Enhancing ASCVD Risk Prediction in NASH/NAFLD Patients

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Background

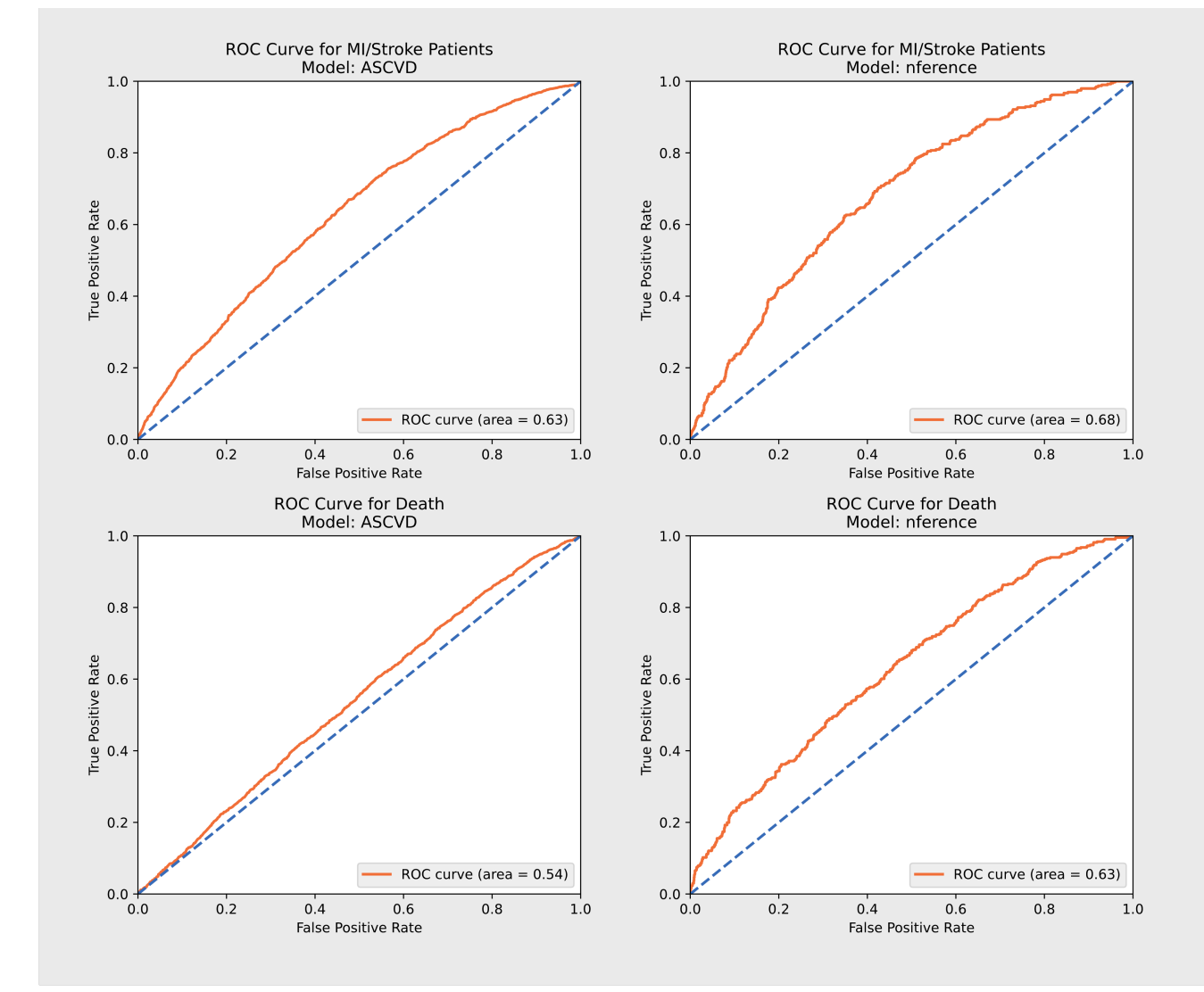
- Non-alcoholic steatohepatitis (NASH), also known as metabolic dysfunction-associated steatohepatitis (MASH), and non-alcoholic fatty liver disease (NAFLD), also known as metabolic dysfunction-associated steatotic liver disease (MASLD), are burgeoning health concerns in the United States, linked to increased cardiovascular diseases (CVD) risks.¹
- The leading cause of morbidity and mortality outside of liver disease is CVD among the NASH population.¹
- The American Heart Association's (AHA) Atherosclerotic Cardiovascular Disease

Table 1. Baseline Demographics and **Clinical Characteristics**

Cohort Statistics									
	Count (%)								
Sample Size									
Total Number	9185.0 (100.0%)								
Past Medical History									
Hypertension	6831.0 (74.4%)								
Type 2 DM	206.0 (2.2%)								
Smoker									
Smoker	1803.0 (19.6%)								
Non-Smoker	7382.0 (80.4%)								
Gender									
Female	4589.0 (50.0%)								
Male	4596.0 (50.0%)								
Race									
*White / Asian / Unknown / Other	9012.0 (98.1%)								
*Black or AA	173.0 (1.9%)								
Observed Outcomes									
Decompensated Cirrhosis	4383.0 (47.7%)								
MI / Stroke	1904.0 (20.7%)								
Death Within 10 yrs	2139.0 (23.3%)								
Data Coverage									
Mean Years After Diagnosis (Std. Dev.)	8.18 (5.9)								
*Race was grouped into categories used in A	ASCVD REP Equation								

Figure 1. Comparison of ROC Curves between ASCVD REP and Our Models

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(ASCVD) Risk Estimator Plus lacks validated efficacy for mortality prediction in NASH/NAFLD patients suggesting that additional variables should be considered to enhance the accuracy.^{2,3,4}

Objective

This study aims to augment the predictive accuracy of ASCVD risk and death among NASH patients by proposing alternative logistic regression models.

Methods

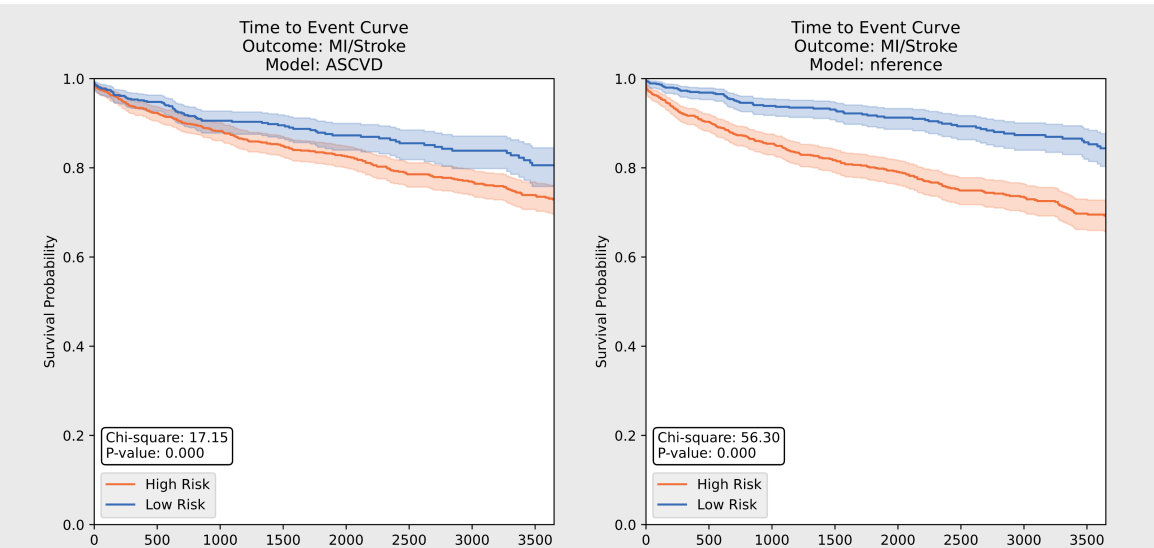
- A retrospective EHR dataset from a large US integrated delivery network health system encompassing multiple sites within the US was curated. A 9,185 patient cohort of biopsy confirmed NASH patients was curated using structured and unstructured data including ICD codes and Clinical Notes. Patients with a history of MI/Stroke before their diagnosis were excluded.
- The variables were constrained to be within 30 days of the NASH diagnosis, and the outcomes were limited to within 10 years and censored by the last available data.

Table 2. Comparison of Variables for Models

	MI/Stroke	Death	ASCVI
Universal			
Age			
Hypertension PMH			
Smoker			
MI/Stroke and Death Models			
International Normalized Ratio (INR)			×
Prothrombin time			×
Total Bilirubin			×
Alanine Aminotransferase			×
Alkaline Phosphatase, Serum			×
Aspartate Aminotransferase			×
Direct Bilirubin			×
Hemoglobin A1c, Blood			•
Exclusive MI/Stroke			
Platelet Count		×	×
Sodium [Moles / Volume] in Serum, Plasma or Blood		×	×
Gamma globulin [Mass/volume] in Serum or Plasma by Electrophoresis		×	×
HDL		×	
Exclusive Death			
LDL Cholesterol [Mass/volume] in Serum or Plasma	×		×
Systolic Blood Pressure	×		
Total Iron binding capacity (TIBC) [Moles/volume] in Serum or Plasma	×		×
Gender	×		
Exclusive ASCVD			
Race	×	×	
Total Cholesterol	×	×	
Diastolic Blood Pressure	×	×	
History of DM		<u> </u>	

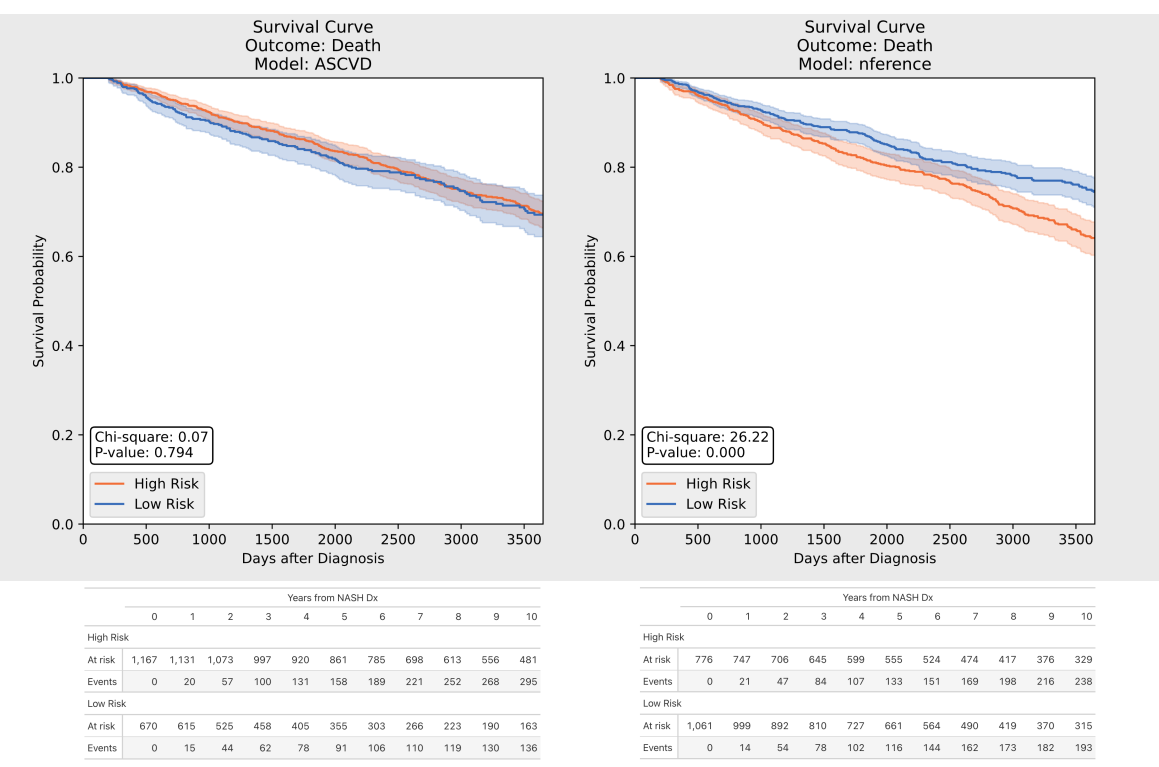
The AUROC curves increased when incorporating additional liver specific variables (0.63 -> 0.68 and 0.54 -> 0.63 for MI/Stroke and Death Outcomes, respectively)

Figure 2. Time to Event and Survival Curves for ASCVD and Our Models



- Demographic, clinical, and laboratory variables underwent a recursive feature elimination process.
- We created two logistic regression models to predict MI/Stroke and Death. An optimal threshold was determined using Youden's J for the output classes of High or Low risk.
- The performance of the AHA's ASCVD Risk Estimator Plus (REP) was compared against our models, which incorporated alternative predictors. A REP score ≥7.5% was considered High Risk. Area Under the Receiver Operating Characteristic curve (AUC) was employed to evaluate and compare both models.
- Survival and time to event curves were generated using the statistical python package lifelines and evaluated for statistical significance at a predetermined alpha of 0.05.

0	500	10	000	150 Days		2000 Diagn		500	3000 3500			0	500	10	000 [150 Days a		2000 Diagn		00	300	0	3500
	Years from NASH Dx														Years f	rom NAS	H Dx						
	0	1	2	3	4	5	6	7	8	9	10		0	1	2	3	4	5	6	7	8	9	10
High Ris	sk											High Ri	sk										
At risk	1,040	939	867	788	713	659	590	526	464	419	357	At risk	917	818	745	682	617	572	507	451	400	360	309
Events	8	68	102	126	145	160	177	195	203	215	227	Events	11	76	114	138	157	172	189	205	212	223	234
Low Ris	k											Low Ris	k										
At risk	623	540	448	386	341	297	259	222	184	161	135	At risk	746	661	570	492	437	384	342	297	248	220	183
Events	6	30	47	54	57	63	66	70	74	75	80	Events	3	22	35	42	45	51	54	60	65	67	73



The time to event and survival curves should significant differences in outcomes for both nference models. The difference was statistically significant for the ASCVD MI/Stroke outcome, however was not significant for the death outcome



The ASCVD Risk Estimator Plus demonstrated suboptimal predictive accuracy for mortality and myocardial infarction (MI) / stroke events in NASH/NAFLD patients, with an AUC of 0.63 for MI/Stroke events and an even lower AUC of 0.54 for mortality (figure 1). In contrast, our logistic regression models exhibited higher AUCs, with 0.68 for MI/Stroke events and 0.63 for mortality, indicating enhanced predictive accuracy for both outcomes (figure 1).

- The inclusion of liver-specific markers, such as alanine aminotransferase, and other novel predictors, such as INR, helped to achieve greater accuracy. A Kaplan-Meier time to event analysis for 10-year MI/Stroke risk was generated to evaluate the prediction over 10 years, and a similar survival analysis was conducted for mortality. Pearson's chi-squared tests yielded significant statistics of 56.30 (p=0.000) for MI/Stroke and 26.22 (p=0.000) for mortality.
- These findings underscore the potential limitations of the ASCVD Risk Estimator Plus in predicting outcomes for NASH/NAFLD patients and highlight the significance of incorporating liver-specific markers such as ALT and INR and other novel predictors for more accurate risk assessment, which could lead to improved clinical decision-making and patient management strategies in this population.

Our findings underscore the necessity of revisiting the current CV risk models for NASH/NAFLD patients to incorporate more holistic and disease-specific variables. Conclusion

• The proposed logistic regression model that includes liver-specific biomarkers, such as ALT and INR, improves the prediction of cardiovascular mortality and MI events, thereby facilitating better clinical decision-making and patient-centered care.

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