PM 592 Regression Analysis for Public Health Data Science

Week 10

Predictive Modeling

Predictive Modeling

Introduction to Prediction Models Predictive Model Building Predictive Power Optimizing Classification

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Lecture Objectives

- > Implement a complete prediction model-building method
- > Explain how to diagnose the predictive ability of these models
- > Describe the ROC curve and its metrics
- $\, \boldsymbol{\succ} \,$ Determine the best cut point for a prediction model

1. Review	4
✓ Assumptions in logistic regression – similarities and differences from OLS regression	_
✓ Goodness-of-fit measures	_
✓ Diagnosing outliers and influential values	_
✓ Automated selection procedures	
natemated solution procedures	-
	_
	_
	」 _
2. Introduction to Predictive Modeling	ī
There are two ways to approach the development of regression models:	1 _
Testing a hypothesized association	
"Does a violence prevention program in high schools successfully reduce the chance that students will experience bullying?"	_
Must consider potential confounders and effect modifiers.	-
2. Developing a prediction model	

"What are the factors that contribute to developing coronary heart disease?"

Confounders aren't of interest, as there is no specific association we are

We just want a model that has a successful prediction rate.

(Framingham coronary risk model)

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2. Introduction to Predictive Modeling			7
The model determines how variables, as a	TABLE 6. β-Coefficients Underlying Using TC Categories	CHD Prediction Sh	ieets
predict the outcome.	Variable Age, y Age squared, y	0.04826	Women 0.33766 -0.00268
	TC, mg/dL <160 160–199	Referent	-0.26138 Referent
	200-239 240-279 ≥280	0.17692 0.50539	0.20771 0.24385 0.53513
	HDL-C, mg/dL <:35 35–44	0.49744	0.84312 0.37796
(Equation 1): L_Chol _{men} =0.04826×age-0.65945 (if cholester	45-49 50-59	Referent -0.05107	0.19785 Referent
<160) +0.0 (if cholesterol 160 to 199) +0.17692 (if cholesterol 20 to 239) +0.50539 (if cholesterol 240 to 279) +0.65713 (if chole terol ≥280) +0.49744 (if HDL-C<35) +0.24310 (if HDL-C 35	Blood pressure Optimal	-0.00226	-0.42951 -0.53363
44) +0.0 (if HDL-C 45 to 49) -0.05107 (if HDL-C 50 to 5 -0.48660 (if HDL-C ≥60) -0.00226 (if blood pressure [B.	High normal Stage I hypertension	0.28320 - 0.52168	Referent -0.06773 0.26288
optimal) +0.0 (if BP normal) +0.28320 (if BP high normal) +0.52168 (if BP stage I hypertension) +0.61859 (if BP stage I hypertension) +0.42830 (if diabetes present) +0.0 (if diabetes)	I Diabetes Smoker	0.42839 0.52337	0.46573 0.59626 0.29246
hypertension) +0.42839 (if diabetes present) +0.0 (if diabetes n present) +0.52337 (if smoker) +0.0 (if not smoker).	Baseline survival function at 10 years, S(I)	0.90015	0.96246
I			
2. Introduction to Predictive Modeling			8
Which independent variables should be	considered?		
Anything that may help predict the outcome.	ome.		
Many independent variables can be incl			
The variables don't have to be etiological			
It is not necessary to consider confound	ing.		
0			
8			
2. Introduction to Predictive Modeling			9
How many variables should be in the fin	al model?		
Enough to predict the outcome well, but		اد	
 Rule of thumb: there should be at least 			
Y=1) for each predictor in the model (so	•		
well-powered and not biased)			

	_
2. Introduction to Predictive Modeling	
Parsimony	
Every model is a simplification of reality (parsimony).	
 If two models fit the data equally well, the more parsimonious model is the one with fewer predictors. 	
 Models that are very complex may "over-fit" the data; providing very good prediction for our current sample but may not be 	
 generalizable to other samples (lacks external validity). Models that are very complex are difficult to interpret. 	
10	
2. Introduction to Predictive Modeling	
Validation	
To ensure the model isn't over-fit to the data, development of a	
prediction model is usually done in two steps	
Develop a model with a training data set	
2. Validate the model with a testing data set	
 The training and testing sets should be independent (i.e., don't validate on the training data) 	-
The validation component will provide evidence for external validity.	
If you split a larger dataset into two sets for this purpose, generally 70% (see to 200%) of the data is used for this purpose,	
70% (up to 80%) of the data is used for training.	
11	
2. Introduction to Predictive Modeling	
<u> </u>	
Recap	
 Prediction models develop the best prediction of outcome and are not concerned with any particular association of interest 	
 Good prediction models are as simple as possible while having predictive ability 	
We will additionally need to validate a prediction model against an	
independent data set	

2. Introduction to Predictive Modeling	
Recap	
> Explain the differences in the approach for prediction modeling vs.	
models of association	
13	
	•
3. Steps in Predictive Model Building	
Overview	
Univariate analysis Variable selection for explainments and all	
Variable selection for multivariate model Preliminary main effects model (does each variable retain	
significance?)	
4. Main effects model (check linearity, scale of variables)	
5. Preliminary final model (check for interactions)	
6. Final model (check model fit and adequacy)	
14	
3. Steps in Predictive Model Building	1
Just like in cooking, a recipe is only as good as the ingredients you put into it.	
Prediction models are only as good as the variables you put into it.	
Check your variables and data before beginning regression modeling.	

3. Steps in Predictive Model Building **Example** Schools in rural areas face increased risk of mental health issues. Researchers want to determine whether characteristics of students' friendship networks, in addition to demographic variables, can be used to identify students at risk of suicide attempt. > with(sos_dat, + gmodels::CrossTable(s_att)) Cell Contents

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3. Steps in Predictive Model Building **Example** We will predict suicide attempt using several variables: male – male gender age – student age grade – student grade in school (9-12) odg – out-degree (# of friends student named) dens – density of the ego's friendship network (range from 0 to 1) recip – reciprocity of friendship nominations (range from 0 to 1) tatot – number of trusted adults named by the student (0-7) bullied - student was bullied at school bully - student bullied others at school

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3. Steps in Predictive Model Building

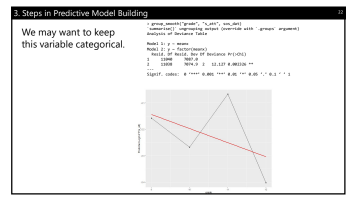
1. Determine the Form of the Outcome

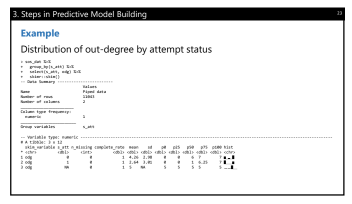
- For a binary outcome, you may use:
 - · Logistic regression
 - Probit regression (not covered in this course)
- For a non-binary outcome, you may use:
 - · Linear regression
 - Poisson regression
 - Others
- For continuous/non-binary outcomes, it helps to evaluate any potential transformations with the entire set of possible X variables.

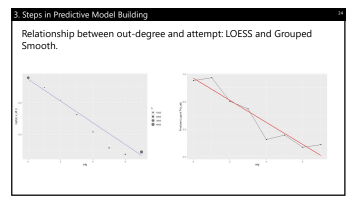
2. Univariate Analyses Categorical Predictors Examine the distribution of X for Y=0 and Y=1 Get univariate odds ratios Pay attention to empty (zero) cells – we will need to deal with these somehow Continuous Predictors: Examine the distribution of X for Y=0 and Y=1 Get univariate odds ratios The x² test from the contingency table is asymptotically equivalent to the LR x² test from the logistic regression model. The 2²-sample t-test is is asymptotically equivalent to the x² tests from the logistic regression model. The 2-sample t-test is is asymptotically equivalent to the x² tests from the logistic regression model. Assess the linearity assumption (grouped smooth, LOESS, fractional polynomials)

3. Steps in Predictive Model Buil	ding					
F	> with(sos_dat	;, gmodels::0	CrossTable(g	rade, s_att, p	prop.chisq=F, prop.t=H	, chisq=T))
Example	Cell Conter	nts				
		N	1			
Relationship between	N .	/ Row Total	į			
grade and attempt:	N .	/ Col Total	-			
		s_att				
contingency table.	grade	0	1	Row Total		
	9	2475	286	2761		
		0.896 0.249	0.104	0.250		
		8.249	0.264			
	10	2502	259	2761		
		0.906 0.251	0.094	0.250		
				i		
	11	2450 0.888	310 0.112	2760 0.250		
		0.888	0.112	0.250		
	12	2531 0.917	0.083	2760 0.250		
		0.254	0.003	0.230		
	Column Total	9958	1084			
	Column local	0.902	0.098	11042		
1	Pearson's Chi-	squared test				
1						
I	Chi^2 = 14.9	5095 d.f.	. = 3 p	= 0.00185904	19	

	> summary(univ_grade.m)	
Example	<pre>Call: glm(formula = s_att ~ factor(grade), family = binomial, data = sos_dat)</pre>	
Relationship between grade and attempt:	Deviance Residuals: Min 10 Median 30 Max -0.4881 -0.4677 -0.4439 -0.4162 2.2313	
ogistic regression.	Coefficients: Estimate Std. Error z value Pr(> z) (Intercept) -2.15800 0.06245 -34.553 < 2e-16 *** factor(grade)10 -0.1801 0.09884 -1.218 0.22331	
	factor(grade)11 0.09073 0.08680 1.045 0.29589 factor(grade)12 -0.24464 0.09307 -2.629 0.00858 ** Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1	
	<pre>> anova(univ_grade.m, test="LRT") Analysis of Deviance Table</pre>	
	Model: binomial, link: logit	
	Response: s_att Terms added sequentially (first to last)	
	Df Deviance Resid. Df Resid. Dev Pr(>Chi) NULL 11041 7690.0	
	factor(grade) 3 15.055 11038 7074.9 0.00177 ** Signif. codes: 0 (***) 0.001 (**) 0.01 (*) 0.05 (.) 0.1 (.) 1	







Steps in Predictive Model Building	25
Relationship between out-degree and attempt: MFP.	
> mfp(_att - fp(odg), family = binomial, data = sos_dat) (all: sfp(formula = s_att - fp(odg), data = sos_dat, family = binomial)	
Deviance table: Resid. Dev Null model 7089.952 Linear model 6812.877 Final model 6812.877	
fractional polynomials: ff.initial select alpha of.final power2 power2 odg 4 1 0.05 1 1 .	
Transformations of covariates: formula odg I(((ods:1)409'1)	
Re-Scaling: Non-positive values in some of the covariates. No re-scaling was performed.	
Coefficients: Intercept odg.1 -1.438 -1.754	
Degrees of Freedom: 11041 Total (i.e. Null); 11040 Residual Null Deviance: 7890 Alc: 6817	

3. Steps in Predictive Model Building

3. Variable Selection

After all variables have been examined univariately, include

- All variables with "clinical importance"
- If this produces too many variables, include all with a univariate p<.25
- Constrain the total number of variables to follow the sample size rule of thumb, either by redefining the definition of "clinically important" or by choosing a lower p-value threshold for inclusion.

We use a less strict p-value to include variables as they may become significant later (in combination with others).

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3. Steps in Predictive Model Building

4. Preliminary Modeling

Use purposeful model building to examine several different models.

- Be careful of automated selection procedures (stepwise regression)
- Forward selection produces more "noise" variables
- Some good models cannot be found with automated selection
- $\bullet\hspace{0.4cm}$ In general automated procedures do not do well with correlated predictors
- Automated selection discourages thinking about the actual problem
- Automated procedures can be helpful for hypothesis-generating analyses (after the primary analysis is done)

3. Steps in Predictive Model Building

4. Preliminary Modeling

Be skeptical of p-values.

- P-values are only valid for testing pre-specified hypotheses
- Since we are screening variables, p-values only indicate relative importance among all variables included
- The larger candidate pool of variables, the more variables will appear significant when they in fact aren't related to outcome

Scrutinize the models for biological plausibility.

Choose the "best" multivariate model

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3. Steps in Predictive Model Building

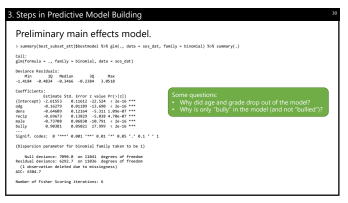
Verify Each Variable

Verify, delete, refit, etc. until you're satisfied that:

- All important variables are included in the model
- All excluded variables are clinically or statistically unimportant

Then you will have the **preliminary main effects model**.

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3. Steps in Predictive Model Building

5. Refine your model

Re-assess linearity for continuous variables.

- This assumption is usually not critical in the variable selection stages; the model-building process is quite forgiving for modest violations of linearity (except for U-shaped relationships)
- Scatterplots (e.g., LOESS) are not easily extended to multivariable models
- Grouped-smooth and fractional polynomials are good approaches to assess linearity in multivariable models

Make sure your model makes sense clinically and scientifically.

You now have the main effects model.

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3. Steps in Predictive Model Building

6. Check for interactions

List all pairs of variables that have scientific plausibility for interaction and add them to the model.

- May need to discuss "plausibility" with your co-investigators with content expertise
- Add one at a time to the main effects model
- Include interactions significant at p<.05 (we're stricter here because nonsignificant interactions tend to inflate the standard errors of beta coefficients).

You now have the preliminary final model.

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3. Steps in Predictive Model Building

7. Check Model Fit

Fit Statistics

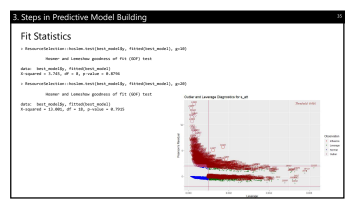
- · Pearson's GOF
- · Hosmer-Lemeshow

Model Diagnostics

- · Closely examine influential points
- Do NOT exclude influential points simply to get better fit
- Consult with investigators and content experts to see if there is a reason why points might be excluded

You now have the final model.

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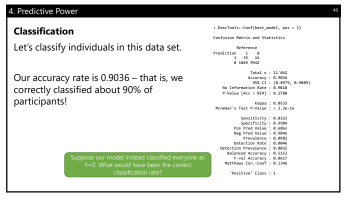
35

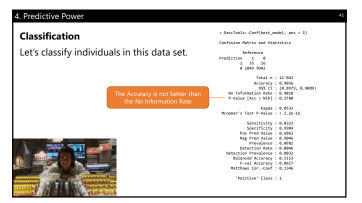
3. Steps in Predictive Model Building

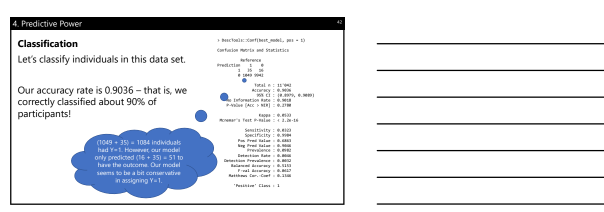
Recap

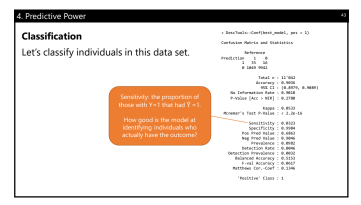
- Model building requires statistical knowledge but is also part art; think of yourself as crafting a model and getting to know the variables along the way
- Good model-building is driven by theory as well; check to make sure your results make sense along the way, and there is theoretical justification for how you handle the variables
- There may be a lot of trial-and-error and refinement in this process

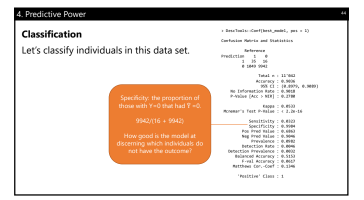
Implement the methods in this section to build a prediction model	3. Steps in Predictive Model Building	
4. Predictive Power From Probability to Outcome	Recap	
4. Predictive Power From Probability to Outcome	> Implement the methods in this section to build a prediction model	
4. Predictive Power From Probability to Outcome		
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From Probability to Outcome ————————————————————————————————————	31	
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From Probability to Outcome		
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From Probability to Outcome ————————————————————————————————————		
From Probability to Outcome Recall that logistic regression provides logit values, which are converted to $\hat{\pi}$, the estimated probability that Y=1.	4. Predictive Power	58
Recall that logistic regression provides logit values, which are converted to $\hat{\pi}$, the estimated probability that Y=1.	From Probability to Outcome	
to $\hat{\pi}$, the estimated probability that Y=1.	Recall that logistic regression provides logit values, which are converted	
	to $\hat{\pi}$, the estimated probability that Y=1.	
38	38	
4. Predictive Power	4 Productive Power	19
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Generally, we classify people into predicted outcome status by:		
i) Fitting the logistic model		
ii) Obtaining the predicted probabilities for each subject ($\widehat{\pi}$)		
iii) Choosing a cutpoint c (usually c=0.5)		
iv) Classifying individuals into an estimated outcome based on their predicted probability and c, such that:	predicted probability and c, such that:	
If $\widehat{\pi}_i > c$, $\widehat{Y}_i = 1$		
If $\widehat{\pi}_i < c$, $\widehat{Y}_i = 0$		
		-
39	39	

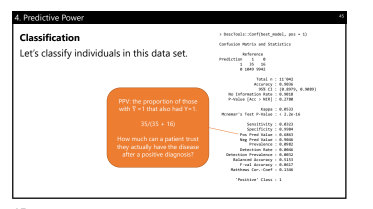












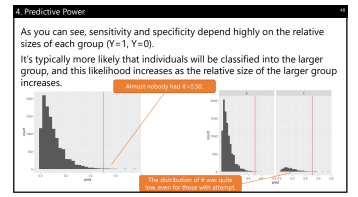
4. Predictive Power	46
Classification	<pre>> DescTools::Conf(best_model, pos = 1) Confusion Matrix and Statistics</pre>
Let's classify individuals in this data set.	Reference Prediction 1 0 1 35 16 0 1049 9942
NPV: the proportion of those with \hat{Y} = 0 that also had Y=0. 9942/(1049+9942) How much can a patient trust they don't have the disease after receiving a negative diagnosis?	Total n: 11'042 Accuracy: 0.0826 Accuracy: 0.0826 No Information & Let (0.8989 0.9089) No Information & Let (0.8989 0.9089) Pollume (Rec > NIR]: 0.2780 Expan: 0.0533 Penemar's Test Publum: 0.2893 Sentitivity: 0.8934 Pos Pred Value: 0.0866 Prevalence: 0.0866 Prevalence: 0.0866 Detection Prevalence: 0.0802 Balanced Accuracy: 0.3539 Balanced Accuracy: 0.3136 Protitive Class: 1

4. Predictive Power

As you can see, sensitivity and specificity depend highly on the relative sizes of each group (Y=1, Y=0).

It's typically more likely that individuals will be classified into the larger group, and this likelihood increases as the relative size of the larger group increases.

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4. Predictive Power	
Classification also depends on how similar individuals in the population are (with respect to $\hat{\pi}_l$):	
1) In a homogenous population, many individuals will have $\hat{\pi}_i$ close to the classification threshold.	
This may be problematic as observations with similar $\hat{\pi}_i$ are forced into discrete outcome categories.	
E.g., A subject with $\hat{\pi}_i$ =0.495 will be classified as no-outcome, and a subject with $\hat{\pi}_i$ =0.505 will be classified as having outcome.	
2) In a polarized population, the $\hat{\pi}_l$ are distributed at the extremes.	
49	
4. Predictive Power 50	
In both cases, we shouldn't expect perfect fit! • If most individuals have $\hat{\pi}_i$ close to 0.5, then we should expect about 50%	
misclassification. • If most individuals have $\hat{\pi}_i$ close to 0.05 or 0.95, then we should expect about 5%	
 misclassification. Classification measures (e.g. sensitivity, specificity) depend on the distribution of π̂_i in the sample and, therefore, are not absolute measures of goodness of 	
classification.	
50	
4. Predictive Power	
We can also change the cutpoint to make it easier or harder to classify $Y=1$.	
This should be done depending on the research question.	
	-

4. Predictive Power 52	
Example	
We want to use this diagnostic tool in order to identify students at school	
who may be particularly at-risk for suicide attempt.	
We will use the tool to discreetly contact students and refer them to resources at school that can help them.	
How would we change the cut point in this case?	
·	_
52	
4. Predictive Power 53	
Example	
We would lower it. • When you raise the cutpoint:	
sensitivity decreases – $P(\hat{Y} = 1 Y = 1)$	
specificity increases – $P(\hat{Y} = 0 Y = 0)$	-
 When you lower the cutpoint: sensitivity increases – P(Ŷ = 1 Y = 1) 	
specificity decreases – $P(\hat{Y} = 0 Y = 0)$	
Therefore we would be able to detect more students with attempt, at the cost of classifying some without attempt as having it.	
, g	
53	
4. Predictive Power 54	•
Sensitivity vs. Specificity – Advantages	
Sensitive models are helpful to identify those who actually have the	
disease, even at the cost of misdiagnosing some individuals without the disease.	
Screening tests with the opportunity for further follow-up	
Examples: mammograms, HIV screening, airport security	
Specific models should be used when we want to verify that an	
individual does not have the disease, even at the cost of misdiagnosing some individuals who actually have the disease.	
 Useful after preliminary screening when being diagnosed "positive" has large 	
risk of physical, emotional, or monetary harm (e.g., biopsies).	

4. Predictive Power

Predictive Value

- Sensitivity and specificity are more useful for clinicians & researchers, when considering diagnosing individuals at the population-level.
- PPV and NPV are more useful to the patient

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4. Predictive Power Changing the Cutpoint Here we change the cutpoint of $\hat{\pi}$ for classifying $\hat{Y}=1$ vs. $\hat{Y}=0$. We graph the accuracy over all possible cutpoints. Note that the accuracy doesn't change much, likely because of the small number of individuals classified with Y=1.

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4. Predictive Power

Recap

- Sensitivity, specificity, positive predictive value, and negative predictive value are all ways of evaluating the predictive power of a model.
- The cutpoint for classifying Y=1 can be changed, and will alter the specificity and sensitivity depending on the goal of the prediction model
- These metrics are highly confounded with the proportion of individuals in the sample with Y=0, Y=1.

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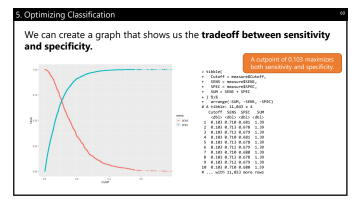
- > Explain the concepts of sensitivity, specificity, positive predictive value, and negative value
- ${\pmb\succ}$ Explain how these measures are affected by the overall proportion of individuals with the outcome

5. Optimizing Classification

We can see that the cutpoint we choose may depend on the nature of the diagnostic tool you'd like to create.

Is there another way to optimize how individuals are classified?

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ROC (receiver operating characteristic) curves are another tool used to optimize classification, and provide a more complete assessment of classification accuracy.

The area under these curves is indicative of how good a model fits.

In world war 2, radar operators had to interpret blips on radar screens as either friendly, hostile, or noise. The blip was the "signal", and an ROC curve was one way of "signal detection," or a set of ways to measure/study how, and to what extent, the receivers could make sense of the signal.

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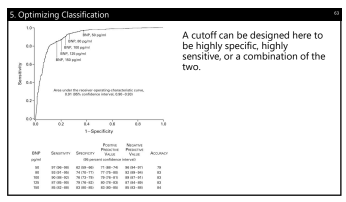
5. Optimizing Classification

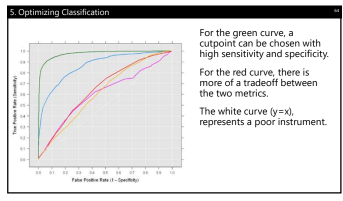
Example

BNP is a cardiac peptide secreted in the heart in response to volume expansion. Therefore, BNP levels can be used to detect cardiac problems. Furthermore, patients presenting with dyspnea (difficult breathing) may be experiencing this symptom due to congestive heart failure

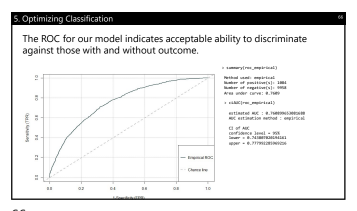
BNP is a potential diagnostic tool for congestive heart failure. (Florkowski 2008)

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5. Optimizing Classification						
General Guideline for an instrument's discriminative ability.						
	AUDOC (A. III I. BOC)	ci :c. :	ı			
	AUROC (Area Under ROC)	Classification				
	0.5	Useless (essentially a coin flip)				
	0.5-0.7	Poor				
	0.7-0.8	Acceptable-Good				
	0.8-0.9	Excellent				
	0.9-1.0	Nearly perfect				



5. Optimizing Classification	67
We can also examine optimal cutpoint	s using differing criteria:
) optimal.cutpoints("-"core", strue ""class") ata east rame(cods locate) esthods = ("Youden", "Rasspie", "RassProdSpie"), ta call: outpoints.default(" "score", strue ""class", tag.healty = orentos < ("Youden", "Arasspie"), data data ("tag.healty = orentos < ("Youden", "Arasspie"), "aras="orentos", tag.healty = orentos < ("Youden", "Arasspie"), "aras="orentos", data data."	e,
Optimal cutoffs: Youdon MasSide MakProdSpds 1 0-4897 0-4898 0-4895 Area under the ROC curve (AUC): 0.761 (0.747, 0.775	Youden = max(Sp + Se - 1) MaxSpSe = max(min(Sp, Se)) MaxProdSpSe = max(Sp*Se)

5. Optimizing Classification

Recap

- Discrimination is another tool for assessing prediction models, in addition to correct classification rates.
- Models that best discriminate between those with Y=0 and Y=1 maximize both sensitivity and specificity.
- \bullet ROC curves show the tradeoff between sensitivity and specificity for different cutpoints.
- Higher area under the ROC curve (AUROC) indicates a model with better discrimination.

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5. Optimizing Classification

Recap

> Use AUC as a metric to explain a model's discriminant ability

6. Recap	
Additional Reading	
More on the Youden Index:	
https://onlinelibrary.wiley.com/doi/pdf/10.1002/bimj.200410135	
nteps,//orininensiary.wiley.com/ doi/ pai/ 10.1002/ birrig.200 110 100	
70	
6. Recap	7
o. Recap	
Packages and Functions	
DescTools::Conf()	
ROCit::measureit()ROCit::rocit()	-
• ROCIE: rock()	
OptimalCutpoints::optimal.cutpoints()	
• plotROC	