Asthma and COPD are both common chronic respiratory diseases affecting approximately 1 in 12 people worldwide. The two conditions are generally considered different disease entities with unique pathophysiological mechanisms and characteristic clinical features. The underlying pathophysiology in COPD is characterized predominantly by neutrophilic inflammation, whereas in asthma the inflammatory pattern is mostly due to eosinophilic inflammation. Asthma typically presents with intermittent respiratory symptoms caused by airflow obstruction predominantly due to bronchial hyperresponsiveness. Asthma is often presented at younger age as part of an atopic constitution, but can also be diagnosed in adulthood. In contrast, COPD is a slowly progressive lung disease with persistent respiratory symptoms and airflow obstruction. In high-income countries like the Netherlands COPD usually presents in patients older than forty who are generally current or former smokers. Patients with asthma or COPD are most diagnosed and managed by primary care clinicians.

Looking at the classic pathophysiological and clinical presentations, the distinction between asthma and COPD seems clear, but in clinical practice patients often show features of both diseases. These similarities make it difficult for clinicians to distinguish between asthma and COPD, especially in older and more diverse patient populations encountered in primary care. However, differentiating between the two respiratory conditions is important as they have different pharmacotherapeutic regimens. In patients with asthma, inhaled corticosteroids (ICS) are highly effective in reducing symptoms and reducing the risk of asthma-related mortality. In contrast, patients with COPD respond poorly to ICS and are mainly treated with (long-acting) bronchodilators to relieve symptoms. In addition to this, misdiagnosing asthma for COPD could lead to serious health risks considering that monotherapy with long-acting bronchodilators is contra-indicated in asthmatics since it increases the risk of severe exacerbation. On the other hand, (unnecessary) treatment with ICS may cause pneumonia and increased risk of osteoporosis.

Thus, establishing a correct diagnosis is essential for optimal treatment of asthma and COPD, but this can be challenging for primary care clinicians. Supporting them in the diagnostic process seems therefore essential, but this also depends on the availability of diagnostic tools. Although quality spirometry has shown to be feasible in primary care settings, there is substantial room for improvement of its use to accurately diagnose chronic respiratory diseases. Thus, the first aim of our current study was to establish which patient characteristics distinguish between patients diagnosed with asthma or COPD. The second and main aim was to establish the added value of spirometry and more advanced lung function measurements to differentiate between these two chronic airways diseases.

**METHODS**

**Study design and population**

In this observational multi-centre cross-sectional study, we compared patients diagnosed with asthma, patients diagnosed with COPD, and subjects without underlying chronic obstructive...
random sample of 1,749 adult subjects (20–70 years old) were included in the study. At the start of the program, patients with pre-existing asthma, COPD or another airway disease were excluded. In 2007, ten years after the start of the initial DIMCA program, all subjects (now aged 30–80 years) received an invitation for a comprehensive respiratory assessment consisting of extensive lung function measurements and a myriad of medical history questions. A total of 532 subjects agreed to participate in this follow-up study. The results of the respiratory assessment of these subjects were submitted to two experienced chest physicians who assessed if a chronic airways disease (i.e., COPD or asthma) was present or absent using guideline criteria, their expert knowledge, and their clinical expertise. Study subjects were randomly assigned to the chest physicians in a 1:1 ratio. If a subject was diagnosed with a chronic airways disease by the assigned chest physician, the subject's data was also presented to the other chest physician and a final joint diagnosis was established. To standardize the diagnostic process, a decision tree (Fig. 1) was created based on international clinical guideline criteria for diagnosing asthma (GINA guideline, 2007 update) and COPD (GOLD guideline, 2006 update) that applied at the time of the study (see below). The results of the chest physicians’ assessments were used as the golden standard in the current study.

The study was approved by the medical ethics review board CMO Regio Arnhem – Nijmegen (https://www.radboudumc.nl/over-het-radboudumc/kwaliteit-en-veiligheid/commissie-mensgebonden-onderzoek; file number: 2002/028). Participants provided written informed consent to take part in the study.

Measurements
Study participants were instructed to interrupt the use of any bronchodilators they might use for a specified number of hours before their visit to the pulmonary function laboratory. Lung function testing involved pre- and postbronchodilator spirometry (both static and dynamic) and measurement of carbon monoxide diffusion capacity (DLCO) and bronchial hyperresponsiveness (BHR). Aerosolized salbutamol 800 μg and/or ipratropium 160 μg were used as bronchodilators and were administered by volume spacer. Postbronchodilator forced expiratory volume in one second (FEV1) was measured 15 min after salbutamol and 45 min after ipratropium. Bronchodilator reversibility was defined as an increase in FEV1 after bronchodilation by at least 12% and 200 mL. BHR was assessed by histamine challenge test and considered positive in case of a >20% drop in FEV1 at a provocative dose histamine of ≤8 mg/mL (PC20). All lung function tests were conducted by certified lung function technicians in a hospital-based pulmonary function laboratory and were performed in accordance with the 1994 American Thoracic Society standards. Predicted normal lung function values for FEV1 were calculated using European Community for Coal and Steel reference values. Following lung function testing, subjects were interviewed by the lung function technician regarding respiratory symptoms, smoking behaviour, presence of allergies and eczema, respiratory problems triggered by environmental exposures, and family history of COPD or asthma.

Diagnostic assessment
Based on the results of the respiratory assessment the chest physicians assessed if a chronic airways disease (i.e., asthma or COPD) was present or absent using guideline criteria, their expert knowledge, and their clinical expertise. Study subjects were randomly assigned to the chest physicians in a 1:1 ratio. If a subject was diagnosed with a chronic airways disease by the assigned chest physician, the subject's data was also presented to the other chest physician and a final joint diagnosis was established. To standardize the diagnostic process, a decision tree (Fig. 1) was created based on international clinical guideline criteria for diagnosing asthma (GINA guideline, 2007 update) and COPD (GOLD guideline, 2006 update) that applied at the time of the study, in co-operation with the two chest physicians. In case of uncertainty about the respiratory diagnosis the chest physicians could request additional diagnostic tests (i.e., allergy skin testing, peak expiratory flow (PEF) monitoring) in order to maximize their diagnostic certainty. Because the concept of asthma-COPD overlap (ACO) was introduced after the current study was conducted, the chest physicians did not consider a diagnosis of ACO as a part of their assessment. They were instructed to, based on their systematic assessment of all diagnostic information.

Fig. 1 Decision tree used by the chest physicians to support their assessment of chronic lung disease diagnoses based on GOLD and GINA guidelines. Postbronchodilator forced expiratory volume. Postbronchodilator vital capacity. 12% change in FEV1 (after bronchodilation), with a change of at least 200 mL. BHR was assessed by histamine challenge test and considered positive in case of a >20% drop in FEV1 at a provocative dose histamine of ≤8 mg/mL (PC20). All lung function tests were conducted by certified lung function technicians in a hospital-based pulmonary function laboratory and were performed in accordance with the 1994 American Thoracic Society standards. Predicted normal lung function values for FEV1 were calculated using European Community for Coal and Steel reference values. Following lung function testing, subjects were interviewed by the lung function technician regarding respiratory symptoms, smoking behaviour, presence of allergies and eczema, respiratory problems triggered by environmental exposures, and family history of COPD or asthma.
Fig. 2 Schematic illustration of the spectrum of chronic obstructive airways disease diagnoses. The current study focuses on the parts to the left and right of the vertical dotted lines as indicated by the arrows. ACO asthma-COPD overlap, COPD chronic obstructive pulmonary disease.

Table 1. Clinical features and lung function values of patients diagnosed with asthma and patients diagnosed with COPD.

|                        | Chronic airways disease | No chronic airways disease | p-value |
|------------------------|-------------------------|----------------------------|---------|
| **Demographic characteristics** |                         |                            |         |
| Age (mean, SD)          | 52.0 (11.4)             | 57.8 (10.0)                | <0.001  |
| Median (IQR)            | 49.8 (14.6)             | 57.0 (15.2)                | <0.001  |
| Range (youngest, oldest) | 36.6-78.9               | 36.9-80.5                  | <0.001  |
| Gender (% female, n)    | 59.5 (50)               | 44.2 (61)                  | 0.027   |
| BMI (mean, SD)          | 27.4 (4.3)              | 26.7 (4.0)                 | 0.22    |
| Smoking behaviour       |                         |                            |         |
| Ever smoking (% n)      | 56.0 (47)               | 81.0 (111)                 | <0.001  |
| Current smoking (% n)   | 17.9 (15)               | 39.1 (54)                  | 0.001   |
| Pack-year (mean, SD)    | 8.9 (14.4)              | 21.3 (19.5)                | <0.001  |
| Atopy (% n)             |                         |                            |         |
| Ever allergy a          | 70.2 (59)               | 19.6 (27)                  | <0.001  |
| Ever eczema             | 26.2 (22)               | 26.1 (36)                  | 0.99    |
| Hyperresponsiveness (% n) |                        |                            |         |
| Respiratory symptoms triggered by cold air smoke or (exhaust) fumes | 71.4 (60) | 59.4 (82) | 0.071 | 22.6 (70) |
| Family history b (% n)  |                         |                            |         |
| Asthma                  | 19.0 (16)               | 15.9 (22)                  | 0.32    |
| COPD                    | 29.8 (25)               | 36.2 (50)                  | 0.65    |
| Current respiratory medication c (% n) |               |                            |         |
| Bronchodilator(s)       | 20.2 (17)               | 16.7 (23)                  | 0.502   |
| Inhaled corticosteroid  | 13.1 (11)               | 9.4 (13)                   | 0.392   |
| Respiratory symptoms (% n) |                        |                            |         |
| Cough d                 | 20.2 (17)               | 26.1 (36)                  | 0.32    |
| Wheeze e                | 46.4 (39)               | 27.5 (39)                  | 0.006   |
| Phlegm f                | 11.9 (10)               | 19.6 (27)                  | 0.14    |
| Breathlessness g        | 40.5 (34)               | 30.4 (42)                  | 0.13    |
| Spirometry:             |                         |                            |         |
| PostBD FEV1/FVC (mean, SD) | 74.2 (4.9)       | 63.3 (6.3)                 | <0.001  |
| PostBD FEV1/ FVC < 0.70 (% n) | 15.7 (13)         | 97.8 (135)                 | <0.001  |
| PostBD FEV1 % predicted ECCS (mean, SD) | 98.9 (13.9) | 88.2 (16.2) | <0.001 | 107.1 (14.0) |

*p-values are for the comparison between the two diagnostic subgroups. Data of patients with no chronic airways disease as presented in the table serve as a general reference, but were not part of the current analysis.

ECCS European Community of Coal and Steel, GINA global initiative for asthma, GLI global lung function initiative, LLN lower limit of normal based in GLI prediction equations, RV residual volume, SD standard deviation, TLC total lung capacity.

*Allergic to pollen, animals, dust mites or seasonal symptoms.

†First degree relatives.

‡As prescribed by the patient’s general practitioner and/or pulmonologist.

§Chronic cough in winter.

¶Wheeze with or without breathlessness (in previous 12 months).

‖Phlegm after getting out of bed (in previous 12 months).

*Based on GLI reference equations (http://gli-calculator.ersnet.org/index.html). The % predicted FEV1 values as considered by the two chest physicians in the study were based on the 1993 ECCS reference equations. The GLI-based % predicted FEV1 values were not used by the two chest physicians.

GINA (2021) states that confidence regarding presence of bronchodilator reversibility is greater if the increase is >15% and >400mls (1).

Fisher’s exact test because one cell had an expected count <5.

Decrease in FEV1 by >20% at provocative dose histamine of ≤8 mg/ml (PC20).

Available, assign one single preferred diagnosis (i.e., either asthma or COPD) that fitted best according to their expert opinion. Figure 2 illustrates the spectrum of chronic obstructive airways disease diagnoses and the parts of the spectrum on which the current study focuses. Strictly for the purpose of describing the study population and its diagnostic subgroups (see Table 1) the Global Lung function Initiative (GLI) reference equations were applied at the time of the data analysis for the current paper.

Categorization of variables

In the present study, we categorized all items of the respiratory assessment in three subsections based on their availability in different healthcare settings, i.e., public health, primary care, and secondary care (Table 2). Subsection 1 consists of items that are available in any public health or healthcare setting since they require no measurements or testing equipment but only medical history questions (i.e., respiratory symptoms, smoking behaviour, body mass index (BMI)). Subsection 2 contains lung function test results that are available to primary care clinicians (i.e., spirometry and reversibility testing) in countries with well-developed healthcare systems.
Regression models were run for three scenarios based on diagnostic availability and multivariable logistic regression analysis from the patient assessment were categorized in three subgroups (i.e., Subsections 1 plus 2 plus 3). Only items with a $p$-value ≤0.20 in the univariate analysis were considered relevant as predictors and were included in the respective models. In each scenario, the item with the highest $p$-value was manually removed from the model after which the logistic model was re-run ("backward selection"). This step was repeated until only variables with $p$-values < 0.10 remained in the model for each scenario. Odds ratios for diagnosing asthma were calculated with COPD as reference group and vice versa. For each scenario a receiver operator characteristics (ROC) curve was created and the percentage explained variance (Nagelkerke R square) determined. Area under the curve (AUC) values from the ROC curves of the three scenarios were statistically compared using a non-parametric approach for correlated ROC curves. SPSS statistics version 25.0 and SAS version 9.4 were used for the analyses. Missing data were not imputed. Two-sided $p$-values < 0.05 were considered statistically significant, except for the testing of the AUC values between Scenarios 1 and 2 and Scenarios 2 and 3, respectively, in which multiple testing was taken into account by using $p < 0.025$ to define statistical significance (i.e., Bonferroni correction: $p = 0.05/2 = 0.025$).

Finally, Subsection 3 contains results from more advanced diagnostic tests as performed mainly in lung function laboratories in hospital care settings. These tests include measurement of static lung volumes, diffusion capacity, and histamine challenge testing.

**Statistical analysis**

Demographic characteristics, clinical features and lung function values were univariately compared between the subgroups of patients diagnosed with asthma and COPD using independent t-tests and Chi-square tests. The further analysis focussed on assessing the ability to differentiate between these chronic obstructive lung diseases in different healthcare settings. Since physicians are not limited to asking a single medical history question or to conducting a single diagnostic test, we used multivariable logistic regression analysis to construct predictive models based on the data of the subjects who were diagnosed with asthma or COPD by the chest physicians (i.e., the binary outcome measure for this analysis was to have a diagnosis of asthma or a diagnosis of COPD). As described above, the items from the patient assessment were categorized in three subsections based on diagnostic availability and multivariable logistic regression models were run for three ‘scenarios’ (Table 2). In the first scenario, we only used the medical history items from Subsection 1 in the model. In the second scenario, we added diagnostic items available to primary care clinicians (i.e., Subsections 1 plus 2) to the model. In the third scenario, we added diagnostic items available to secondary care clinicians to the model (i.e., Subsections 1 plus 2 plus 3). Only items with a $p$-value ≤0.20 in the univariate analysis were considered relevant as predictors and were included in the respective models. In each scenario, the item with the highest $p$-value was manually removed from the model after which the logistic model was re-run ("backward selection"). This step was repeated until only variables with $p$-values < 0.10 remained in the model for each scenario. Odds ratios for diagnosing asthma were calculated with COPD as reference group and vice versa. For each scenario a receiver operator characteristics (ROC) curve was created and the percentage explained variance (Nagelkerke R square) determined. Area under the curve (AUC) values from the ROC curves of the three scenarios were statistically compared using a non-parametric approach for correlated ROC curves. SPSS statistics version 25.0 and SAS version 9.4 were used for the analyses. Missing data were not imputed. Two-sided $p$-values < 0.05 were considered statistically significant, except for the testing of the AUC values between Scenarios 1 and 2 and Scenarios 2 and 3, respectively, in which multiple testing was taken into account by using $p < 0.025$ to define statistical significance (i.e., Bonferroni correction: $p = 0.05/2 = 0.025$).

| Table 2. Categorization of variables in three subsections based on diagnostic availability and multivariable logistic regression analysis for the three scenarios. |
|---|
| **Subsection 1 – Patient history questions:** |
| - Demographic characteristics $^a$ |
| - Smoking behaviour $^b$ |
| - Atopy $^c$ |
| - Hyperresponsiveness $^d$ |
| - Family history $^e$ $^f$ |
| - Respiratory symptoms $^f$ |
| **Subsection 2 – Lung function tests available in primary care settings:** |
| - Spirometry $^i$ |
| - Reversibility testing $^*$ |
| **Subsection 3 – Lung function tests available in secondary care settings:** |
| - RV/TLC $^h$ |
| - Bronchial hyperresponsiveness $^*$ |
| - Diffusion capacity |

$^a$Age, gender and BMI.  
$^b$Ever and current smoking, packyears.  
$^c$Ever allergy, ever eczema.  
$^d$Respiratory symptoms triggered by cold air, smoke or (exhaust) fumes.  
$^e$First degree relative with asthma or COPD.  
$^f$Cough, wheeze, phlegm, breathlessness.  
$^g$Postbronchodilator FEV1 and FEV1/FVC.  
$^h$Residual volume/total lung capacity.  
$^i$Not included in multivariable logistic regression as $p$ was <0.20 in univariate analysis.

**RESULTS**

**Study population**

In the total sample of 532 study subjects (all Caucasians), 84 (16%) were diagnosed with asthma, 138 (26%) were diagnosed with COPD, and in 310 subjects (58%) no chronic airways disease was diagnosed (Table 1). Compared to patients with COPD the patients diagnosed with asthma were significantly younger (mean age 50.2 (SD 11.4) versus 57.8 (SD 10.0); $p < 0.001$) and more likely to be female (59.5% versus 44.2%; $p = 0.027$). There was no statistically significant difference in BMI between the two diagnostic subgroups ($p = 0.22$).
Differences and similarities in clinical features and lung function

Table 1 gives an overview of the differences and similarities in demographic characteristics, clinical features and lung function values between patients with asthma and patients with COPD. Patients diagnosed with COPD were significantly more likely to be former or current smokers and had more packyears compared to patients with asthma (21.3 (SD 19.5) versus 9.1 (14.4); \( p < 0.001 \)). Patients with asthma were significantly more likely to have allergies compared to patients with COPD (\( p < 0.001 \)) but there was no difference in the prevalence of eczema between the subgroups (\( p = 0.99 \)). Patients with asthma had significantly more often symptoms of wheezing (\( p = 0.006 \)) compared to patients with COPD. The prevalence of having chronic cough, phlegm or breathlessness was not significantly different between the groups. Patients with COPD had significantly lower % predicted post-bronchodilator FEV1 values (88.2% versus 98.9%; \( p < 0.001 \)) compared to patients with asthma. There were no differences in the presence of reversibility (\( p = 0.75 \)) or bronchial hyperresponsiveness (\( p = 0.68 \)) between the two subgroups. No additional diagnostic tests were requested by the two chest physicians.

Differentiating ability of diagnostic items

Demographic characteristics, clinical features and lung function tests yielded a total of 21 diagnostic variables (Table 1). Excluding items with p-values of >0.20 in the univariate analysis resulted in twelve items that were considered as relevant discriminants to be entered in the multivariable logistic models: age, gender, packyears, wheeze, phlegm, breathlessness, allergy, respiratory symptoms triggered by environmental exposures, post-bronchodilator FEV1 % predicted, postbronchodilator FEV1/FVC <0.70, RV/TLC and diffusion capacity.

Table 3 shows an overview of the differentiating ability of all relevant items. In Scenario 1 (only medical history questions), eight items were included in the model, four of which showed a statistically significant relationship when differentiating between asthma and COPD: packyears, wheeze, phlegm and allergy. In Scenario 2, ten items were included in the model, six of which showed a significant relation in differentiating between asthma and COPD: age, wheeze, breathlessness, allergy, FEV1 % predicted and FEV1/FVC. In Scenario 3, twelve items were included in the model, six showing statistical significance when differentiating between asthma and COPD: age, wheeze, breathlessness, allergy, FEV1 predicted and FEV1/FVC. Independent of the scenario, postbronchodilator FEV1/FVC was an important discriminant.

In Scenario 1 the logistic model showed a percentage explained variance of 41% and ROC characteristics showed an area under the curve (AUC) of 0.84 (95% confidence interval (CI): 0.78–0.89) (Fig. 3). By adding diagnostic variables available to primary care (i.e., spirometry) in Scenario 2, the explained variance increased to 54% and AUC increased to 0.89 (95% CI 0.84–0.93). Finally, by adding more advanced diagnostic tests available to secondary care in Scenario 3, the explained variance increased to 56% but AUC remained 0.89 (95% CI 0.85–0.94). Statistical testing showed a statistically significant difference between the AUCs of Scenarios 2 and 1 (\( p = 0.020 \)) but no such difference between the AUCs of Scenarios 3 and 2 (\( p = 0.967 \), see Table 3).

DISCUSSION

In this study, we looked at which patient characteristics distinguish between patients diagnosed with asthma or COPD, and established the added value of spirometry and of more advanced lung function measurements when differentiating between the two chronic airways diseases. Although asthma and COPD are both heterogeneous conditions with multiple overlapping features, there are important clinical differences as well.

We observed that in the scenario using only medical history questions, it is already possible to reliably distinguish between asthma and COPD. The most important factors to aid differentiation are smoking behaviour, certain respiratory symptoms and the presence of allergies. The use of postbronchodilator spirometry provided important additional discriminative power in correctly labelling a patient as having asthma or COPD. More advanced diagnostic tests that are mainly used in secondary care, such as measuring bronchial hyperresponsiveness and diffusion capacity, did not provide a better differentiation in this primary care study population.

In the present study, both bronchodilator reversibility and bronchial hyperresponsiveness had a similar prevalence in patients diagnosed with asthma and COPD. This finding is noteworthy, as the current GINA guideline refers to reversibility testing and bronchial hyperresponsiveness as criteria supporting the diagnosis of asthma. However, our finding is not unique as previous studies have concluded that solely the presence of reversibility or bronchial hyperresponsiveness does not distinguish between the two obstructive airways diseases. Besides these similarities, there were several clinical features that were statistically different between the two diagnostic subgroups and for that reason, these features can aid primary care clinicians when differentiating between asthma and COPD. Using only medical history questions in the logistic model (Scenario 1) already showed rather good differentiating ability (AUC = 0.84). These findings are in line with other studies that assessed the ability of solely using medical history questionnaires to distinguish between asthma and COPD. Beeh et al. concluded that with only medical history questions, it is possible to distinguish between asthma and COPD for the majority of patients with suspected or established obstructive lung disease. Likewise, in their study Tinkelmann et al. reported that a simple self-administered questionnaire can facilitate differentiation between obstructive lung diseases. However, these studies did not look at the additional use of spirometry or more advanced diagnostic tests to discriminate between asthma and COPD nor did they quantify this in, for instance, an area under the curve analysis like we did. In the present study we found that postbronchodilator spirometry was important when differentiating the two conditions and together with medical history questions, the discriminating ability of the model improved (from AUC = 0.84 in Scenario 1 to AUC = 0.89 in Scenario 2). In contrast, more advanced diagnostic tests did not provide a better diagnostic differentiation (AUC remained 0.89 in Scenario 3). This does not mean that these tests are useless, as they have an important role in evaluating the presence and severity of structural lung damage (like, for instance, in emphysema and bronchiectasis) and in differentiating obstructive lung disease from other aetiologies in selected patients.

A particular strength of our study is that we used standardized methods to conduct the lung function testing and to obtain the respiratory diagnoses. All questionnaires and lung function tests were standardized and prospectively collected, were supervised by certified lung function technicians, and the lung function tests met established quality standards.

Given the central role of general practice in the Dutch healthcare system, nearly all inhabitants are registered in a general practice of their own choice. Therefore, the subjects who participated in the initial DIMCA program and provided for the sample in the current analysis can be seen as representative for the adult Dutch population. On top of this, our study is original in categorizing diagnostic variables based on their availability in different healthcare settings.

However, there were limitations as well. We only looked at the diagnosis itself and did not consider the severity of the diagnosed chronic airways diseases. Because each subject was initially assessed by only one of the chest physicians we were not able to look at the interobserver agreement. Subjects who were
considered to have no asthma or COPD were not mutually discussed by the chest physicians to reach a maximum substantiated outcome. However, given that the aim of our study was to differentiate between asthma and COPD and not to distinguish between being ‘respiratory healthy’ or not, we do not consider this to be a relevant limitation of the study.

In some cases the chest physicians’ assessment may have led to false positive diagnoses of COPD, as some subjects who had a post-BD FEV1/FVC value >0.70 (n = 3; see Table 2) or reported to never have smoked (n = 27) were assigned a COPD diagnosis nonetheless. Unfortunately, we cannot in retrospect ascertain the chest physicians’ specific considerations for assigning this diagnosis in these cases.

Whereas the data collection and diagnostic approach in the DIMCA study by Albers et al.22 were conducted in a prospective manner, our study was retrospective in design and we were limited to using a pre-existing list of diagnostic items. The data collection dates from more than a decade ago and therefore several more recent diagnostic tests were not included. For instance, several recent studies have shown that the underlying type of inflammation in patients with asthma and COPD is markedly different3,41. Tests like sputum cell count, peripheral

![Table 3. Differentiating abilities of relevant items and overall model performance.](https://example.com/table3.png)

| Subsection                        | Scenario 1 | Scenario 2 | Scenario 3 |
|----------------------------------|------------|------------|------------|
|                                  | Asthma     | COPD       | Asthma     | COPD       | Asthma     | COPD       |
|                                  | p-value    | p-value    | p-value    | p-value    | p-value    | p-value    |
| Medical history questions        |            |            |            |            |            |            |
| Age                              | 0.97       | 1.03       | 0.96       | 1.05       | 0.93       | 1.08       |
|                                  | (0.94, 1.01)| (1.00, 1.06)| (0.92, 0.99)| (1.01, 1.09)| (0.88, 0.97)| (1.03, 1.13)|
| Gender (female)                  | x          |            | x          |            | x          |            |
| Packyears                        | 0.97       | 1.03       | 0.98       | 1.02       | x          |            |
|                                  | (0.95, 0.99)| (1.01, 1.06)| (0.96, 1.00)| (1.00, 1.05)|            |            |
| Wheezeb                          | 2.76       | 0.36       | 3.62       | 0.28       | 2.79       | 0.36       |
|                                  | (1.33, 5.57)| (0.17, 0.75)| (1.52, 8.59)| (0.12, 0.66)| (1.15, 6.75)| (0.15, 0.87)|
| Phlegmc                          | 0.33       | 2.99       | x          |            |            |            |
|                                  | (0.12, 0.90)| (1.11, 8.08)|            |            |            |            |
| Breathlessnessd                  | x          |            |            |            | x          |            |
|                                  |            |            |            |            |            |            |
| Lung function tests available to primary care |            |            |            |            |            |            |
| FEV1 % predicted ECCSg           | 1.07       | 0.94       | 1.08       | 0.93       |            |            |
|                                  | (1.03, 1.10)| (0.91, 0.97)| (1.04, 1.11)| (0.90, 0.96)|            |            |
| FEV1/FVC< 0.70                   | 0.14       | 7.25       | 0.11       | 8.81       |            |            |
|                                  | (0.04, 0.52)| (1.92, 27.45)| (0.03, 0.44)| (2.27, 34.18)|            |            |
| Lung function tests available to secondary care |            |            |            |            |            |            |
| Diffusion capacityi              |            |            |            |            | x          |            |
|                                  |            |            |            |            |            |            |
| Explained variancej              | 0.41       | 0.54       | 0.56       |            |            |            |
|                                  |            |            |            |            |            |            |
| AUCf (95%CI)                     | 0.84       | 0.89       | 0.89       |            |            |            |
|                                  | (0.78-0.89)| (0.84-0.93)| (0.85-0.94)|            |            |            |
| p-value for differencek          |            |            |            |            |            |            |
| between AUCs                      | 0.020g     |            | 0.967g     |            |            |            |
|                                  |            |            |            |            |            |            |

Odds ratios (95% confidence intervals) for diagnosing asthma or COPD together with corresponding p-values are calculated for the three different scenarios based on the items available.

AUC area under the curve, ECCS European community of coal and steel, FEV1 forced expiratory volume in 1 s, FVC forced vital capacity, ROC receiver operator characteristics, RV residual volume, TLC total lung capacity.

*aPackyears were missing in 2 subjects, RV/TLC in 3 subjects; there were no further missings.

*bWheeze with or without breathlessness (in previous 12 months).

*cPhlegma after getting out of bed (in previous 12 months).

*dBreathlessness on exertion (in previous 12 months).

*eAllergic to pollen, animals, dust mites or seasonal symptoms.

*fRespiratory symptoms triggered by cold air, smoke or (exhaust)fumes.

*gPostbronchodilator FEV1 as % of predicted value.

*hPostbronchodilator FEV1/FVC.

*iDiffusion capacity in mmol/kPa/min.

*jNagelkerke R square.

kAUC of ROC curve with COPD as reference group.

l*x refers to variables manually removed from the model as p-values were >0.10.

mFor the difference between asthma and COPD diagnoses within each scenario separately.

nFor the difference between Scenarios 2 and 1.

oFor the difference between Scenarios 3 and 2.
A final limitation of the study that should be mentioned is that younger adults (i.e., those aged 18–30) were not included in the study. However, as the aim of the study was to differentiate between asthma and COPD and a diagnosis of COPD below the age of 30 is highly unlikely, we do not think this has had a relevant impact on the findings as reported.

Besides the good discriminating ability of solely using anamnestic questions, our results emphasize the importance of postbronchodilator spirometry in distinguishing asthma from COPD and vice versa. However, it is important to realise that the lung function tests in our study were conducted by well-trained staff in a pulmonary function laboratory and interpreted by experienced chest physicians. To translate these results to the real-life setting, it requires standardized procedures, quality assurance and trained clinicians to interpret the spirometry data accurately and this may be difficult to achieve in primary care.46–48. However, previous studies have shown that it is feasible to conduct reproducible and clinically meaningful spirometry tests in primary care and that primary care clinicians can interpret spirometry test results correctly.29,49,50. Even while in our study bronchial hyperresponsiveness testing did not improve diagnostic differentiation, it has been shown that bronchial challenge testing is safe and feasible in a suitably equipped primary care diagnostic centre.51. Referral to secondary care is indicated in the few cases in which it is not possible to establish a diagnosis on the basis of thorough medical history taking and well-conducted spirometry alone, or to exclude other possible underlying conditions.

In conclusion, primary care clinicians should be able to reliably differentiate between asthma and COPD with the combination of relevant patient history questions and postbronchodilator spirometry tests for the majority of patients with suspected chronic airways disease. More advanced diagnostic tests used in hospital care settings do not seem to provide a better overall diagnostic differentiation between asthma and COPD in primary care patients. Given the important additional role of postbronchodilator spirometry in this process of differentiating, the implementation of quality-assured spirometry testing and sufficient training should be mandatory in primary care practices. Furthermore, the availability of inflammatory markers in primary care could potentially provide better discriminating diagnostic ability but we did not investigate this in the current study.

DATA AVAILABILITY
The data from the DIMCA study are not made publicly accessible because the variable names, labels and codebook are all in the Dutch language. The dataset can be requested from the corresponding author without restrictions, in which case relevant variables and labels will be translated to English.

CODE AVAILABILITY
Readers can access the code (i.e., SPSS syntax) of the statistical analyses by sending a request to the corresponding author.

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AUTHOR CONTRIBUTIONS

J.D.M.B. and T.R.S. initiated the study, performed the data analysis, and wrote the initial draft version of the paper. E.W.M.A.B. and J.C.C.M.V. critically reviewed the paper. All authors approved the final version of the paper that was submitted.

COMPETING INTERESTS

The authors declare no competing interests.

ADDITIONAL INFORMATION

Supplementary information

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Correspondence and requests for materials should be addressed to Tjard R. Schermer.

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