Introducing a new prognostic instrument for long-term mortality prediction in COPD patients: the CADOT index

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Objectives. The BODE (BMI, Obstruction – FEV\textsubscript{1}, Dyspnoea – mMRC, Exercise – 6-MWT) and the ADO (Age, Dyspnoea – mMRC, Obstruction – FEV\textsubscript{1}) indices are widely used prognosis assessment tools for long-term mortality prediction in COPD patients but subject to limitations for use in daily clinical practice. The aim of this research was to construct a prognostic instrument that prevents these limitations and which would serve as a complementary prognostic tool for clinical use in these patients.

Methods and Participants. The data of 699 COPD subjects were extracted from the Czech Multicentre Research Database (CMRD) of COPD patients (the derivation cohort) and analysed to identify factors associated with the long-term risk of mortality. These were entered into the ROC analysis and reclassification analysis. Those with the strongest discriminative power were used to construct the new index (CADOT). The new index was validated on 187 patients of the CIROCO+ cohort (Netherlands; the validation cohort).

Results. The CADOT was constructed by adding two newly identified prognosis-determining factors, chronic heart failure (CHF) and TL\textsubscript{CO}, to the ADO index. In a head-to-head comparison, the CADOT index showed highest c-statistic values compared to the BODE and ADO indices (0.701 vs 0.677 vs 0.644, respectively). The prognostic power was more definitive when applied to the Dutch validation (CIROCO+) cohort (0.842 vs 0.799 vs 0.825, respectively).

Conclusions. The CADOT index has comparable prognostic power to the BODE and ADO indices. The CADOT is complementary/an alternative to the BODE (if 6-MWT is not feasible) and ADO (with less dependence on the age factor) indices.

Trial registration: ClinicalTrials.gov (NCT01923051).

Key words: COPD, prognostic index, pulmonary function, mortality

INTRODUCTION

According to the latest World Health Organization data, chronic obstructive pulmonary disease (COPD) was the third leading cause of death worldwide, claiming approximately 3 million lives in 2016 (ref.\textsuperscript{1}). In the Czech Republic, the annual COPD-related death rate is around 3,500 events per 10.6 million population\textsuperscript{2}. According to an epidemiological prediction model, COPD prevalence is expected to rise in the coming years\textsuperscript{3}.

COPD is considered a heterogeneous syndrome with inter-individual differences in disease manifestation, co-morbidity and long-term mortality risk\textsuperscript{4,5}. For this reason, accurate tools for estimating the life expectancy of COPD patients are warranted\textsuperscript{6,7}. The BODE (Body mass index, airflow Obstruction, Dyspnoea and Exercise) and the ADO (Age, Dyspnoea and Obstruction) indices are globally the most widely used instruments for long-term mortality assessment\textsuperscript{8,9}. A number of other prediction tools have been constructed, e.g. the e-BODE, BODEx and
COTE (ref. 10,11). Use of the BODE (and derived indices) may be difficult in some patients (e.g. with a disability) or in some outpatient settings where the 6-minute walk test (6-MWT) cannot be performed (e.g. lack of a ~30-meter corridor). Two of the 3 parameters determining the ADO score may be associated with other confounders – the specificity of the mMRC score (alternative causes of dyspnoea – pulmonary “other-than-COPD”, cardiogenic, extrathoracic, neuromuscular, systemic, etc.) and the age factor (mortality risk/rate is strongly determined by age – the “Gompertz-Makeham Law of Mortality”) (ref. 9,12,13).

The Czech Multicenter Research Database of COPD (CMRD) comprises a large number of regularly monitored COPD patients14. Based on an analysis of all-cause mortality in CMRD COPD subjects, the aim of the present research was to construct an alternative long-term prognostic instrument for use in situations where the BODE score cannot be calculated and that would improve the ADO index by augmenting the role of COPD-specific conditions predictive of poorer prognosis.

METHODS

The derivation cohort (Czech Republic)

The data for development of the new scale were extracted from the CMRD Registry14, an observational prospective study with a primary objective to monitor and assess morbidity and all-cause mortality in patients with moderate to very severe COPD (Global Initiative for Obstructive Lung Disease (GOLD) grades II to IV) in the Czech Republic (ClinicalTrials.gov Identifier: NCT01923051). Patients were recruited in 14 centers providing specialised respiratory care between February 2013 and December 2016. Follow-up of patients within the CMRD Registry is still ongoing. Detailed description was published elsewhere14.

At the time of the new prognostic instrument construction (July 2016), the registry included 699 COPD patients. Parameters assessed at enrolment included demographics, patient history data [general practitioners’ (GPs’) and specialists’ records], symptoms [dyspnoea – mMRC score15], COPD Assessment Test (CAT) (ref. 16), quality of life measures [St George’s Respiratory Questionnaire (SGRQ) (ref. 17)], treatment, pulmonary functions and other clinical examinations (chest CT, ECG; blood gases, echocardiography etc.). GPs’ and specialists’ records were also used to identify chronic heart failure (CHF).

The validation cohort (Netherlands)

To validate the new prognostic index, data from the CIROCO+ cohort, an observational single-center study, were used18. Patients with moderate to very severe COPD (GOLD grades II to IV) (ref. 19), aged 40 to 80 years and in a clinically stable condition were prospectively recruited between November 2007 and November 2010 during initial evaluation of a comprehensive pulmonary rehabilitation program at CIRO+ (ref. 20). CHF was identified from patient history (Charlson comorbidity index); FEV1 and TLco values were measured at inclusion20.

Development of the new index

The steps to develop the new index included identification of parameters discriminating between patients who died and those who were alive (Step 1), testing of the discriminatory power of these parameters (Step 2) and refinement with a reclassification analysis (Step 3).

The derivation cohort was separated into patients who died and those who were alive at the time of analysis. Clinical characteristics of these two subgroups were compared using the Mann-Whitney U-test and the Fisher exact test to identify parameters discriminating between the two subgroups (Step 1).

The discriminatory power of the identified parameters (Step 2) was tested with the receiver operating characteristic (ROC) analysis. The ROC analysis included the existing ADO and BODE indices alone and ADO and BODE indices with addition of the risk parameters identified during Step 1. The quality of fit was assessed with the c-statistic that equals the area under the ROC curve. The significance of the differences between the ROC curves was tested with the DeLong test21.

Reclassification analyses (Step 3), namely the NRI and IDI methods22,23, were used for parameters that significantly increased the c-statistics (the ROC analysis) of the ADO/BODE to further refine selection of components for the new prognostic index; only the parameters with significant results in the reclassification analysis were included in the final index.

Logistic regression was used to calculate the risk of death for the newly added parameters and, based on this, patients were divided into risk groups. Each group was assigned risk points that these parameters added to the new scale. The risk points were determined by rounding the ORs from the logistic regression. The new index was then divided into risk categories with similar prognostic power.

To validate the new scale, we calculated long-term survival estimates for the derivation (July 2016) and validation (April 2017) cohorts to assess differences in mortality risk between subgroups assigned to the risk categories. The CMRD Registry is still an ongoing project and the prospective nature of the study enabled us to re-assess the prognostic utility of the CADOT twice more in March 2018 and in January 2020, in order to confirm its unique prognostic properties.

All presented analyses were performed using the IBM SPSS Statistics 24.0 (ref. 24) and R-studio software (ref. 25). All statistical tests used α=0.05 as the level of significance.

RESULTS

Step 1: The derivation cohort (median follow-up 18.5 months) was split into subgroups of 616 alive and 83 dead patients. The two subgroups differed in the total ADO and BODE scores as well as the individual parameters within the two indices. Concurrently, we observed lower values of pulmonary function tests (FEV1/FVC, TLco, Kco) in the subgroup of dead vs living patients (P<0.001 for all). CHF was significantly more frequent in the dead vs alive subgroup (33.7% vs 15.0%). Detailed characteristics of
the derivation cohort and its subgroups are presented in Table 1.

Step 2: Since total ADO/BODE and their individual items discriminated between the two subgroups, the ADO and BODE indices were used as platforms for the development of a new index.

The potential of the newly identified parameters, i.e. CHF, FEV$_1$/FVC, TL$_{CO}$ and K$_{CO}$ to increase the discriminative power for all-cause mortality prediction of the existing risk indices (ADO/BODE) is presented in the Table A1. The DeLong test showed that adding CHF, TL$_{CO}$ and K$_{CO}$ to the ADO platform provided statistically significant differences in mortality prediction. No significant result was identified for any combination of the new parameters and the BODE.

Consequently, CHF, TL$_{CO}$ and K$_{CO}$ were entered in the reclassification analysis (Step 3) using the ADO platform as the cornerstone. The reclassification analysis showed CHF and TL$_{CO}$ to be appropriate for definitive use as additional factors for mortality prediction (Table A2).
Therefore, the development process resulted in a new prognostic index with 5 components, i.e. CHF, Age, Dyspnoea (mMRC score), Obstruction (FEV\textsubscript{1}, \% of predicted value) and TL\textsubscript{CO} (% of predicted value) (CADOT). CHF was assigned 0 points (absent) and 3 points (present; rounded OR = 3). TL\textsubscript{CO} values were categorized into three categories and assigned 0 points (≥425%), 1 point (30-44%) and 2 points (<30%). The rating of the individual items of the CADOT index is described in Table 2. The CADOT scores can range between 0 and 15 points. Detailed characteristics of the CMRD cohort with complete CADOT data are presented in Table A3. Like the BODE index, the individual scores were split into 4 prognostic categories (low risk, intermediate risk, high risk, very high risk of death) with similar prognostic power (Table A3 and Table 3).

### Validation

Characteristics of the validation cohort (median follow-up 26.2 months) are presented in Table A4. The outcomes of the validation are presented in Table 3 (the CADOT index performance), Table 4 (prognostic power of the BODE, ADO and CADOT) and in Fig. 1 (long-term survival estimates).

Estimated two-year survival rates for the low, intermediate, high and very high risk groups were 100%, 92.1%, 84.8% and 54.6%, respectively (P<0.001) in the derivation cohort, 100%, 95.7%, 79.5% and N/A, respectively (P<0.001) in the validation cohort.

### Confirmation of prognostic utility of the CADOT from a long-term perspective

The c-statistic of the CADOT in the March 2018 reassessment was 0.685 (P<0.001). In the last reassessment in January 2020, estimated five-year survival rates for the low, intermediate, high and very high risk groups were 88.9%, 66.7%, 42.6% and 22.9%, respectively (P<0.001). (Fig. A1).

### DISCUSSION

The BODE index is the most widely used and globally accepted instrument for COPD prognosis assessment. We present a complementary/alternative prognostic tool, the CADOT index. Having comparable prognostic power to the BODE and ADO indices, the CADOT has features that may be of special benefit in selected settings, in particular, if the 6-MWT is not practicable. The CADOT index also functioned well in subjects with milder airflow obstruction (Table 4) and its properties were confirmed on an independent validation cohort.

The CADOT addresses some specific weaknesses of the ADO and BODE, such as 6-MWT in BODE or the impact of age on ADO.

Disabled or unfit COPD patients (e.g. severe arthrosis, polyneuropathy, lower limb amputees) may be unable (or unwilling) to undergo the 6-MWT or complete a shorter distance and fall into a BODE poorer prognosis. In addition, not all medical offices are equipped to perform the 6-MWT (e.g. lack of a 30-meter corridor). Consequently, the BODE may be of limited use in COPD populations with disability/immobility. In 2009, Puhan et al. attempted to increase the prognostic accuracy of the BODE and ADO indices\textsuperscript{4}. For BODE (“updated BODE index”), the main difference was the 6-MWT scoring where the different walking distances were assigned 0 (>350 m), 4 (250-349 m), 7 (150-249 m) or 9 points (<150 m), respectively\textsuperscript{9}. However, this modification reinforced the reliance of the BODE on the 6-MWT, since a single one meter difference in walking distance (e.g. 350 vs 349 m) may result in a total score change of up to 4 points\textsuperscript{9}. Since the CADOT does not include the 6-MWT, the issues associated with the 6-MWT are completely avoided.

The ADO index is based on age, FEV\textsubscript{1} and mMRC score assessments\textsuperscript{8}. Up to 50% of the total score is determined by age alone (5 points of the 10-point scale assigned to age >90 years) (ref.\textsuperscript{9}). The 2012 ADO update

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**Table 3.** Prediction of 2-year mortality (95% CI) according to the CADOT total score.

<table>
<thead>
<tr>
<th>Category</th>
<th>Score</th>
<th>Derivation cohort</th>
<th>Validation cohort</th>
</tr>
</thead>
<tbody>
<tr>
<td>CADOT 1</td>
<td>0-2</td>
<td>0.0%</td>
<td>0.0%</td>
</tr>
<tr>
<td>CADOT 2</td>
<td>3-5</td>
<td>7.9% (3.2% – 12.5%)</td>
<td>4.3% (0.2% – 8.5%)</td>
</tr>
<tr>
<td>CADOT 3</td>
<td>6-9</td>
<td>15.2% (9.0% – 21.5%)</td>
<td>20.5% (6.9% – 34.1%)</td>
</tr>
<tr>
<td>CADOT 4</td>
<td>10-15</td>
<td>45.4% (21.0% – 69.9%)</td>
<td>-</td>
</tr>
</tbody>
</table>

CADOT = Chronic heart failure, Age, Dyspnoea, airflow Obstruction, TL\textsubscript{CO} = diffusion capacity (Transfer factor) for Carbon Monoxide; CI = Confidence Interval

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**Table 4.** ROC analysis - index capacity to separate patients according to mortality.

<table>
<thead>
<tr>
<th>Index</th>
<th>Cohort</th>
<th>C-statistics (95% CI)</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>BODE</td>
<td>Derivation cohort</td>
<td>0.677 (0.610–0.744)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td></td>
<td>Validation cohort - NL</td>
<td>0.799 (0.681–0.917)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>ADO</td>
<td>Derivation cohort</td>
<td>0.644 (0.581–0.706)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td></td>
<td>Validation cohort - NL</td>
<td>0.825 (0.735–0.914)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>CADOT</td>
<td>Derivation cohort</td>
<td>0.701 (0.625–0.776)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td></td>
<td>Validation cohort - NL</td>
<td>0.842 (0.755–0.930)</td>
<td>&lt;0.001</td>
</tr>
</tbody>
</table>

ADO = Age, Dyspnoea and airflow Obstruction; BODE = Body-mass index, airflow Obstruction, Dyspnoea, and Exercise; CADOT = Chronic heart failure, Age, Dyspnoea, airflow Obstruction, TL\textsubscript{CO} = diffusion capacity (Transfer factor) for Carbon Monoxide
In COPD patients, reduced TLco usually reflects the presence of emphysema, pulmonary arterial hypertension or CHF that are all associated with increased risk of long-term mortality. In Central Europe, COPD patients are cared for mainly by pulmonary physicians, unlike Western Europe, where GPs are the prevailing caregivers. In the Czech Republic, around 80% of outpatient non-hospital respiratory practices are equipped with (or have access to) a diffusion capacity assessment device. The situation is similar in Germany (personal communication with the Lemon Medical GmBH). Recent analysis of the POPE cohort illustrated that TLco vs 6-MWT data were available in 90% vs 11% of Croatians and 63% vs 20% of Czechs with COPD (ref.30-34). This means that in some regions or countries, TLco assessment may be more accessible than a ~30-meter corridor, i.e. the 6-MWT.

The CADOT index performed equally well in various populations of patients (CMRD and CIROCO). CMRD represents a population of moderate-to-very severe COPD subjects with higher (>17%) prevalence of CHF, while the CIROCO cohort included more patients with milder airflow obstruction (mean FEV1, was 50%). In addition, the CIROCO patients were younger, had higher mean TLco (56%) and CHF was less prevalent (3%). This was consequent to the CIROCO study exclusion criteria (unstable COPD, myocardial infarction in the previous 6 months, asthma history, alpha-1 antitrypsine deficiency, previous lung surgery, malignancy in the previous 5 years) (ref.35). Importantly, the prognostic power (c-value) of the CADOT was higher in the “milder” COPD population of the CIROCO cohort. Since the long-term mortality risk among GOLD I subjects is very low, a large cohort of these subjects and a year-long follow-up would be needed to learn the prognostic properties of CADOT in this category of patients. An easy way for assessing risk among GOLD I patients using CADOT is a periodic (e.g., annual) calculation of the CADOT score where the disease progression/deterioration can be captured.

Our study has limitations. First, the derivation cohort included patients from tertiary and university hospital-based centers and thus, further external validation on larger cohorts with higher proportions of mild COPD patients are desirable. Second, the TLco test may be less available in primary care settings. However, this is strongly region/country-dependent and the availability of TLco in some regions or countries may exceed that of 6-MWT. For example, in Czechia, the availability of TLco for respiratory physicians is more than 80%. Third, the presence of CHF has not been re-assessed at patient inclusion. However, of the 120 CMRD subjects with a history of CHF, 29 had an echocardiographic (ECHO) examination of the heart within the CMRD study protocol (ECHO was a non-mandatory test). Of these, 26 patients (93%) had ECHO signs compatible with left- or right-sided CHF. Of the remaining 94 subjects with a CHF history, 87 (93%) were treated with 1 or more CHF treatments (ACE inhibitors, angiotensin II receptor antagonists, betablockers, diuretics). These data suggest reliability of the patient history data from the CMRD database. Fourth, our cohorts included a lower portion of patients with COPD, GOLD grade I. Though the CADOT performed well on a cohort with milder airflow obstruction, further studies are needed to assess the utility of the CADOT in GOLD grade I subjects. Finally, the majority of both cohorts was composed of men (74% of the derivation cohort and 58% of the validation cohort, respectively) and the applicability to

**Fig. 1.** Long-term survival according to the CADOT index (comparison of the derivation - CZ (A) and validation - NL (B) cohorts). CZ - Czech Republic; NL - Netherlands.
women may be somewhat limited. However, in the BODE and ADO construction studies, the proportion of men and women was unequal as well - in the 2012 study of Puhan et al., the proportion of men was 60%, while in the Puhan study of 2009, men accounted for a 60% (Swiss cohort) and 93% (Spanish cohort) (ref.9,11).

CONCLUSIONS AND IMPLICATIONS

We constructed and validated a new prognostic index (CADOT) that has slightly higher prognostic power than the BODE and ADO indices. The CADOT is complementary (or alternative) to the BODE in situations where 6-MWT is not feasible. The CADOT index improves the ability of respiratory physicians to determine risk for patients with COPD and severe comorbidities.

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Authors contributions: All authors contributed to the data collection, analysis and interpretation and writing of the manuscript. MS, KH, KB, VK: had full access to all CMRD data in the study and take the responsibility for the integrity of the data and the accuracy of the data analysis; MS: carried out the statistical analysis; VK, KH: conceived and designed the CMRD study in the 2012; BN: took the responsibility for CMRD project registration at ClinicalTrials.gov (NCT01923051) and for validation of CMRD data; KB, MP, MS, DE, VK: drafted the manuscript; DE: edited the English language; VK: obtained funding. All authors critically revised the manuscript for important intellectual content and approved the submitted version. All authors are committed to ensuring that questions related to the accuracy and integrity of any part of the work are appropriately investigated and resolved.

Conflict of interest statement: The authors declare there are no conflicts of interest related to the manuscript content.

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REFERENCES


Supplemental Material:
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