Comparison of early warning scores in patients with COPD exacerbation: DECAF and NEWS score

Carlos Echevarría, John Steer, Stephen C Bourke

ABSTRACT

Background The National Early Warning Score 2 (NEWS2) includes two oxygen saturation scales; the second adjusts target saturations to 88%–92% for those with hypercapnic respiratory failure. Using this second scale in all patients with COPD exacerbation ("NEWS2\_AI\_COPD") would simplify practice, but the impact on alert frequency and prognostic performance is unknown. Admission NEWS2 score has not been compared with DECAF (dyspnoea, eosinopenia, consolidation, acidaemia, atrial fibrillation) for inpatient mortality prediction.

Methods NEWS, NEWS2 and NEWS2\_AI\_COPD and DECAF were calculated at admission in 2645 patients with COPD exacerbation attending consecutively to one of six UK hospitals, all of whom met spirometry criteria for COPD. Alert frequency and appropriateness were assessed for all NEWS iterations. Prognostic performance was compared using the area under the receiver operating characteristic (AUROC) curve. Missing data were imputed using multiple imputation.

Findings Compared with NEWS, NEWS2 reclassified 3.1% patients as not requiring review by a senior clinician (score\geq5). NEWS2\_AI\_COPD reduced alerts by 12.6%, or 16.1% if scoring for injudicious use of oxygen was exempted. Mortality was low in reclassified patients, with no patients dying the same day as being identified as low risk. NEWS2\_AI\_COPD was a better prognostic score than NEWS (AUROC 0.72 vs 0.65, p<0.001), with similar performance to NEWS2 (AUROC 0.72 vs 0.70, p=0.090). DECAF was superior to all scores (validation cohort AUROC 0.82) and offered a more clinically useful range of risk stratification (DECAF=1.2%–25.5%; NEWS2=3.5%–15.4%).

Conclusion NEWS2\_AI\_COPD safely reduces the alert frequency compared with NEWS2. DECAF offers superior prognostic performance to guide clinical decision-making on admission, but does not replace repeated measures of NEWS2 during hospitalisation to detect the deteriorating patient.

BACKGROUND

The National Early Warning Score (NEWS) facilitates prompt recognition of clinical deterioration in patients hospitalised with a wide range of conditions. It is composed of bedside indices (respiratory rate, oxygen saturations, systolic blood pressure, pulse, level of consciousness and temperature) with clear thresholds commanding the urgency of clinical response. Early warning scores, such as NEWS, are intended to be scored repeatedly during the inpatient stay to monitor patients for deterioration.

Although NEWS is widely used in the UK, two key limitations have been highlighted. First, patients with COPD and chronic hypoxaemia raise false alerts, which may lead to alert fatigue and complacency. Second, in severe COPD exacerbations (ECOPD), excess oxygen is associated with increased need for ventilation and mortality, but this risk is not identified by NEWS. The recently updated NEWS2 includes a second oxygen saturation scale which is intended for patients with hypercapnic respiratory failure (table 1). New onset of confusion has also been added to the level of consciousness index. The new oxygen saturation scale encourages the prescription and delivery of oxygen to target saturations of 88%–92% in those with hypercapnia, while the original oxygen saturation scale is retained for patients without hypercapnia. This incorrectly assumes that patients with ECOPD without hypercapnia are at risk of harm from excess oxygen;
Normocapnia will remain at risk throughout. The BTS guidelines recommend reassessing risk until blood gas analysis and clinical review, while those with hypercapnia will be at risk for deterioration throughout admission. However, admission NEWS2 has not been compared with DECAF. The UK National Asthma and COPD Audit Programme currently relies on admission NEWS2 to allow case-mix adjustment for mortality risk; the superiority of DECAF would support its routine use for this purpose. Furthermore, if admission NEWS2 offered similar performance to DECAF in ECOPD, then the principle of parsimony would favour reliance on NEWS2 alone and render the DECAF score redundant.

Despite clear guidelines by the British Thoracic Society (BTS) promoting target saturations of 88%–92% in most patients with COPD, this is frequently not achieved in clinical practice. By assigning risk points to those with excess oxygen, NEWS2 will encourage the correct titration of oxygen in patients identified as at risk. Unfortunately, this is limited to patients confirmed to have hypercapnic respiratory failure on blood gas analysis and with clinician approval. Patients with hypercapnia will be at risk until blood gas analysis and clinical review, while those with normocapnia will remain at risk throughout. The BTS guidelines limit risk by recommending initial target saturations of 88%–92%, but risk is not avoided as the target can be increased to 94%–98% following confirmation of normocapnia. In unselected patients with a clinical diagnosis of ECOPD, there is randomised controlled trial (RCT) evidence that target saturations of 88%–92% improve mortality. This favours adopting the second NEWS2 oxygen scale (88%–92%) in all patients with ECOPD, which would simplify the pathway facilitating implementation, and should greatly reduce the risk of excess oxygen use.

The DECAF (dyspnoea, eosinopenia, consolidation, acidemia, atrial fibrillation) score is a risk stratification tool intended to be scored on admission and accurately predicts risk of death. It can be easily calculated at the bedside to guide treatment, such as hospital at home for low-risk patients. DECAF and NEWS2 serve separate purposes in clinical practice: the former to risk assess patients accurately at admission and the latter to monitor patients for deterioration throughout admission. However, NEWS2 was initially calculated using scale 1 and scale 2 for normocapnic and hypercapnic patients, respectively (‘NEWS2’). NEWS2 was then re-calculated with the assumption that all patients with COPD should have oxygen saturations of 88%–92% (‘NEWS2\textsubscript{AI COPD?’}). Patients were assigned to standard risk groups for each early warning score. For NEWS, NEWS2 and NEWS2\textsubscript{AI COPD}, a score of 0 to 4 is low risk, 5 or 6 is moderate risk, and 7 or more is high risk. A patient with a NEWS of 5 or more should have at least hourly observations and be reviewed by a clinician with competencies in dealing with acute illness, such as a ward doctor. A score of 7 or more should prompt an emergency review, with continuous monitoring of observations. For the DECAF score, 0 to 1 equates to a low in-hospital mortality risk, 2 is moderate risk, and 3 or more is high risk. Alert frequencies (NEWS, NEWS2, NEWS2\textsubscript{AI COPD}) and other proportions were compared with Fisher’s exact test. The performance of NEWS, NEWS2, NEWS2\textsubscript{AI COPD} and DECAF for the prediction of mortality was assessed using receiver operating characteristic (ROC) curves and area under the curve (AUC) statistics. The comparison of NEWS2\textsubscript{AI COPD} with NEWS2 and DECAF was pre-specified in the original study protocols. The first set of NEWS indices was used, either from the accident and emergency (A+E) department, or the medical admissions unit if the patient bypassed A+E. Missing data were handled using multiple imputation as described previously. NEWS2 was initially calculated using scale 1 and scale 2 for normocapnic and hypercapnic patients, respectively (‘NEWS2’). NEWS2 was then re-calculated with the assumption that all patients with COPD should have oxygen saturations of 88%–92% (‘NEWS2\textsubscript{AI COPD}’).

### METHODS

The DECAF derivation and validation cohorts are composed of 2645 consecutive admissions of unique patients with ECOPD to six UK hospitals with predmission obstructive spirometry, age of 35 or older, and smoking history of 10 or more cigarette pack-years (ISRCTN13946813 and ISRCTN29082260).

Collection of the DECAF and NEWS score indices, including confusion, was pre-specified in the original study protocols. The first set of NEWS indices was used, either from the accident and emergency (A+E) department, or the medical admissions unit if the patient bypassed A+E. Missing data were handled using multiple imputation as described previously. NEWS2 was initially calculated using scale 1 and scale 2 for normocapnic and hypercapnic patients, respectively (‘NEWS2’). NEWS2 was then re-calculated with the assumption that all patients with COPD should have oxygen saturations of 88%–92% (‘NEWS2\textsubscript{AI COPD}’).

**Table 1** National Early Warning Score 2 (NEWS2) with risk score for oxygen saturations alone shown for scale 1 (normocapnic patients) and for scale 2 (hypercapnic patients)

<table>
<thead>
<tr>
<th>NEWS2 scale 1 and original NEWS (normocapnic patients)</th>
<th>NEWS2 scale 2 (hypercapnic patients)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Oxygen saturations, %</td>
<td>Score</td>
</tr>
<tr>
<td>91 or less</td>
<td>3</td>
</tr>
<tr>
<td>92–93</td>
<td>2</td>
</tr>
<tr>
<td>94–95</td>
<td>1</td>
</tr>
<tr>
<td>96 or more</td>
<td>0</td>
</tr>
<tr>
<td>93 or more</td>
<td>0</td>
</tr>
<tr>
<td>95–96 on O\textsubscript{2}</td>
<td>2</td>
</tr>
<tr>
<td>97 or more on O\textsubscript{2}</td>
<td>3</td>
</tr>
</tbody>
</table>

Scale 1 is the same as the original NEWS oxygen saturation index and is recommended for most patients including patients with COPD without hypercapnic respiratory failure. Scale 2 is recommended for patients with hypercapnic respiratory failure, whose target saturations are set lower (ie, 88%–92%).

**Table 2** Indices that comprise news and DECAF scores, and rates of missing data in each cohort

<table>
<thead>
<tr>
<th>Indices that comprise news and DECAF scores, and rates of missing data in each cohort</th>
<th>Derivation cohort (n=920)</th>
<th>Validation cohort (n=1725)</th>
</tr>
</thead>
<tbody>
<tr>
<td>RESPIRATORY RATE 0.33%</td>
<td>1.8%</td>
<td>1.8%</td>
</tr>
<tr>
<td>OXYGEN SATURATION 0%</td>
<td>1.8%</td>
<td>0%</td>
</tr>
<tr>
<td>SUPPLEMENTAL OXYGEN 1.5%</td>
<td>0%</td>
<td>0%</td>
</tr>
<tr>
<td>SYSTOLIC BLOOD PRESSURE 0%</td>
<td>1.7%</td>
<td>1.5%</td>
</tr>
<tr>
<td>PULSE 0%</td>
<td>1.8%</td>
<td>0%</td>
</tr>
<tr>
<td>LEVEL OF CONSCIOUSNESS 0%</td>
<td>0%</td>
<td>2.6%</td>
</tr>
<tr>
<td>CONFUSION (NEWS2 only) 0%</td>
<td>2.6%</td>
<td>0%</td>
</tr>
<tr>
<td>TEMPERATURE 0.54%</td>
<td>0%</td>
<td>2.6%</td>
</tr>
<tr>
<td>DECAF SCORE eMRCD score 0%</td>
<td>0%</td>
<td>0%</td>
</tr>
<tr>
<td>EOSINOPENIA 0%</td>
<td>0%</td>
<td>0%</td>
</tr>
<tr>
<td>CHEST X-RAY CONSOLIDATION 0%</td>
<td>0%</td>
<td>0%</td>
</tr>
<tr>
<td>ACIDEMIA (pH &lt;7.3) 0%</td>
<td>0%</td>
<td>0%</td>
</tr>
<tr>
<td>ATRIAL FIBRILLATION 0%</td>
<td>0%</td>
<td>0%</td>
</tr>
</tbody>
</table>

*In patients without an arterial blood gas, it was assumed that none had an acidemia of <7.30 provided oxygen saturations were 93% or more while breathing room air. This was based on data from the derivation cohort of 118 patients meeting these criteria, none had an arterial pH of <7.30.

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inpatient mortality was assessed and compared using the area under the receiver operating characteristic (AUROC) curve.\textsuperscript{17} Calibration was assessed using the Hosmer-Lemeshow (HL) test and by visual assessment in calibration plots.\textsuperscript{18} Analyses were performed on IBM SPSS Statistics V.22 and SigmaPlot V.12.3.

**RESULTS**

**Population and missing data**

In 2645 patients, the mean (SD) age was 73.1 (10.2), 54% were women and the mean (SD) FEV\textsubscript{1} was 45.1% (SD 18.2) predicted. Almost half were unable to leave the house unassisted (Extended Medical Research Council Dyspnoea score 5a or 5b=47%), and 29.8% had consolidation on the admission chest X-ray. Furthermore, 17.9% of patients were acidaemic on the admission arterial blood gas (ABG) analysis, and 10.1% had a pH of less than 7.3. Comorbid conditions were common; 55.4% had a Charlson index of 2 or more, and 28.9% scored 3 or more. Overall inpatient mortality was 8.6%; 96 in 920 (10.4%) in the derivation cohort and 132 in 1725 (7.7%) in the validation cohort.

A more detailed breakdown of patient characteristics by site and cohort is published elsewhere.\textsuperscript{11} \textsuperscript{12} There were no missing data for the primary outcome, inpatient mortality. Rates of missing data for indices were low and are shown in table 2.

**Risk groups and alerts**

Inpatient mortality by risk group is shown in table 3. The percentage of patients classified as requiring an urgent review (score of 5 or more) for NEWS\textsubscript{2 All COPD}, NEWS\textsubscript{2} and NEWS was 62.1%, 71.6% and 74.7% (middle column figure 1). NEWS\textsubscript{2 All COPD} resulted in a shift of total scores towards the lower risk range compared with NEWS\textsubscript{2} (figure 2). NEWS\textsubscript{2 All COPD} identified fewer patients requiring medical review relative to NEWS\textsubscript{2} (9.5% absolute reduction), but the risk of death in the low-risk group was similar (3.5% and 3.1%, p=0.686).

A patient receiving excess oxygen may not necessarily require an urgent clinical review and should first have their oxygen titrated down to target range. For NEWS\textsubscript{2}, if excess oxygen scores were discounted, there was a significant reduction in the frequency of alerts for patients with a score of 5 or more (71.6% to 68.0%, p=0.005). For NEWS\textsubscript{2 All COPD}, discounting excess oxygen reduced reviews in this group from 62.1% to 58.6% (p=0.010), with little difference in the percentage of deaths in the low-risk patient group (counting excess oxygen=3.5% vs discounting excess oxygen=3.6%, p=0.905). Of those patients who were assigned a low-risk NEWS\textsubscript{2} or NEWS\textsubscript{2 All COPD} score, none died that same day. This suggests that all of these patients were correctly categorised as low risk at the time; early warning scores are repeated with each set of observations and are likely to have risen in those who subsequently died.

**Performance of admission DECAF and early warning scores for inpatient mortality**

For all patients (n=2645), NEWS\textsubscript{2 All COPD} was a stronger mortality predictor than NEWS (AUROC NEWS\textsubscript{2 All COPD}=0.72, 95%CI 0.68 to 0.76, vs NEWS=0.65, 95%CI 0.61 to 0.68; p<0.001). NEWS\textsubscript{2 All COPD} showed a trend towards superiority compared with NEWS\textsubscript{2} (AUROC NEWS\textsubscript{2 All COPD}=0.72, 95%CI 0.68 to 0.76, vs NEWS\textsubscript{2}=0.70, 95%CI 0.67 to 0.74; p=0.090).

Of interest, the addition of confusion to the ‘level of consciousness’ component of NEWS\textsubscript{2} accounted for part of the improved performance in NEWS\textsubscript{2} compared with NEWS. For example,
Chronic obstructive pulmonary disease

**Table 4** Sensitivity, specificity, negative predictive value (NPV) and positive predictive value (PPV) at NEWS thresholds of 5 and 7

<table>
<thead>
<tr>
<th>Low-risk group (cut-off less than 5 for NEWS)</th>
<th>Low and moderate risk group combined (cut-off less than 7 for NEWS)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sensitivity</td>
<td>Specificity</td>
</tr>
<tr>
<td>NEWS2_{\text{All COPD}}</td>
<td>0.85</td>
</tr>
<tr>
<td>NEWS2</td>
<td>0.90</td>
</tr>
<tr>
<td>NEWS</td>
<td>0.89</td>
</tr>
<tr>
<td>DECAF</td>
<td>0.93</td>
</tr>
</tbody>
</table>

When scored without confusion, the AUROC for NEWS2_{\text{All COPD}} was only 0.68 (95% CI 0.64 to 0.71) which is significantly worse than NEWS2_{\text{All COPD}} scored with confusion (p<0.001).

DECAF provided optimal prediction of inpatient mortality, and was superior to NEWS2 and NEWS2_{\text{All COPD}} in terms of performance as shown in figure 3 (further data in online supplementary table 1).

Table 4 shows the performance of early warning scores at low-risk and high-risk NEWS thresholds (5 and 7), with the DECAF score the gold standard. The positive predictive value (PPV) and negative predictive value (NPV) most inform the clinician: the PPV is the chance of a patient dying if they score above the threshold, while the NPV is the chance of a patient surviving if they score below the threshold. Sensitivity is the proportion of those who died that were correctly identified as at risk (above the threshold), while the specificity looks at the proportion of those who survived, identified as not at risk (below the threshold).

In the low-risk group, a higher sensitivity and NPV is desirable. With regards to sensitivity, NEWS2 performed optimally, while both NEWS2 scores performed equally well for NPV. In the high-risk group, a high specificity and PPV is desirable, for which NEWS2_{\text{All COPD}} performed optimally.

Calibration describes how well individuals’ predicted risks from a logistic regression model agree with the observed risk in terms of the proportion of individuals affected. The goodness of fit can be measured with the HL test, with poor fit indicated by a small calculated p value. DECAF, NEWS2_{\text{All COPD}} and NEWS were well calibrated with HL statistics of p=0.209, p=0.237 and p=0.736. NEWS2 was not well calibrated, with an HL statistic of p=0.048.

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Calibration curves are shown in figure 4. A good prognostic tool should cover a range of risks, have an intercept close to zero and a slope of one, and have the intersection of predicted and observed risk close to the line (shown by the dots). NEWS2_{\text{All COPD}} was well calibrated and covered a larger range of risks than other early warning scores.

**Discussion**

Patients admitted with ECOPD have a modest in-hospital mortality rate (3.9%–4.3%), but frequently trigger alerts on the National Early Warning Scale (NEWS). This can place a substantial burden on healthcare professionals, create complacency and promote excess oxygen use in those who are at particular risk of its harmful effects. In this study, NEWS2 showed superior discrimination for mortality to NEWS, but only

![Figure 3](image3.png) **Figure 3** Receiver operator curves for the DECAF score, NEWS2_{\text{All COPD}} and NEWS all cohorts. Figure shows the performance of the DECAF, NEWS2_{\text{All COPD}}, NEWS2 and NEWS scores. A higher area under the receiver operating characteristic curve (’A’) shows better prediction. DECAF is not included in the derivation cohort, as this would unfairly favour its performance.

![Figure 4](image4.png) **Figure 4** Calibration curves for DECAF, NEWS2_{\text{All COPD}}, NEWS2 and NEWS in all cohorts. Figure shows calibration curves for NEWS2_{\text{All COPD}}, NEWS2 and NEWS in all patients. DECAF is shown in the validation cohort, as inclusion of the derivation cohort would unfairly favour its performance. Predicted risk is shown on the y-axis which was calculated using logistic regression analysis to give individual’s mortality risk as a percentage. The x-axis shows the observed risk, which is a proportion of the number of patients who died within an assigned risk range. The slope of the equation should be near 1 (eg, for DECAF, the slope is 1.038), and the intercept should be near 0 (for DECAF, this is 0.0022). The R^2 represents the correlation between the predicted risk and observed risk using Pearson’s correlation, with scores closest to 1 showing maximum correlation.

reclassified 3.1% as not requiring a review, based on an alert trigger of 5 or more. NEWS2_{AE,COPD} adopting target saturations of 88%–92% and scale 2 of the NEWS2 tool for all patients, led to an absolute reduction in alert frequency of 12.6%. In clinical practice, titrating down oxygen prior to alerts may be appropriate; this approach decreased alerts by 16.1%. The performance of NEWS2_{All,COPD} was similar to NEWS2, with no increased risk of death in the low-risk group. Of key importance, there were no deaths on the same day that patients were classified as not requiring review.

DECAF was prognostically superior to all scores, showed the most consistent performance and was well calibrated. Accurate mortality prediction informs clinical decision-making; the low-risk DECAF group was larger with a lower risk of death than seen with other scores, which makes DECAF suitable for the identification of patients for early discharge or hospital at home services. Additionally, in the National Asthma and COPD Audit, baseline mortality risk is being quantified by the NEWS2 score, but our results show NEWS2 should not be used in place of the DECAF score for this purpose.

Eligibility criteria were broad and rates of missing data were low, with robust data analyses, including methods to deal with missing data. The assessment of the NEWS score was pre-specified, as was the assessment of confusion; therefore, there is no bias in the subsequent re-coding of NEWS2. Furthermore, in both cohorts, the inclusion of NEWS indices was pre-specified, and data were collected prior to and/or without knowledge of the (objective) outcome. For the AUROC curve comparison, DECAF was compared with NEWS iterations only in the validation cohort; comparing performance in the derivation cohort would have introduced bias in favour of the DECAF score. A key strength is that our population was well described, with respiratory specialist confirmation of exacerbation and spirometric confirmation of COPD. These results may not apply to those with presumed ECOPD without obstructive spirometry (patients misdiagnosed with COPD). In our study, considerable efforts were made to obtain previous lung function results: we checked primary and secondary paper and electronic records before concluding spirometry was unavailable.

Our study has the following limitations. Scores were compared using admission data only. For the comparison with the DECAF score, using admission data is appropriate as we are assessing the tools’ performance as a guide to inform clinical decision-making at the point of admission. However, for NEWS the inclusion of multiple measures would have added further useful data.

‘NEWS2-without confusion’ has been compared with NEWS by Hodgson and others, in a large cohort of patients. They reported concerns that high-risk patients were inappropriately downgraded by NEWS2. However, NEWS2 was calculated from NEWS, which does not include confusion. This is an important omission as we have shown that confusion offers additional prognostic benefit, and inclusion of confusion may have resulted in fewer patients being downgraded. Furthermore, time to death was not reported: in our study, no low-risk patients died the same day, which supports correct risk categorisation. In common with other studies assessing similar early warning scores (such as Salford NEWS), Hodgson et al used ICD-10 codes to identify patients, which are known to be inaccurate.

A recent retrospective study by Pimentel et al offers the advantage of capture of sequential early warning scores in 251 266 acute adult admissions. Again, reliance on coding limits data quality in terms of case ascertainment. Again, confusion was not included, and therefore NEWS2 was not correctly scored. In 48 898 patients with conditions that were associated with risk of hypercapnia based on ICD-10 codes, ‘NEWS2-without confusion’ was slightly inferior to NEWS for identifying death within 24 hours. In a subgroup of 1394 patients with confirmed hypercapnia, ‘NEWS2-without confusion’ offered similar discrimination and a higher PPV than NEWS. In common with our study, ‘NEWS2-without confusion’ generated fewer alerts, and mortality rates were low on the same day as patients having low-risk assessments. Our study differs in that we only included patients with confirmed COPD, NEWS2 was correctly scored and case ascertainment was robust.

While no patient should receive a diagnosis of COPD without obstructive spirometry, we acknowledge that this does occur and that our results may not be generalisable to such patients. However, we highlight that the Austin RCT showed that target SpO2 88%–92% in all patients with a presumed diagnosis of ECOPD at ambulance pick-up was associated with lower mortality, and this included patients without a spirometry-confirmed diagnosis of COPD.

An important limitation of this and other studies of NEWS2 is that we did not observe NEWS2 in practice. Real-time recording of NEWS2 may result in changes in clinical behaviour. For example, while the performance of NEWS2_{AE,COPD} and NEWS2 was similar in our study, the reduction in false alerts with NEWS2_{AE,COPD} could result in the more timely clinical assessment of other patients and hence a positive clinical outcome. Prospective randomised studies are required to address these points and to see if the use of NEWS2_{AE,COPD} leads to improvements in mortality by preventing excess oxygen prescribing, but would be challenging to perform. Mortality was high in this study, though appropriate given the higher levels of comorbidity, pneumonia and frailty in our cohort compared with UK national audit data collected at a similar time.

The current BTS oxygen guidelines recommend initial target saturations of 88%–92% in patients with COPD. NEWS2 requires confirmation of hypercapnia before the alternative oxygen saturation scale is activated and should only be instigated by a clinician. Of concern, the NEWS2 implementation guidance and NEWS2 chart may result in many patients receiving excess oxygen prior to ABG analysis or clinician review. RCT data show a mortality benefit in those treated with oxygen target saturations 88%–92% at the point of ambulance pick-up. These patients did not have baseline ABG analysis prior to oxygen treatment, and this approach was beneficial even though a substantial proportion of patients were subsequently described as not having COPD. RCTs have identified increases in mortality with excess oxygen in patients with a range of acute medical presentations even where there is no risk of hypercapnia.

Using NEWS2_{ul,COPD} in all patients with ECOPD could offer improved prognostic performance and calibration, with significantly fewer alerts, and lead to a reduction in harm from excess oxygen delivery in this at-risk population.

Contributors The study was conceived by SCB and designed by CE, JS and SCB. CE performed the statistical analyses. All authors contributed to data analysis and interpretation, and all were involved in drafting the manuscript and approving the final version. The corresponding author attests that all listed authors meet authorship criteria and that no others meeting the criteria have been omitted.

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Patient consent for publication Not required.

Ethics approval The study was approved by NRES Committee North East, UK (REC reference: 12/NE/0379).

Provenance and peer review Not commissioned; externally peer reviewed.

Data availability statement Data are available on reasonable request.

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