Prediction models for the mortality risk in chronic dialysis patients: a systematic review and independent external validation study

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Objective: In medicine, many more prediction models have been developed than are implemented or used in clinical practice. These models cannot be recommended for clinical use before external validity is established. Though various models to predict mortality in dialysis patients have been published, very few have been validated and none are used in routine clinical practice. The aim of the current study was to identify existing models for predicting mortality in dialysis patients through a review and subsequently to externally validate these models in the same large independent patient cohort, in order to assess and compare their predictive capacities.

Methods: A systematic review was performed following the preferred reporting items for systematic reviews and meta-analyses (PRISMA) guidelines. To account for missing data, multiple imputation was performed. The original prediction formulae were extracted from selected studies. The probability of death per model was calculated for each individual within the Netherlands Cooperative Study on the Adequacy of Dialysis (NECOSAD). The predictive performance of the models was assessed based on their discrimination and calibration.

Results: In total, 16 articles were included in the systematic review. External validation was performed in 1,943 dialysis patients from NECOSAD for a total of seven models. The models performed moderately to well in terms of discrimination, with C-statistics ranging from 0.710 (interquartile range 0.708–0.711) to 0.752 (interquartile range 0.750–0.753) for a time frame of 1 year. According to the calibration, most models overestimated the probability of death.

Conclusion: Overall, the performance of the models was poorer in the external validation than in the original population, affirming the importance of external validation. Floege et al’s models showed the highest predictive performance. The present study is a step forward in the use of a prediction model as a useful tool for nephrologists, using evidence-based medicine that combines individual clinical expertise, patients’ choices, and the best available external evidence.

Keywords: external validation, prediction, nephrology, dialysis, mortality

Introduction
End-stage renal disease (ESRD) is a large public health problem. Treatment options for ESRD consist of transplantation, conservative treatment, or dialysis, of which there are two main forms: hemodialysis (HD) and peritoneal dialysis (PD). Dialysis patients are a very vulnerable population, with 3-year cumulative survival of approximately 50%, considerably lower than survival rates for cancer and heart-failure patients.1,2 Many different patient characteristics have been reported to influence this poor survival, and there is wide interindividual variability in prognosis.3–5

Identifying dialysis patients with an increased risk of dying through prediction models could offer valuable information on prognosis to patients, as well as help clinicians...
recognize these high-risk patients and apply more intensive care and supportive therapies when needed.\(^5,6,7\) Furthermore, a risk-stratification model could be used as an important research tool when adjusting for baseline risk in comparative studies, targeting specific interventions to high- or low-risk patients or even provide tools to compare mortality risks among different centers, regions, or countries.\(^3,7,8\)

Various models to predict mortality in dialysis patients have been published.\(^7-13\) However, no review of these articles exists, and only a fraction of these prediction models or risk scores have been externally validated.\(^7\) The few studies that have externally validated models to predict mortality in dialysis patients did so for not more than one or two models at a time.\(^7,12,13\) Differences in cohorts used to validate various models make it difficult to compare predictive performances fairly; ideally, one would externally validate all available models in the same cohort. External validation entails that a model is tested in other data than that in which the model was derived, in order to test the generalizability of the tool on the targeted patient population.\(^14\) Since the performance of a prediction model is generally lower in new individuals than in the population in which the model was developed,\(^15\) these models cannot be recommended for clinical use before external validity is established, as is also stressed in the TRIPOD (transparent reporting of a multivariable prediction model for individual prognosis or diagnosis) statement for reporting prediction models.\(^14-20\) Though the importance of combining evidence is recognized, many more models have been developed in the medical research field than are used in clinical practice or even tested on different populations.\(^17,21,22\) To combat this research waste, the realization of more external validation studies is imperative and increasingly in demand.\(^14,17,20,23-25\)

A direct comparison of the predictive performance of existing models in the same external-validation cohort could help bridge the gap between the development of models and their clinical application. The aim of the current study was to perform a systematic literature review to identify existing models for predicting the risk of death in dialysis patients and subsequently to externally validate these models in the same large independent patient cohort, in order to assess and compare their predictive capacities.

**Methods**

**Systematic review**

The PRISMA (preferred reporting items for systematic reviews and meta-analyses) guidelines were followed for the review process.\(^26\) A systematic literature search was performed in PubMed to identify articles that described prediction models for risk of death in dialysis patients. PubMed was searched on October 28, 2016 using the search string presented in the Supplementary materials. Additional relevant articles were identified by checking the references of the selected articles from the PubMed search. Titles and abstracts were screened, and full texts of potentially eligible articles were read carefully.

Studies were included in the systematic review if they met the following criteria: 1) the study contained at least one formal, original prediction model from which individual risk estimates could be calculated; 2) the study population consisted of chronic dialysis patients or patients with ESRD; 3) the end point of the prediction was all-cause mortality for a defined time frame, studied in a longitudinal design; and 4) the goal of the study was to predict mortality to the best of the authors’ ability, and not to research whether a particular comorbidity score or single variable was predictive of mortality.

Titles, abstracts, and full papers were reviewed by two authors independently (CLR and PWMV), and any discrepancies in the full-text review were solved by consulting a third party (MvD). For the articles included, two investigators (CLR and PWMV) independently extracted the main characteristics and necessary data from the main paper and any accompanying supplemental material, as recommended by the PRISMA guidelines.\(^26\)

**Validation cohort**

NECOSAD (Netherlands Cooperative Study on the Adequacy of Dialysis) was a multicenter, prospective cohort study, in which 38 dialysis centers throughout the Netherlands participated. Incident dialysis patients were included at the start of dialysis treatment if they were aged ≥18 years and had no history of previous renal replacement therapy. Patient characteristics on dialysis modality, comorbidities, laboratory investigations, and clinical parameters were recorded. High-sensitivity CRP was not assessed in standard fashion, but retrospectively determined in 2002 from frozen blood samples for all patients included till then. Patients were followed till time of death or censored due to kidney transplantation or loss to follow-up. Inclusion of patients took place between 1997 and 2007, and follow-up data on death were available until February 1, 2015. For more information on study design, please refer to one of the hallmark NECOSAD articles published in the *Lancet*.\(^27\) The study protocol was fully approved by the Amsterdam Medical Centre ethics committee, and subsequently all local medical ethics committees gave a declaration of agreement. All patients gave written informed consent. The NECOSAD data set will be made available upon request.
For the current study, baseline measurements were taken 3 months after dialysis initiation, with the exception of predialysis serum creatinine. This was primarily done to ascertain that the dialysis patients included had chronic renal failure and the choice of treatment modality would be more definite. Furthermore, laboratory investigations were collected at this time, since the clinical condition of the patient was more stabilized. Patients who had a follow-up within NECOSAD of less than 90 days were thus excluded from the current study.

Studies selected for validation
Predictors from the studies included in the review were assessed and matched to variables available in NECOSAD. For predictors that were not recorded in NECOSAD, a panel of researchers and nephrologists was consulted on the possible use of proxies. When the panel considered it appropriate, original predictors were replaced with proxies, in order to allow validation of the model. If these studies presented more than one formal prediction model, their final model or the model with the best predictive performance was selected. The original prediction models with regression coefficients per predictor and intercept and/or baseline hazard were extracted from the selected studies and used as published to evaluate predictive performance within our external population. If the full prediction formula was not provided in the article or its supplementary material, the authors were approached multiple times by email or phone with a request to supply this formula.

Statistical analysis
Baseline characteristics of the patients in NECOSAD are presented as mean values with standard deviations (SD) or as median values with interquartile ranges (IQRs) for continuous variables (depending on the normality of the distribution), and as percentages for categorical variables. Baseline characteristics included all the predictors from the validated studies, and were stratified per treatment modality.

To account for missing data on predictors, multiple imputation was performed using the fully conditional specification. The imputation model contained all predictor variables, outcome variables, and any variables that could help impute the missing data. Normally distributed variables were log-transformed. Outcomes from the ten imputed data sets were combined into one overall estimate, with associated variance that incorporated both within- and between-imputation variability. This was done through applying the rules described by Marshall et al, which are an extension of Rubin’s rules aimed specifically at imputation in prognostic modeling studies.

Probabilities of death predicted by each validated model were calculated for each individual within the validation cohort, using the original model formulae and the time frame used in the original papers. The predictive performance of the models was assessed based on outcomes for discrimination and calibration, and these measures are described later (see Supplementary materials for a full description of the model formulae and predictors/proxies used).

Discrimination is a measure for how well a model is able to distinguish between people who will or will not have the event (death) within the given time frame. It can be assessed with the C-statistic, which indicates the probability that the model will assign a higher risk to a random case than to a random non-case. The C-statistic lies between 0.5 and 1, where 1 is perfect, 0.7 is considered reasonable, and 0.8 is considered good discrimination. For a logistic regression model, the C-statistic is equivalent to the area under the receiver-operating curve. For Cox proportional-hazard models, Harrell’s C-index of concordance is presented. This outcome measure takes censoring into account. The C-statistic was assessed per model for multiple prediction time frames and stratified by dialysis modality. Additionally, we computed Kaplan–Meier survival curves for four risk groups, based on quartiles of prognostic indices. These survival curves can provide informal evidence of discrimination, and allow for a visual interpretation of model performance over time: the further the survival curves lie from one another, the better the discrimination.

Calibration describes how accurately the predicted risks from a model reflect the mortality risks in the observed data. This is best represented visually in a calibration plot, in which observed mean mortality risk is plotted against predicted mean mortality risk by a tenth of the predicted risk. The calibration plot can be characterized by a slope and calibration-in-the-large. The calibration-in-the-large indicates to what extent the model may systematically over- or underestimate the probability of death. For a perfectly calibrated risk score, the calibration plot would lie on the 45° line and have a slope of 1, and the calibration-in-the-large would be identical to the observed risk. In the current study, calibration was assessed per model by presenting a calibration plot, calibration slope, and calibration-in-the-large for multiple prediction time frames (if possible) and stratified by dialysis modality.

To test the additional benefit of recalibration, we updated the models in a sensitivity analysis. For the Cox models, this was done by adjusting the baseline hazard of death to the hazard of death observed within the NECOSAD population in the model.
calculating a correction factor for the intercept based on the incidence of death in the validation cohort, as described by Janssen et al. Recalibration can improve calibration measures, but does not affect the discrimination (C-statistic) of the model performance. All statistical analyses were performed in SPSS version 23.

**Results**

**Systematic review**

Using the PubMed search strategy (Supplementary materials), 508 articles were identified and their titles screened. From these articles, 112 were selected for abstract review. From the reviewed abstracts, 42 articles were selected for full-text review and three that had been identified through references of other articles were added to the full-text review selection. After the full review 16 articles were included in the systematic review, based on the predefined inclusion criteria as described in the “Methods” section. The study-inclusion flowchart is presented in Figure 1. The main reasons for exclusion were that no formal prediction model was presented or that the goal of the model was not to predict mortality in individual patients. Characteristics from the selected studies were collected by two independent researchers, and are summarized in Table 1. Most of the studies were performed on HD patients, as opposed to PD patients, and made use of Cox models. Many studies used predictors not routinely

![Flowchart of study selection for systematic review and external validation.](figure1.png)

**Figure 1** Flowchart of study selection for systematic review and external validation.

**Abbreviations:** ESRD, end-stage renal disease; NECOSAD, Netherlands Cooperative Study on the Adequacy of Dialysis.
collected by nephrologists; these were: socioeconomic status, fasting blood glucose, left ventricular ejection fraction, surprise question, plasma S100A12, peritoneal equilibration test, and ventilation.10,11,37–41

As shown in Table 1, a discrimination value was presented for 13 of the 16 models, and ranged from 0.71 to 0.87 in the derivation cohorts. For 12 models, information on calibration in the form of a plot or table was present. A full prediction formula, including baseline hazard and/or intercept and regression coefficients, was available in ten of 16 studies. The baseline hazard function, with which one can calculate predicted probabilities for any time frame in a Cox model, was only offered in one study by Geddes et al.3 Multiple studies offered their estimated baseline hazard for multiple time frames, allowing the model to be used to predict mortality over different prediction horizons.

Selection for external validation

All studies included in the systematic review were considered carefully for validation. A number of studies used very specific predictors that could not be replaced by a proxy. These studies (n=7) were excluded from the validation (see Supplementary material for the specific variables).10,11,37–41 Two studies were excluded because they were (partly) developed on NECOSAD patients.42,43 Finally, one study was excluded because the authors explicitly stated that they had recorded all their predictors before the initiation of dialysis and the values of these predictors change notably after dialysis initiation.44 As shown in Figure 1, a total of six studies presenting seven final-prediction models were finally selected for external validation. Of these studies, four were developed in HD patients and two in dialysis patients in general (HD and PD). The statistical model used was a Cox model in all but one of the validated models, the exception being Mauri et al.45 Upon request, the full prediction formula was made available for all studies.

Baseline characteristics of the validation cohort

Baseline patient characteristics of the NECOSAD patients included at 3 months after dialysis initiation stratified for dialysis modality are summarized in Table 2. The mean time between dialysis initiation and baseline measurement was 93 (SD 13) days. As expected, some marked differences between HD and PD patients were present. For example, the median age was much lower in PD patients. In Table S1, baseline characteristics are given for each predictor/proxy used for validation. In Table S2, baseline characteristics from the derivation cohorts of the validated models are shown. In the first year after baseline (at approximately 15 months after dialysis initiation), 12.6% of the patients (n=244) died. After 2 years 23.3% (n=452) had passed away, and this was 30.2% (n=586) after 3 years and 40.5% (n=787) after 5 years.

Performance of validated models

Table 3 shows the discrimination of the validated models presented as the pooled mean Harrell’s C-index from the ten imputation sets with IQRs. In Table 4, the discrimination is presented when stratified for dialysis modality. The models performed moderately to well in terms of discrimination, with C-statistics ranging from 0.71 (IQR 0.708–0.711) to 0.752 (IQR 0.750–0.753) for a time frame of 1 year. Models tested for different time frames performed consistently more poorly when the prediction horizon increased. All models performed better in the PD population than in the HD population. The model of Wagner et al8 showed the highest discriminatory ability for 1 year, with a C-statistic of 0.752 (IQR 0.750–0.753), while models by Floege et al7 showed similarly high discrimination, with highest discriminatory ability for a 2-year prediction time frame with a C-statistic of 0.740 (IQR 0.737–0.742). To assist comparison, C-statistics found through current external validation were plotted against original C-statistics (if available; Figure 2). As one can tell from this figure, all studies that presented a C-statistic performed more poorly in external validation. In Figure 3, Kaplan–Meier curves are shown per model for quartiles of the prognostic indices. This clarifies the models’ capability of distinguishing between patients with high or low mortality rates, and gives insight on how the discrimination capacity changes over time.

The overall calibration of the models can best be judged by examining the calibration plots presented in Figure 4 and taking the calibration slope and calibration-in-the-large into account, as shown in Tables 5 and 6. Calibration plots are presented for single time frames and combined for HD and PD patients (see Figure S1 for calibration plots stratified by dialysis modality and for various prediction time frames). Though the model tended to overestimate slightly, the calibration of Holme et al’s5 model was fairly good, as can be seen in the plot, and was exemplified by a calibration slope close to 1. Floege et al’s1–2 and 2-year models also showed fairly good calibration. The models of Geddes et al3 and Hutchinson et al15 both overestimated considerably, especially in the higher risk ranges. The model of Wagner et al8 significantly underestimated the probability of mortality, and the model of Mauri et al45 slightly underestimated and had a poor
slope of 0.27. After adjustment for differences in mortality rates between NECOSAD and the development cohorts as a sensitivity analysis, overall the calibration plots improved (Table S3 and Figure S2).

### Discussion

In this study, which included a systematic review and external validation, it was shown that there are 16 published models predicting death in dialysis patients, which previously have barely been validated and have not been implemented in clinical practice on a large scale. Validating them in a large independent cohort showed that they all performed more poorly on new patients, demonstrating the importance of external validation before clinical implementation in the shared decision-making process.

The systematic review exemplified that many articles describing the development of a prediction model used rare predictors that are difficult to validate. Besides complicating validation, the use of rare predictors makes these tools extremely impractical for nephrologists and large-scale clinical implementation highly improbable. The systematic review also showed that many studies were incomplete in providing all information necessary for complete external validation or described outcome-assessment measures differently from

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**Table 1** General characteristics of models to predict risk of all-cause mortality in dialysis patients

| Reference | Events:sample size | Study population | Patient inclusion time span | Outcome, prediction horizon | Statistical model |
|-----------|--------------------|------------------|---------------------------|----------------------------|------------------|
| Hemke et al, Netherlands, “easy” model | 1,225 | Incident dialysis patients | 1997–2007 | 3-, 5-, 10-year mortality | Cox |
| Cao et al, China | 371,5405 | Incident CAPD patients | 2012 | 1-year mortality | Cox |
| Doi et al, Japan | 62,688 | Incident HD patients | 2006–2011 | 1-year mortality | Logistic |
| Floege et al, Germany | 1,060,1,714.722 | Incident and prevalent HD patients | 2007–2009 | 1- and 2-year mortality | Cox |
| Chua et al, Singapore | 169,983 | Incident dialysis patients | 2005–2010 | 1-year mortality | Logistic and linear |
| Zhao et al, China, model 3 | 175,903 | Incident CAPD patients | 2006–2011 | 2-year mortality | Cox |
| Hemke et al, The Netherlands | --, 3502, 4549** | Incident RRT patients | 1995–2005 | 3-, 5-, 10-year mortality | Cox |
| Shiotsu et al, Japan | 50,550 | Prevalent HD patients | 2009 | 2-year mortality | Cox |
| Holme et al, Norway | 880,1,868 | Prevalent HD patients | 2003–2004 | 3-year mortality | Cox |
| Quinn et al, Canada | 1,326,8,103 | Incident dialysis patients | 1998–2005 | 1-year mortality | Logistic |
| Wagner et al, UK, model 2* | 1,078,3,631 | Incident dialysis patients | 2002–2004 | 3-year mortality | Cox |
| Cohen et al, USA | 43,512 | Prevalent HD patients | 2006–2007 | 6-month mortality | Cox |
| Mauri et al, Spain | 558,3,455 | Incident HD patients | 1997–2003 | 1-year mortality | Logistic |
| Geddes et al, UK, MVR model\* | 351, 837** | Incident RRT patients | 1997 | 1- and 5-year mortality | Cox |
| Foley, Canada | 73,325 | Incident dialysis patients | 1980–1991 | 6-month mortality (logistic) | Logistic and Cox model |
| Hutchinson et al, Canada | 101** | Incident HD patients | 1970–1975 | 5-year mortality | Cox |

**Notes:** *Externally validated in the current independent study; **deducted/calculated based on information available in article; ***baseline hazard provided upon request; ****shrinkage factor provided upon request; ^f^3, 5-year mortality, respectively; ^f^1 and 2-year mortality events respectively; ^f^3, 5 and 10 year mortality events, respectively; "", 1- and 5-year mortality events, respectively; "" not provided.

**Abbreviations:** CAPD, continuous ambulatory peritoneal dialysis; HD, hemodialysis; HR, hazard ratio; MVR, multivariate regression; RRT, renal replacement therapy.
### Validation

| Validation                     | Predictors, n | Discrimination, C-statistic (CI) (of derivation) | Discrimination, C-statistic (CI) (of validation) | Calibration plot/table, slope | Presentation of model |
|-------------------------------|---------------|--------------------------------------------------|--------------------------------------------------|------------------------------|-----------------------|
| Internal split-sample validation (610 patients) | 10            | –                                                | 0.79, 0.79, 0.78e | Yes                           | Full prediction formula provided |
| External validation (1,764 patients)                | 6             | 0.71 (0.6–0.83)                                  | 0.72 (0.63–0.81)                                 | –                            | /-value per predictor given |
| Internal bootstrap validation                                   | 6             | 0.83                                             | 0.83 (0.79–0.89)                                 | Yes                          | /-value per predictor given and score chart |
| External validation (10,615 patients)                   | 12, 13a       | –                                                | 0.73, 0.72a                                      | Yes                          | Full prediction formulae provided |
| –                                                   | 7             | 0.74, 0.74, 0.75 (logistic, linear, score)       | –                                                | –                            | Logistic, linear model, and score chart presented |
| Internal split-sample validation (451 patients)         | 10            | 0.79                                             | 0.76 (0.68–0.84)                                 | Yes                          | Full prediction formula provided |
| Internal split-sample validation (6,934 patients)        | 4             | –                                                | 0.71 (0.7–0.72), 0.72 (0.7–0.72), 0.72 (0.71–0.73)c | Yes                          | Full prediction formula provided |
| Bootstrapping and external validation (303 patients)     | 4             | 0.73 (0.66–0.8)                                  | 0.72 (0.63–0.82)                                 | Yes                          | /-value per predictor given and score chart |
| Internal split-sample validation (905 patients)          | 5             | 0.73 (0.71–0.75)                                 | 0.72 (0.69–0.76)                                 | Yes                          | Full prediction formula provided |
| Internal split-sample validation, twice (4,052 and 4,051 patients) and bootstrapping | 15            | 0.77                                             | 0.76 and 0.76                                   | Yes                          | Full prediction formula provided and score chart |
| Internal split-sample validation (1,816 patients)        | 11            | 0.75 (0.73–0.77)                                 | 0.73 (0.71–0.76)                                 | Yes                          | HR per predictor given*** |
| External validation (514 patients)                      | 5             | 0.87 (0.82–0.92)                                 | 0.8 (0.73–0.88)                                  | –                            | HR per predictor given |
| Internal split-sample validation (2,283 patients)        | 10            | 0.78                                             | 0.78                                             | Yes                          | Full prediction formula provided |
| Internal split-sample validation (1,171 patients)        | 17            | –                                                | –                                                | Yes                          | Full prediction formula provided*** |
| –                                                   | 14            | –                                                | –                                                | Yes                          | Full prediction formula (logistic), HRs for Cox and score chart |
| –                                                   | 3             | –                                                | –                                                | Yes                          | Full prediction formula and score chart |

#### Notes:

- **Hutchinson et al,55**
- **Foley,41**
- **Canada 73:325 Incident dialysis patients 1980–1991 6-month mortality**
- **Geddes et al,3 UK, MVR**
- **Mauri et al,45 Spain* 558:3,455 Incident HD patients 1997–2003 1-year mortality Logistic**
- **Cohen et al,38 USA 43:512 Prevalent HD patients 2006–2007 6-month mortality Cox**
- **Wagner et al,8 UK, model**
- **Quinn et al,10 Canada 1,326:8,103 Incident dialysis patients 1998–2005 1-year mortality Logistic**
- **Holme et al,9 Norway* 880:1,868 Prevalent HD patients 2003–2004 3-year mortality Cox**
- **Shiotsu et al,39 Japan 50:550 Prevalent HD patients 2009 2-year mortality Cox**
- **Hemke et al,42 The**
- **Zhao et al,11 China,**
- **Chua et al,37 Singapore 169:983 Incident dialysis patients 2005–2010 1-year mortality Logistic and**
- **Floege et al,7 Germany* 1,060, 1,714:9,722b Incident and prevalent HD**
- **Doi et al,44 Japan 62:688 Incident HD patients 2006–2011 1-year mortality Logistic**

#### Reference Events:

- Sample size
- Study population
- Patient inclusion

#### General characteristics of models to predict risk of all-cause mortality in dialysis patients

1. **Discrimination**
2. **Calibration**
3. **Presentation of model**

#### Statistical methods:

- Cox model
- Logistic and
- Linear
- Score chart
- Slope
- HR per predictor given
- Beta-value per predictor given
- Full prediction formula
- Full prediction formula and score chart
- Score chart

#### Challenges:

- **Discrimination and calibration**
- **Comparison of different models and comparison of model performance in different populations.**
- **Similar problems**
- **Different time frames**
- **Comparing different models and comparing model performance**
- **Standard discrimination and calibration**
- **Complicating comparison of different models**
- **Comparison of model performance in different populations.**

#### Model performance:

- **Wagner et al**'s model seems to be best suited for distinguishing high- and low-mortality-risk patients in a population of HD and PD patients over a time frame of 1 year in this validation cohort. The models of Floege et al also showed high discriminatory ability, with the highest C-statistics for a 2-year time frame. For predicting 3-year probability of mortality, Holme et al’s model performed best on discrimination. As far as calibration goes, most models showed poor calibration when eyeing the calibration plots, and calibration seemed to be best in the models of Floege et al and Holme et al.
When comparing the predictive performances found in this external validation to the performance reported in the original articles (Figure 2), we can state that overall performance was poorer in the validation (as expected), especially when looking at performance in only HD patients. In conclusion, the models presented by Floege et al seem to be best suited to their goal, and showed results in validation that were very consistent with their original findings. This might be due to the fact that they presented predictive performances that were already based on external validation (as well as internal validation), which they had performed on their model in the original publication, but also could have been influenced by the high resemblance among the Western European patient populations in which the model was derived and validated. In addition, parsimony of a model may weigh against a slight loss in performance. Since the models compared differed greatly in number of predictors included, one could imagine that a model with fewer predictors, such as that of Holme et al, which uses only five variables, might be preferred in

### Table 2 Baseline characteristics of external validation cohort (NECOSAD) at 3 months after dialysis initiation

| Characteristics               | Missing (%) | Total (n=1,943) | HD patients (n=1,266) | PD patients (n=677) |
|-------------------------------|-------------|-----------------|-----------------------|---------------------|
| Age, years                    | 0.2%        | 62.8 (49.8–72.2)| 66.7 (55.6–74.4)      | 54.3 (43.0–65.1)    |
| Sex (% male)                  | 0.2%        | 62.3%           | 59.4%                 | 67.8%               |
| Current smokers               | 10.2%       | 22.5%           | 21.2%                 | 24.8%               |
| Ethnicity (% Caucasian)       | 0.2%        | 91.9%           | 92.0%                 | 91.6%               |
| Primary kidney disease        | 0.2%        |                 |                       |                     |
| Diabetes mellitus             |             | 14.4%           | 14.1%                 | 15.1%               |
| Glomerulonephritis            |             | 12.4%           | 9%                    | 18.8%               |
| Renal vascular disease        |             | 16.9%           | 19.5%                 | 11.8%               |
| Other                         |             | 56.3%           | 57.4%                 | 54.3%               |
| **Comorbidities**             |             |                 |                       |                     |
| Cardiovascular disease        | 9.8%        | 35.3%           | 40.9%                 | 25.3%               |
| Diabetes mellitus             | 0.2%        | 5.6%            | 6.8%                  | 3.3%                |
| Cancer                        | 10.8%       | 9.6%            | 11.9%                 | 5.5%                |
| **Medication**                |             |                 |                       |                     |
| ACE inhibitors                | 0.6%        | 24.5%           | 22.4%                 | 28.6%               |
| Calcium antagonists           | 0.6%        | 34.1%           | 32.4%                 | 37.3%               |
| β-blockers                    | 0.6%        | 31.7%           | 30.3%                 | 34.2%               |
| Diuretics                     | 0.6%        | 27.3%           | 27.0%                 | 28.0%               |
| Erythropoietin                | 1.5%        | 85.3%           | 91.7%                 | 73.2%               |
| **Laboratory**                |             |                 |                       |                     |
| Albumin, g/L                  | 5.8%        | 36.0 (5.3)      | 35.8 (5.3)            | 36.2 (5.3)          |
| mGFR, mL/min/1.73 m²         | 21.0%       | 3.4 (1.8–5.4)   | 3.1 (1.6–4.9)         | 4.1 (2.2–6.3)       |
| Hemoglobin, mmol/L            | 1.6%        | 6.9 (1.0)       | 6.7 (0.9)             | 7.4 (1.0)           |
| HS-CRP mg/L                   | 56.6%       | 5.0 (3.0–14.0)  | 6.0 (3.0–16.0)        | 4.0 (3.0–9.0)       |
| BMI (kg/m²)                   | 10.1%       | 24.7 (4.2)      | 24.7 (4.4)            | 24.8 (3.9)          |
| Vascular access mode          | 4.6%        | NA              | NA                    | NA                  |
| Catheter                      |             | NA              | 21.0%                 | NA                  |
| Graft/fistula                 |             | NA              | 79.0%                 | NA                  |

**Notes:** Continuous variables presented as mean values with standard deviations for normally distributed variables. Continuous variables presented as median values with interquartile ranges for non-normal distributed variables. Categorical variables presented as valid percentages.

**Abbreviations:** NECOSAD, Netherlands Cooperative Study on the Adequacy of Dialysis; HD, hemodialysis; PD, peritoneal dialysis; mGFR, measured glomerular filtration rate (combined creatinine and urea clearance in 24-hour urine samples at 3 months); BMI, body-mass index; NA, not applicable; HS-CRP, high sensitivity C-reactive protein.

### Table 3 Discrimination results of external validation for HD and PD patients

| Reference       | Time frame | Original population | Discrimination: C-statistic,* |
|-----------------|------------|---------------------|-------------------------------|
|                 |            | HD and PD           | HD and PD                     |
| Floege et al7   | 1 year     | HD                  | 0.740 (0.738–0.742)           |
| Floege et al7   | 2 years    | HD                  | 0.740 (0.737–0.742)           |
| Holme et al9    | 3 years    | HD                  | 0.734 (0.730–0.737)           |
| Wagner et al8   | 1 year     | HD and PD           | 0.752 (0.750–0.753)           |
|                 | 2 years    | HD and PD           | 0.738 (0.737–0.738)           |
|                 | 3 years    | HD and PD           | 0.730 (0.729–0.731)           |
| Mauri et al45   | 1 year     | HD                  | 0.728 (0.724–0.730)           |
| Geddes et al3   | 1 year     | HD and PD           | 0.721 (0.717–0.723)           |
|                 | 2 years    | HD and PD           | 0.706 (0.704–0.708)           |
|                 | 3 years    | HD and PD           | 0.705 (0.702–0.707)           |
|                 | 5 years    | HD and PD           | 0.698 (0.695–0.700)           |
| Hutchinson et al1 | 1 year | HD                  | 0.710 (0.708–0.711)           |
|                 | 2 years    | HD                  | 0.702 (0.701–0.706)           |
|                 | 3 years    | HD                  | 0.700 (0.699–0.702)           |
|                 | 5 years    | HD                  | 0.697 (0.696–0.699)           |

**Note:** *C-statistics presented as medians and interquartile ranges from pooled imputation results.

**Abbreviations:** HD, hemodialysis; PD, peritoneal dialysis.
Prediction models for mortality risk in chronic dialysis

By evaluating the different models in the same data set, this study provides a far greater amount of information than separate validation studies would, as attested by Collins and Moons and recently demonstrated in a study by Abbasi et al, who systematically validated prediction models for the risk of developing diabetes.47,48 Unfortunately, a lack of well-conducted external-validation studies of prognostic models was found, and many validation studies handle missing data inadequately or do not report measures of model calibration.18,22 These are issues that we have accounted for. Still, there are a number of limitations to our study.

First, our results were influenced by differences in case-mix and mortality rates between the derivation cohorts and our validation cohort. Inherently, some of the derivation cohorts will resemble our validation cohort more than others, since patient characteristics can differ strongly per country.

Table 4 Discrimination results of external validation stratified for dialysis modality

| Reference     | Time frame | Original population | Discrimination: C-statistic* |
|---------------|------------|---------------------|------------------------------|
|               |            |                     | HD                           |
| Floege et al  | 1 year     | HD                  | 0.701 (0.699–0.709)          |
| Floege et al  | 2 years    | HD                  | 0.711 (0.705–0.718)          |
| Holme et al   | 3 years    | HD                  | 0.705 (0.700–0.708)          |
| Wagner et al  | 1 year     | HD and PD           | 0.712 (0.711–0.715)          |
|               | 2 years    |                      | 0.712 (0.709–0.713)          |
|               | 3 years    |                      | 0.703 (0.702–0.706)          |
| Mauri et al   | 1 year     | HD                  | 0.670 (0.668–0.675)          |
| Geddes et al  | 1 year     | HD and PD           | 0.676 (0.671–0.677)          |
|               | 2 years    |                      | 0.671 (0.668–0.674)          |
|               | 3 years    |                      | 0.667 (0.664–0.669)          |
|               | 5 years    |                      | 0.663 (0.660–0.665)          |
| Hutchinson et al | 1 year | HD                  | 0.667 (0.666–0.671)          |
|               | 2 years    |                      | 0.667 (0.666–0.669)          |
|               | 3 years    |                      | 0.663 (0.661–0.665)          |
|               | 5 years    |                      | 0.663 (0.662–0.665)          |
| Floege et al  | 1 year     | PD                  | 0.778 (0.768–0.780)          |
| Floege et al  | 2 years    | PD                  | 0.775 (0.767–0.777)          |
| Holme et al   | 3 years    | PD                  | 0.767 (0.765–0.768)          |
| Wagner et al  | 1 year     | HD and PD           | 0.709 (0.789–0.793)          |
|               | 2 years    |                      | 0.770 (0.770–0.775)          |
|               | 3 years    |                      | 0.763 (0.762–0.767)          |
| Mauri et al   | 1 year     | PD                  | 0.800 (0.791–0.808)          |
| Geddes et al  | 1 year     | HD and PD           | 0.722 (0.769–0.774)          |
|               | 2 years    |                      | 0.748 (0.746–0.749)          |
|               | 3 years    |                      | 0.750 (0.748–0.753)          |
|               | 5 years    |                      | 0.737 (0.735–0.740)          |
| Hutchinson et al | 1 year | HD                  | 0.767 (0.763–0.771)          |
|               | 2 years    |                      | 0.752 (0.748–0.754)          |
|               | 3 years    |                      | 0.749 (0.748–0.753)          |
|               | 5 years    |                      | 0.742 (0.741–0.745)          |

* C-statistics presented as medians and interquartile ranges from pooled imputation results.

Abbreviations: HD, hemodialysis; PD, peritoneal dialysis.

Figure 2 C-statistics reported in the original study compared to those found through external validation (for the original time frame and original dialysis modality).

Note: The validation in the original study of Floege et al was an external validation also.
per time span of patient inclusion, or per chosen method for patient inclusion.

Furthermore, percentage of missing data of predictors can differ between derivation and validation, especially for CRP, which had a large missing rate in our cohort. However, this was dealt with by using multiple imputation, under the assumption that these data were missing at random. Baseline mortality, which can largely influence calibration results, differs strongly per time period and also per country and setting, such as an inpatient clinic, hospital, or outpatient clinic. The marked overestimation of mortality in the model of Hutchinson et al can be accounted for by the dramatic change in mortality rates of dialysis patients between the 1970s and late 1990s, and exemplifies the fact that recalibration over time is important to adjust models to new event rates.

The importance of differences in patient characteristics was also seen in the large difference between results in HD and PD patients. It seems counterintuitive that all models showed better discrimination for PD patients, even though they were not designed for this group of patients. However, this can be accounted for by a larger case-mix heterogeneity within the PD-patient population, specifically in age, allowing

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**Figure 3** Kaplan–Meier survival curves by quartile of the prognostic index.
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Figure 4 Calibration plots per model.

Abbreviations: HD, hemodialysis; PD, peritoneal dialysis.

for better discrimination, which is a population characteristic and has little to do with model fit. Due to such differences in case-mix and mortality rates, this validation study can be seen as a true test of the transportability of the models to patients from different source populations, in which population differences on a geographical and temporal level can be a great strength. If a model performs well across a spectrum of different populations, a universally applicable model may be within reach.

Another factor that complicates comparison is the different prediction time frames used by the studies. The larger the time frame, the more difficult it is to predict a patient’s death. For a better comparison and validation, we urge studies to include the baseline hazard function of a Cox prediction model or at least the baseline hazard value for clinically relevant time points, as specified in the TRIPOD guidelines.

To conclude, we warn against drawing general conclusions on which validated model is “the best”, as the different models were tested on transportability to a different degree and reproducibility was not always apparent. Clinical expertise remains leading when interpreting the differences between derivation and validation cohorts and differences between model performance.

This study contains a number of great strengths. To our knowledge, this is the first study to conduct a systematic review of existing models for predicting mortality in dialysis.
patients and subsequently validate and compare the existing models. To guarantee study reproducibility and transparency to the greatest extent possible, the PRISMA and TRIPOD guidelines were followed. Furthermore, missing data were accounted for by performing multiple imputations. This allowed us to test each model on the same patient population, and decreased the chance of selection bias.28 Additionally, NECOSAD is a highly representative large cohort of incident dialysis patients, and the large number of events within NECOSAD provide us with unbiased and precise estimations of calibration and discrimination performance measures.51,52

We have chosen to perform very conservative model updating as a sensitivity analysis only, and recommend that our main (unupdated) results be used when interpreting the performance of the validated models. Since validation means assessing the performance of a predefined model with new data, one could wonder whether model updating leads to a new model, which would then need external validation itself.34,53 If the validation results demonstrate that a model is not transportable to a specific (sub)population, this could give incentive for model updating in the context of an impact study or during clinical implementation.23

It is clear that before clinical implementation of prognostic models is warranted, extensive validation is necessary. The aim of clinical implementation is for such models to function as a useful tool for nephrologists, using evidence-based

| Reference | Time frame | Original population | Calibration: HD and PD | Calibration: PD |
|-----------|------------|---------------------|------------------------|----------------|
|           |            |                     | Calibration slope (SE) | Calibration slope (SE) |
|           |            |                     | Calibration-in-the-large* | Calibration-in-the-large* |
| Floege et al7 | 1 year | HD | 0.71 (0.04) | 14.0% vs 13.5% |
| Floege et al7 | 2 years | HD | 0.83 (0.04) | 22.1% vs 26.9% |
| Holme et al7 | 3 years | HD | 0.96 (0.05) | 47.6% vs 37.2% |
| Wagner et al7 | 1 year | HD and PD | 0.82 (0.04) | 4.0% vs 13.5% |
|             | 2 years | HD and PD | 0.82 (0.04) | 8.9% vs 26.9% |
|             | 3 years | HD and PD | 0.82 (0.04) | 14.2% vs 37.2% |
| Mauri et al45 | 1 year | HD | 0.42 (0.04) | 12.0% vs 12.6% |
| Geddes et al3 | 1 year | HD and PD | 0.82 (0.05) | 41.7% vs 13.5% |
|             | 2 years | HD and PD | 0.82 (0.05) | 59.3% vs 26.9% |
|             | 3 years | HD and PD | 0.82 (0.05) | 69.7% vs 37.2% |
|             | 5 years | HD and PD | 0.82 (0.05) | 80.9% vs 58.2% |
| Hutchinson et al55 | 1 year | HD | 0.50 (0.03) | 45.3% vs 13.5% |
|             | 2 years | HD | 0.50 (0.03) | 51.4% vs 26.9% |
|             | 3 years | HD | 0.50 (0.03) | 60.3% vs 37.2% |
|             | 5 years | HD | 0.50 (0.03) | 70.8% vs 58.2% |

Note: *Predicted vs observed risk of death.
Abbreviations: HD, hemodialysis; PD, peritoneal dialysis; SE, standard error.

| Reference | Time frame | Original population | Calibration: HD | Calibration: PD |
|-----------|------------|---------------------|----------------|----------------|
|           |            |                     | Calibration slope (SE) | Calibration slope (SE) |
|           |            |                     | Calibration-in-the-large* | Calibration-in-the-large* |
| Floege et al7 | 1 year | HD | 0.67 (0.04) | 16.7% vs 16.9% |
| Floege et al7 | 2 years | HD | 0.79 (0.05) | 26.2% vs 30.9% |
| Holme et al7 | 3 years | HD | 0.92 (0.06) | 52.8% vs 42.2% |
| Wagner et al7 | 1 year | HD and PD | 0.85 (0.05) | 5.0% vs 16.9% |
|             | 2 years | HD and PD | 0.85 (0.05) | 11.1% vs 30.9% |
|             | 3 years | HD and PD | 0.85 (0.05) | 17.6% vs 42.2% |
|             | 5 years | HD and PD | 0.85 (0.05) | 25.0% vs 52.8% |
| Mauri et al45 | 1 year | HD | 0.34 (0.05) | 16.9% vs 15.7% |
| Geddes et al3 | 1 year | HD and PD | 0.75 (0.06) | 47.6% vs 16.9% |
|             | 2 years | HD and PD | 0.75 (0.06) | 66.0% vs 30.9% |
|             | 3 years | HD and PD | 0.75 (0.06) | 76.1% vs 42.2% |
|             | 5 years | HD and PD | 0.75 (0.06) | 86.1% vs 64.2% |
| Hutchinson et al55 | 1 year | HD | 0.46 (0.03) | 51.6% vs 16.9% |
|             | 2 years | HD | 0.46 (0.03) | 58.1% vs 30.9% |
|             | 3 years | HD | 0.46 (0.03) | 67.3% vs 42.2% |
|             | 5 years | HD | 0.46 (0.03) | 77.5% vs 64.2% |

Note: *Predicted vs observed risk of death.
Abbreviations: HD, hemodialysis; PD, peritoneal dialysis; SE, standard error.
medicine that combines individual clinical expertise, patients’ choices, and the best external evidence.5,4

For future research, we suggest that more effort is put into validating, building upon, and implementing existing models in different populations. In the emerging field of prediction modeling, it is essential to provide summaries and evaluations of previously conducted studies through external validation, in order to move forward and prevent research waste. This study brings us a step closer to the use of a model with which personalized information on prognosis can be given to dialysis patients, aiding patient-centered decision making.

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