Biostatistics

Applications in Medicine

Syllabus

1. General review

- a. What is Biostatistics?
- b. Population/Sample/Sample size
- c. Type of Data quantitative and qualitative variables
- d. Common probability distributions
- e. Work example Malaria in Tanzania

2. Applications in Medicine

- a. Construction and analysis of diagnostic tools Binomial distribution, sensitivity, specificity, ROC curve, Rogal-Gladen estimator
- b. Estimation of treatment effects generalized linear models
- c. Survival analysis Kaplan-Meier curve, log-rank test, Cox's proportional hazards model

3. Applications in Genetics, Genomics, and other 'omics data

- a. Genetic association studies Hardy-Weinberg test, homozygosity, minor allele frequencies, additive model, multiple testing correction
- b. Methylation association studies M versus beta values, estimation of biological age
- c. Gene expression studies based on RNA-seq experiments Tests based on Poisson and Negative-Binomial

4. Other Topics

- a. Estimation of Species diversity Diversity indexes, Poisson mixture models
- b. Serological analysis Gaussian (skew-normal) mixture models
- c. Advanced sample size and power calculations

What is the test perfect for diagnosis?

What is the test perfect for diagnosis?

Positive for all true cases

P(+|true case)=1

Sensitivity=1

Negative for all true non-cases

P(-|true non case)=1

Specificity=1

(Almost) Perfect diagnostic = Gold Standard

Hypothetical data

Disease status	Positive	Negative
True Cases	n_1	0
True Non-cases	0	n_2

Some mathematics (sampling model)

True cases

$$x_1 \mid n_1, \pi_{Se} \rightsquigarrow \text{Binomial}(n_1; \pi_{Se})$$

True non-cases

$$x_2 \mid n_2, \pi_{Sp} \rightsquigarrow \text{Binomial}\left(n_2; \pi_{Sp}\right)$$

What are the assumptions?

Random versus deterministic

$$H_0$$
: Sensitivity < 1 vs H_1 : Sensitivity = 1

$$H_0: \pi_{Se} < 1 \text{ vs } H_1: \pi_{Se} = 1$$

$$H_0$$
: Specificity < 1 vs H_1 : Specificity $= 1$

$$H_0: \pi_{Sp} < 1 \text{ vs } H_1: \pi_{Sp} = 1$$

Technical problem I

$$\pi_{Se} \in (0,1) \text{ and } \pi_{Sp} \in (0,1)$$

Practical Solution

$$H_0: \pi_{Se} = 0.999 \text{ vs } H_1: \pi_{Se} > 0.999$$

$$H_0: \pi_{Sp} = 0.999 \text{ vs } H_1: \pi_{Sp} > 0.999$$

Simple test: calculate confidence intervals and check whether 0.999 is within it.

Should the confidence intervals be one-tail or two-tail?

Random versus deterministic

$$IC_{95\%}(\pi_{Se}) = \hat{\pi}_{Se} \pm 1.96 \times se(\hat{\pi}_{Se})$$

$$IC_{95\%}(\pi_{Se}) = (\hat{\pi}_{Se} - 1.65 \times se(\hat{\pi}_{Se}); 1)$$

Technical problem II (standard error undefined)

$$se\left(\hat{\pi}_{Se}\right) = \sqrt{\frac{\pi_{Se}\left(1 - \pi_{Se}\right)}{n}}$$

$$se\left(\hat{\pi}_{Se}\right) = \sqrt{\frac{\hat{\pi}_{Se}\left(1 - \hat{\pi}_{Se}\right)}{n}}$$

Solution: use an alternative confidence interval

Clopper-Pearson exact interval

Exact version

Over-coverage

Wilson score interval

Better approximations to the Normal distributions, especially when the true proportion is close to 0 or 1

Agresti-Coull interval

Arcsin approximation interval

Under-coverage



Disease status	Positive	Negative
True Cases	30	0
True Non-cases	0	40

Calculate one-tailed 95% confidence intervals and test the null hypothesis

Use Wilson score test (prop.test)

Sample size determination

What is the sample size required for us to be confidence we are close to a deterministic situation?



- 1. Use prop.test function
- 2. Calculate (one tailed test) 95% confidence intervals for perfect sensitivity/specificity with different methods
- 3. Determine the minimum sample size required to obtain the one-tailed 95% confidence interval that does not contain 99.9%

You observe x positive tests in a sample of n individuals

The diagnostic test is imperfect

What is the estimate for the proportion of true positive individuals?

$$x \mid n, \pi \rightsquigarrow \text{Binomial}(n; \pi)$$

$$\pi$$
 = probability of a positive test

$$\pi = \pi_{Se}\pi_{+} + (1 - \pi_{Sp})\pi_{-}$$

$$= \pi_{Se}\pi_{+} + (1 - \pi_{Sp})(1 - \pi_{+})$$

$$\pi_{Se} = Sensitivity$$

$$\pi_{Sp} = Specificity$$

 π_{+} = proportion of sampling a true positive

Can you estimate π_{Se} , π_{Sp} , and π_{+} from the data?

$$x \mid n, \pi \rightsquigarrow \text{Binomial}(n; \pi)$$

$$\pi$$
 = probability of a positive test

$$\pi = \pi_{Se}\pi_{+} + (1 - \pi_{Sp})\pi_{-}$$

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$$\pi_{Se} = Sensitivity$$

$$\pi_{Sp} = Specificity$$

 π_{+} = proportion of sampling a true positive

Can you estimate π_{Se} , π_{Sp} , and π_{+} from the data?

No, overparametrization and lack of identifiability!

$$x \mid n, \pi \rightsquigarrow \text{Binomial}(n; \pi)$$

$$\pi$$
 = probability of a positive test

$$\pi = \pi_{Se}\pi_{+} + (1 - \pi_{Sp})\pi_{-}$$

$$= \pi_{Se}\pi_{+} + (1 - \pi_{Sp})(1 - \pi_{+})$$

$$\pi_{Se} = Sensitivity$$

$$\pi_{SD}$$
 = Specificity

 π_{+} = proportion of sampling a true positive

Can you estimate π_{Se} , π_{Sp} , and π_{+} from the data?

No, overparametrization and lack of identifiability!

Do you have any solution?

$$x \mid n, \pi \rightsquigarrow \text{Binomial}(n; \pi)$$

$$\pi$$
 = probability of a positive test

$$\pi = \pi_{Se}\pi_{+} + (1 - \pi_{Sp})\pi_{-}$$

$$= \pi_{Se}\pi_{+} + (1 - \pi_{Sp})(1 - \pi_{+})$$

$$\pi_{Se} = Sensitivity$$

$$\pi_{Sp} = Specificity$$

 π_{+} = proportion of sampling a true positive

Can you estimate π_{Se} , π_{Sp} , and π_{+} from the data?

No, overparametrization and lack of identifiability!

Do you have any solution? Use external estimates for π_{Se} and π_{Sp}

$$\hat{\pi}_{+} = \frac{\hat{\pi} + \hat{\pi}_{Sp} - 1}{\hat{\pi}_{Se} + \hat{\pi}_{Sp} - 1}$$

$$\hat{\pi} = \frac{x}{n}$$

Method of moments

$$\hat{\pi} = \hat{\pi}_{Se}\pi_{+} + (1 - \hat{\pi}_{Sp})(1 - \pi_{+})$$

$$\hat{\pi} = (\hat{\pi}_{Se} + \hat{\pi}_{Sp} - 1)\hat{\pi}_{+} + (1 - \hat{\pi}_{Sp})$$

$$\hat{\pi} + \pi_{Sp} - 1 = (\pi_{Se} + \pi_{Sp} - 1)\pi_{+}$$

$$\pi_{+} = \frac{\hat{\pi} + \hat{\pi}_{Sp} - 1}{\hat{\pi}_{Se} + \hat{\pi}_{Sp} - 1}$$

Invariant property of MLE

Wald's Confidence interval

$$IC_{95\%}(\pi_{+}) = \hat{\pi}_{+} \pm 1.96 \times se(\hat{\pi}_{+})$$

$$\hat{\pi} = \frac{x}{n}$$

$$\operatorname{Var}\left[\hat{\pi}_{+} \,|\, \hat{\pi}_{Se}, \hat{\pi}_{Sp}\right] = \operatorname{Var}\left[\frac{\hat{\pi} + \hat{\pi}_{Sp} - 1}{\hat{\pi}_{Se} + \hat{\pi}_{Sp} - 1} \,|\, \hat{\pi}_{Se}, \hat{\pi}_{Sp}\right] = \frac{\operatorname{Var}\left[\hat{\pi}\right]}{\left(\hat{\pi}_{Se} + \hat{\pi}_{Sp} - 1\right)^{2}} = \frac{\pi\left(1 - \pi\right)}{n\left(\hat{\pi}_{Se} + \hat{\pi}_{Sp} - 1\right)^{2}}$$

$$\hat{\text{Var}} \left[\hat{\pi}_{+} | \hat{\pi}_{Se}, \hat{\pi}_{Sp} \right] = \frac{\hat{\pi} (1 - \hat{\pi})}{n \left(\hat{\pi}_{Se} + \hat{\pi}_{Sp} - 1 \right)^{2}} \qquad se(\hat{\pi}_{+}) = \sqrt{\hat{\text{Var}} \left[\hat{\pi}_{+} | \hat{\pi}_{Se}, \hat{\pi}_{Sp} \right]}$$

Limitation and extension

 $\hat{\pi}_{Se}$

are assumed to be known constant

 $\hat{\pi}_{Sp}$

 $\hat{\pi}_{Se}$

are assumed to be estimators

$$\hat{\pi}_{Sp}$$

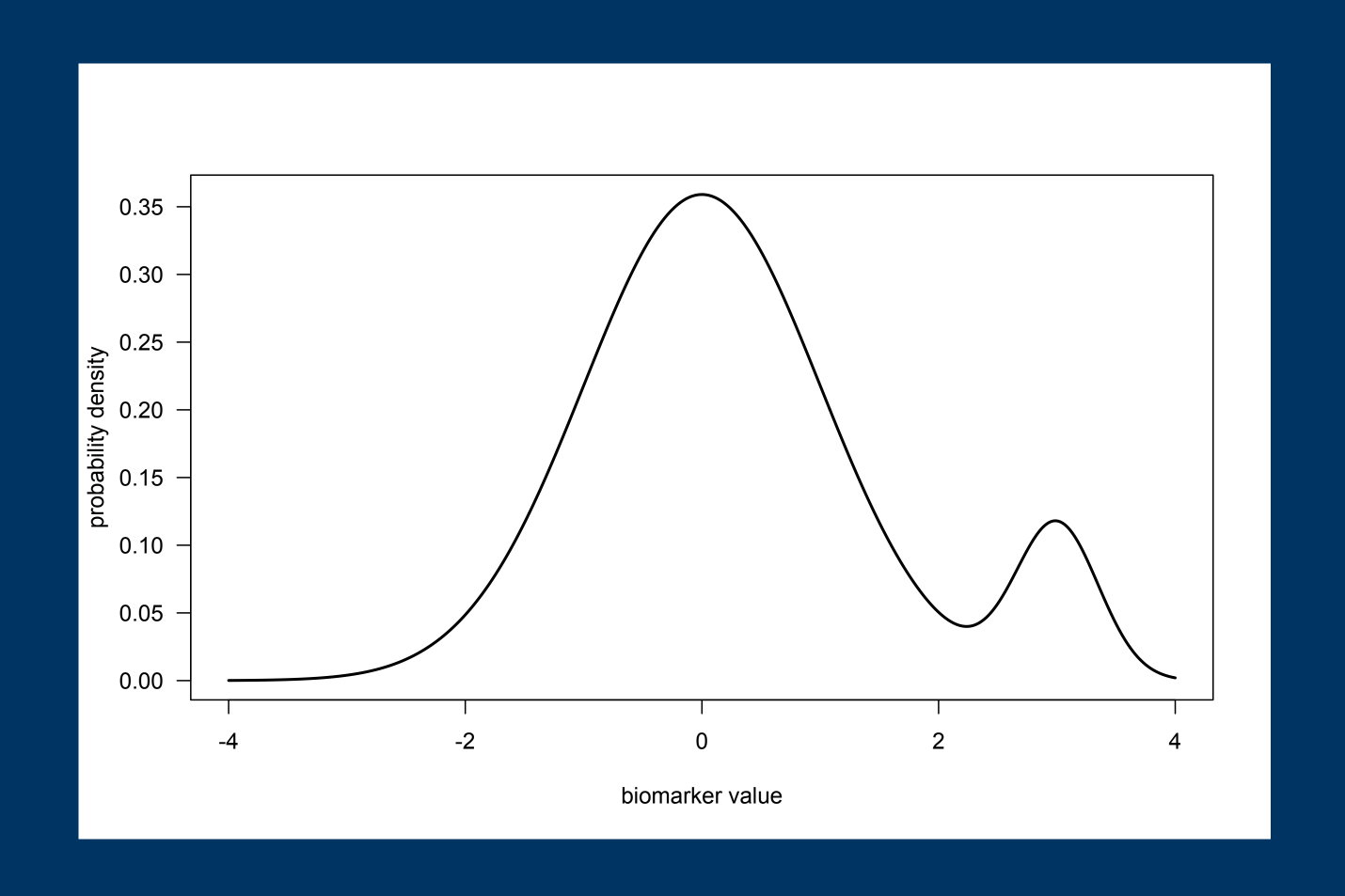
$$\hat{\text{Var}}\left[\hat{\pi}_{+}\right] = E\left[Var\left[\hat{\pi}_{+} | \hat{\pi}_{Se}, \hat{\pi}_{Sp}\right]\right] + \text{Var}\left[E\left[\hat{\pi}_{+} | \hat{\pi}_{Se}, \hat{\pi}_{Sp}\right]\right]$$

Exercise

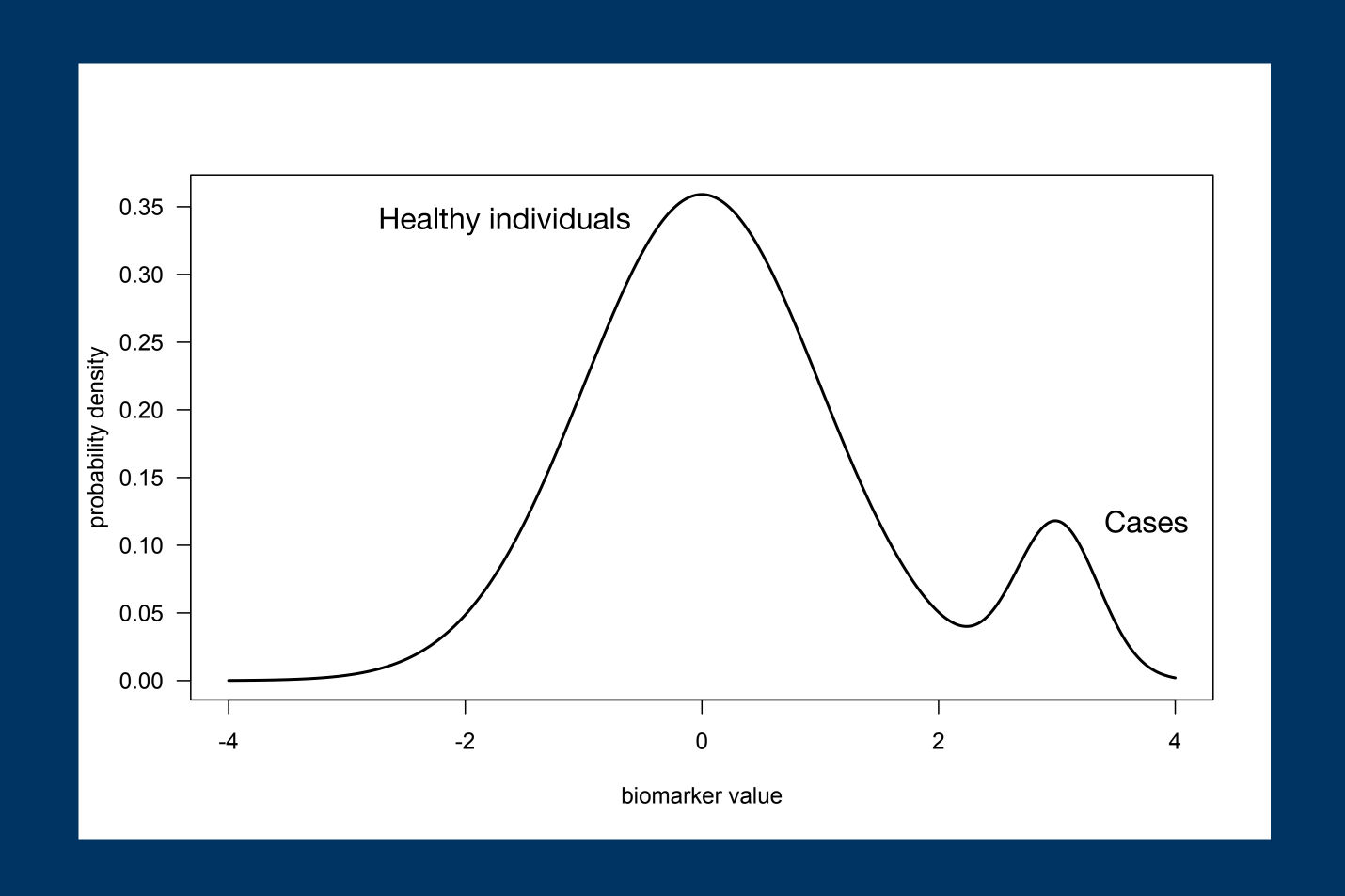
	Positive	n	Sensitivity	Specificity
Anti-S antibodies	1271	2474	0.665 (India)	0.99 (India)
Anti-N antibodies	848	2474	0.995 (manufacturer)	0.997 (manufacturer)

Estimate the proportion of true positive using the Rogan-Gladen estimator and its 95% confidence interval

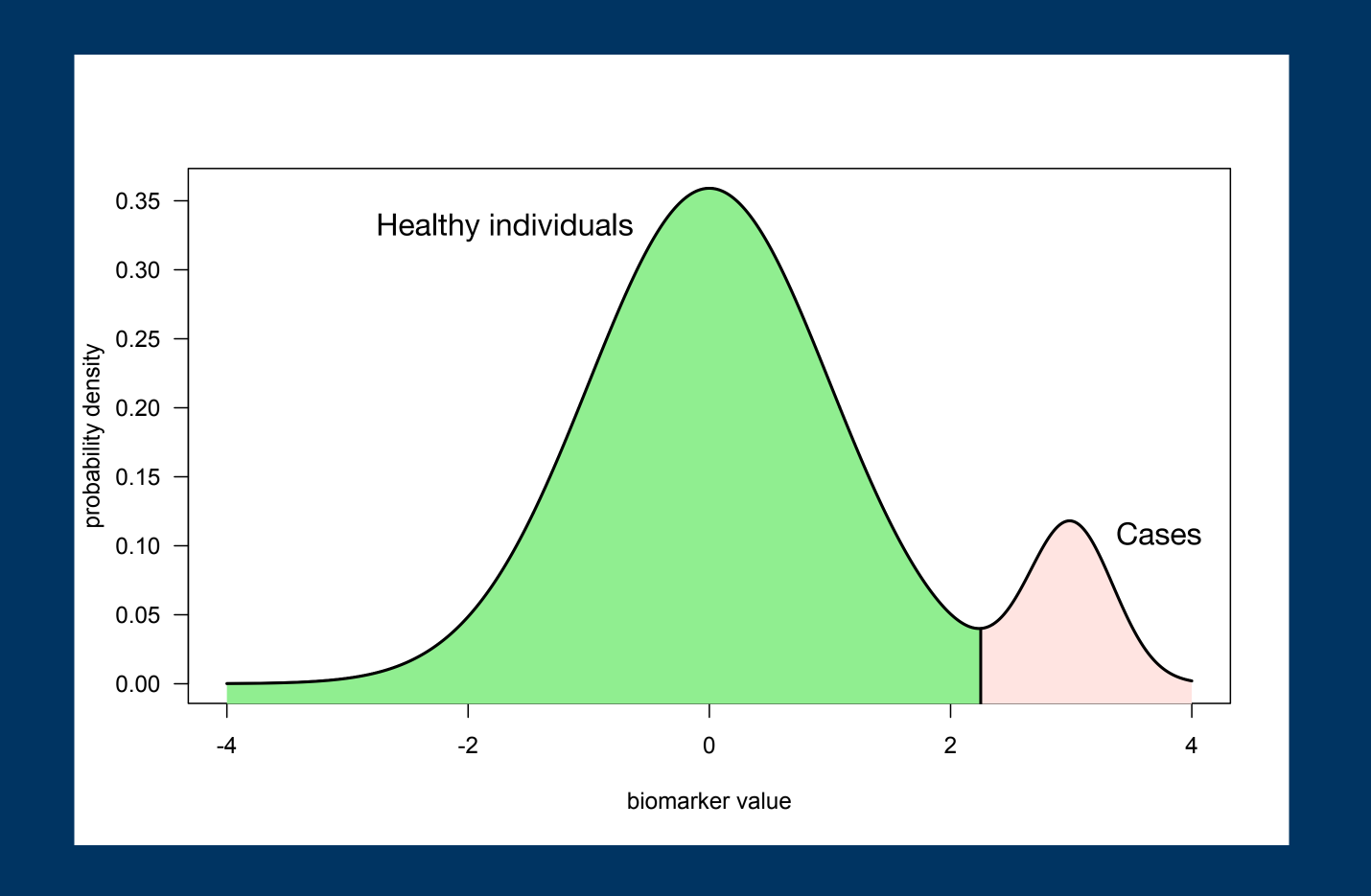
Constructing a diagnostic tool based on quantitative biomarker



Constructing a diagnostic tool based on quantitative biomarker



Constructing a diagnostic tool based on quantitative biomarker



How to estimate the optimal cutoff to distinguish cases from healthy controls

Evaluating the diagnostic potential of a biomarket

Two samples

True cases True healthy

Estimate sensitivity and specificity using all possible cutoffs for the biomarker

Evaluating the diagnostic potential of a biomarket

Two samples

True cases True healthy

Estimate sensitivity and specificity using all possible cutoffs for the biomarker

$$x_1 \mid n_1, \pi_{Se} \rightsquigarrow \text{Binomial}(n_1; \pi_{Se})$$

$$\pi_{Se}(\kappa) = P\left[X \ge \kappa \mid n_1, \text{True case}\right]$$

$$\hat{\pi}_{Se}(\kappa) = \frac{\text{\# individuals with X} \geq \kappa}{n_1}$$

Evaluating the diagnostic potential of a biomarker

Two samples

True cases

True healthy

Estimate sensitivity and specificity using all possible cutoffs for the biomarker

$$x_1 \mid n_1, \pi_{Se} \rightsquigarrow \text{Binomial}(n_1; \pi_{Se})$$

$$x_2 \mid n_2, \pi_{Sp} \rightsquigarrow \text{Binomial}\left(n_2; \pi_{Sp}\right)$$

$$\pi_{Se}(\kappa) = P\left[X \ge \kappa \mid n_1, \text{True case}\right]$$

$$\pi_{Sp}(\kappa) = P\left[X < \kappa \mid n_2, \text{True healthy}\right]$$

$$\hat{\pi}_{Se}(\kappa) = \frac{\text{\# individuals with X} \ge \kappa}{n_1}$$

$$\hat{\pi}_{Sp}(\kappa) = \frac{\text{\# individuals with X} < \kappa}{n_2}$$

Evaluating the diagnostic potential of a biomarket

Two samples

True cases True healthy

Estimate sensitivity and specificity using all possible cutoffs for the biomarker

Construct a plot with the respective estimated sensitivities and 1-specificities

Receiver Operating Characteristic (ROC) curve

Real world data



Journal of Psychosomatic Research

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Measuring fatigue in clinical and community settings

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Abstract

Objective: The Chalder Fatigue Scale (CFQ) is a widely used instrument to assess fatigue in both clinical and nonclinical settings. Psychometric properties of the scale and discriminative abilities were examined. Methods: A total of 361 patients with CFS and 1615 individuals in the community were assessed with the CFQ. Principal component analysis (PCA) was used to explore the structure of the scale. Receiver-operating characteristic curve (ROC) was used to investigate the discriminative properties.

Results: Two components, physical and mental fatigue, were identified in the CFS patient group and in the general population samples. Area under the curve for ROC was .91. The fatigue scale effectively discriminates, at high scores, between CFS patients and the general population. Conclusion: Physical and mental fatigue are clearly separable components of fatigue. The CFQ can discriminate reliably between clinical and nonclinical conditions. © 2010 Elsevier Inc. All rights reserved.

Keywords: CFS; Chalder fatigue scale; Chronic fatigue syndrome; Fatigue; ROC

Table 3								
Sensitivity	and	1-	specificity	for	the	fatigue	scales	scores

0 1.000 1.000 1 1.000 .999 2 .997 .998 3 .997 .993 4 .997 .993 5 .994 .985 6 .989 .979 7 .989 .940 8 .989 .940 9 .989 .913 10 .978 .866 11 .966 .637 12 .966 .565 13 .958 .484 14 .936 .408 15 .922 .330 16 .899 .260 17 .888 .207 18 .854 .167 19 .824 .136 20 .773 .107 21 .728 .084 22 .658 .062 23 .605 .043 24 .546 .029 25 .473 .018 26 .4	CFQ Score	Sensitivity	1-Specificity
2 .997 .997 3 .997 .993 4 .997 .993 5 .994 .985 6 .989 .979 7 .989 .964 8 .989 .940 9 .989 .913 10 .978 .866 11 .966 .637 12 .966 .565 13 .958 .484 14 .936 .408 15 .922 .330 16 .899 .260 17 .888 .207 18 .854 .167 19 .824 .136 20 .773 .107 21 .728 .084 22 .658 .062 23 .605 .043 24 .546 .029 25 .473 .018 26 .415 .014 27 .339 .010 28 .28	0	1.000	1.000
3 .997 .993 4 .997 .993 5 .994 .985 6 .989 .979 7 .989 .964 8 .989 .940 9 .989 .913 10 .978 .866 11 .966 .637 12 .966 .565 13 .958 .484 14 .936 .408 15 .922 .330 16 .899 .260 17 .888 .207 18 .854 .167 19 .824 .136 20 .773 .107 21 .728 .084 22 .658 .062 23 .605 .043 24 .546 .029 25 .473 .018 26 .415 .014 27 .339 .010 28 .286 .007 29 .2	1	1.000	.999
4 .997 .993 5 .994 .985 6 .989 .979 7 .989 .964 8 .989 .940 9 .989 .913 10 .978 .866 11 .966 .637 12 .966 .565 13 .958 .484 14 .936 .408 15 .922 .330 16 .899 .260 17 .888 .207 18 .854 .167 19 .824 .136 20 .773 .107 21 .728 .084 22 .658 .062 23 .605 .043 24 .546 .029 25 .473 .018 26 .415 .014 27 .339 .010 28 .286 .007 29 .202 .004 30 .	2	.997	.998
5 .994 .985 6 .989 .979 7 .989 .964 8 .989 .940 9 .989 .913 10 .978 .866 11 .966 .637 12 .966 .565 13 .958 .484 14 .936 .408 15 .922 .330 16 .899 .260 17 .888 .207 18 .854 .167 19 .824 .136 20 .773 .107 21 .728 .084 22 .658 .062 23 .605 .043 24 .546 .029 25 .473 .018 26 .415 .014 27 .339 .010 28 .286 .007 29 .202 .004 30 .151 .000 31	3	.997	.997
6 .989 .979 7 .989 .964 8 .989 .940 9 .989 .913 10 .978 .866 11 .966 .637 12 .966 .565 13 .958 .484 14 .936 .408 15 .922 .330 16 .899 .260 17 .888 .207 18 .854 .167 19 .824 .136 20 .773 .107 21 .728 .084 22 .658 .062 23 .605 .043 24 .546 .029 25 .473 .018 26 .415 .014 27 .339 .010 28 .286 .007 29 .202 .004 30 .151 .000 31 .095 .000 32 <td< td=""><td>4</td><td>.997</td><td>.993</td></td<>	4	.997	.993
7 .989 .940 8 .989 .940 9 .989 .913 10 .978 .866 11 .966 .637 12 .966 .565 13 .958 .484 14 .936 .408 15 .922 .330 16 .899 .260 17 .888 .207 18 .854 .167 19 .824 .136 20 .773 .107 21 .728 .084 22 .658 .062 23 .605 .043 24 .546 .029 25 .473 .018 26 .415 .014 27 .339 .010 28 .286 .007 29 .202 .004 30 .151 .000 31 .095 .000 32 .062 .000	5	.994	.985
8 .989 .940 9 .989 .913 10 .978 .866 11 .966 .637 12 .966 .565 13 .958 .484 14 .936 .408 15 .922 .330 16 .899 .260 17 .888 .207 18 .854 .167 19 .824 .136 20 .773 .107 21 .728 .084 22 .658 .062 23 .605 .043 24 .546 .029 25 .473 .018 26 .415 .014 27 .339 .010 28 .286 .007 29 .202 .004 30 .151 .000 31 .095 .000 32 .062 .000	6	.989	.979
9 .989 .913 10 .978 .866 11 .966 .637 12 .966 .565 13 .958 .484 14 .936 .408 15 .922 .330 16 .899 .260 17 .888 .207 18 .854 .167 19 .824 .136 20 .773 .107 21 .728 .084 22 .658 .062 23 .605 .043 24 .546 .029 25 .473 .018 26 .415 .014 27 .339 .010 28 .286 .007 29 .202 .004 30 .151 .000 31 .095 .000 32 .062 .000	7	.989	.964
10 .978 .866 11 .966 .637 12 .966 .565 13 .958 .484 14 .936 .408 15 .922 .330 16 .899 .260 17 .888 .207 18 .854 .167 19 .824 .136 20 .773 .107 21 .728 .084 22 .658 .062 23 .605 .043 24 .546 .029 25 .473 .018 26 .415 .014 27 .339 .010 28 .286 .007 29 .202 .004 30 .151 .000 31 .095 .000 32 .062 .000	8	.989	.940
11 .966 .565 12 .966 .565 13 .958 .484 14 .936 .408 15 .922 .330 16 .899 .260 17 .888 .207 18 .854 .167 19 .824 .136 20 .773 .107 21 .728 .084 22 .658 .062 23 .605 .043 24 .546 .029 25 .473 .018 26 .415 .014 27 .339 .010 28 .286 .007 29 .202 .004 30 .151 .000 31 .095 .000 32 .062 .000	9	.989	.913
12 .966 .565 13 .958 .484 14 .936 .408 15 .922 .330 16 .899 .260 17 .888 .207 18 .854 .167 19 .824 .136 20 .773 .107 21 .728 .084 22 .658 .062 23 .605 .043 24 .546 .029 25 .473 .018 26 .415 .014 27 .339 .010 28 .286 .007 29 .202 .004 30 .151 .000 31 .095 .000 32 .062 .000	10	.978	.866
13 .958 .484 14 .936 .408 15 .922 .330 16 .899 .260 17 .888 .207 18 .854 .167 19 .824 .136 20 .773 .107 21 .728 .084 22 .658 .062 23 .605 .043 24 .546 .029 25 .473 .018 26 .415 .014 27 .339 .010 28 .286 .007 29 .202 .004 30 .151 .000 31 .095 .000 32 .062 .000	11	.966	.637
14 .936 .408 15 .922 .330 16 .899 .260 17 .888 .207 18 .854 .167 19 .824 .136 20 .773 .107 21 .728 .084 22 .658 .062 23 .605 .043 24 .546 .029 25 .473 .018 26 .415 .014 27 .339 .010 28 .286 .007 29 .202 .004 30 .151 .000 31 .095 .000 32 .062 .000	12	.966	.565
15 .922 .330 16 .899 .260 17 .888 .207 18 .854 .167 19 .824 .136 20 .773 .107 21 .728 .084 22 .658 .062 23 .605 .043 24 .546 .029 25 .473 .018 26 .415 .014 27 .339 .010 28 .286 .007 29 .202 .004 30 .151 .000 31 .095 .000 32 .062 .000	13	.958	.484
16 .899 .260 17 .888 .207 18 .854 .167 19 .824 .136 20 .773 .107 21 .728 .084 22 .658 .062 23 .605 .043 24 .546 .029 25 .473 .018 26 .415 .014 27 .339 .010 28 .286 .007 29 .202 .004 30 .151 .000 31 .095 .000 32 .062 .000	14	.936	.408
17 .8888 .207 18 .854 .167 19 .824 .136 20 .773 .107 21 .728 .084 22 .658 .062 23 .605 .043 24 .546 .029 25 .473 .018 26 .415 .014 27 .339 .010 28 .286 .007 29 .202 .004 30 .151 .000 31 .095 .000 32 .062 .000	15	.922	.330
18 .854 .167 19 .824 .136 20 .773 .107 21 .728 .084 22 .658 .062 23 .605 .043 24 .546 .029 25 .473 .018 26 .415 .014 27 .339 .010 28 .286 .007 29 .202 .004 30 .151 .000 31 .095 .000 32 .062 .000	16	.899	.260
19 .824 .136 20 .773 .107 21 .728 .084 22 .658 .062 23 .605 .043 24 .546 .029 25 .473 .018 26 .415 .014 27 .339 .010 28 .286 .007 29 .202 .004 30 .151 .000 31 .095 .000 32 .062 .000	17	.888	.207
20 .773 .107 21 .728 .084 22 .658 .062 23 .605 .043 24 .546 .029 25 .473 .018 26 .415 .014 27 .339 .010 28 .286 .007 29 .202 .004 30 .151 .000 31 .095 .000 32 .062 .000	18	.854	.167
21 .728 .084 22 .658 .062 23 .605 .043 24 .546 .029 25 .473 .018 26 .415 .014 27 .339 .010 28 .286 .007 29 .202 .004 30 .151 .000 31 .095 .000 32 .062 .000	19	.824	.136
22 .658 .062 23 .605 .043 24 .546 .029 25 .473 .018 26 .415 .014 27 .339 .010 28 .286 .007 29 .202 .004 30 .151 .000 31 .095 .000 32 .062 .000	20	.773	.107
23 .605 .043 24 .546 .029 25 .473 .018 26 .415 .014 27 .339 .010 28 .286 .007 29 .202 .004 30 .151 .000 31 .095 .000 32 .062 .000	21	.728	.084
24 .546 .029 25 .473 .018 26 .415 .014 27 .339 .010 28 .286 .007 29 .202 .004 30 .151 .000 31 .095 .000 32 .062 .000	22	.658	.062
25 .473 .018 26 .415 .014 27 .339 .010 28 .286 .007 29 .202 .004 30 .151 .000 31 .095 .000 32 .062 .000	23	.605	.043
26 .415 .014 27 .339 .010 28 .286 .007 29 .202 .004 30 .151 .000 31 .095 .000 32 .062 .000	24	.546	.029
27 .339 .010 28 .286 .007 29 .202 .004 30 .151 .000 31 .095 .000 32 .062 .000	25	.473	.018
28 .286 .007 29 .202 .004 30 .151 .000 31 .095 .000 32 .062 .000	26	.415	.014
29 .202 .004 30 .151 .000 31 .095 .000 32 .062 .000	27	.339	.010
30 .151 .000 31 .095 .000 32 .062 .000	28	.286	.007
31 .095 .000 32 .062 .000	29	.202	.004
32 .062 .000	30	.151	.000
	31	.095	.000
33 .000 .000	32	.062	.000
	33	.000	.000

ROC curve



Journal of Psychosomatic Research

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Measuring fatigue in clinical and community settings

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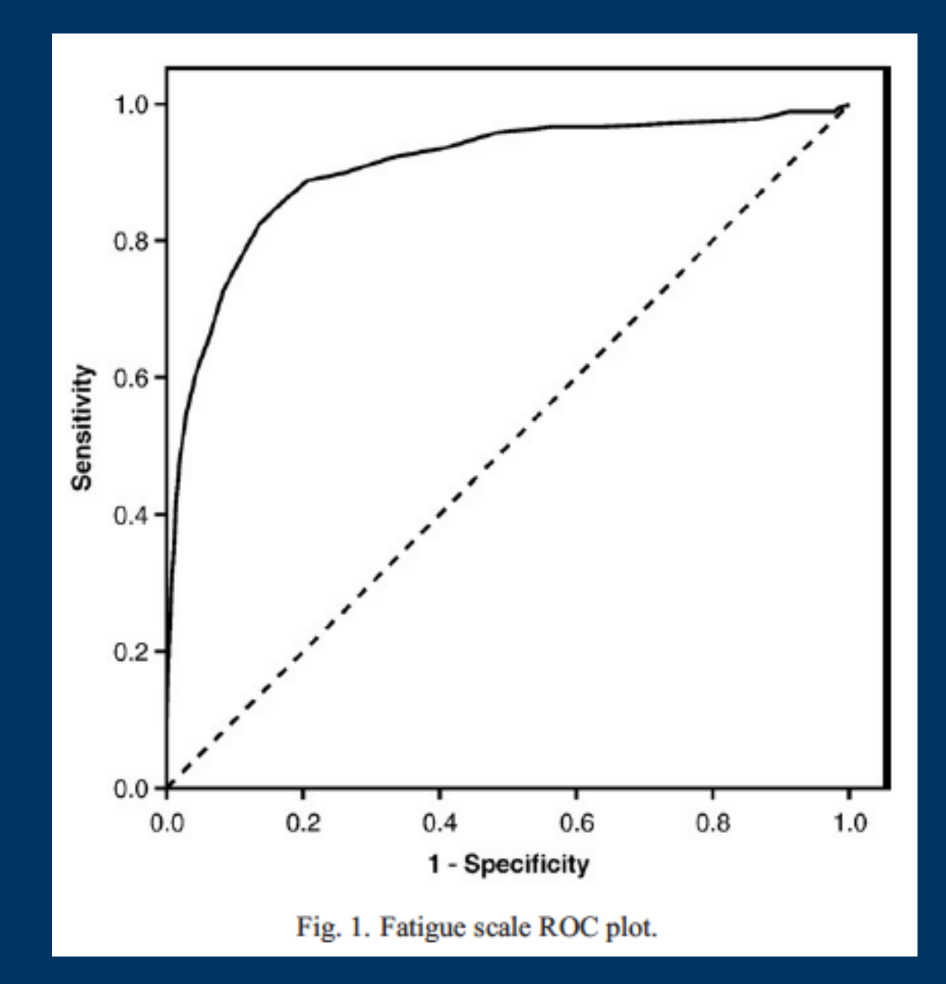
Received 11 June 2009; received in revised form 8 October 2009; accepted 13 October 2009

Abstract

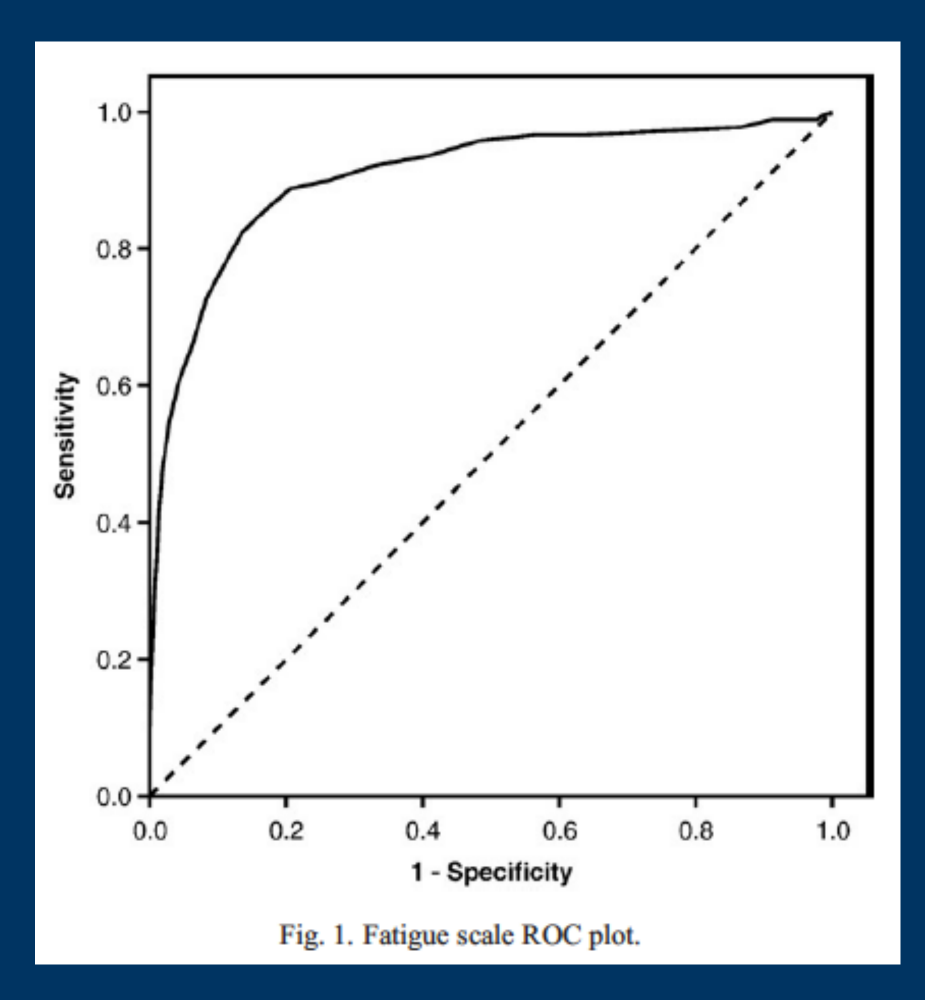
Objective: The Chalder Fatigue Scale (CFQ) is a widely used instrument to assess fatigue in both clinical and nonclinical settings. Psychometric properties of the scale and discriminative abilities were examined. Methods: A total of 361 patients with CFS and 1615 individuals in the community were assessed with the CFQ. Principal component analysis (PCA) was used to explore the structure of the scale. Receiver-operating characteristic curve (ROC) was used to investigate the discriminative properties.

Results: Two components, physical and mental fatigue, were identified in the CFS patient group and in the general population samples. Area under the curve for ROC was .91. The fatigue scale effectively discriminates, at high scores, between CFS patients and the general population. Conclusion: Physical and mental fatigue are clearly separable components of fatigue. The CFQ can discriminate reliably between clinical and nonclinical conditions. © 2010 Elsevier Inc. All rights reserved.

Keywords: CFS; Chalder fatigue scale; Chronic fatigue syndrome; Fatigue; ROC

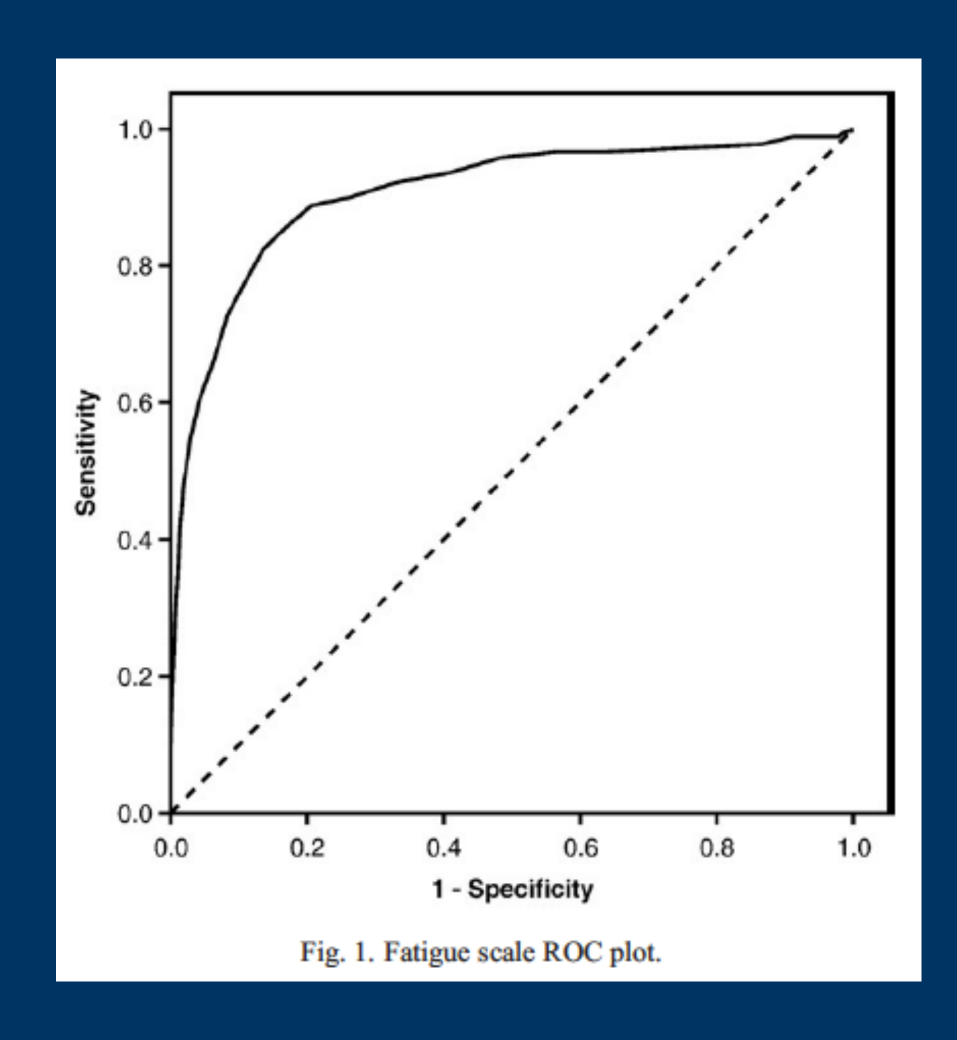


Real practical questions



How good is a diagnostic tool?

What is the best cutoff for discriminating cases from controls?



Area under the curve (AUC)

$$AUC \in [0,1]$$

AUC = $\pi_{Se} \times \pi_{Sp}$ for a binary predictor

Warm-up Exercise

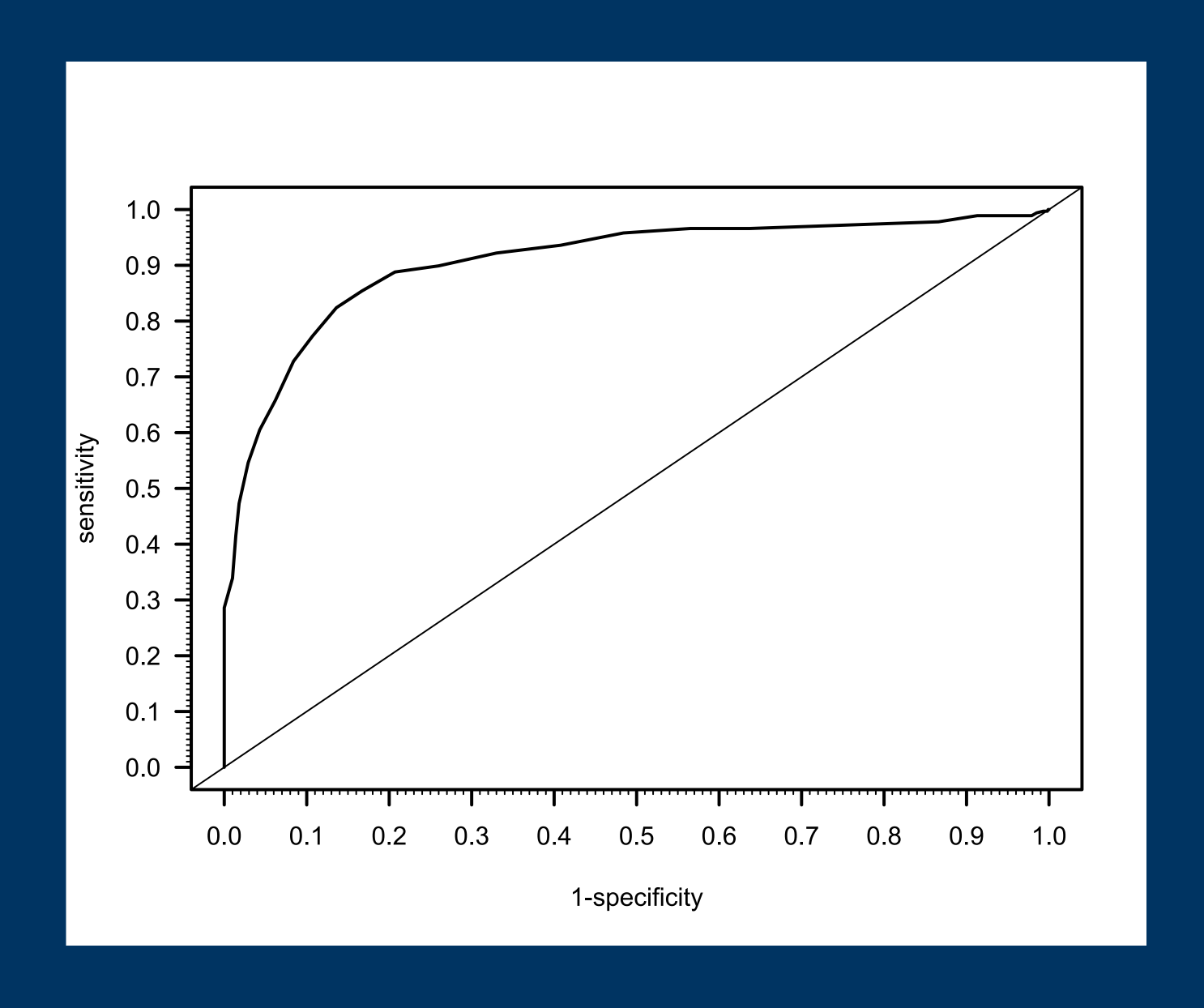
$$AUC = 0 \qquad AUC = 0.5$$

$$AUC = 1$$

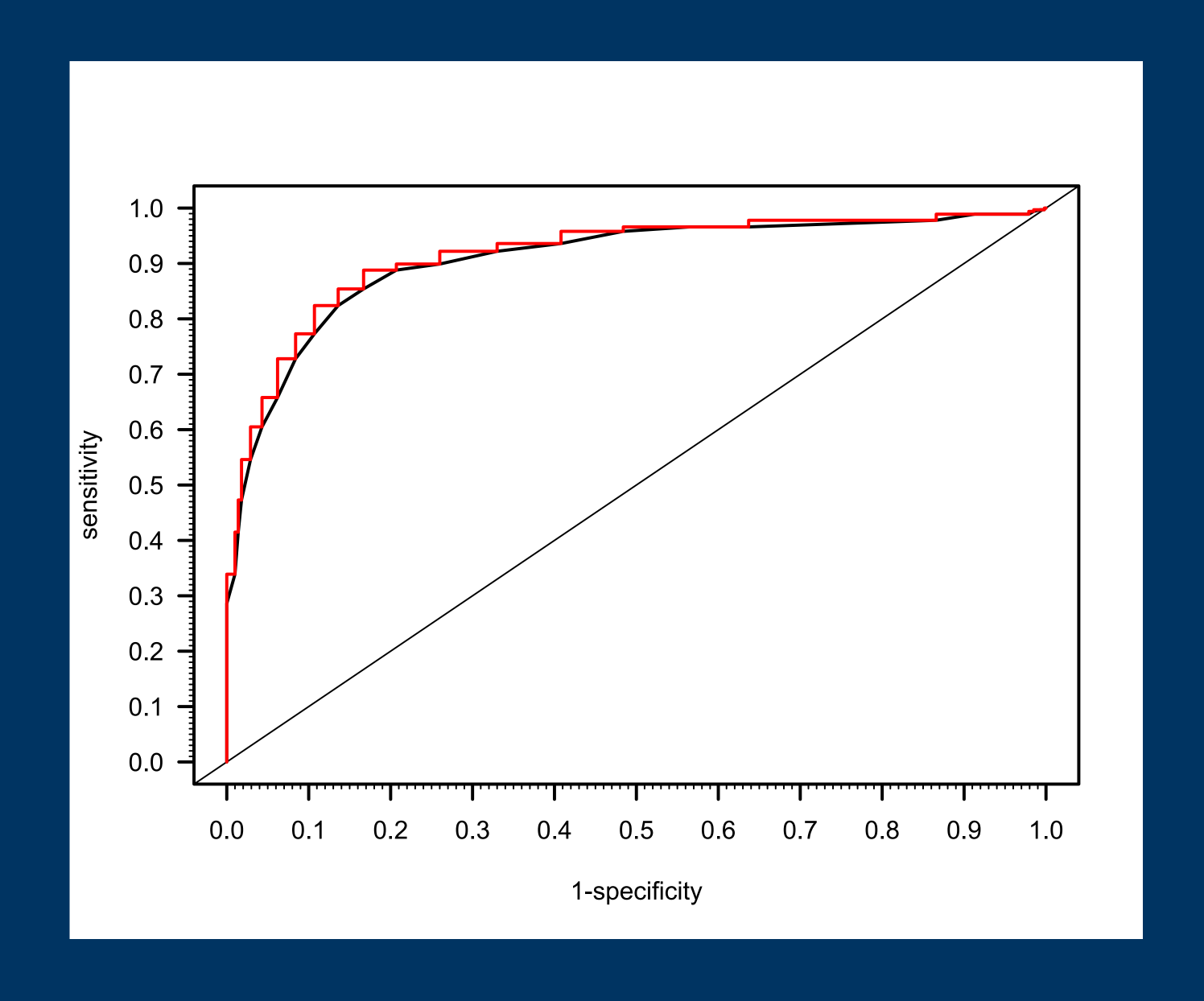
Draw the respective ROC curves

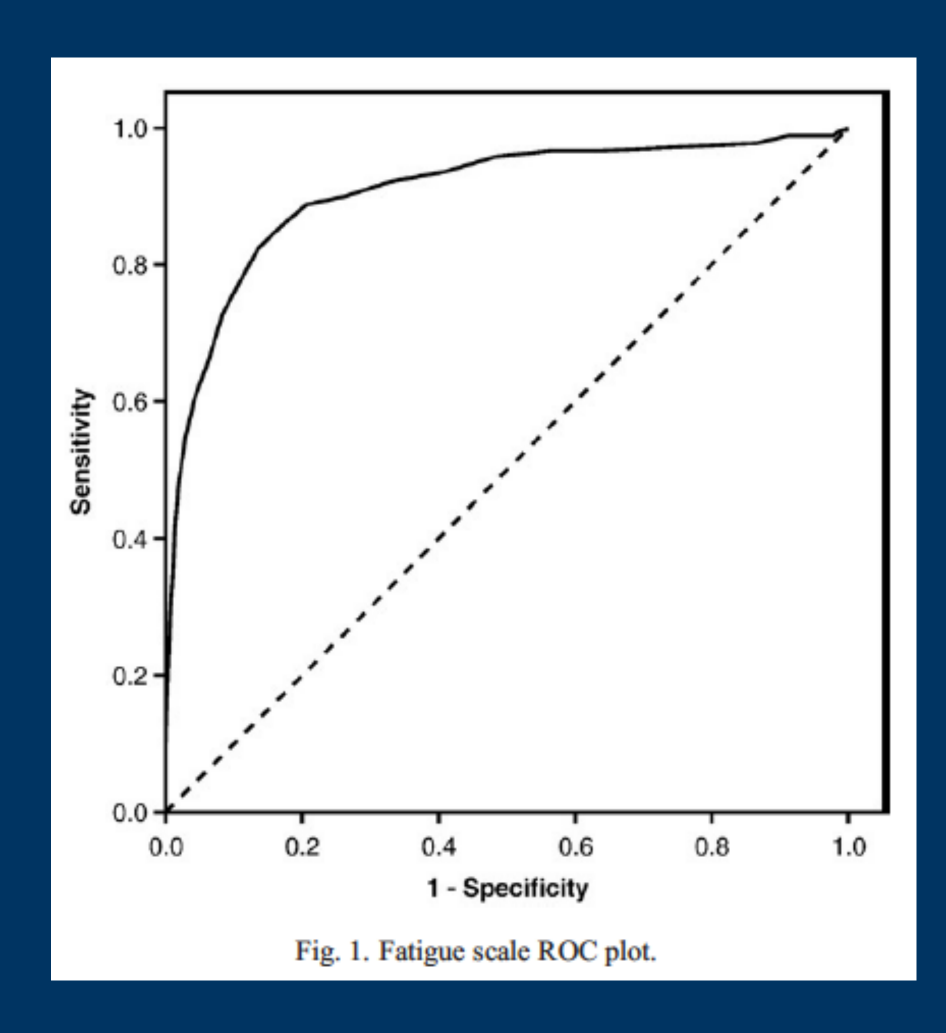
What is the respective interpretation?

A nuance conversation about ROC curve visualisation and AUC

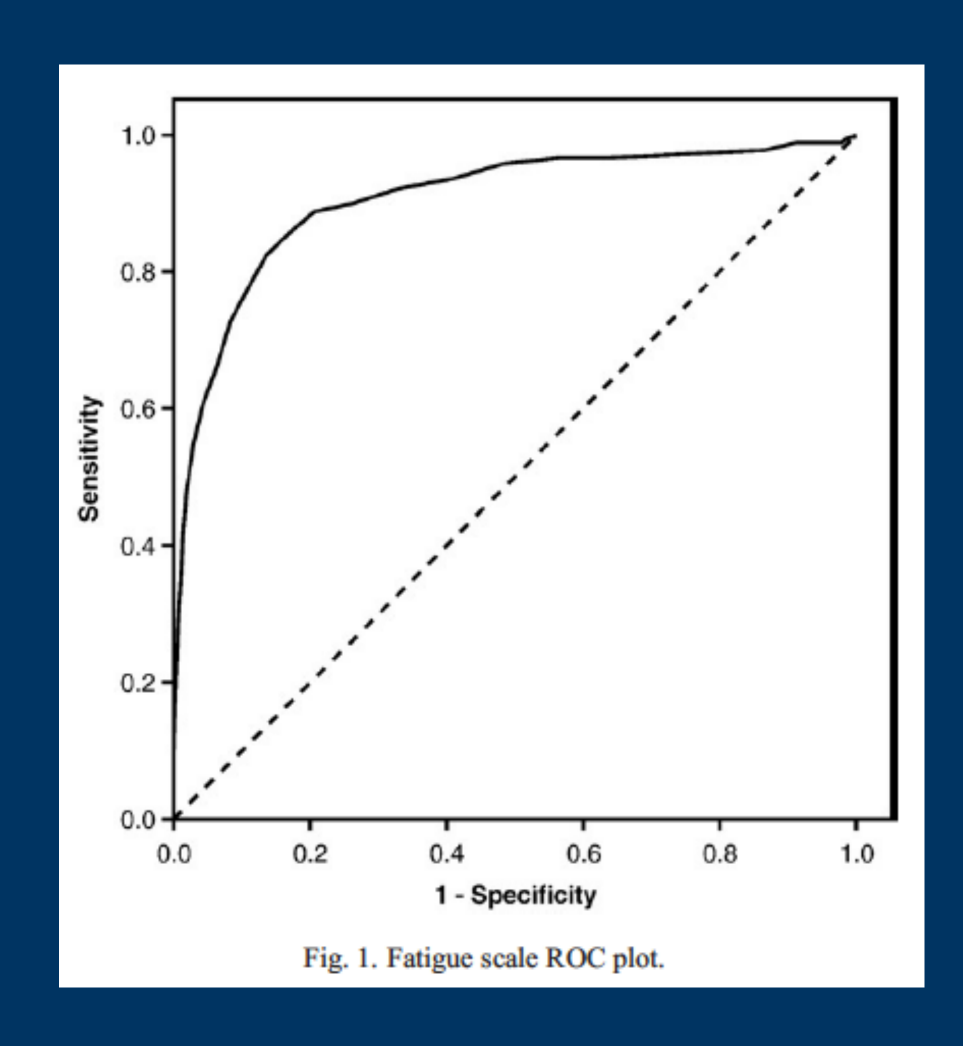


A nuance conversation about ROC curve visualisation and AUC





In practice, what is the most important hypothesis to test against the data?



In practice, what is the most important hypothesis to test against the data?

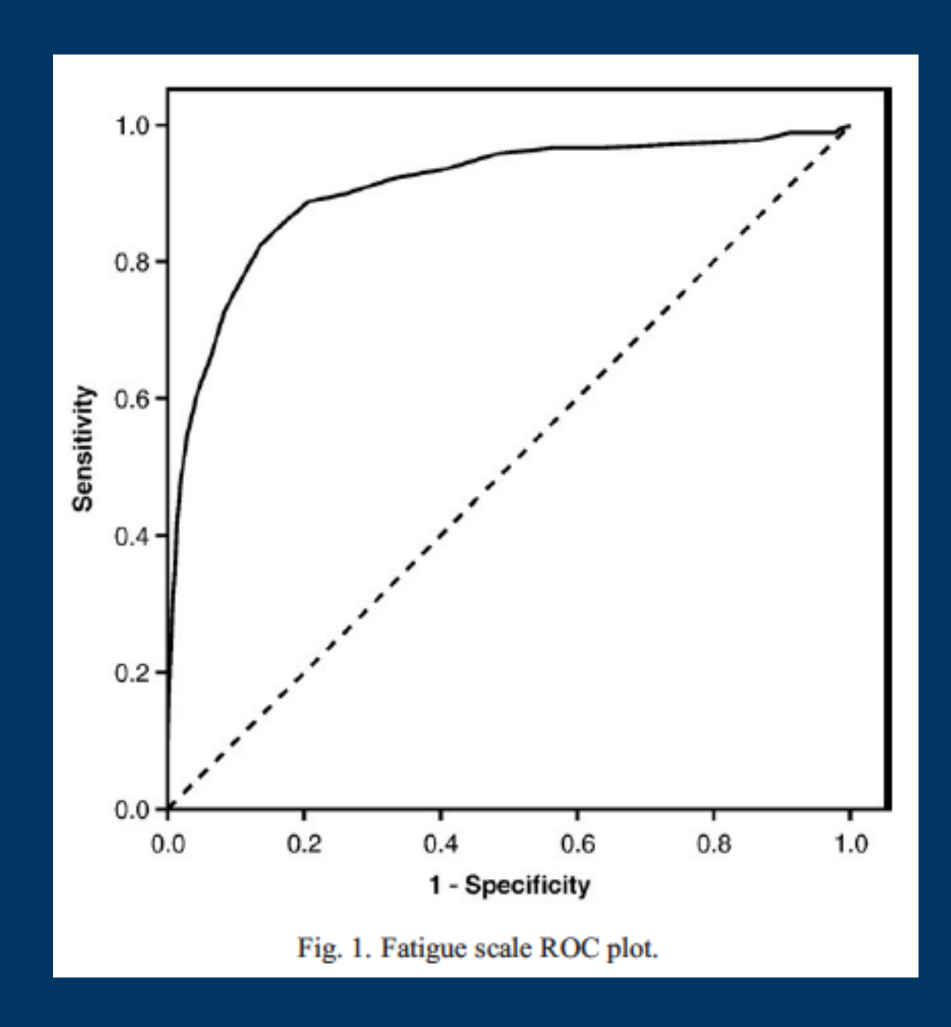
 $\overline{H_0}$: AUC = 0.5 versus AUC > 0.5

Wald's Confidence interval

$$IC_{95\%}$$
 (AUC) = $\hat{AUC} \pm 1.96 \times se$ (AÛC)

Accept H_0 if $0.5 \in IC_{95\%}$ (AUC)

Reject H_0 , otherwise

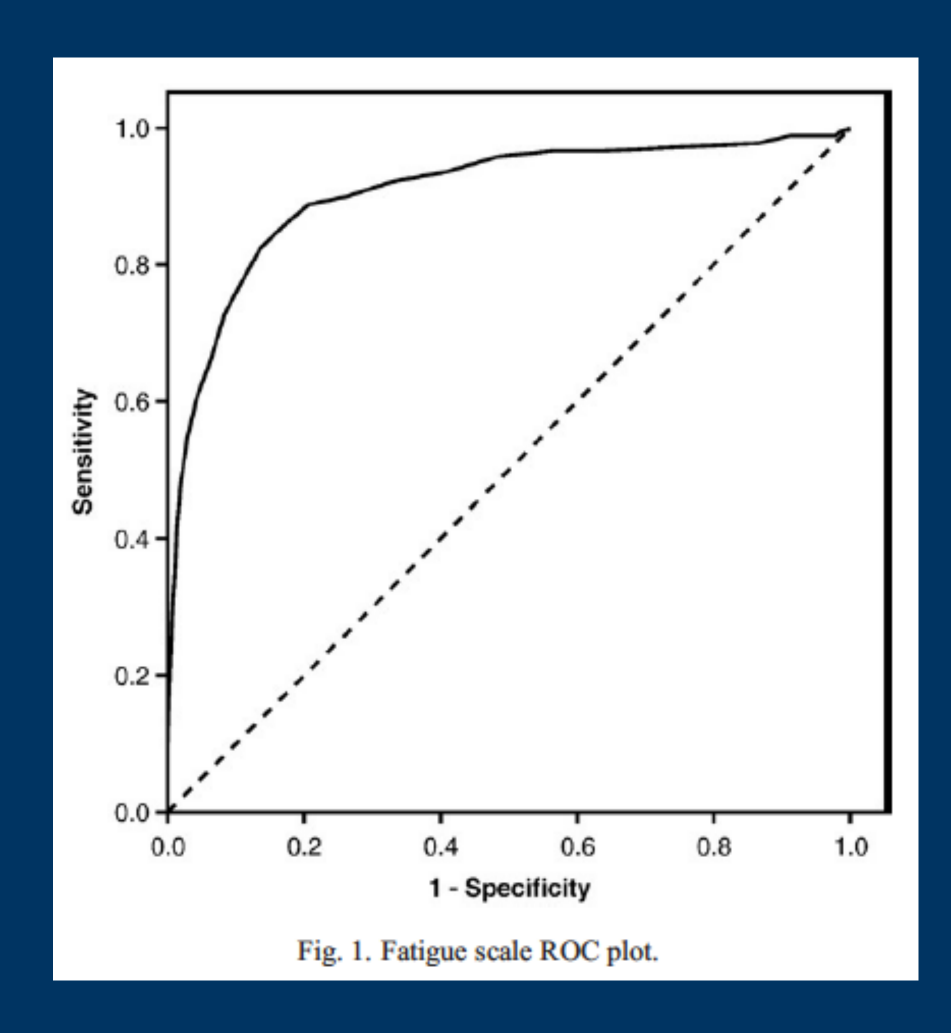


Wald's Confidence interval

$$IC_{95\%}$$
 (AUC) = $\hat{AUC} \pm 1.96 \times se$ (AÛC)

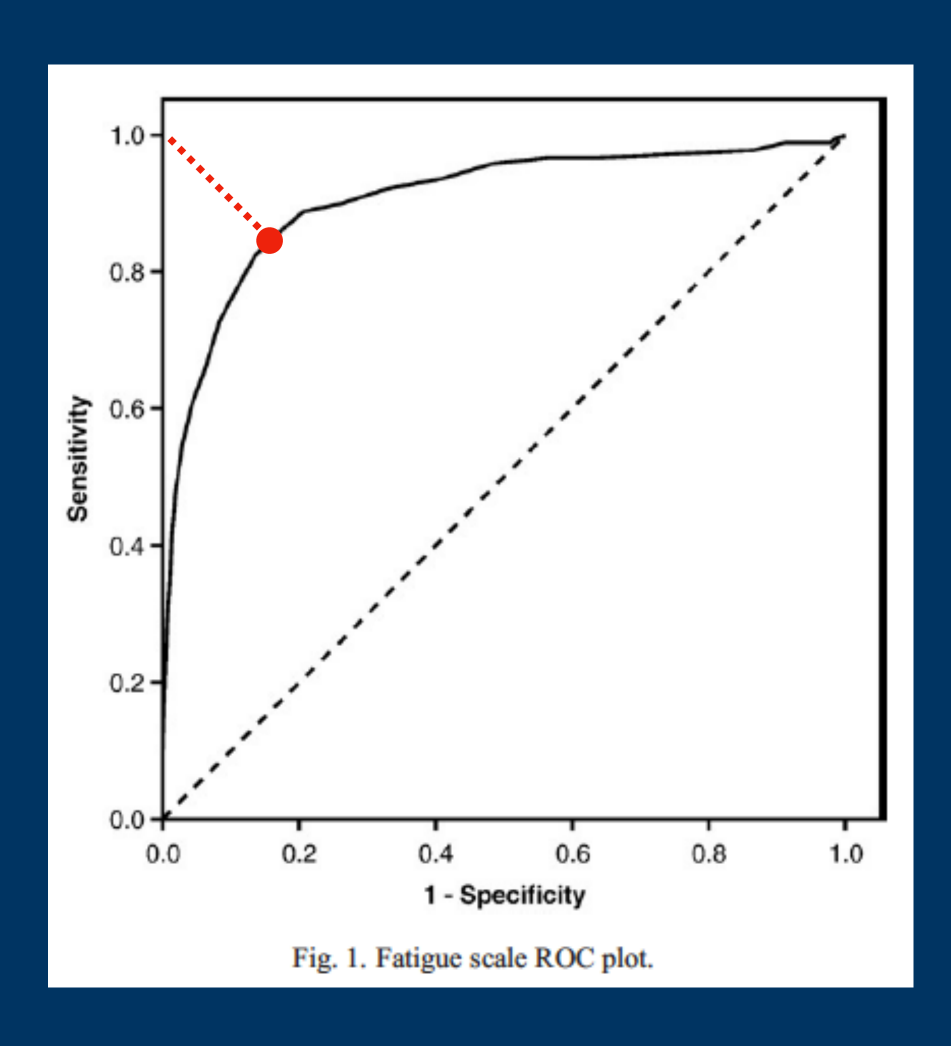
Problem: the upper limit might be larger than 1

Solution: use bootstrap (beyond the scope of the course).



Which point choose we choose from the ROC curve?

Which possible criteria can we use?

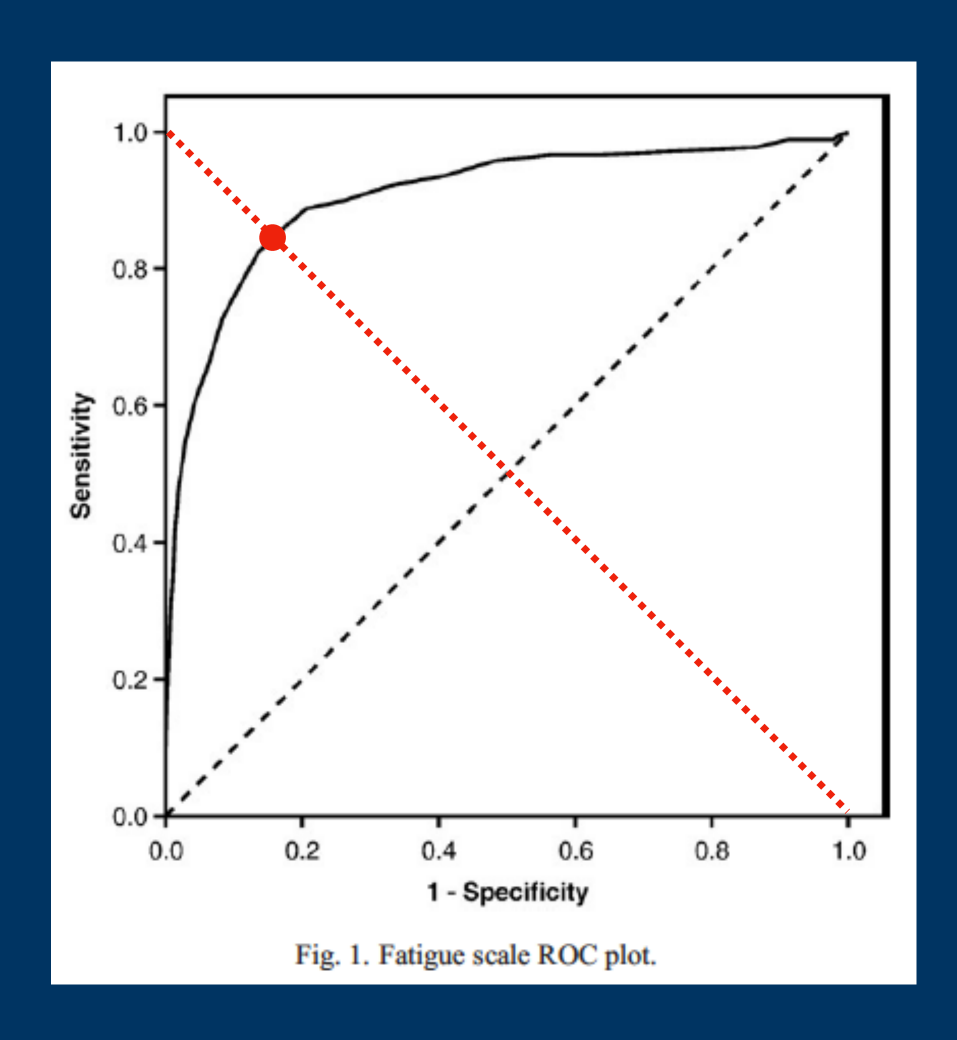


Which point choose we choose from the ROC curve?

Which possible criteria can we use?

Criterion I: ROC01

Point that is the closest to perfect classification (X=0,Y=1)



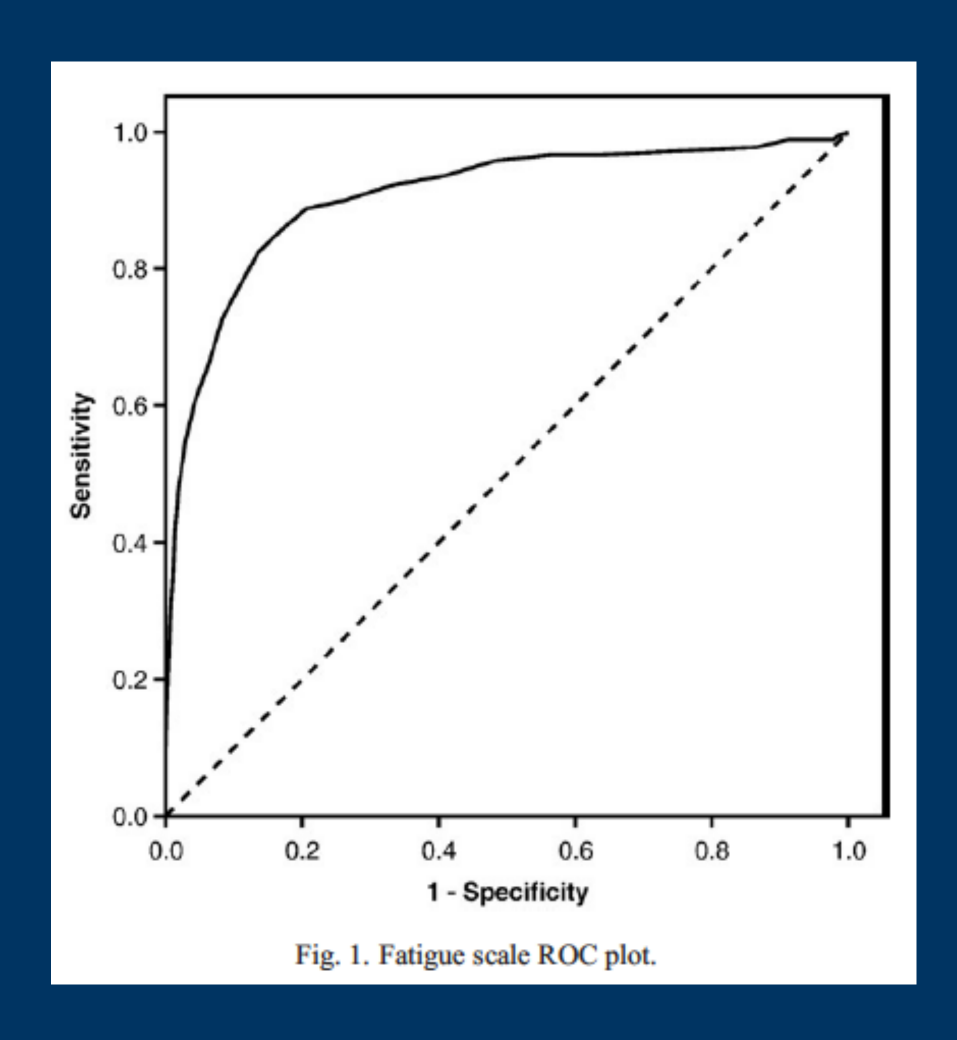
Which point choose we choose from the ROC curve?

Which possible criteria can we use?

Criterion II: SeEqualSp

Point where the sensitivity and specificity are the same

Point where the distance between sensitivity and specificity is minimized



Which point choose we choose from the ROC curve?

Which possible criteria can we use?

Criterion III: maxEfficiency

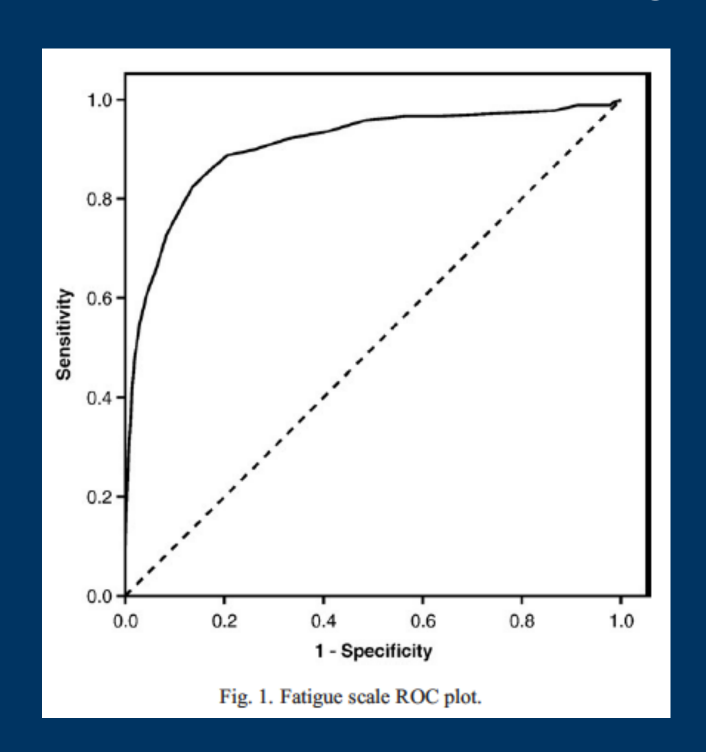
Point where the sensitivity and specificity provide the best accuracy

Exercise

Dataset: data_CFQ_extended.csv

Patients with chronic fatigue syndrome versus individuals from the general population





CFQ Score	Sensitivity	1-Specificity
0	1.000	1.000
1	1.000	.999
2	.997	.998
3	.997	.997
4	.997	.993
5	.994	.985
6	.989	.979
7	.989	.964
8	.989	.940
9	.989	.913
10	.978	.866
11	.966	.637
12	.966	.565
13	.958	.484
14	.936	.408
15	.922	.330
16	.899	.260
17	.888	.207
18	.854	.167
19	.824	.136
20	.773	.107
21	.728	.084
22	.658	.062
23	.605	.043
24	.546	.029
25	.473	.018
26	.415	.014
27	.339	.010
28	.286	.007
29	.202	.004
30	.151	.000
31	.095	.000
32	.062	.000
33	.000	.000

Calculate AUC (pROC package) and its 95% confidence interval.

Estimate the optimal cutoff for CFQ score according to ROC01, SeEqualSp and maxEfficiency (OptimalCutpoints package).

What are your conclusions?

Note

When the diagnostic tools comes in a form of estimated probability (e.g., ML applications), it is usually to use 0.5 as the cutoff for the probability. Accuracy is then estimated.

This cut-off might not be optimal for accuracy.

So we can use previous criteria to select a different cutoff for the estimated probabilities.

Exercise

Dataset: data_ML_diagnostic_malaria_exposure.csv

ML Diagnostic tool (randomForest) to identify individuals previously exposed to malaria parasites (these individuals would go to prophylactic treatment).

Calculate ROC curve.

Calculate the accuracy using 0.5 as the cutoff for the probability.

Estimate the optimal cutoff for CFQ score according to maxEfficiency criterion and calculate accuracy (OptimalCutpoints package).