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Step-up Therapy for Children with Uncontrolled Asthma Receiving Inhaled Corticosteroids

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ABSTRACT

BACKGROUND

For children who have uncontrolled asthma despite the use of low-dose inhaled corticosteroids (ICS), evidence to guide step-up therapy is lacking.

METHODS

We randomly assigned 182 children (6 to 17 years of age), who had uncontrolled asthma while receiving 100 μ g of fluticasone twice daily, to receive each of three blinded step-up therapies in random order for 16 weeks: 250 μ g of fluticasone twice daily (ICS step-up), 100 μ g of fluticasone plus 50 μ g of a long-acting beta-agonist twice daily (LABA step-up), or 100 μ g of fluticasone twice daily plus 5 or 10 mg of a leukotriene-receptor antagonist daily (LTRA step-up). We used a triple-cross-over design and a composite of three outcomes (exacerbations, asthma-control days, and the forced expiratory volume in 1 second) to determine whether the frequency of a differential response to the step-up regimens was more than 25%.

RESULTS

A differential response occurred in 161 of 165 patients who were evaluated (P<0.001). The response to LABA step-up therapy was most likely to be the best response, as compared with responses to LTRA step-up (relative probability, 1.6; 95% confidence interval [CI], 1.1 to 2.3; P=0.004) and ICS step-up (relative probability, 1.7; 95% CI, 1.2 to 2.4; P=0.002). Higher scores on the Asthma Control Test before randomization (indicating better control at baseline) predicted a better response to LABA step-up (P=0.009). White race predicted a better response to LABA step-up, whereas black patients were least likely to have a best response to LTRA step-up (P=0.005).

CONCLUSIONS

Nearly all the children had a differential response to each step-up therapy. LABA step-up was significantly more likely to provide the best response than either ICS or LTRA step-up. However, many children had a best response to ICS or LTRA step-up therapy, highlighting the need to regularly monitor and appropriately adjust each child's asthma therapy. (ClinicalTrials.gov number, NCT00395304.)

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NCONTROLLED ASTHMA OCCURS IN many children who receive treatment with low-dose inhaled corticosteroids. In the Pediatric Asthma Controller Trial (PACT),¹ administration of 100 μ g of fluticasone twice daily was the most effective therapy, but uncontrolled asthma occurred in more than 50% of the children, and 39% of the children had at least one asthma exacerbation that was treated with oral corticosteroids during a 48-week period. The few data that are available to guide practitioners about how to treat children whose asthma is poorly controlled while they are receiving low-dose inhaled corticosteroids2-8 are inconsistent. The discrepancies may be related to the choice of a single primary outcome measure (e.g., peak expiratory flow); to differences in the duration of treatment, eligibility criteria, or choice of comparison groups; and to evaluation of the responses of groups rather than of individual patients. Evidence from comparison studies is needed to establish which step-up therapy is best and whether there are phenotypic or genotypic characteristics that can be used to predict whether a child will have a better response to one particular step-up treatment than to another.

In this study, called the Best Add-on Therapy Giving Effective Responses (BADGER) trial, we assessed the frequency of differential responses to three blinded step-up treatments in children who had uncontrolled asthma while receiving lowdose inhaled corticosteroids. We used a threeway crossover design with a composite of outcomes (asthma exacerbations, asthma-control days, and the forced expiratory volume in 1 second [FEV₁]), which allowed us to determine the probability that a given treatment would produce the best response. We also determined whether specific baseline characteristics could be used to predict the response to step-up treatment.

METHODS

STUDY PATIENTS

From March 2007 through July 2008, we recruited patients, 6 to 17 years of age, at Childhood Asthma Research and Education (CARE) Network centers. Inclusion criteria were mild-to-moderate asthma diagnosed by a physician on the basis of criteria recommended by the National Asthma Education and Prevention Program,⁹ an ability to perform reproducible spirometry, an FEV₁ of at least 60% before bronchodilation, and an in-

crease in the FEV_1 of at least 12% (bronchodilator reversibility) or a methacholine provocation concentration causing a 20% fall (PC₂₀) in the FEV₁ of 12.5 mg per milliliter or less.

Each center's institutional review board approved the study, and parents or guardians provided written informed consent. In addition, children under 7 years of age provided oral assent, and older children provided written assent.

STUDY PROTOCOL

The protocol that we used to guide study procedures is provided in the Supplementary Appendix, available with the full text of this article at NEJM .org. Briefly, all patients were enrolled in a run-in period of 2 to 8 weeks to determine whether their asthma was poorly controlled while they were receiving 100 μ g of fluticasone twice daily. The run-in period could be shortened by up to 1 week for safety reasons in case of a sudden worsening in symptoms. Patients or their parents or guardians recorded symptoms and peak-flow determinations in a diary each day. A patient became eligible for randomization after the documentation of uncontrolled asthma, which was defined as the occurrence of at least one of the following for more than 2 days per week on average during a 2-week period: diary-reported symptoms (coughing rated as moderate or severe or wheezing rated as mild, moderate, or severe), rescue use of an inhaled bronchodilator with two or more puffs per day, or peak flows under 80% of the predetermined reference value.

Patients then entered a randomized, doubleblind, three-treatment, three-period crossover trial for a total of 48 weeks. During each 16-week period, patients received 250 μ g of fluticasone (Flovent Diskus, GlaxoSmithKline) twice daily (inhaled-corticosteroid [ICS] step-up therapy), 100 μ g of fluticasone plus 50 μ g of the longacting beta-agonist salmeterol (Advair Diskus, GlaxoSmithKline) twice daily (LABA step-up therapy), or 100 μ g of fluticasone twice daily plus 5 or 10 mg of the leukotriene-receptor antagonist montelukast (Singulair, Merck) daily (LTRA stepup therapy). The drug assignments were masked with the use of placebo tablets and dummy disk devices that discharged powder without the active drug. The initial 4 weeks of the last two 16-week periods were considered to be the active washout from the previous period.

Patients received an open-label metered-dose inhaler of albuterol (Ventolin HFA, GlaxoSmith-

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Kline), prednisone, and a customized written action plan to guide use. A standardized course of prednisone treatment was initiated for an asthma exacerbation if predetermined clinical criteria were met. Asthma characteristics were assessed by means of CARE Network procedures that have been described previously.¹⁰ Patients were evaluated at 4-week intervals (Fig. 1). Measurements of the fraction of exhaled nitric oxide and methacholine bronchoprovocation were performed as described previously.1 We administered the validated Pediatric Asthma Quality of Life Questionnaire¹¹ to assess patients' impairment from asthma, with scores ranging from 1 to 7 and higher scores indicating less impairment (with a minimally important difference [MID] of 0.5); the Asthma Control Test^{12,13} (for children 12 years of age or older), with scores ranging from 5 to 25 and higher scores indicating greater control (with an MID of 3.0); and the Childhood Asthma Control Test¹⁴ (for children between the ages of 4 and 11 years), with scores ranging from 0 to 27 and higher scores indicating greater control (with an undefined MID).

OUTCOME MEASURES

The primary outcome was the differential response to each of the three step-up therapies on the basis of fixed threshold criteria for the following three asthma-control measures: the need for treatment with oral prednisone for acute asthma exacerbations, the number of asthmacontrol days, and the FEV₁. One treatment period was ranked as better than another if the total amount of prednisone received during the period was at least 180 mg less, if the number of annualized asthma-control days during the final 12 weeks of the period was increased by at least 31 days, or if the FEV₁ at the end of the period was at least 5% higher. If the prednisone threshold was met, then we ignored the number of asthmacontrol days and the FEV₁. If the threshold for asthma-control days was met, then we ignored the FEV₁. Otherwise, the order of response was determined by the FEV₁. If none of the thresholds were met, then we gave the same rank to each treatment period and the patient was considered not to have had a differential response. A patient was considered to have had a differential response if at least one treatment period was ranked as better than another.

Oral prednisone was administered according to a protocol that has been described previously.¹

An asthma-control day, as documented in each patient's diary, was a day during which there was no use of albuterol rescue (excluding the use of albuterol as preexercise treatment), no use of a nonstudy asthma medication, no daytime or nighttime asthma symptoms, no unscheduled visit to a health care provider for asthma, and no peak expiratory flow of less than 80% of the predetermined reference value. We calculated annualized asthma-control days as 365 times the proportion of asthma-control days during the final 12 weeks of the period, which were adjusted for seasonal differences.

STUDY OVERSIGHT

Fluticasone and salmeterol were donated by the manufacturer, GlaxoSmithKline, which also donated the albuterol that was provided to patients; montelukast was donated by Merck. The manufacturers had no role in the study design, the accrual or analyses of data, or the preparation of the manuscript. The authors vouch for the completeness and accuracy of the data.

STATISTICAL ANALYSIS

The primary analysis involved two stages: a onesided exact test for binomial proportions to test the null hypothesis that the percentage of patients with a differential response (i.e., those for whom the three treatments were not ranked the same) would be less than or equal to 25% at the 0.01 significance level, and if there was a significant result in stage one, a rank-ordered logistic regression¹⁵ to test whether four prespecified baseline covariates would predict distinct patterns of differential responses. Each covariate was tested at the 0.01 significance level so that the overall type I error rate for the primary analysis was 0.05. Secondary analyses tested the predictive values of other baseline covariates at the 0.05 significance level.

The number of patients who were enrolled in the study provided adequate power for both stages of the primary analysis. A total sample of 180 patients, with an assumed 15% loss to follow-up, would provide a power of 90% for the first stage if the true percentage of patients with a differential response was at least 40%, and would provide the same power for each of the four prespecified covariates in the second stage if the covariate predicted a change of more than 15% in the probability of a best response for each treatment. We constructed rank-ordered logit models and calculated bootstrap-based 95% confidence intervals with the use of Stata software, version 11. Classification analysis was used to examine the goodness of fit of the model. Model results are presented for dichotomized covariates to simplify interpretation of continuous predictors. Sensitivity analyses also modeled them as ordinal covariates (quartiles) and continuous covariates to confirm that dichotomization did not mask relationships. Sensitivity analyses assessed the possible influence of treatment carryover effects by testing for period effect and treatment-by-period interaction effect in the regression models.

RESULTS

STUDY PATIENTS

Of the 480 patients who were enrolled, 298 were excluded during the run-in phase. Thus, 182 patients underwent randomization, of whom 157 completed all three study periods (Fig. 1). A total of 165 patients completed two study periods, permitting assessment of the differential response. Table 1 presents baseline demographic and physiological data, stratified according to age group. (Baseline data stratified according to the studygroup assignment during period 1 are available in Table E1 in the Supplementary Appendix.) Patients completed 90% of the study visits and provided sufficient data in the symptom diaries to determine control status on 96% of study days. The rate of adherence to study medication was 84% for study tablets (as measured by an electronic monitor in the bottle cap) and 87% for study inhalers (as measured by a disk counter).

DIFFERENTIAL RESPONSE TO THE THREE STEP-UP THERAPIES

A differential response occurred in 161 of 165 patients (98%), a proportion that was significantly greater than the one-sided null hypothesis of 25% (P<0.001). The percentage of asthma-control days differed according to season in all study groups, ranging from 71% in winter months to 79% in summer months. Asthma exacerbations were most frequent during winter months. The average FEV₁ (measured as a percentage of the predicted value) varied by less than 1% across seasons. Seasonal variation affected only 12% of patients for whom the number of annualized asthma-control days determined the differential response. Sensitivity analysis indicated that seasonal differences in the FEV_1 had no significant effect on the results, whereas seasonal differences in exacerbations had a small effect that was neither statistically nor qualitatively significant.

Subsequent analyses were performed to characterize the pattern of differential response. In pairwise comparisons, the proportion of patients who had a better response to LABA step-up was higher than the proportion with a better response to LTRA step-up (52% vs. 34%, P=0.02), and the proportion with a better response to LABA stepup was higher than the proportion with a better response to ICS step-up (54% vs. 32%, P=0.004), whereas the responses to LTRA and ICS step-up therapies were similar (Fig. 2A). The primary outcome of the trial, a three-way comparison of step-up therapy with the use of rank-ordered logistic regression, predicted that the response to LABA step-up was significantly more likely to be the best response, as compared with the response to LTRA step-up (relative probability, 1.6; 95% confidence interval [CI], 1.1 to 2.3; P=0.004) and the response to ICS step-up (relative probability, 1.7; 95% CI, 1.2 to 2.4; P=0.002) (Fig. 2B).

PRIMARY ANALYSIS OF PREDICTORS OF DIFFERENTIAL RESPONSE

The ability of the four preselected factors to predict patterns of differential response is shown in Figure 3. Patterns of differential response were not predicted by methacholine PC20 values, either dichotomized at the median baseline value (P=0.2) or examined as a continuous covariate (P=0.37) (Fig. 3A), or by the fraction of exhaled nitric oxide, either dichotomized at the median baseline value (P=0.7) or examined as a continuous covariate (P=0.52) (Fig. 3B). Baseline scores on the Asthma Control Test and the Childhood Asthma Control Test that were dichotomized at validated numerical scores indicative of acceptable control (>19) or unacceptable control (\leq 19) significantly predicted patterns of differential response (P=0.009), with higher scores predicting a greater probability that the best response would be to LABA step-up (Fig. 3C). Finally, the genotype at position 16 of the β_2 -adrenergic receptor did not predict patterns of response (P=0.49) (Fig. 3D).

SECONDARY ANALYSIS OF PREDICTORS OF DIFFERENTIAL RESPONSE

We evaluated other factors post hoc, including demographic and physiological characteristics and asthma history, as predictors of a differential re-

STEP-UP THERAPY FOR CHILDREN WITH UNCONTROLLED ASTHMA

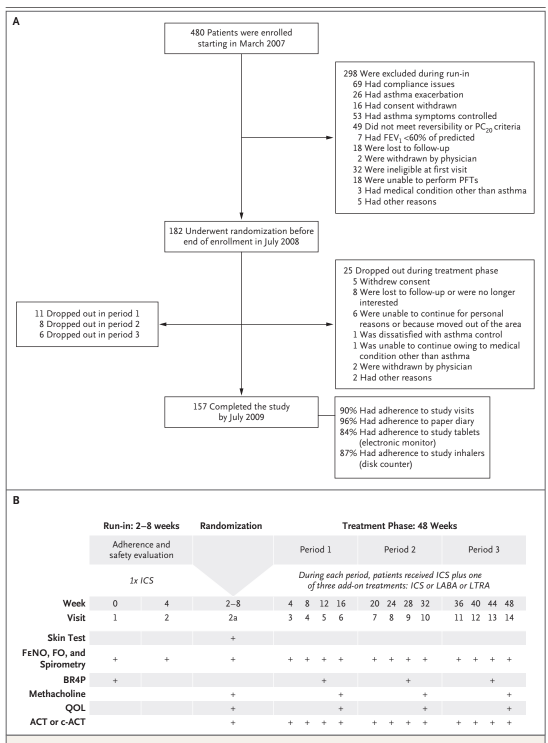


Figure 1. Enrollment and Outcomes and Schedule of Evaluations.

Panel A shows the number of patients who enrolled in the study, underwent randomization, and completed the study. FEV_1 denotes forced expiratory volume in 1 second, PC_{20} provocation concentration of methacholine causing a 20% fall in FEV₁, and PFT pulmonary-function test. Panel B shows the schedule for various tests and procedures and the administration of the Pediatric Asthma Quality of Life Questionnaire to assess asthma impairment at specific study visits. Visit 2a was the randomization visit. ACT denotes Asthma Control Test, BR4P bronchodilator response to four puffs of albuterol, c-ACT Childhood Asthma Control Test, FENO fraction of exhaled nitric oxide, FO forced oscillometry, and QOL quality-of-life questionnaire.

Characteristic	Age Group	
	6-11 Yr (N=126)	12–17 Yr (N=56)
Age — yr	9.1±1.5	14.7±1.7
Male sex — no. (%)	83 (66)	36 (64)
Race or ethnic group — no. (%)†		
Hispanic or Latino	38 (30)	22 (39)
Non-Hispanic white	54 (43)	20 (36)
Black	37 (29)	12 (21)
Hispanic white	28 (22)	15 (27)
Other	7 (6)	9 (16)
Height — cm	134.3±10.8	164.2±11.0
Weight — kg	36.1±12.7	63.4±17.2
Body-mass index	19.6±4.5	23.3±4.8
Age at asthma diagnosis — yr	3.3±2.2	4.7±4.1
Age at onset of asthma symptoms — yr	2.4±2.2	3.8±3.6
Family history of asthma — no. (%)		
Father	33 (26)	17 (30)
Mother	44 (35)	12 (21)
Eczema — no. (%)	69 (55)	24 (43)
Positive aeroallergen skin test		
Any — no.	2.6±2.1	3.7±2.4
≥1 — no./total no. (%)	95/122 (78)	48/55 (87)
Positive perennial skin test	/ (-/	
Any — no.	1.4±1.3	2.0±1.4
≥1 — no./total no. (%)	82/122 (67)	48/55 (87)
Serum IgE — IU/ml	493.4±690.5	530.5±589.0
Blood eosinophils — %	5.1±3.7	5.3±4.8
Eligibility classification — no. (%)‡	0111011	0.01
Step-up	35 (28)	14 (25)
Step-neutral	16 (13)	11 (20)
Step-down	75 (60)	31 (55)
Medication use in previous year — no. (%)	, 5 (00)	51 (55)
Inhaled or nebulized corticosteroid	82 (65)	39 (70)
Leukotriene modifier	46 (37)	14 (25)
Salmeterol	5 (4)	5 (9)
Theophylline	0	0
Cromolyn or nedocromil	0	
Salmeterol plus fluticasone, or budesonide plus formoterol	41 (33)	1 (2) 18 (32)
Prednisone (≥ 1 courses) Prebronchodilator FEV ₁ — % of predicted value	56 (44) 98.5±13.1	25 (45) 95.0±14.8
Prebronchodilator FEV ₁ :FVC ratio	81.6±7.1	78.2±7.1
Bronchodilator response, 4 puffs — % Asthma-control days during worst 2 weeks of run-in period — %	11.2±11.2 30±21	13.5±10.2 36±23

Table 1. (Continued.)		
Characteristic	Age Group	
	6–11 Yr (N=126)	12–17 Yr (N=56)
Score on Asthma Control Test or Childhood Asthma Control Test§	20.5±3.8	19.8±3.4
Exhaled nitric oxide — ppb		
Median	8.5	17.7
Interquartile range	5.8-13.1	11.4–26.2
Methacholine PC ₂₀ — mg/ml		
Median	1.24	2.06
Interquartile range	0.56-3.83	0.70-5.44
β_2 -adrenergic-receptor genotype — no. (%)¶		
Arg/Arg	19 (15)	10 (18)
Arg/Gly	71 (57)	31 (55)
Gly/Gly	34 (27)	15 (27)

* Plus-minus values are means ±SD. The body-mass index is the weight in kilograms divided by the square of the height in meters. Percentages may not total 100 because of rounding. FEV₁ denotes forced expiratory volume in 1 second, FVC forced vital capacity, and PC₂₀ provocation concentration causing a 20% fall in FEV₁.

† Race or ethnic group was self-reported.

 \ddagger The run-in medication was 100 μ g of fluticasone twice daily. The category for patients who were taking an equivalent dose of an inhaled corticosteroid at baseline was termed "step-neutral," the category for those who were not receiving an inhaled corticosteroid or were taking a lower dose at baseline was termed "step-up," and the category for those who were taking a higher dose of an inhaled corticosteroid or combination therapy at baseline was termed "step-down."

§ Scores on the Asthma Control Test (for patients 12 years of age or older) are measured on a scale of 5 to 25, with higher scores indicating greater control. Scores on the Childhood Asthma Control Test (for children between the ages of 4 and 11 years) are measured on a scale from 0 to 27, with higher scores indicating greater control.

¶ Data were missing for two patients between the ages of 6 and 11 years.

sponse. Figure 4 shows the clinical relevance of three of these factors: race or ethnic group, the presence or absence of eczema, and age. Race or ethnic group significantly predicted patterns of differential response (P=0.005 without correction for multiple testing), with Hispanic and non-Hispanic white patients most likely to have a best response to LABA step-up and least likely to have a best response to ICS step-up, and black patients equally likely to have a best response to LABA or ICS step-up therapy and less likely to have a best response to LTRA step-up (Fig. 4A). There were no differences in the patterns of differential response according to age, whether age was examined as a dichotomous covariate (5 to 11 years or \geq 12 years) or as a continuous covariate (Fig. 4B). Finally, patients who did not have eczema were most likely to have a best response to LABA stepup (P=0.006 without correction for multiple testing) (Fig. 4C).

Of the seven other factors examined post hoc to predict patterns of differential response, none were significant as either dichotomous or continuous covariates. Factors that were evaluated

were sex (P=0.10), presence or absence of perennial allergen sensitization (P=0.16), baseline prebronchodilator FEV₁ of 90% or less or more than 90% (P=0.70), bronchodilator reversibility (>10% vs. \leq 10%) (P=0.08), previous use or nonuse of a controller medication (P=0.90), the baseline number of asthma-control days (P=0.08), and the number of recent asthma exacerbations (0 vs. \geq 1) (P=0.90).

The rank-ordered logit model that included only the significant predictors correctly classified the observed ranks 68% of the time. The interaction between period and treatment according to period was not significant, indicating that it was unlikely that effects of treatment carryover could have biased the results.

ADVERSE EVENTS

There were seven serious adverse events, one during the run-in period (asthma exacerbation) and six during the treatment phase (five of which required hospitalization): two each in the LABA step-up group (asthma exacerbation and appendicitis), the LTRA step-up group (asthma exacer-

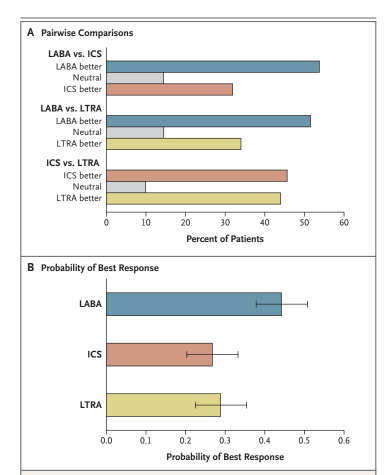


Figure 2. Pairwise Comparison of Three Step-up Therapies and the Overall Probability of Best Response.

Panel A shows pairwise comparisons of the three step-up therapies. The proportion of patients with a best response to a long-acting beta-agonist (LABA) step-up was higher than the proportion with a best response to a leukotriene-receptor antagonist (LTRA) step-up (52% vs. 34%, P=0.02) or an inhaled corticosteroid (ICS) step-up (54% vs. 32%, P=0.004); the best-response results for LTRA step-up were similar to those for an ICS step-up. Panel B shows the pattern of differential response according to the probability that the best response to a treatment occurred during the period in which that treatment was received; LABA step-up was most likely to provide the best response. Covariate-adjusted estimates were obtained from rank-ordered logistic-regression models; I bars indicate bootstrap-based 95% confidence intervals.

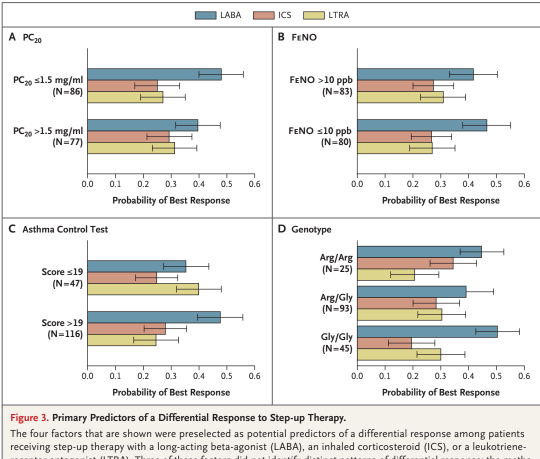
bation and need for tonsillectomy–adenoidectomy), and the ICS step-up group (asthma exacerbation and behavioral issues not requiring hospitalization). Thrush that was classified as probably related to the study drug was reported in one patient during treatment with LABA step-up and in two patients during treatment with ICS step-up. A total of 25 treatment failures occurred: 3 hospitalizations for asthma-related symptoms (1 during each study treatment) and 22 involving the need for a second prednisone burst (3 during treatment with LABA, 8 during treatment with ICS, and 11 during treatment with LTRA). A total of 120 prednisone bursts were prescribed for asthma exacerbations (30 during treatment with LABA, 47 during treatment with ICS, and 43 during treatment with LTRA) (for details, see Table E2 in the Supplementary Appendix).

DISCUSSION

In this trial, we examined step-up strategies for children who had uncontrolled asthma while receiving low-dose inhaled corticosteroids. When the three step-up treatments were compared, 98% of the children had a differential response, with the best response during LABA step-up occurring significantly more frequently than during LTRA or ICS step-up. The direction of this best response should not have been biased by the entry criteria, because 20% of the children qualified for the study on the basis of bronchodilator reversibility alone, 58% qualified on the basis of methacholine hyperresponsiveness alone, and 22% met both criteria. Moreover, FEV1 values after bronchodilation were not significant predictors of the response to LABA step-up, providing further evidence against this bias.

Our findings suggest that there is a ceiling effect of low-dose inhaled corticosteroids in many, though not all, children. Our data show that to achieve improvements in asthma control, the addition of a different class of medication is often required. Furthermore, although LABA step-up therapy was most likely to provide the best response, some children had a best response to one of the other step-up therapies. For the most part, baseline characteristics did not predict the therapy most likely to be best for an individual patient. We found that baseline scores on the Asthma Control Test and the Childhood Asthma Control Test indicating better control were associated with a higher probability that LABA would be the best add-on therapy. The predictive value of other characteristics may have been limited by run-in treatment — for example. baseline values for the fraction of exhaled nitric oxide and PC20 were obtained after at least 2 weeks of low-dose inhaled corticosteroids, unlike the predictive value of these measures in children who had not received inhaled corticosteroids, as reported in PACT.¹⁶

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receiving step-up therapy with a long-acting beta-agonist (LABA), an inhaled corticosteroid (ICS), or a leukotrienereceptor antagonist (LTRA). Three of these factors did not identify distinct patterns of differential response: the methacholine provocation concentration causing a 20% fall (PC₂₀) in the forced expiratory volume in 1 second (Panel A), the fraction of exhaled nitric oxide (FENO) (Panel B), and the genotype for the β_2 -adrenergic receptor (Panel D). The fourth factor — the baseline score on the Asthma Control Test (for children 12 years of age or older, measured on a scale of 5 to 25) or the Childhood Asthma Control Test (for children between the ages of 4 and 11 years, measured on a scale from 0 to 27, with higher scores on both tests indicating greater control) — did identify distinct patterns of differential response (P=0.009), with scores of more than 19 predicting an increased likelihood of a best response to LABA step-up therapy (Panel C). Estimates were obtained from rank-ordered logistic-regression models; I bars indicate bootstrap-based 95% confidence intervals.

In secondary analyses, findings regarding self-reported race or ethnic group were noteworthy. Black patients were equally likely to have a best response to LABA or ICS step-up therapy and were least likely to have a best response to LTRA step-up. In contrast, in white subjects, LABA step-up was clearly most likely to provide the best response. A number of studies have reported a differential response to asthma therapy on the basis of race or ethnic group.^{17,18} These differences appear to primarily involve responsiveness to beta-agonists, which may be related to genetic variations¹⁹ and gene-by-gene interactions.²⁰ The observed best overall performance of LABA therapy should be weighed against the potential increased risks of such therapy that are cited in the Food and Drug Administration's approved labeling.²¹ Our study was not designed or powered to evaluate the long-term safety of LABA in children. Although we found that LABA step-up therapy was associated with the smallest number of treatment failures and exacerbations, the between-group differences were not significant, and we emphasize that the duration of our trial and the size of our sample preclude statements regarding long-term risks. Clinicians who prescribe LABAs (never to be used as monothera-

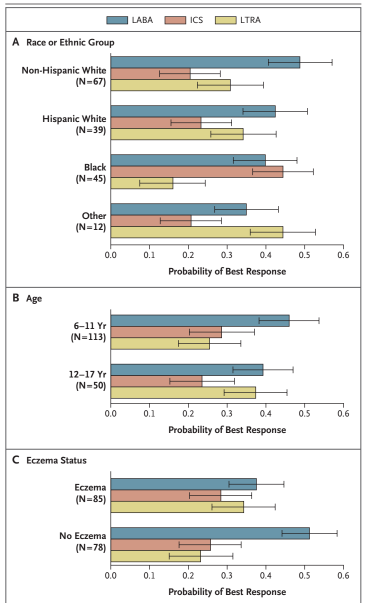


Figure 4. Secondary Predictors of a Differential Response to Step-up Therapy. Shown is the probability of a best response to step-up therapy with a longacting beta-agonist (LABA), an inhaled corticosteroid (ICS), or a leukotriene-receptor antagonist (LTRA), according to race or ethnic group, age, and the presence or absence of eczema, factors that were evaluated in post hoc analysis. Race or ethnic group was a significant predictor (P=0.005), with Hispanic and non-Hispanic white patients most likely to have a best response to LABA step-up therapy and black patients equally likely to have a best response to LABA or ICS step-up and least likely to have a best response to LTRA step-up (Panel A). Age was not a significant predictor of a differential response (Panel B). Patients who did not have eczema were most likely to have a best response to LABA step-up therapy (P=0.006) (Panel C). Estimates were obtained from rank-ordered logistic-regression models; I bars indicate bootstrap-based 95% confidence intervals. P values have not been corrected for multiple testing. py^{22,23}) in combination with inhaled corticosteroids should continue to evaluate risk–benefit ratios.

Despite step-up in daily therapy, 120 exacerbations requiring the use of oral corticosteroids occurred during the treatment periods. Day-today asthma control, as reflected by the number of asthma-control days, was quite good with all three step-up therapies. However, none of the step-up therapies completely prevented asthma exacerbations. These findings underscore the need for new therapies aimed at preventing exacerbations.

In summary, our study showed a clinically significant differential response in nearly all the children and identified several characteristics of the children that predicted the direction of differential responses, including race or ethnic group and two readily available clinical attributes: asthma control, as indicated by the score on the Asthma Control Test, and the presence or absence of eczema. More expensive and labor-intensive measures of physiological factors (e.g., methacholine PC₂₀) and biomarkers (e.g., the fraction of exhaled nitric oxide) did not have predictive value. Although LABA step-up was significantly more likely to provide the best response than either ICS or LTRA step-up, many children had a best response to ICS or LTRA step-up, highlighting the need to regularly monitor and appropriately adjust each child's asthma therapy within this level of care before further step-up.

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