Bronchodilator Responsiveness Defined by the 2005 and 2021 ERS/ATS Criteria in Patients with Asthma as Well as Chronic Obstructive Pulmonary Disease

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Background: In the 2021 ERS/ATS interpretive strategies for routine lung function tests, a positive bronchodilator response (BDR) was updated as a change of >10% relative to the predicted value in forced expiratory volume in 1 second (FEV\textsubscript{1}) or forced vital capacity (FVC). We aimed to explore the differences between the 2005 and 2021 ERS/ATS criteria applied to patients with asthma as well as chronic obstructive pulmonary disease (COPD).

Methods: BDR test data about asthma patients aged 6–80 years and COPD patients aged 18–80 years were derived from the National Respiratory Medicine Center, First Affiliated Hospital of Guangzhou Medical University, from January 2017 to March 2022. BDR results defined by the 2005 and 2021 ERS/ATS criteria were named 2005-BDR and 2021-BDR, respectively. We compared differences between 2005-BDR and 2021-BDR and analyzed the trend in the proportion of positive BDR (BDR+) with the level of airflow obstruction.

Results: A total of 4457 patients with asthma and 7764 patients with COPD were included in the analysis. The percentages of 2005-BDR+ and 2021-BDR+ were 63.32% and 52.84% for asthma, 30.92% and 22.94% for COPD, respectively. Of patients with 2005-BDR+, 81.86% for asthma and 70.18% for COPD showed 2021-BDR+ results, and these patients had higher FEV\textsubscript{1}\%pred, FVC\%pred (all P<0.05). Whichever BDR criterion was adopted, the proportion of BDR+ had an upward linear trend with the increased degree of airflow obstruction in COPD, but exhibited an approximate inverted U-shaped curve in asthma. In COPD, the proportion of BDR\textsubscript{FEV1} was negatively associated with the degree of airflow obstruction, while BDR\textsubscript{FVC} was positively associated (all P<0.05).

Conclusion: Compared with 2005-BDR+, the proportion of 2021-BDR+ reduced markedly in patients with asthma and COPD, but their trends with the degree of airflow obstruction did not change. Patients with consistent BDR+ had higher initial FEV\textsubscript{1}\%pred and FVC\%pred.

Keywords: pulmonary function test, bronchodilator responsiveness testing, asthma, chronic obstructive pulmonary disease, European Respiratory Society, ERS, American Thoracic Society, ATS

Introduction

Pulmonary function tests (PFTs) have played a vital role in the screening, diagnosis and management of respiratory diseases. In order to improve the quality control and interpretation of PFTs, a series of technical documents have been jointly published by the American Thoracic Society (ATS) and European Respiratory Society (ERS).\textsuperscript{1–7} Bronchodilator responsiveness (BDR) testing, widely used to assess the degree of volume and airflow improvement in response to an...
inhaled short-acting bronchodilator, is considered to be a key diagnostic criterion for asthma, as well as differentiating asthma from chronic obstructive pulmonary disease (COPD).

However, recommendations on how to express a bronchodilator response are still controversial. Three approaches are commonly used to express BDR, including an absolute change from the initial value, a relative change to the initial value, and a change relative to the predicted value. The 2005 ERS/ATS interpretive strategies for routine lung function tests recommended that a change in forced expiratory volume in 1 second (FEV$_1$) or forced vital capacity (FVC) ≥12% and ≥200mL of the initial value was defined as significant BDR. Nevertheless, the absolute and relative changes in FEV$_1$ and FVC are closely associated with initial lung function, and biased toward height, age and sex in both health and disease. Furthermore, requiring a fixed minimum change in FEV$_1$ ≥200mL seems unrealistic, since in many subjects with a low baseline FEV$_1$, especially children and elder adults, the FEV$_1$ increased >12% of baseline but <200mL.

Previous studies showed that BDR expressed as the percent change relative to the individual’s predicted value might minimize sex and size bias. In a study involving 3922 healthy individuals, the 95th percentile of BDR was estimated to be 10.0% (9.5% to 10.5%) of predicted value for FEV$_1$ and 9.2% (7.9% to 10.5%) of predicted value for FVC. In another study of 2371 healthy non-smokers, the 95th percentiles of BDR in FEV$_1$ and FVC were 11.6% and 10.2%, respectively. For the preschool children aged 3–5 years, the 95th percentiles of the change in FEV$_1$, FEF$_{25–75}$%, FEV$_{0.75}$, and FEV$_{0.5}$ relative to the predicted value following bronchodilator were 11.6%, 16.0%, 8.5%, and 35.5%, respectively. Therefore, based on these considerations, the 2021 ERS/ATS interpretive strategies updated the positive response as a change of >10% relative to the predicted value for FEV$_1$ or FVC. For example, a man has a pre-bronchodilator FEV$_1$ of 2.0L, a post-bronchodilator FEV$_1$ of 2.4L and his predicted FEV$_1$ is 3.2L, then the BDR = (2.4−2.0)*100/3.2=12.5%, which is >10% and classified as a positive response.

However, there are limited data about making a comparison between the two criteria applied to clinical practice. Since BDR is mostly used in the diagnosis of obstructive pulmonary diseases, the present study aimed to compare the differences between the two BDR criteria in a large sample of patients with asthma as well as COPD.

**Materials and Methods**

**Methods**

This was a retrospective study and was performed in accordance with the Declaration of Helsinki and approved by the Ethics Committee of First Affiliated Hospital of Guangzhou Medical University (2020–124).

**Data Resource and Study Population**

The data were derived from the Respiratory Health Big Database of the National Respiratory Medicine Center, First Affiliated Hospital of Guangzhou Medical University. BDR testing reports from January 2017 to March 2022 were extracted from the database and then patients were preliminarily screened according to the International Classification of Diseases Volume 10 (ICD-10) and Systematized Nomenclature of Medicine Clinical Terms (SNOMED) standard terminology. The rationality of the diagnosis was checked by reviewing the electronic medical records. Patients with asthma were clinically diagnosed according to the guideline of the Global Initiative for Asthma (GINA) and aged 6–80 years old. Patients with COPD were diagnosed according to the guideline of the Global Initiative for Chronic Obstructive Lung Disease (GOLD) and aged 18–80 years old. Excluded criteria were as followings: patients having a history of interstitial lung diseases or pulmonary tuberculosis or severe bronchiectasis; those with ambiguous diagnosis; those diagnosed with obliterative bronchiolitis; those with missing important parameters such as age, weight, FEV$_1$, FVC and FEV$_1$/FVC. For the subjects who have performed BDR testing more than once, only the initial report was selected.

**Spirometry Tests**

Spirometry tests were performed by trained technicians according to the relevant guidelines by ERS/ATS and Pulmonary Function Group, Respiratory Diseases Society of Chinese Medical Association. All the subjects signed consent form before tests. At least three acceptable curves were needed. Each subject inhaled 100μg/puff sprays of salbutamol (Ventolin, Glaxo Wellcome Products, France) from an inhaler for totally 4 puffs and repeated spirometry after
15 to 30 minutes. Severity of lung function impairment was assessed by FEV₁%pred with 4 critical values of 70%, 60%, 50% and 35%, namely mild: FEV₁%pred ≥70%, moderate: 60% ≤ FEV₁%pred < 70%, moderate-severe: 50% ≤ FEV₁%pred < 60%, severe: 35% ≤ FEV₁%pred < 50%, extremely severe: FEV₁%pred < 35%.

Lung Function Indices and Variable Definitions
Lung function indices were recorded at pre- and post-bronchodilator. The regular indices included FEV₁, FEV₁%pred, FVC, FVC%pred and FEV₁/VCmax (%). The differences between the optimal value of baseline and post-bronchodilator in FEV₁ (or FVC) were called ΔFEV₁, ΔFVC. To simplify the expression, BDR results defined by the 2005 and 2021 ERS/ATS criteria were named 2005-BDR and 2021-BDR, and the positive and negative BDR were named BDR+ and BDR-, respectively. Patients who had the same 2005-BDR and 2021-BDR results were named consistent group, and those who had reversed results were named inconsistent group. Moreover, three BDR+ subgroups were defined as 2005-BDR_{FEV1}, 2005-BDR_{FVC} and 2005-BDR_{Both}, which met the requirement of ΔFEV₁%init ≥ 12% and ΔFEV₁ ≥ 200mL, ΔFVC%init ≥ 12% and ΔFVC ≥ 200mL, and the both of above, respectively. 2021-BDR_{FEV1}, 2021-BDR_{FVC} and 2021-BDR_{Both} were defined as ΔFEV₁%pred > 10%, ΔFVC%pred > 10%, and the both of above, respectively.

Statistical Analysis
Differences were compared between asthma and COPD, between the percentages of 2005-BDR+ and 2021-BDR+, between BDR- and BDR+ groups, between inconsistent and consistent groups. The statistical analysis was performed using SPSS 23 (SPSS, Chicago, Illinois). Results are presented as mean and standard deviation or quartiles for continuous variables and frequency and percentage for categorical variables. Differences in the percentages of 2005-BDR+ and 2021-BDR+ were compared using McNemar test. Differences in baseline characteristics and lung function indices were compared between two groups using a two sample t-test (or Wilcoxon rank sum test, if normality was not met) for continuous variables and a Chi-square test for categorical variables. The continuous variables of three or more groups were compared by one-way analysis of variance as appropriate. The trend in the proportion of BDR+ with the level of airflow obstruction was studied using linear by linear association. The P value <0.05 was considered statistically significant.

Results
Differences in Baseline Characteristics Between Asthma and COPD
A total of 23,072 BDR reports were initially extracted from the database. After rigorous screening, 4457 patients with asthma (male 2201; female 2256) aged 6–80 years and 7764 patients with COPD (male 6817; female 947) aged 18–80 years were included in the final analysis (Figure 1). Compared with asthma patients, COPD patients were greater in percentage of males, age, weight, height [49.4% vs 87.8%, (35, 58) (y) vs (59, 70) (y), 58.39 ± 14.92 (Kg) vs 59.67 ± 10.98 (Kg), 159.12 ± 11.59 (cm) vs 163.68 ± 7.16 (cm), all P<0.05], less in FEV₁, FEV₁%pred, FVC%pred and FEV₁/VCmax [1.57 ± 0.65 (L) vs 1.34 ± 0.66 (L), 58.85 ± 18.81 (%) vs 50.85 ± 22.25 (%), 83.29 ± 18.00 (%) vs 79.66 ± 20.44 (%), 56.03 ± 12.08 (%) vs 48.05 ± 13.00 (%), all P<0.05], and no significant difference in FVC [2.68 ± 0.92 (L) vs 2.65 ± 0.83 (L), P=0.127].

Differences Between the Percentages of 2005-BDR+ and 2021-BDR+
The percentages of 2005-BDR+ and 2021-BDR+ were 63.32% (2822/4457) and 52.84% (2355/4457) for asthma, and 30.92% (2401/7764) and 22.94% (1781/7764) for COPD, respectively. Compared with 2005-BDR+, the percentage of 2021-BDR+ reduced significantly in both asthma and COPD (P<0.001, Table 1). Both the percentages of 2005-BDR+ and 2021-BDR+ were higher in asthma patients than those in COPD patients (P<0.001). When asthma patients were divided into children group aged 6–17 years and adult group aged 18–80 years, the percentage of 2021-BDR+ (64.6%, 306/474) was similar to that of 2005-BDR+ (63.1%,299/474) in children but decreased in adults (51.4% vs 63.3%, P<0.001, Table 2).
Differences Between BDR- and BDR+, Between Inconsistent and Consistent Groups

As shown in Table 3, compared with 2005-BDR-group, the 2005-BDR+ group had higher percentage of males, height, FEV₁pred, ΔFEV₁%init, ΔFEV₁%pred, FVCpred, ΔFVC%init and ΔFVC%pred but less FEV₁, FEV₁%pred, FVC%pred and FEV₁/VCmax (all P<0.05). Conversely, 2021-BDR+ group had lower FEV₁pred, FVCpred than 2021-BDR- group.

**Table 1** The Number of BDR+ and BDR- Defined by Two Criteria in Asthma and COPD

|                | 2021-BDR+ | 2021-BDR- | Total | P value | 2021-BDR+ | 2021-BDR- | Total | P value |
|----------------|-----------|-----------|-------|---------|-----------|-----------|-------|---------|
| Asthma (n=4457)| 2310      | 512       | 2822  | <0.001  | 1685      | 716       | 2401  | <0.001  |
| COPD (n=7764)  | 2005-BDR- | 45        | 1590  | 1635    | 96        | 5267      | 5363  |         |
| 2005-BDR+      | 2355      | 2102      | 4457  |         | 1781      | 5983      | 3764  |         |

*Table 2* The Number of BDR+ and BDR- Defined by Two Criteria in Patients with Asthma

|                | 2021-BDR+ | 2021-BDR- | Total | P value | 2021-BDR+ | 2021-BDR- | Total | P value |
|----------------|-----------|-----------|-------|---------|-----------|-----------|-------|---------|
| Children (n=474)| 284       | 15        | 299   | 0.324   | 2026      | 497       | 2523  | <0.001  |
| Adults (n=3983)| 2005-BDR- | 22        | 153   | 175     | 23        | 1437      | 1460  |         |
| 2005-BDR+      | 306       | 168       | 474   |         | 2049      | 1934      | 3983  |         |
Table 3 Differences in Pulmonary Function Indices Between 2005-BDR- and 2005-BDR+ Groups

|                  | Asthma (n=4457) | COPD (n=7764) | t or χ² or Z | P value |
|------------------|-----------------|---------------|-------------|---------|
|                  | 2005-BDR-(n=1635) | 2005-BDR+(n=2822) | t or χ² or Z | P value |
| Male, n (%)      | 697 (42.6)      | 1504 (53.3)   | 47.11       | <0.001  |
| Female, n (%)    | 938 (57.4)      | 1318 (46.7)   | −1.661      | 0.097   |
| Age(year)        | (33.00, 58.00)  | (35.00, 59.00) | −4.369      | <0.001  |
| Weight(Kg)       | 57.16±14.49     | 59.10±14.21   | −6.199      | <0.001  |
| Height(cm)       | 158.07±11.92    | 159.73±11.35  | −4.403      | <0.001  |
| FEV₁ pred (L)    | 3.14±0.86       | 3.27±0.87     | −5.743      | <0.001  |
| FEV₁ (L)         | 2.61±0.71       | 2.71±0.71     | −4.043      | <0.001  |
| FEV₁ %pred       | 62.99±21.52     | 65.45±16.58   | −6.148      | <0.001  |
| ΔFEV₁ (mL)       | 91.53±88.65     | 136.18±190.33 | −65.148     | <0.001  |
| ΔFEV₁%init       | 6.13±6.55       | 26.01±15.00   | −61.046     | <0.001  |
| ΔFEV₁%pred       | 3.64±3.48       | 13.69±6.33    | −68.313     | <0.001  |
| FVC pred (L)     | 3.14±0.86       | 3.27±0.87     | −5.029      | <0.001  |
| FVC(L)           | 2.64±0.95       | 2.70±0.90     | −2.042      | 0.041   |
| FVC%pred         | 84.39±19.93     | 82.65±16.75   | 2.963       | 0.003   |
| ΔFVC (mL)        | 57.05±121.84    | 332.29±42.92  | −50.259     | <0.001  |
| ΔFVC%init        | 2.47±5.37       | 13.87±11.15   | −45.964     | <0.001  |
| ΔFVC%pred        | 1.86±4.07       | 10.48±7.26    | −50.761     | <0.001  |
| FEV₁/VCmax (%)   | 59.26±13.30     | 54.16±10.88   | 13.157      | <0.001  |

(Table 4). As shown in Table 5, in asthma, compared with inconsistent group with 2005-BDR+ and 2021-BDR- (18.14%, 512/2822), consistent group with 2005-BDR+ and 2021-BDR+ (81.86%, 2310/2822) had lower percentage of males, weight, height, FVC, FEV₁ pred and FVCpred, but higher FEV₁ %pred, FVC %pred (all P<0.05). In COPD, compared with inconsistent group (29.82%, 716/2401), consistent group (70.18%, 1685/2401) had lower percentage of males, height, FEV₁ pred and FVC pred, but higher age, FEV₁, FVC, FEV₁ %pred, FVC %pred (all P<0.05). Additionally, of 1635 asthma patients with 2005-BDR-, 45 (2.75%) converted to 2021-BDR+; of 5363 COPD patients with 2005-BDR-, 96 (1.79%) converted to 2021-BDR+.

The Trend in the Proportion of BDR+ with the Level of Airflow Obstruction

As shown in Figure 2, with the increasing degree of airflow obstruction, the percentage of 2005-BDR+ increased linearly in COPD (P<0.001) but exhibited an approximate inverted U-shaped curve in asthma (P=0.097). Similar but more flat trends were observed in 2021-BDR (P<0.001, P=0.949). As shown in Figure 3, in asthma, the proportion of 2005-BDR_FEV₁ was negatively associated with the level of airflow obstruction, while 2005-BDR_FVC was positively associated (both P<0.001), and the 2005-BDR_both had an increased trend but decreased at extremely severe obstruction (P=0.244). In COPD, the proportion of 2005-BDR_FEV₁ was negatively associated with the degree of airflow obstruction, while 2005-BDR_FVC was positively associated (both P<0.001), and the 2005-BDR_both had an increased trend but decreased at extremely severe obstruction (P=0.585). Similar results were observed in 2021-BDR, but the percentage of 2021-BDR_FEV₁ decreased significantly in both asthma and COPD (all P<0.05). The component percentages of BDR+ among different levels of airflow obstruction are shown in Supplementary Figure 1 and Supplementary Table 1. The component percentage of 2005-BDR_FEV₁, 2005-BDR_FVC, 2005-BDR_both, 2021-BDR_FEV₁, 2021-BDR_FVC and 2021-BDR_both, were 62.1%, 11.5%, 26.4%, 42.8%, 14.8%, 42.4% for asthma, 35.5%, 42.4%, 22.0%, 15.4%, 55.3%, 29.3% for COPD, respectively. More details are found in Supplementary Figure 2 and Supplementary Table 2.
To the best of our knowledge, there is a paucity of data exploring the differences in BDR between the 2005 and 2021 ERS/ATS criteria applied to a large sample of patients with asthma as well as COPD. The main findings of our study...
were the decreased proportion of 2021-BDR+ in asthma and COPD and the similar trends in the proportion of BDR+ with the level of airflow obstruction.

The present study showed the percentage of 2021-BDR+ reduced by 10.48% and 7.98% in asthma and COPD, respectively, being lower than a previous study in which the percentage of responders reduced by about 50% when BDR expressed as a percentage of the predicted value or a change in $z$ score.\(^\text{11}\) On the contrary, Bhatt reported the prevalence of BDR+ in COPD increased from 32.5% (2005-BDR+) to 44.6% (2021-BDR+).\(^\text{19}\) The different results may be partly due to the heterogeneity in subjects. Interestingly, we found that the percentage of BDR+ increased slightly in children with asthma, which may be attributed to the lower predicted FEV\(_1\) or FVC in children. As we know, BDR+ is often misinterpreted as a hallmark of asthma. Herein, approximately 20–30% and 50–60% of BDR+ occurred in COPD and asthma, respectively. As reported previously, 39% of COPD patients had a $\geq 10\%$ absolute increase in percent predicted FEV\(_1\) values;\(^\text{20}\) 52.7% COPD patients exhibited BDR.\(^\text{21}\) Undoubtedly, it is quite difficult to distinguish asthma from

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**Figure 2** The trend in the proportion of BDR+ with the increased degree of airflow obstruction in asthma (A) and COPD (B).
COPD based merely on BDR test in clinical practice. In addition, in our study, whichever criterion was adopted, BDR+ group had lower FEV$_1$, FVC, FEV$_1$%pred, FVC%pred than BDR-group. In contrast, patients with consistent BDR+ had higher initial FEV$_1$%pred, FVC%pred, indicating that consistent BDR+ was associated with better lung function. Notably, some patients changed from 2005-BDR- to 2021-BDR+, and their characteristics need further exploration.

With the deterioration of respiratory function, the percentage of BDR+ increased linearly in COPD, but exhibited an approximate inverted U-shaped curve in asthma. This reveals that patients with severe asthma patients may respond poorly to bronchodilator. As reported previously, severity of asthma had a significant influence on BDR and more severe asthma predicted a smaller BDR. An explanation account for the above observations might be the irreversible airflow limitation in severe asthma, which was associated with airway inflammation, airway wall thickening, smooth muscle.

**Figure 3** The trend in the proportion of BDR+ subgroups with the increased degree of airflow obstruction in asthma (A) and COPD (B).
hypertrophy, and hyperinflation.\textsuperscript{23,24} We also found that in COPD patients, the proportion of BDR\textsubscript{FEV1} was negatively associated with the degree of airflow obstruction while BDR\textsubscript{FVC} was positively associated, indicating BDR\textsubscript{FEV1} occurred more frequently in mild COPD while BDR\textsubscript{FVC} occurred more often in advanced COPD. This finding was consistent with that previously reported by Barjaktarevic et al.\textsuperscript{21} Previous studies showed that BDR\textsubscript{FVC} rather than BDR\textsubscript{FEV1} better reflected the physiological processes of air trapping\textsuperscript{25–28} and BDR\textsubscript{FVC} was demonstrated to be more prevalent in subjects with greater emphysema and gas trapping.\textsuperscript{21,29} As is known, in advanced COPD, emphysema with loss of alveolar attachments, distal airway remodeling and mucus hypersecretion may lead to air trapping and dynamic hyperinflation. Bronchodilator administration can partly reduce lung hyperinflation, resulting in a more significant impact on the change in FVC in comparison to that in FEV\textsubscript{1}. Meanwhile, we noted that BDR\textsubscript{Both} had a quite component percentage among three BDR+ subgroups, but its clinical significance was not fully understood. Fortis and coworkers reported that BDR\textsubscript{Both} was associated with less emphysema and indicated a COPD phenotype with asthma-like characteristics.\textsuperscript{30}

In the present study, we mainly addressed the differences in baseline pulmonary indices, the proportion of BDR+ and the trend with respiratory impairment, but did not include other indices associated with clinical outcomes such as dyspnea, exercise capacity, radiological airway measures, exacerbations and mortality. However, an ideal criterion for BDR should be integrated with clinical data to estimate prognosis. Patients with BDR\textsubscript{FEV1}>8% of predicted were reported to have an optimal survival advantage compared with those with BDR\textsubscript{FEV1} ≤ 8% of predicted.\textsuperscript{10} In another study, neither the 2005 nor the 2021 BDR criterion predicted exacerbations or mortality in COPD when adjusted for the severity of lung disease.\textsuperscript{19} Furthermore, whether an acute response can predict long-term outcomes other than survival remains unclear.\textsuperscript{20,31,32} On the other side, some researchers argued that reversibility of airway obstruction in response to a bronchodilator was not a dichotomous trait but a continuous variable, so any cut-off level of a positive BDR might be arbitrary.\textsuperscript{9} In addition to binary BDR categorization, a novel classification with five distinct categories was put forward, including negative, minimal, mild, moderate or marked, which was based on the changes in FEV\textsubscript{1} relative to the predicted values with the following intervals ≤0%, 0–2%, 2–4%, 4–8% and >8%.\textsuperscript{33}

The strengths of our study include the stringent quality control of data processing and a large dataset, but it has several limitations. First, as mentioned earlier, whether 2021-BDR outweighs 2005-BDR deserves further study and could not be judged from the available results. Second, the presence of BDR was reported to be variable at follow-up visits,\textsuperscript{30} and the results from a single BDR test in this retrospective study should be interpreted with caution. Third, BDR results may be influenced by a variety of factors, such as smoking index, disease phenotype, comorbidities and the administration protocol of bronchodilators, more relevant factors should be considered in our future analysis.

**Conclusions**

When BDR was defined by the 2021 ERS/ATS criterion, the proportion of BDR+, especially BDR\textsubscript{FEV1} decreased in asthma patients and COPD patients. The proportion of BDR+ had an upward linear trend with the increased degree of airflow obstruction in COPD but exhibited an approximate inverted U-shaped curve in asthma. Patients with consistent BDR+ have higher initial FEV\textsubscript{1},%pred and FVC%pred. However, the clinical significance of the new criterion in the evaluation of obstructive lung diseases warrants for further study.

**Data Sharing Statement**

The data presented in this study are available on request from the corresponding author.

**Author Contributions**

All authors made a significant contribution to the work reported, whether that is in the conception, study design, execution, acquisition of data, analysis and interpretation, or in all these areas; took part in drafting, revising or critically reviewing the article; gave final approval of the version to be published; have agreed on the journal to which the article has been submitted; and agree to be accountable for all aspects of the work.
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Disclosure
The authors report no conflicts of interest in this work.

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