Predicting 6-Year Mortality Risk in Patients With Type 2 Diabetes

Brian J. Wells, MD1
Anil Jain, MD2
Susana Arrigain, MA1
Changhong Yu, MS1
Wayne A. Rosenkrans, Jr., PhD3
Michael W. Kattan, PhD1

OBJECTIVE — The objective of this study was to create a tool that predicts the risk of mortality in patients with type 2 diabetes.

RESEARCH DESIGN AND METHODS — This study was based on a cohort of 33,067 patients with type 2 diabetes identified in the Cleveland Clinic electronic health record (EHR) who were initially prescribed a single oral hypoglycemic agent between 1998 and 2006. Mortality was determined in the EHR and the Social Security Death Index. A Cox proportional hazards regression model was created using medication class and 20 other predictor variables chosen for their association with mortality. A prediction tool was created using the Cox model coefficients. The tool was internally validated using repeated, random subsets of the cohort, which were not used to create the prediction model.

RESULTS — Follow-up in the cohort ranged from 1 day to 8.2 years (median 28.6 months), and 3,661 deaths were observed. The prediction tool had a concordance index (i.e., c statistic) of 0.752.

CONCLUSIONS — We successfully created a tool that accurately predicts mortality risk in patients with type 2 diabetes. The incorporation of medications into mortality predictions in patients with type 2 diabetes should improve treatment decisions.

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Patients with type 2 diabetes have an increased risk of mortality (1–4), and tools for predicting overall mortality in diabetic patients are lacking. Tools for predicting the risk of cardiovascular disease (CVD) have been created and assist physicians in the prevention of CVD among patients with type 2 diabetes (5–8). However, these tools do not consider the risk of all-cause mortality. Preventing a myocardial infarction is important but not if the patient dies prematurely from kidney disease. Treating each disease in isolation is a strategy that fails to consider the overall effect to the patient. Scores have been created to adjust for diabetes severity and other comorbid conditions when mortality is studied (9,10). However, these indexes do not provide overall mortality predictions and therefore are of limited use in routine clinical practice. Furthermore, existing prediction tools do not adjust for a patient’s current therapy. Specific diabetes medications may decrease the risk of one complication while increasing the risk of other outcomes. Tools are needed to help clinicians tailor therapy to individual patients to minimize mortality risk on the basis of characteristics of the patient, his or her disease, and the available treatment options. The purpose of this study was to create a mortality risk calculator for patients with type 2 diabetes that can be used to aid in treatment decisions.

RESEARCH DESIGN AND METHODS — This study was conducted on a retrospective cohort of patients with type 2 diabetes whose data were collected for clinical and administrative purposes in the electronic health record (EHR) at Cleveland Clinic. The cohort began in 1998, and follow-up data were obtained through 2006.

Eligibility criteria at baseline
Baseline was defined as the date of the first prescription for a qualifying oral antidiabetic agent in an eligible patient. Because we are interested in patients with type 2 diabetes, the cohort was limited to patients at least 18 years of age with a diagnosis of diabetes. Patients with diabetes were identified if they had a single diagnosis of diabetes in the “History” or “Problem List” sections of their chart. To reduce the chance of misclassification due to “rule out” diagnoses, we required two occurrences of “diabetes” for patients with diabetes identified from the “encounter diagnosis” section. The following codes from the ICD-9 were used to identify patients with diabetes: 250–250.99, 357.2, 362.01, 362.02, 366.41, and 648.01–648.04.

The cohort was further limited to patients who were prescribed a single one of the four most common types of oral hypoglycemic agents: sulfonylureas (SFUs), meglitinides (MEGs), biguanides (BIGs), or thiazolidinediones (TZDs). Patients prescribed α-glucosidase inhibitors were excluded because of an inadequate sample (n = 149). Patients with prescriptions for multiple oral agents at baseline were excluded because of the substantial number of possible two- and three-drug (10) combinations.

Exclusions
Polycystic ovarian syndrome is sometimes treated with a biguanide and could be confused with diabetes. Noninsulin injectable medications are used infrequently in the treatment of type 2 diabetes, and the patients receiving these medications may be fundamentally different. Therefore, patients with polycystic ovarian syndrome (ICD-9 256.4) and patients prescribed noninsulin injectable diabetes medications at baseline were excluded.

Outcome
Mortality was determined in the EHR and with linkage to the Social Security Death...
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Index (SSDI). Patients classified as deceased per the SSDI or the EHR but who continued to have vital statistics entered into the EHR were counted as alive.

**Predictor variables**
The following variables were included in the model because of their independent associations with mortality in the literature: estimated glomerular filtration rate (GFR), A1C, BMI, systolic blood pressure (SBP), diastolic blood pressure (DBP), HDL and LDL cholesterol, triglycerides, history of congestive heart failure (CHF), history of coronary heart disease, smoking status, use of concomitant medications (insulin, ACE inhibitor/angiotensin receptor blocker [ARB], aspirin, clopidogrel, or lipid-lowering drug), new diabetes, sex, race, age, and oral medication class. Values for all of the predictor variables were extracted from the EHR. GFR was calculated from serum creatinine using the simplified equation from the Modification of Diet in Renal Disease Study Group (11). BMI was calculated using the standard equation (weight in kilograms divided by the square of height in meters). The baseline date was defined as the date of the first prescription for an oral hypoglycemic agent in a qualified individual. The baseline value for the predictor variables was defined as the value on the baseline date or the most recent historical value. If no historical lab or vital value was available then the value closest to the baseline date up to 21 days into the future was defined as the baseline. Patients were considered to be newly diabetic if they had been seen before their baseline date by either an endocrinologist or primary care physician at Cleveland Clinic and did not have a diagnosis of diabetes entered in the EHR at that time.

**Interactions**
A limited number of interactions were explored for inclusion in the predictive model but for parsimony were only included if they were statistically significant ($P < 0.05$). The following interactions were investigated in the model:

Medication class $\times$ GFR and medication class $\times$ age. The medication class interactions with GFR and age were included because of the precautions advised for use of biguanides in older adults and in patients with renal dysfunction (12).

Sex $\times$ race. The interaction for sex and race was explored because of the higher risk of coronary heart disease seen in black women (13).

Medication class $\times$ CHF. The interaction between medication class and heart failure was explored because of the precautions advised for using TZDs and BIGs in patients with CHF (12).

**Statistical methods**
The analytic dataset was built using SAS, version 9.1. Missing values were imputed using Multiple Imputation by Chained Equations package, version 1.16, for R (14). Imputation was performed to maximize the available information and to reduce the potential bias introduced by deleting incomplete records. The imputation was performed using regression techniques that include all baseline patients and all baseline variables as predictors and without knowledge of the outcome. Linearity assumptions of the ordinal and continuous variables were relaxed by fitting the model using restricted cubic splines.

A Cox proportional hazards regression model was created with the predictor variables and interactions listed above with time to death as the outcome. Statistical analyses were performed using R for Unix, version 2.3.1 (15). The coefficients from the fitted Cox model were also used to develop an interactive Web-based tool that calculates the predictions automatically. The Web-based calculator is available from http://www.clinicriskcalculators.org.

Tenfold cross-validation was used to validate our modeling approach. With this method, the data are randomly partitioned into 10 equal segments. One segment (10% of the data) is extracted before the prediction model is fitted. The model is fitted with the remaining 90% of the data, and the prediction accuracy is evaluated on the outcomes observed in the 10% subsample. The process is repeated (10 times) until each segment of the data has been used to assess the prediction accuracy. A calibration curve was created by plotting the quintiles of the predicted probabilities on the observed estimates for the entire cohort. The model was also evaluated through the use of the concordance index (i.e., c statistic). In this process, the model is graded on its ability to differentiate between all possible discordant pairs of patients. For example, patient 1 dies after 6 months in the cohort. Patient 2 dies after 3 years in the cohort. Does the predictive model correctly show that patient 1 had a higher risk of death?

Concordance indexes can vary between 0.5 (chance) and 1.0 (perfect prediction).

**RESULTS** — The final cohort had a sample size of 33,067 patients. Table 1 shows the characteristics of the cohort by drug class. Overall, the patients were predominantly white with a similar proportion of men and women. Patients taking BIGs were younger, had less heart failure, and were more likely to have newly diagnosed diabetes compared with patients taking the other drugs. Patients taking TZDs were the most likely to be using insulin, whereas patients taking MEGs had the lowest levels of LDL cholesterol and triglycerides.

Cumulative mortality by drug class is shown in Fig. 1. There were 1,958 patients followed for at least 5 years. The median length of follow-up was 28.6 months (range 1 day–8.2 years). The number of deaths per drug class were 799, 135, 2,220, and 507 for BIGs, MEGs, SFUs, and TZDs, respectively.

The interaction between race and sex was found not to be statistically significant and was removed from the model for parsimony. The other interactions (medication $\times$ GFR, medication $\times$ age, and medication $\times$ CHF) were all statistically significant. The medication $\times$ CHF interaction was the least significant of these interactions and was removed from the final model to reduce the size of the final nomogram. Figure 2A and B shows the paper-based nomogram for predicting 6-year survival (model coefficients are available in the supplemental table in an online appendix, available at http://dx.doi.org/10.2337/dc08-1047).

The BIGs were associated with the lowest risk of mortality in younger patients. Because of the interactions between medication with GFR and medication with age, the medication associated with the highest probability of survival varied according to individual patient characteristics. As expected, smoking and high levels of A1C were associated with lower survival. LDL had a U-shaped relationship with mortality. The lowest risk of mortality was associated with an LDL of 150 mg/dl and increased with higher or lower LDL levels.

Figure 3 shows the calibration curve for the mortality prediction. The predicted survival was quite accurate at all quintiles of mortality risk. The concordance index for the survival prediction was 0.752, which indicates that the no-
program was correct 75.2% of the time in identifying which patient had the highest risk among all possible discordant patient pairs.

CONCLUSIONS — The prediction tool created in this study was accurate in predicting 6-year mortality risk among patients with type 2 diabetes. The c statistic of 0.752 indicates good discrimination ability of this tool, and the calibration curve shows that the prediction does not significantly overestimate or underestimate risk. If the current prediction tool performs well in other cohorts of patients with type 2 diabetes, it offers clinicians a tool for tailoring antidiabetes treatments with the aim of improving survival among patients with type 2 diabetes.

We are not aware of other tools designed to predict mortality in patients with type 2 diabetes in the clinical setting. It is hoped that the recently published Diabetes Complications Severity Index (DCSI) will improve the ability to adjust for the severity of type 2 diabetes in future regression models predicting mortality (9). However, the DCSI is not a stand-alone prediction tool and is designed to be used as a covariate in a larger model. We did not have all of the predictors necessary to add the DCSI score to our model.

The widespread use of this tool may be limited by the number of predictor variables that we included in the model and the requirement of GFR. We did not attempt to reduce the model using variable selection techniques (e.g., stepwise regression) because the omission of insignificant predictors tends to harm predic-
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A
Points

Age (GFR=30 Medication=IG)
Age (GFR=60 Medication=IG)
Age (GFR=90 Medication=IG)
Age (GFR=30 Medication=MEG)
Age (GFR=60 Medication=MEG)
Age (GFR=90 Medication=MEG)
Age (GFR=30 Medication=SFU)
Age (GFR=60 Medication=SFU)
Age (GFR=90 Medication=SFU)
Age (GFR=30 Medication=TZD)
Age (GFR=60 Medication=TZD)
Age (GFR=90 Medication=TZD)

Hemoglobin A1c
Body Mass Index
Systolic Blood Pressure
Diastolic Blood Pressure
High Density Lipoprotein
Low Density Lipoprotein

B
Points

Triglycerides
Gender
Caucasian
History of Heart Disease?
History of Heart Failure?
Smoking Status
Is Pt on Insulin?
Is Pt on an Ace Inhibitor or ARB?
New Diabetic?
Is Pt on Aspirin?
Is Pt on Clopidogrel?
Is Pt on a Lipid Lowering Drug?
Total Points
6-Year Survival Prob.
We hope that the online version of the calculator will make the model more user-friendly by calculating GFR automatically from serum creatinine and eliminating the need for the paper nomogram. The online calculator is available as mentioned earlier under statistical methods.

This study has several other weaknesses that should be discussed. First, there is the potential for misclassification bias surrounding the designation of the baseline oral medication. Prescriptions outside Cleveland Clinic are unknown, and we did not assess for medication changes within our system. However, in a separate analysis of this cohort, we found that ~75% of patients were treated with the same oral diabetes medication throughout their time in the cohort. A second weakness of the study involves the substantial amount of missing data for some predictor variables. However, the imputation techniques used should help to limit the potential bias caused by simply eliminating incomplete records. Third, we were unable to accurately define the exact duration of type 2 diabetes. Patients whose diabetes was diagnosed on their first visit to Cleveland Clinic will not be recognized as newly diabetic. However, duration of diabetes is probably captured by the presence or absence of comorbidities and other variables such as GFR. Diabetes complications may be the most accurate reflection of diabetes duration because the disease frequently goes undetected for years after it first appears (17). Young et al. (9) found that duration of diabetes was not independently predictive of mortality. In addition, the presence of comorbid conditions such as heart failure and stroke may be underdocumented in the EHR. However, there is no reason to suggest that the documentation would vary according to drug class. To maximize the presence or absence of baseline conditions, we included information beyond documentation of ICD-9 codes. For instance, patients with a documentation of a coronary revascularization procedure were considered to have a history of heart disease. Despite any possible lack of documentation in the EHR, the model performed very well. A final weakness could be the loss to follow-up that is inevitable in this type of study. Fortunately, we were able to link participants with the SSDI, which should capture deaths of patients regardless of loss to follow-up at Cleveland Clinic.

The major strengths of the study include the large sample size and cohort design. The good prediction accuracy obtained through the cross-validation of the model suggests strong internal validity of these results. The Web-based version of this calculator provides the tool in an easily accessible format for clinical use. The current model requires external validation before the applicability of this model to other patient populations will be known.
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