Analysis on the Cessation Effect on Smoking among Patients with MDD with the Combination Treatment of Behavioral Activation and Varenicline

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Introduction

Major Depressive Disorder (MDD) and smoking demonstrate a significant bidirectional relationship, with MDD patients showing higher probability of heavy smoking patterns, elevated nicotine dependence levels, and intensified withdrawal symptoms during cessation attempts. While previous studies have documented varenicline's effectiveness in smoking cessation treatment, and Behavioral Activation (BA) has emerged as a promising intervention for depression-related symptoms, we still lack comprehensive understanding of their combined therapeutic effects in treating smokers with MDD. In a recent randomized, placebo-controlled, 2×2 factorial trial comparing BA for smoking cessation (BASC) and standard behavioral treatment (ST), with and without varenicline administration, researchers found no significant advantage of BASC over ST in adult smokers with current or past MDD diagnosis.

In this report, we aim to extend the previous findings by analyzing the potential moderating effects of baseline variables on treatment outcomes and examining the predictive factors for smoking abstinence, while controlling for both behavioral intervention approaches and pharmacotherapy. We investigate multiple potential moderators, including depression-related variables (e.g., anhedonia severity, current MDD status, depression intensity), smoking-related characteristics (e.g., nicotine dependence level, daily cigarette consumption, cessation readiness), and demographic factors. Through this analysis, we expect to identify specific patient characteristics that may predict better responses to particular treatment approaches, ultimately contributing to the development of more individualized and effective smoking cessation strategies for this high-risk population.

Data

The dataset used in this analysis contains 300 observations with 25 variables (including 1 identification column) collected from a randomized, placebo-controlled, 2×2 factorial trial involving adult smokers diagnosed with current or past major depressive disorder (MDD). In order to ensure data quality and reliability of our analysis, we firstly conducted data preprocessing to handle missing values. After removing observations with incomplete records, our final analytical sample consisted of 241 participants.

The data documented participants' demographic information, depression-related variables, and smoking-related characteristics measured at baseline, as well as their treatment assignment and outcomes. The participants were randomized to receive either Behavioral Activation for smoking cessation (BASC) or standard behavioral treatment (ST), combined with either varenicline or placebo, forming a 2×2 factorial design. The completion of treatment and follow-up assessments were carefully monitored throughout the trial period.

Variable	Description
abst	Smoking Abstinence
Var	Pharmacotherapy (Varenicline)
BA	Psychotherapy (Behavioral Activation)
age_ps	Age at phone interview
sex_ps	Sex at phone interview
NHW	Non-Hispanic White indicator
Black	Black indicator
Hisp	Hispanic indicator
inc	Income (ordinal categorical, low to high)
edu	Education (ordinal categorical, low to high)
$ftcd_score$	FTCD score at baseline
ftcd.5.mins	Smoking within 5 mins of waking up
bdi_score_pq1	BDI score at baseline (a measure of depression)
cpd_ps	Cigarettes per day at baseline phone survey
crv_total_pq1	Cigarette reward value at baseline
$hedonsum_n_pq1$	Pleasurable Events Scale at baseline – substitute reinforcers
$hedonsum_y_pq1$	Pleasurable Events Scale at baseline – complementary reinforcers
$shaps_score_pq1$	Anhedonia
otherdiag	Other lifetime DSM-5 diagnosis
antidepmed	Taking antidepressant medication at baseline
$\mathrm{mde}_\mathrm{curr}$	Current vs past MDD
NMR	Nicotine Metabolism Ratio
Only.Menthol	Exclusive Mentholated Cigarette User
readiness	Baseline readiness to quit smoking

Analysis of the baseline characteristics by treatment groups reveals several distinctive patterns that need attention. When comparing the varenicline and placebo groups, we observe a substantial disparity in abstinence rates, with the varenicline group achieving 28% success compared to 8.4% in the placebo group (p < 0.001), indicating varenicline's potential therapeutic effectiveness. However, other baseline characteristics maintained balanced distributions between these pharmacotherapy groups, suggesting successful randomization. Similarly, when comparing behavioral activation (BA) and standard treatment groups, most characteristics were comparably distributed, with one notable exception: the proportion of participants using antidepressant medication differed significantly between groups (35% in BA group versus 19% in standard treatment group, p = 0.004).

Table 2: Data Summary Table

	Grouped by Pharmacotherapy Status		Grouped by F	sychotherapy Sta	tus	
Characteristic	No Varenicline	Varenicline	p-value	No Psychotherapy	Psychotherapy	p-value
abst			< 0.001			0.7
0	98 (92%)	96 (72%)		93 (79%)	101 (81%)	
1	9 (8.4%)	38 (28%)		24 (21%)	23 (19%)	
BA	, ,	, ,	0.6	,	, ,	
0	54 (50%)	63 (47%)				
1	53 (50%)	71 (53%)				
age_ps	52(45, 59)	52 (40, 59)	0.4	51 (43, 58)	54 (41, 60)	0.2
sex_ps			0.4			> 0.9
1	45 (42%)	63 (47%)		52 (44%)	56 (45%)	
2	62 (58%)	71 (53%)		65 (56%)	68 (55%)	
NHW	, ,	, ,	0.7	, ,	, ,	0.5
0	70 (65%)	85 (63%)		78 (67%)	77 (62%)	
1	37 (35%)	49 (37%)		39 (33%)	47 (38%)	
Black	. ,		0.2			0.6

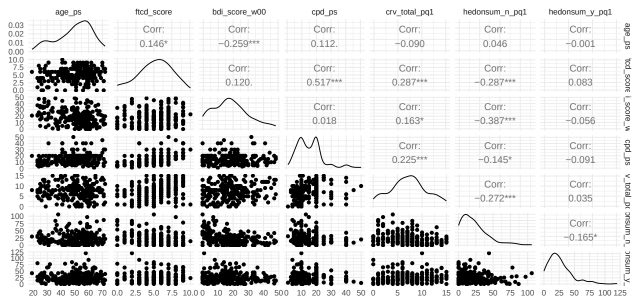
Table 2: Data Summary Table (continued)

	Grouped by Pharmacotherapy Status		Grouped by Psychotherapy Status			
Characteristic	No Varenicline	Varenicline	p-value	No Psychotherapy	Psychotherapy	p-value
0	47 (44%)	70 (52%)		55 (47%)	62 (50%)	
1	60 (56%)	64 (48%)		62 (53%)	62 (50%)	
Hisp	, ,	, ,	0.7	, ,	, ,	0.5
0	100 (93%)	127 (95%)		109 (93%)	118 (95%)	
1	7 (6.5%)	7 (5.2%)		8 (6.8%)	6 (4.8%)	
inc	, ,	, ,	0.6	, ,	, ,	0.3
1	38 (36%)	47 (35%)		40 (34%)	45 (36%)	
2	25 (23%)	31 (23%)		28 (24%)	28 (23%)	
3	18 (17%)	20 (15%)		21 (18%)	17 (14%)	
4	16 (15%)	15 (11%)		10 (8.5%)	21 (17%)	
5	10 (9.3%)	21 (16%)		18 (15%)	13 (10%)	
edu		(, , ,	0.2	(0.6
1	1 (0.9%)	0 (0%)		0 (0%)	1 (0.8%)	
2	3 (2.8%)	9 (6.7%)		4 (3.4%)	8 (6.5%)	
3	29 (27%)	34 (25%)		30 (26%)	33 (27%)	
4	46 (43%)	45 (34%)		48 (41%)	43 (35%)	
5	28 (26%)	46 (34%)		35 (30%)	39 (31%)	
ftcd_score	6 (4, 7)	5 (4, 7)	0.3	6 (4, 7)	5 (4, 7)	0.4
ftcd.5.mins	51 (48%)	57 (43%)	0.4	57 (49%)	51 (41%)	0.2
bdi score w00	18 (9, 27)	18 (9, 24)	0.9	18 (12, 25)	18 (9, 26)	0.6
cpd_ps	15 (10, 20)	15 (10, 20)	0.5	15 (10, 20)	15 (10, 20)	>0.9
crv total pq1	7 (5, 9)	7 (5, 10)	0.8	7 (5, 9)	8 (5, 10)	0.4
hedonsum_n_pq1	17 (9, 30)	20 (9, 35)	0.3	18 (9, 32)	20 (9, 31)	>0.9
hedonsum_y_pq1	24 (12, 36)	19 (13, 31)	0.2	21 (13, 35)	22 (12, 33)	>0.9
shaps_score_pq1	1 (0, 4)	1(0,3)	>0.9	1 (0, 4)	1 (0, 3)	0.5
otherdiag	1 (0, 1)	1 (0, 0)	0.6	1 (0, 1)	1 (0, 0)	0.7
0	58 (54%)	77 (57%)	0.0	64 (55%)	71 (57%)	0.1
1	49 (46%)	57 (43%)		53 (45%)	53 (43%)	
antidepmed	10 (1070)	0. (10/0)	0.3	33 (1370)	00 (10/0)	0.004
0	74 (69%)	101 (75%)	0.0	95 (81%)	80 (65%)	0.001
1	33 (31%)	33 (25%)		22 (19%)	44 (35%)	
mde curr	00 (0170)	00 (2070)	0.6	22 (1070)	11 (0070)	0.8
0	59 (55%)	69 (51%)	0.0	61 (52%)	67 (54%)	0.0
1	48 (45%)	65 (49%)		56 (48%)	57 (46%)	
NMR	0.32 (0.20, 0.45)	0.31 (0.21, 0.50)	0.8	0.32 (0.20, 0.46)	0.32 (0.22, 0.48)	0.7
Only.Menthol	5.52 (5.25, 5.45)	0.01 (0.21, 0.00)	0.7	0.02 (0.20, 0.40)	0.02 (0.22, 0.40)	>0.7
0	41 (38%)	55 (41%)	0.1	47 (40%)	49 (40%)	/0.0
1	66 (62%)	79 (59%)		70 (60%)	75 (60%)	
readiness	7.00 (6.00, 8.00)	7.00 (6.00, 8.00)	0.2	7.00 (6.00, 8.00)	7.00 (6.00, 8.00)	0.9
Var	1.00 (0.00, 0.00)	1.00 (0.00, 0.00)	0.2	1.00 (0.00, 0.00)	1.00 (0.00, 0.00)	0.6
0				54 (46%)	53 (43%)	0.0
1				63 (54%)	71 (57%)	
¹ n (%); Median (IC				00 (04/0)	11 (01/0)	

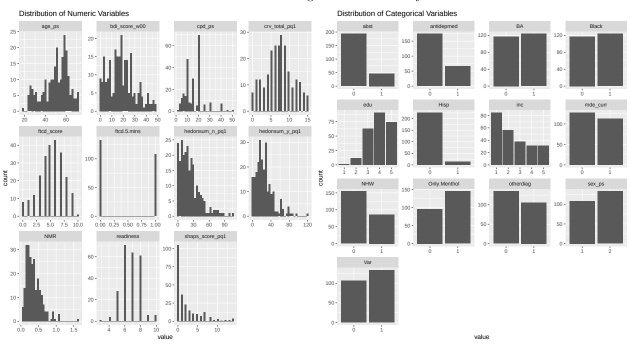
² Pearson's Chi-squared test; Wilcoxon rank sum test; Fisher's exact test

In our examination of the relationships among continuous variables in the dataset, we identified several significant correlations that deserve attention. The daily cigarette consumption (cpd_ps) exhibited a moderate positive correlation with the Fagerström Test for Cigarette Dependence score (FTCD score) (r=0.517, p<0.001). This correlation aligns with theoretical expectations, as both measures are designed to assess different aspects of smoking dependency. Furthermore, we observed that the Beck Depression Inventory score (bdi_score_w00) was negatively correlated with both measures of hedonic capacity (hedonsum_n_pq1 and hedonsum_y_pq1), a finding that supports the established relationship between increased depression severity and diminished ability to experience pleasure.

Correlation Matrix of Baseline Predictors

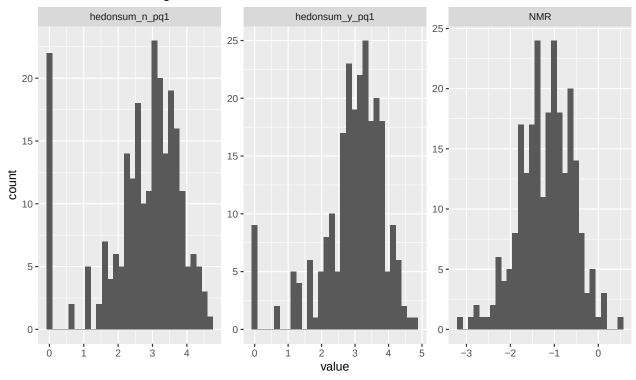


Our distribution analysis of numeric variables in the dataset identified notable right-skewed patterns in several key measures, particularly in both hedonic capacity indices (hedonsum_n_pq1, hedonsum_y_pq1) and the nicotine metabolite ratio (NMR). To address these violations of normality assumptions, we implemented log transformations for the affected variables. Post-transformation examination revealed substantially improved symmetry and reduced skewness in these distributions, ensuring their suitability for subsequent statistical analyses. We found that the remaining numeric variables, including participant age, Fagerström Test for Cigarette Dependence score, and depression-related measures, demonstrated approximately normal distributions and were therefore retained in their original scale for analysis.



This is the distribution of hedonsum n pq1, hedonsum y pq1, and NMR after log transformation.

Distribution after Log Transformation



Regression Analysis

Basic Models

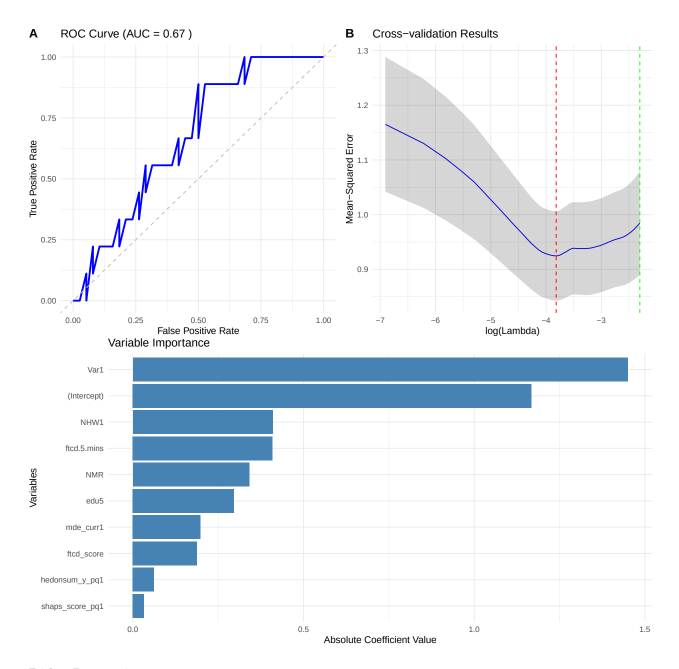
We first implemented two fundamental regularized regression models (Ridge and LASSO regression without interaction terms) to examine the relationship between smoking abstinence and our predictors.

Lasso Regression

Our LASSO regression model demonstrated strong predictive performance, achieving an Area Under the Curve (AUC) of 0.841 on the training dataset and 0.705 on the test dataset. Through its inherent variable selection mechanism, the LASSO model identified 10 key predictors of smoking cessation success.

Among these predictors, varenicline treatment emerged as the most influential factor, showing the strongest positive effect (coefficient = 0.802) on abstinence probability. This coefficient indicates that varenicline administration increases the log-odds of abstinence by 0.802, which translates to an odds ratio of $\exp(0.802)$ = 2.23. In the social domain, we found that both high income level (inc5: coefficient = 0.226, odds ratio = $\exp(0.226) = 1.25$) and antidepressant medication use (coefficient = 0.237, odds ratio = $\exp(0.237) = 1.27$) were associated with increased likelihood of abstinence.

The impact of education levels showed heterogeneous effects in our analysis. While edu4 demonstrated a negative association with abstinence (coefficient = -0.342, reducing odds by 29% as $\exp(-0.342) = 0.71$), edu5 showed a positive relationship (coefficient = 0.236, increasing odds by 27% as $\exp(0.236) = 1.27$). Regarding clinical measures, we observed moderate negative associations with abstinence: each unit increase in Fagerström Test score (ftcd_score) was associated with a 16% reduction in abstinence odds (coefficient = -0.169, odds ratio = $\exp(-0.169) = 0.84$), and similarly, each unit increase in positive hedonic capacity (hedonsum_y_pq1) corresponded to a 24% decrease in abstinence odds (coefficient = -0.280, odds ratio = $\exp(-0.280) = 0.76$).

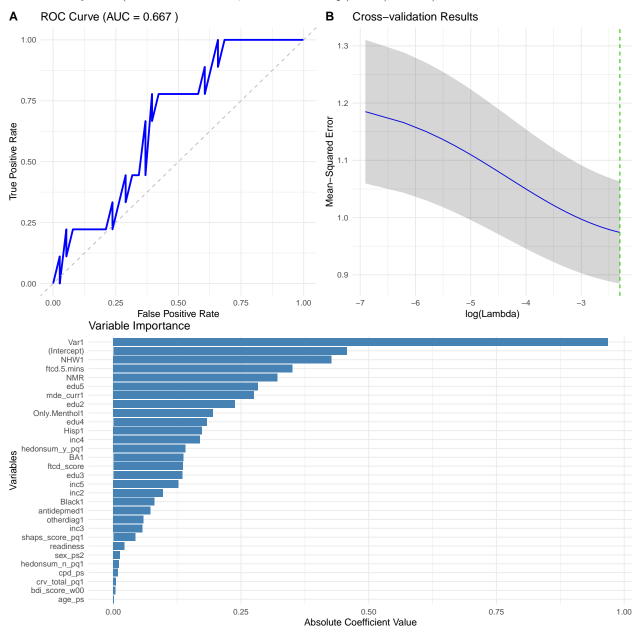


Ridge Regression

The Ridge regression model demonstrated stronger performance on the training set with an AUC of 0.878, though its test set performance (AUC = 0.651) was slightly lower than the LASSO model. While Ridge regression retains all variables in the model, the variable importance analysis revealed patterns similar to our LASSO findings, with varenicline treatment emerging as one of the strongest positive predictors (coefficient = 0.740, odds ratio = $\exp(0.740) = 2.10$). Notably, both antidepressant medication use (coefficient = 0.480, odds ratio = $\exp(0.480) = 1.62$) and Hispanic ethnicity (coefficient = 0.534, odds ratio = $\exp(0.534) = 1.71$) showed substantial positive associations with abstinence outcomes.

Among the clinical predictors, we observed moderate positive associations for both nicotine metabolism ratio (NMR: coefficient = 0.372, increasing odds by 45% as $\exp(0.372) = 1.45$) and menthol cigarette use (coefficient = 0.377, increasing odds by 46% as $\exp(0.377) = 1.46$). In contrast, depression-related measures demonstrated consistent negative associations with abstinence probability. Specifically, current MDD status decreased the odds of abstinence by 26% (coefficient = -0.299, odds ratio = $\exp(-0.299) = 0.74$), and higher

scores on the positive hedonic capacity measure (hedonsum_y_pq1) were associated with reduced odds of abstinence by 28% (coefficient = -0.336, odds ratio = $\exp(-0.336) = 0.72$).



The comparison of the Ridge model and the Lasso model is shown below. The table of coefficients are also included.

Table 3: Comparison of Basic Ridge and Lasso Regression Models

Variable	Ridge Regression Model	Lasso Regression Model
(Intercept) Var1 BA1	-0.457 0.968 -0.137	-1.168 1.45 Eliminated
age_ps sex_ps2 NHW1	0.001 0.013 0.427	Eliminated Eliminated 0.41
Black1	-0.081	Eliminated

Table 3: Comparison of Basic Ridge and Lasso Regression Models (continued)

Variable	Ridge Regression Model	Lasso Regression Model
Hisp1	0.174	Eliminated
inc2	-0.097	Eliminated
inc3	0.057	Eliminated
inc4	-0.170	Eliminated
inc5	0.128	Eliminated
edu2	0.238	Eliminated
edu3	-0.135	Eliminated
edu4	-0.183	Eliminated
edu5	0.283	0.296
$ftcd_score$	-0.136	-0.188
${ m ftcd.5.mins}$	0.351	0.409
bdi_score_w00	-0.004	Eliminated
$\operatorname{cpd}\mathtt{_ps}$	-0.009	Eliminated
crv_total_pq1	0.005	Eliminated
$hedonsum_n_pq1$	0.011	Eliminated
$hedonsum_y_pq1$	-0.141	-0.062
$shaps_score_pq1$	-0.043	-0.033
otherdiag1	-0.059	Eliminated
antidepmed1	0.073	Eliminated
mde_curr1	-0.275	-0.198
NMR	0.321	0.341
Only.Menthol1	0.195	Eliminated
readiness	-0.022	Eliminated

Table 4: Results of Basic Ridge and Lasso Regression Models

Metric	Ridge Regression Model	Lasso Regression Model
AUC (Train)	0.831	0.824
AUC (Test)	0.667	0.670
Accuracy (Train)	0.809	0.814
Accuracy (Test)	0.809	0.787
Lambda Min	0.100	0.022

Model with Interaction Terms

Our investigation of interaction effects progressed through two stages, each guided by specific theoretical considerations.

Basic Interaction Model

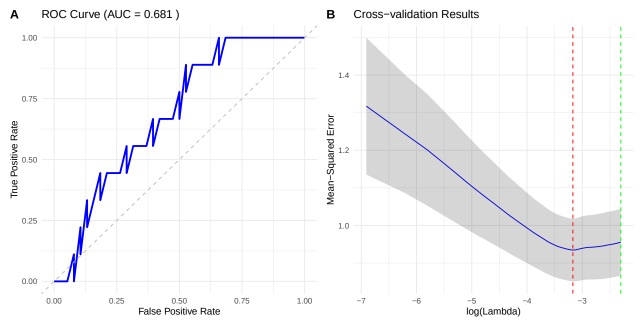
In constructing our basic interaction model, we prioritized the examination of treatment-specific interactions that aligned with our primary research objectives regarding treatment effectiveness. For Behavioral Activation (BA), we specifically focused on its interactions with depression-related variables, including current MDD status (mde_curr), Beck Depression Inventory score (bdi_score_w00), and Snaith-Hamilton Pleasure Scale score (shaps_score_pq1). This selection was theoretically driven, as BA was explicitly designed to target depression symptoms, with particular emphasis on addressing anhedonic symptoms. Additionally, we incorporated the interaction between BA and current MDD status to evaluate whether BA's therapeutic effectiveness varies between patients with current versus past MDD diagnoses.

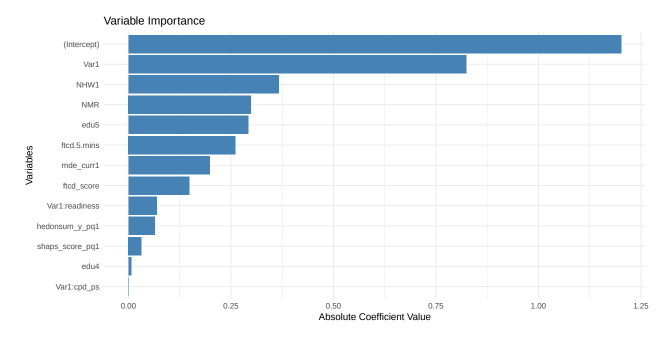
Meanwhile, our analysis of varenicline interactions concentrated on smoking-related characteristics, specifically the Fagerström Test score (ftcd_score), cigarettes per day (cpd_ps), and readiness to quit measures. This selection was informed by previous research findings suggesting that pharmacotherapy outcomes may be moderated by baseline smoking intensity and motivational factors. These interaction terms were essential

for understanding the differential effects of varenicline across varying levels of nicotine dependence and quit motivation.

Our initial interaction model demonstrated moderate predictive performance, achieving an AUC of 0.685 on the test dataset. The variable importance analysis revealed that varenicline maintained its position as a strong predictor, showing a substantial main effect (coefficient = 0.309) that increased the base odds of abstinence by $\exp(0.309) = 1.36$ times. However, we found that varenicline's effectiveness exhibited variation across different baseline smoking characteristics. Particularly noteworthy was the positive interaction with daily cigarette consumption (coefficient = 0.021, odds ratio = $\exp(0.021) = 1.02$), indicating an enhanced treatment effect with higher baseline smoking intensity. To illustrate, for participants reporting 20 cigarettes per day, varenicline's effectiveness showed an enhancement factor of $\exp(0.021 * 20) = 1.52$ compared to its baseline effect.

In examining behavioral treatment interactions, we observed that the relationship between Behavioral Activation (BA) and current MDD status revealed a subtle but important pattern. The negative interaction coefficient (-0.013) suggests that for patients with current MDD, BA's effectiveness was marginally reduced (odds ratio = $\exp(-0.013) = 0.987$, approximately 1.3% reduction in abstinence odds). This finding underscores the particular challenges in treating this specific patient subgroup. Additionally, our analysis revealed that educational level served as a significant moderator of treatment effectiveness, with contrasting effects observed at different levels: edu4 was associated with reduced odds of abstinence (coefficient = -0.250, odds ratio = $\exp(-0.250) = 0.779$, representing a 22.1% reduction), while edu5 showed increased odds (coefficient = 0.215, odds ratio = $\exp(0.215) = 1.24$, indicating a 24% increase).





Advanced Interaction Model

In developing our advanced interaction model, we expanded the scope of our analysis to incorporate a more comprehensive set of theoretically grounded interactions, focusing on three fundamental mechanisms. Firstly, we investigated the interaction between depression symptoms, specifically examining the relationship between Beck Depression Inventory score and Snaith-Hamilton Pleasure Scale score (bdi_score_w00 \times shaps_score_pq1). This interaction was selected based on the hypothesis that concurrent presence of severe depression and pronounced anhedonia might create uniquely challenging conditions for achieving smoking cessation.

Secondly, we examined potential interactions between smoking behavior characteristics and quit motivation (ftcd_score \times readiness). This investigation was driven by the theoretical premise that elevated levels of nicotine dependence, as measured by the Fagerström Test score, might attenuate the typically positive influence of quit motivation on cessation outcomes. Finally, we explored social interactions, particularly focusing on education level's interaction with quit readiness (edu \times readiness) and income level's interaction with antidepressant medication use (inc \times antidepmed). These interactions were chosen to investigate how educational background might moderate the translation of motivation into successful cessation, and how social factors might influence medication effectiveness through differential access to healthcare resources and support systems.

Our advanced interaction model demonstrated superior predictive performance, achieving the highest test set Area Under the Curve (AUC) of 0.711. This improvement in predictive accuracy suggests that the incorporation of comprehensive interaction terms successfully captured important complex relationships in the data. The variable importance analysis revealed several notable interaction patterns that warrant detailed examination.

The most prominent finding emerged from the social domain, where we observed a strong interaction between middle income level and antidepressant medication use (coefficient = 1.204, odds ratio = $\exp(1.204)$ = 3.33). This substantial effect indicates that participants in the middle income group experienced more than triple the effectiveness from antidepressant medication compared to other income groups. Treatment-related interactions also demonstrated meaningful patterns: varenicline's effectiveness showed positive interaction with both daily cigarette consumption (coefficient = 0.019, odds ratio per cigarette = $\exp(0.019)$ = 1.019) and readiness to quit (coefficient = 0.017, odds ratio per unit = $\exp(0.017)$ = 1.017). To illustrate, for participants smoking 20 cigarettes per day, the varenicline treatment effect was enhanced by a factor of $\exp(0.019 * 20) = 1.46$.

The model additionally captured significant behavioral interactions that provide insights into treatment mechanisms. We found that increased nicotine dependence slightly attenuated the benefit of quit readiness (coefficient = -0.011, odds ratio = $\exp(-0.011) = 0.989$ per unit increase in dependence score). Similarly, educational background emerged as a modifier of quit readiness effects, with edu4 level showing a modest negative interaction (coefficient = -0.014, odds ratio = $\exp(-0.014) = 0.986$ per unit increase in readiness score). These findings suggest that both nicotine dependence and educational background influence how effectively quit motivation translates into successful abstinence outcomes.

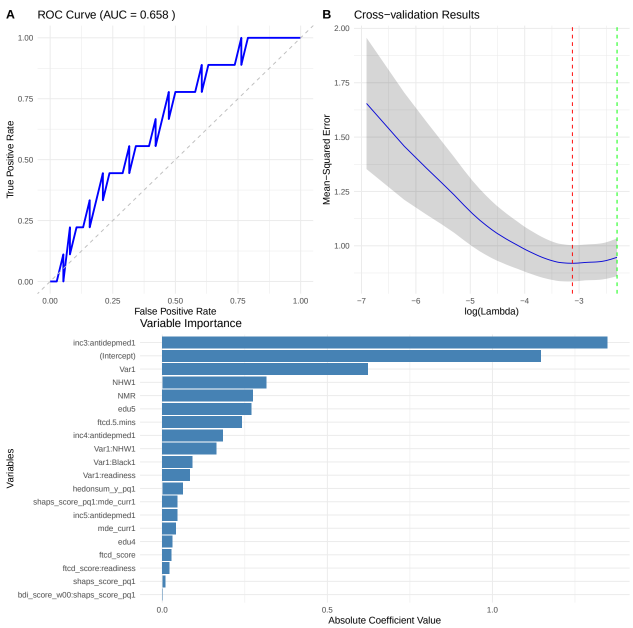


Table 5: Comparison of Models with Interaction Terms

Variable	Model with Basic Interaction Term	Model with Advanced Interaction Terms
(Intercept)	-1.202	-1.147
BA1	Eliminated	Eliminated
Var1	0.824	0.623
age_ps	Eliminated	Eliminated

Table 5: Comparison of Models with Interaction Terms (continued)

Variable	Model with Basic Interaction Term	Model with Advanced Interaction Terms
sex_ps2	Eliminated	Eliminated
NHW1	0.367	0.315
Black1	Eliminated	Eliminated
Hisp1	Eliminated	Eliminated
inc2	Eliminated	Eliminated
inc3	Eliminated	Eliminated
inc4	Eliminated	Eliminated
inc5	Eliminated	Eliminated
edu2	Eliminated	Eliminated
edu3	Eliminated	Eliminated
edu4	-0.007	-0.031
edu5	0.292	0.27
bdi_score_w00	Eliminated	Eliminated
cpd _ps	Eliminated	Eliminated
$ftcd_score$	-0.148	-0.028
ftcd.5.mins	0.261	0.241
shaps_score_pq1	-0.032	-0.009
mde_curr1	-0.198	-0.041
crv total pq1	Eliminated	Eliminated
hedonsum_n_pq1	Eliminated	Eliminated
hedonsum_y_pq1	-0.064	-0.062
otherdiag1	Eliminated	Eliminated
antidepmed1	Eliminated	Eliminated
NMR	0.299	0.275
Only.Menthol1	Eliminated	Eliminated
readiness	Eliminated	Eliminated
BA1:shaps_score_pq1	Eliminated	Eliminated
BA1:mde_curr1	Eliminated	Eliminated
BA1:bdi_score_w00	Eliminated	Eliminated
Var1:ftcd score	Eliminated	Eliminated
Var1:ncd_score Var1:cpd_ps	0.001	Eliminated
-		
Var1:readiness	0.069	0.084
BA1:Black1	NA	Eliminated
BA1:NHW1	NA	Eliminated
Var1:Black1	NA	0.092
Var1:NHW1	NA	0.164
$bdi_score_w00:shaps_score_pq1$	NA	0
$shaps_score_pq1:mde_curr1$	NA	-0.046
$\operatorname{cpd}\operatorname{_ps:ftcd}\operatorname{_score}$	NA	Eliminated
ftcd_score:readiness	NA	-0.022
inc2:antidepmed1	NA	Eliminated
inc3:antidepmed1	NA	1.348
inc4:antidepmed1	NA	-0.184
inc5:antidepmed1	NA	-0.046
edu2:readiness	NA	Eliminated
edu3:readiness	NA	Eliminated
edu4:readiness	NA	Eliminated
edu5:readiness	NA	Eliminated
ftcd_score:Only.Menthol1	NA	Eliminated
NMR:Only.Menthol1	NA	Eliminated
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Table 6: Results of Models with Interaction Terms

Metric	Model with Basic Interaction Term	Model with Advanced Interaction Terms
AUC (Train)	0.821	0.839
AUC (Test)	0.681	0.658
Accuracy (Train)	0.809	0.820
Accuracy (Test)	0.809	0.787
Lambda Min	0.042	0.044

Discussion

In this study, we employed a series of regularized regression models to investigate the effectiveness of behavioral activation and varenicline for smoking cessation among individuals with current or past MDD. Through progressively complex modeling approaches, from basic LASSO and Ridge regression to interaction models, we gained several key insights about treatment effects and their moderators.

Our analytical models demonstrated consistently robust predictive performance, with test AUC values ranging from 0.651 to 0.711. The advanced interaction model achieved the highest predictive accuracy, highlighting the importance of considering treatment effect heterogeneity in our analysis. Notably, the relatively small gap between training and test performance (AUC difference < 0.15) suggests our models successfully avoided severe overfitting, which we attribute to the effective implementation of regularization strategies.

Regarding treatment effectiveness, varenicline emerged as a consistently strong predictor of smoking cessation success across all model specifications. The basic LASSO model identified a substantial positive effect (coefficient = 0.802), which maintained its robustness even after accounting for various interaction terms. In contrast, behavioral activation demonstrated more modest and variable effects, with the lack of a strong main effect aligning with previous trial findings where BASC showed no significant advantage over standard behavioral treatment.

Our interaction models revealed significant heterogeneity in treatment effects across different patient subgroups. For varenicline, we observed enhanced effectiveness among heavier smokers (Var \times cpd_ps: 0.019) and participants with higher readiness to quit (Var \times readiness: 0.017), suggesting particular benefit for more severely dependent smokers who demonstrate strong quit motivation. For behavioral activation, the negative interaction with current MDD status (BA \times mde_curr: -0.013) indicated potentially reduced effectiveness among currently depressed patients, underscoring the challenges in treating this specific subgroup.

The advanced interaction model uncovered important social dimensions of treatment response. We identified a strong positive interaction between middle income level and antidepressant medication use (inc3 \times antidepmed: 1.204), suggesting significant social moderation of depression treatment effectiveness. Additionally, the negative interaction between education level and quit readiness (edu4 \times readiness: -0.014) indicated that the translation of quit motivation into successful cessation varies across educational backgrounds.

Several limitations should be noted in our analysis. First, our relatively small sample size (n=241 after removing missing values) may have limited our ability to detect smaller interaction effects. Second, our binary outcome measure may not fully capture the complexity of smoking cessation trajectories. Third, our models did not account for potential time-varying treatment effects or the dynamic nature of depression symptoms during quit attempts.

Future research directions should address several key areas. Larger sample sizes would enable more precise estimation of interaction effects and potentially reveal additional treatment effect moderators. The incorporation of longitudinal measures for both smoking behavior and depression symptoms could provide valuable insights into temporal dynamics of treatment effects. Additionally, more sophisticated modeling approaches, such as Bayesian methods or advanced machine learning techniques, might better capture the complex relationships between patient characteristics and treatment outcomes.

Despite these limitations, our findings yield important clinical implications. The consistent effectiveness of varenicline, particularly among heavier smokers, supports its continued use as a first-line treatment. However,

the variable effects of behavioral activation and the significant role of social factors suggest the need for more personalized treatment approaches. Clinicians should carefully consider both clinical and social factors when developing smoking cessation interventions for patients with MDD.

References

1. Hitsman, Brian, George D. Papandonatos, Jacqueline K. Gollan, Mark D. Huffman, Raymond Niaura, David C. Mohr, Anna K. Veluz-Wilkins, et al. "Efficacy and Safety of Combination Behavioral Activation for Smoking Cessation and Varenicline for Treating Tobacco Dependence among Individuals with Current or Past Major Depressive Disorder: A 2 × 2 Factorial, Randomized, Placebo-controlled Trial." Addiction 118, no. 9 (September 2023): 1710–25. https://doi.org/10.1111/add.16209.

Code Appendix

```
knitr::opts_chunk$set(echo = FALSE)
knitr::opts_chunk$set(message = FALSE)
knitr::opts_chunk$set(warning = FALSE)
library(mice, warn.conflicts = FALSE)
library(naniar)
library(ggplot2)
library(dplyr)
library(readr)
library(tidyr)
library(readxl)
library(ggpubr)
library(gtsummary)
library(GGally)
library(ggcorrplot)
library(knitr)
library(kableExtra)
library(lubridate)
library(patchwork)
library(introdataviz)
library(glmnet)
library(gt)
library(L0Learn)
library(gridExtra)
library(purrr)
library(pROC)
library(caret)
# Load the data
data <- read.csv("../Data/project2.csv")</pre>
# Transform data type
data$abst <- as.factor(data$abst)</pre>
data$Var <- as.factor(data$Var)</pre>
data$BA <- as.factor(data$BA)</pre>
data$sex_ps <- as.factor(data$sex_ps)</pre>
data$NHW <- as.factor(data$NHW)</pre>
data$Black <- as.factor(data$Black)</pre>
data$Hisp <- as.factor(data$Hisp)</pre>
data$inc <- as.factor(data$inc)</pre>
data$edu <- as.factor(data$edu)</pre>
data$otherdiag <- as.factor(data$otherdiag)</pre>
data$antidepmed <- as.factor(data$antidepmed)</pre>
data$mde curr <- as.factor(data$mde curr)</pre>
data$Only.Menthol <- as.factor(data$Only.Menthol)</pre>
data$readiness <- as.numeric(data$readiness)</pre>
dim(data)
summary(data)
data <- na.omit(data)</pre>
```

```
summary(data)
dim(data)
summary_var1 <- tbl_summary(data %>% select(-id), by = Var, type = list(readiness ~ "continuous")) %>%
  add_p() %>%
  modify header(stat 1 = "**No Varenicline**", stat 2 = "**Varenicline**")
summary_var2 <- tbl_summary(data %>% select(-id), by = BA, type = list(readiness ~ "continuous")) %>%
  add p() %>%
  modify_header(stat_1 = "**No Psychotherapy**", stat_2 = "**Psychotherapy**")
tbl_merge(
 tbls = list(summary_var1, summary_var2),
  tab_spanner = c("**Grouped by Pharmacotherapy Status**", "**Grouped by Psychotherapy Status**")
) %>%
  as_kable_extra(booktabs = TRUE,
caption = "Data Summary Table",
longtable = TRUE, linesep = "") %>%
kableExtra::kable_styling(font_size = 8, latex_options = c("repeat_header", "HOLD_position"))
continuous_vars <- data[, c("age_ps", "ftcd_score", "bdi_score_w00", "cpd_ps", "crv_total_pq1", "hedons</pre>
corr_plot <- ggpairs(continuous_vars, title = "Correlation Matrix of Baseline Predictors") +</pre>
  theme_minimal()
ggsave("../Figures/corr_plot.png", plot = corr_plot, dpi = 300)
corr plot
dist numeric <- data %>%
  select(-id) %>%
  keep(is.numeric) %>%
  gather() %>%
  ggplot(aes(value)) +
    facet_wrap(~ key, scales = "free") +
    geom_histogram() +
    ggtitle("Distribution of Numeric Variables")
dist_cat <- data %>%
  select(-id) %>%
  keep(is.factor) %>%
  gather(key = "key", value = "value") %>%
  ggplot(aes(x = value)) +
    facet_wrap(~ key, scales = "free") +
    geom_bar()+
    ggtitle("Distribution of Categorical Variables")
gridExtra::grid.arrange(dist_numeric, dist_cat, ncol = 2)
data$hedonsum_n_pq1 <- log(data$hedonsum_n_pq1 + 1)</pre>
data$hedonsum_y_pq1 <- log(data$hedonsum_y_pq1 + 1)</pre>
data$NMR <- log(data$NMR)</pre>
dist_after_log <- data %>%
  select(c(hedonsum_n_pq1, hedonsum_y_pq1, NMR)) %>%
```

```
gather() %>%
  ggplot(aes(value)) +
    facet_wrap(~ key, scales = "free") +
    geom_histogram() +
    ggtitle("Distribution after Log Transformation")
ggsave("../Figures/dist_after_log.png", plot = dist_after_log, dpi = 300)
dist_after_log
# Split the data
set.seed(123)
train_index <- createDataPartition(data$abst, p = 0.8, list = FALSE)</pre>
train_data <- data[train_index, ] %>% select(-id)
test_data <- data[-train_index, ] %>% select(-id)
X_train <- model.matrix(abst ~ ., data = train_data)[, -1]</pre>
Y_train <- factor(train_data$abst)</pre>
X_test <- model.matrix(abst ~ ., data = test_data)[, -1]</pre>
Y_test <- factor(test_data$abst)</pre>
model_fit <- function(X_train, Y_train, X_test, Y_test, alpha, lambda_seq = seq(0.001, 0.1, length.out
set.seed(123)
 cv_model <- cv.glmnet(X_train, Y_train, alpha = alpha, family = "binomial", lambda = lambda_seq)</pre>
best_lambda <- cv_model$lambda.min</pre>
lasso_coef <- coef(cv_model, s = best_lambda)</pre>
 # Get predictions for both train and test sets
 train_pred <- predict(cv_model, newx = X_train, s = best_lambda, type = "response")</pre>
 test_pred <- predict(cv_model, newx = X_test, s = best_lambda, type = "response")</pre>
 # Convert predictions to binary using 0.5 threshold
 train_pred_binary <- ifelse(train_pred > 0.5, 1, 0)
 test_pred_binary <- ifelse(test_pred > 0.5, 1, 0)
 # ROC and AUC
train_roc <- roc(Y_train, as.numeric(train_pred))</pre>
 test_roc <- roc(Y_test, as.numeric(test_pred))</pre>
 # Calculate metrics
metrics <- data.frame(</pre>
   Metric = c(
     "AUC (Train)",
     "AUC (Test)",
     "Accuracy (Train)",
     "Accuracy (Test)",
     "Lambda Min"
   ),
   Value = c(
     auc(train_roc),
     auc(test_roc),
     mean(train_pred_binary == Y_train),
```

```
mean(test_pred_binary == Y_test),
    cv_model$lambda.min
  )
) %>%
  mutate(Value = round(Value, 3))
# Create ROC curve data
roc df <- data.frame(</pre>
  FPR = 1 - test_roc$specificities,
  TPR = test roc$sensitivities
)
# Create ROC plot using ggplot2
roc_plot <- ggplot(roc_df, aes(x = FPR, y = TPR)) +</pre>
  geom_line(color = "blue", size = 1) +
  geom_abline(slope = 1, intercept = 0, linetype = "dashed", color = "gray") +
  labs(
    title = paste("ROC Curve (AUC =", round(auc(test_roc), 3), ")"),
   x = "False Positive Rate",
    y = "True Positive Rate"
  ) +
  theme_minimal()
# Create CV plot data
cv_df <- data.frame(</pre>
  lambda = log(cv_model$lambda),
  mean = cv_model$cvm,
  upper = cv_model$cvup,
  lower = cv_model$cvlo
)
# Create CV plot using qqplot2
cv_plot <- ggplot(cv_df, aes(x = lambda)) +</pre>
  geom_line(aes(y = mean), color = "blue") +
  geom_ribbon(aes(ymin = lower, ymax = upper), alpha = 0.2) +
  geom_vline(xintercept = log(cv_model$lambda.min), linetype = "dashed", color = "red") +
  geom_vline(xintercept = log(cv_model$lambda.1se), linetype = "dashed", color = "green") +
    title = "Cross-validation Results",
    x = "log(Lambda)",
   y = "Mean-Squared Error"
  ) +
  theme_minimal()
# Combine plots
combined_plot <- ggarrange(roc_plot, cv_plot,</pre>
                          ncol = 2,
                          labels = c("A", "B"))
# Create coefficient dataframe
coef_df <- data.frame(</pre>
  Variable = rownames(lasso_coef),
  Coefficient = as.vector(lasso_coef)
```

```
) %>%
mutate(Coefficient = ifelse(Coefficient == 0, "Eliminated", round(Coefficient, 3)))
return(list(
   combined_plot = combined_plot,
  coef_df = coef_df,
  metrics = metrics
))
}
plot_variable_importance <- function(coef_df) {</pre>
 # Calculate variable importance by taking absolute values of non-eliminated coefficients
var_importance <- coef_df %>%
  filter(Coefficient != "Eliminated") %>%
  mutate(
     Coefficient = as.numeric(Coefficient),
     Importance = abs(Coefficient)
   ) %>%
   arrange(desc(Importance))
 # Create bar plot of variable importance
 importance_plot <- ggplot(var_importance, aes(x = reorder(Variable, Importance), y = Importance)) +
  geom_bar(stat = "identity", fill = "steelblue") +
   coord_flip() + # Flip coordinates for horizontal bars
   theme minimal() + # Use minimal theme for clean look
  labs(title = "Variable Importance",
       x = "Variables",
        y = "Absolute Coefficient Value")
 # Return both plot and processed data
return(list(
   plot = importance_plot,
   importance_df = var_importance
))
}
lambda_seq <- seq(0.001, 0.1, length.out = 100)</pre>
lasso_model <- model_fit(X_train, Y_train, X_test, Y_test, alpha = 1, lambda_seq = lambda_seq)</pre>
print(lasso_model$combined_plot)
plot_variable_importance(lasso_model$coef_df)$plot
ggsave("../Figures/lasso_model.png", plot = lasso_model$combined_plot, dpi = 300)
ggsave("../Figures/lasso_variable_importance.png", plot = plot_variable_importance(lasso_model$coef_df)
lambda_seq \leftarrow seq(0.001, 0.1, length.out = 100)
ridge_model <- model_fit(X_train, Y_train, X_test, Y_test, alpha = 0, lambda_seq = lambda_seq)
print(ridge_model$combined_plot)
plot_variable_importance(ridge_model$coef_df)$plot
ggsave("../Figures/ridge_model.png", plot = ridge_model$combined_plot, dpi = 300)
ggsave("../Figures/ridge_variable_importance.png", plot = plot_variable_importance(ridge_model$coef_df)
```

```
combined_table <- full_join(ridge_model$coef_df, lasso_model$coef_df, by = "Variable") %>%
  kable(
    col.names = c("Variable", "Ridge Regression Model", "Lasso Regression Model"),
    caption = "Comparison of Basic Ridge and Lasso Regression Models"
  ) %>%
  kable_styling(bootstrap_options = c("striped", "hover"),
                full_width = FALSE, font_size = 8, latex_options = c("repeat_header", "HOLD_position"))
  column spec(1, bold = TRUE)
combined table
combined_result <- full_join(ridge_model$metrics, lasso_model$metrics, by = "Metric") %>%
    col.names = c("Metric", "Ridge Regression Model", "Lasso Regression Model"),
    caption = "Results of Basic Ridge and Lasso Regression Models"
  ) %>%
  kable_styling(bootstrap_options = c("striped", "hover"),
                full_width = FALSE, font_size = 8, latex_options = c("repeat_header", "HOLD_position"))
  column_spec(1, bold = TRUE)
combined_result
X_train_interactions <- model.matrix(</pre>
 ~ BA + Var +
   # Demographic variables
  age ps + sex ps + NHW + Black + Hisp + inc + edu +
   # Clinical and smoking-related variables
  bdi_score_w00 + cpd_ps + ftcd_score + ftcd.5.mins +
   shaps_score_pq1 + mde_curr + crv_total_pq1 +
  hedonsum_n_pq1 + hedonsum_y_pq1 +
  otherdiag + antidepmed + NMR + Only.Menthol + readiness +
   # BA interaction terms
  BA:shaps_score_pq1 + # BA * anhedonia
  BA:mde_curr + # BA * current depression status
BA:bdi_score_w00 + # BA * depression severity
   # Varenicline interaction terms
  Var:ftcd_score +  # Varenicline * nicotine dependence
  Var:cpd_ps +
                         # Varenicline * cigarettes per day
                    # Varenicline * readiness to quit
   Var:readiness,
 data = train_data %>% select(-abst))[, -1]
X_test_interactions <- model.matrix(</pre>
 ~ BA + Var +
   # Demographic variables
  age_ps + sex_ps + NHW + Black + Hisp + inc + edu +
   # Clinical and smoking-related variables
  bdi_score_w00 + cpd_ps + ftcd_score + ftcd.5.mins +
   shaps_score_pq1 + mde_curr + crv_total_pq1 +
  hedonsum_n_pq1 + hedonsum_y_pq1 +
  otherdiag + antidepmed + NMR + Only.Menthol + readiness +
   # BA interaction terms
   BA:shaps_score_pq1 + # BA * anhedonia
  BA:mde_curr + # BA * current depression status
BA:bdi_score_w00 + # BA * depression severity
```

```
# Varenicline interaction terms
  Var:readiness,
                          # Varenicline * readiness to quit
data = test_data %>% select(-abst))[, -1]
lambda_seq \leftarrow seq(0.001, 0.1, length.out = 100)
basic_interaction_model <- model_fit(X_train_interactions, Y_train, X_test_interactions, Y_test, alpha
print(basic_interaction_model$combined_plot)
plot_variable_importance(basic_interaction_model$coef_df)$plot
ggsave("../Figures/basic_interaction_model.png", plot = basic_interaction_model$combined_plot, dpi = 30
ggsave("../Figures/basic_interaction_variable_importance.png", plot = plot_variable_importance(basic_in
# Create model matrix with extended interaction terms
X_train_interactions <- model.matrix(</pre>
 ~ BA + Var +
   # Demographic variables
   age_ps + sex_ps + NHW + Black + Hisp + inc + edu +
   # Clinical and smoking-related variables
   bdi_score_w00 + cpd_ps + ftcd_score + ftcd.5.mins +
   shaps_score_pq1 + mde_curr + crv_total_pq1 +
   hedonsum_n_pq1 + hedonsum_y_pq1 +
   otherdiag + antidepmed + NMR + Only.Menthol + readiness +
   # BA interaction terms
   BA:shaps_score_pq1 + # BA * anhedonia
   BA:mde_curr + # BA * current depression status
BA:bdi_score_w00 + # BA * depression severity
   BA:Black +
                          # BA * race (Black)
   BA:NHW +
                          # BA * race (Non-Hispanic White)
   # Varenicline interaction terms
  Var:ftcd_score +  # Varenicline * nicotine dependence
Var:cpd_ps +  # Varenicline * cigarettes per day
Var:readiness +  # Varenicline * readiness to quit
Var:Black +  # Varenicline * race (Black)
                           # Varenicline * race (Non-Hispanic White)
   # Depression symptom interactions
   bdi_score_w00:shaps_score_pq1 + # Depression severity * anhedonia
   mde_curr:shaps_score_pq1 + # Current depression status * anhedonia
   # Smoking behavior interactions
   ftcd_score:cpd_ps +
                           # Nicotine dependence * cigarettes per day
   ftcd_score:readiness + # Nicotine dependence * readiness to quit
   # social status interactions
   antidepmed:inc +
                                 # Antidepressant medication * income
   edu:readiness +
                                  # Education * readiness to quit
   # Menthol cigarette interactions
   Only.Menthol:ftcd_score +  # Menthol use * nicotine dependence
Only.Menthol:NMR,  # Menthol use * nicotine metabolism ratio
data = train_data %>% select(-abst))[, -1]
X_test_interactions <- model.matrix(</pre>
 ~ BA + Var +
   # Demographic variables
```

```
age_ps + sex_ps + NHW + Black + Hisp + inc + edu +
   # Clinical and smoking-related variables
  bdi_score_w00 + cpd_ps + ftcd_score + ftcd.5.mins +
   shaps_score_pq1 + mde_curr + crv_total_pq1 +
  hedonsum_n_pq1 + hedonsum_y_pq1 +
   otherdiag + antidepmed + NMR + Only.Menthol + readiness +
   # BA interaction terms
  BA:shaps_score_pq1 + # BA * anhedonia
  BA:mde_curr + # BA * current depression status
BA:bdi_score_w00 + # BA * depression severity
  BA:Black +
                         # BA * race (Black)
  BA:NHW +
                          # BA * race (Non-Hispanic White)
   # Varenicline interaction terms
  Var:ftcd_score + # Varenicline * nicotine dependence
  Var:cpd_ps +  # Varenicline * cigarettes per day
Var:readiness +  # Varenicline * readiness to quit
  Var:Black +
                         # Varenicline * race (Black)
  Var:NHW +
                          # Varenicline * race (Non-Hispanic White)
   # Depression symptom interactions
  bdi_score_w00:shaps_score_pq1 + # Depression severity * anhedonia
  mde_curr:shaps_score_pq1 + # Current depression status * anhedonia
   # Smoking behavior interactions
  ftcd_score:cpd_ps +
                          # Nicotine dependence * cigarettes per day
  ftcd_score:readiness +
                               # Nicotine dependence * readiness to quit
   # social status interactions
  antidepmed:inc +
                         # Antidepressant medication * income
  edu:readiness +
                               # Education * readiness to quit
   # Menthol cigarette interactions
  Only.Menthol:ftcd_score + # Menthol use * nicotine dependence
   Only.Menthol:NMR,
                                # Menthol use * nicotine metabolism ratio
data = test_data %>% select(-abst))[, -1]
lambda_seq \leftarrow seq(0.001, 0.1, length.out = 100)
advanced_interaction_model <- model_fit(X_train_interactions, Y_train, X_test_interactions, Y_test, alp
print(advanced interaction model$combined plot)
plot_variable_importance(advanced_interaction_model$coef_df)$plot
ggsave("../Figures/advanced_interaction_model.png", plot = advanced_interaction_model$combined_plot, dp
ggsave("../Figures/advanced_interaction_variable_importance.png", plot = plot_variable_importance(advan
combined_table <- full_join(basic_interaction_model$coef_df, advanced_interaction_model$coef_df, by = "</pre>
 kable(
    col.names = c("Variable", "Model with Basic Interaction Term", "Model with Advanced Interaction Term
    caption = "Comparison of Models with Interaction Terms"
  kable_styling(bootstrap_options = c("striped", "hover"),
                full_width = FALSE, font_size = 8, latex_options = c("repeat_header", "HOLD_position"))
  column_spec(1, bold = TRUE) # Make Variable column bold
combined_table
combined_result <- full_join(basic_interaction_model$metrics, advanced_interaction_model$metrics, by =
 kable(
```