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Empirical Likelihood Based Confidence Intervals for the Difference between Two Sensitivities of Continuous-scale Diagnostic Tests at a Fixed Level of Specificity

Suqin Yao

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Empirical Likelihood Based Confidence Intervals for the Difference between Two Sensitivities of Continuous-scale Diagnostic Tests at a Fixed Level of Specificity

by

SUQIN YAO

Under the Direction of Gengsheng Qin

ABSTRACT

Diagnostic testing is essential to distinguish non-diseased individuals from diseased individuals. The sensitivity and specificity are two important indices for the diagnostic accuracy of continuous-scale diagnostic tests. If we want to compare the effectiveness of two tests, it is of interest to construct a confidence interval for the difference of the two sensitivities at a fixed level of specificity. In this thesis, we propose two empirical likelihood based confidence intervals (HBEI and HBEII) for the difference of two sensitivities at a predetermined specificity level. Simulation studies show that when correlation between the two test results exists, HBEI and HBEII intervals perform better than the existing bootstrap based BCa, BTI and BTII intervals due to shorter interval lengths. However, when there is no correlation, BCa, BTI and BTII intervals outperform HBEI and HBEII intervals due to better coverage probability in most simulation settings.

INDEX WORDS: Empirical likelihood, diagnostic test, sensitivity, specificity
Empirical Likelihood Based Confidence Intervals for the Difference between Two
Sensitivities of Continuous-scale Diagnostic Test at a Fixed Level of Specificity

by

SUQIN YAO

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Empirical Likelihood Based Confidence Intervals for the Difference between Two
Sensitivities of Continuous-scale Diagnostic Test at a Fixed Level of Specificity

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Table of Contents

List of Tables .................................................................................................................................. vi

Chapter 1 Introduction .................................................................................................................... 1

Chapter 2 Existing methods ........................................................................................................... 5
  2.1 Normal-approximation-based interval ................................................................................. 5
  2.2 Bootstrap based intervals .................................................................................................... 7
    2.2.1 Paired uncorrelated samples .................................................................................. 8
    2.2.2 Paired dependent samples ..................................................................................... 10
    2.2.3 New bootstrap intervals for $D(p_o)$ ...................................................................... 10

Chapter 3 Hybrid empirical likelihood based intervals for the difference between two sensitivities ........................................................................................................................................ 12

Chapter 4 Simulation .................................................................................................................... 17

Chapter 5 Dermatoscope example ................................................................................................ 20

Chapter 6 Discussion .................................................................................................................... 22

References ..................................................................................................................................... 23

Appendix I: Simulation tables ..................................................................................................... 25

Appendix II S-plus code for Simulation ....................................................................................... 34
  1. Normal distribution ............................................................................................................. 34
  2. Exponential distribution – no correlation .......................................................................... 40
  3. Exponential distribution – correlation ............................................................................... 45
  4. Dermatoscope example ..................................................................................................... 47
List of Tables

Table a- 1 Level of 95 per cent confidence interval for $D(p)=0$. Bivariate normal distribution with $\rho = 0$ ......................................................................................................................................... 25

Table a- 2 Level of 95 per cent confidence interval for $D(p)=0$. Bivariate normal distribution with $\rho = 0.5$ ........................................................................................................................................ 27

Table a- 3 Level of 95 per cent confidence interval for $D(p)=0$. Bivariate exponential distribution with $\rho = 0$ ......................................................................................................................................... 29

Table a- 4 Level of 95 per cent confidence interval for $D(p)=0$. Bivariate exponential distribution with $\rho \neq 0$ (Using 0.02 to generate diseased random sample) ................................................................. 31

Table a- 5 95 per cent confidence interval for the difference of sensitivities between the two clinical assessments with and without the use of dermatoscopy ......................................................... 33
Chapter 1 Introduction

Diagnostic tests play a key role in modern medicine by screening a specific population for evidence of disease. The interpretation to the result of a diagnostic test depends on the discriminatory accuracy of the test to distinguish diseased patients from non-diseased subjects (Shapiro, 1999). Sensitivity and specificity are two measurements to describe the discriminatory accuracy of a test, which are defined as the probability of the test correctly identifying the diseased and non-diseased subjects respectively.

A diagnostic test is named continuous, dichotomous, or ordinal test depending on whether the test generate a continuous result (e.g. blood pressure), a dichotomous outcome (e.g. positive or negative), or an ordinal conclusion (e.g. confidence rating for presence of disease—definitely, probably, possibly, probably not, definitely not) (Shapiro, 1999). The main focus in this thesis is on continuous-scale diagnostic tests.

In continuous-scale diagnostic tests, it is common to define a threshold or a cut-off point $\gamma$ and classify the subject as diseased if the test result $Y$ is greater than or equal to $\gamma$ and non-diseased if the test result $X$ is less than $\gamma$. Thus, sensitivity and specificity are defined for each cut-off point $\gamma$ as:

$$R = P(Y \geq \gamma) = 1 - G(\gamma),$$

$$Sp = P(X < \gamma) = F(\gamma),$$

respectively, where $G$ and $F$ are the distribution functions of $Y$ and $X$ respectively. Let $X_1, X_2, ..., X_m$ be the test results of a random sample of non-diseased subjects, and $Y_1, Y_2, ..., Y_n$ be the test results of a random sample of diseased subjects. As we can see, when $\gamma$ decreases,
sensitivity increases but specificity decreases; as $\gamma$ increases, specificity increases at the expense of sensitivity. Therefore, there is a compromise between sensitivity and specificity when cut-off point changes, which is accounted for assessing discriminatory accuracy. From equation (1-1), the relationship between sensitivity and specificity can be set up without knowing the exact value of cut-off point $\gamma$. Let the specificity of a test be $p (0 \leq p \leq 1)$, the corresponding sensitivity of the test is

$$R(p) = 1 - G(F^{-1}(p)),$$  

(1-2)

where $F^{-1}$ is the inverse function of $F$.

Using equation (1-2), we can estimate the sensitivity of a test at a fixed level of specificity based on test results from the diseased and non-diseased subjects. It is also of interest to construct confidence intervals for the sensitivity $R(p)$. However, if we have two (or more) continuous-scale diagnostic tests to the same set of subjects, some of whom are non-diseases, some diseased, we may be more interested in knowing which test is better, especially when only a particular value of specificity is relevant (e.g. 70%, 80%, 90%). There are studies in literature for comparing the accuracy of two or more diagnostic tests, including comparing ROC curves and comparing summary accuracy indices (such as AUC, partial AUC, sensitivity and specificity). Some studies used ‘unpaired’ design, in which each diagnostic test is applied to a different group of subjects. The other studies utilized paired design, in which the diagnostic tests are applied to the same subjects (Shapiro, 1999). We focus on the comparison of sensitivities of two tests at a common specificity in this thesis.

Greenhouse and Mantel (1950) provided normal-theory that a diagnostic test has at least a specified sensitivity (e.g. $\geq 0.9$) with specificity higher than a specified value (e.g. $\geq 0.95$). Based on the result of Greenhouse and Mantel, Linnet (1987) proposed both parametric and non-
parametric methods for constructing confidence intervals for the sensitivity of a test at a fixed value of specificity, accounting for the random variation associated with the estimated cut-off point. Wieand et al. (1989) studied asymptotic behaviors of these non-parametric procedures and generalized them to a comparison of two weighted average of sensitivities. Their theory can be used to construct a normal approximation based confidence interval (WGJ interval) for the difference between two sensitivities. Qin et al. (2006) proposed three new bootstrap based intervals (BCa, BTI, BTII) that have better coverage accuracy than the WGJ interval.

Empirical likelihood (EL) (Owen, 1990, 2001) is a popular non-parametric method traditionally used for providing confidence intervals for means. The EL method has many advantages over other non-parametric methods. For example, it has better small sample performance than approaches based on normal approximation; empirical likelihood based confidence regions are range preserving and transformation respecting; the regularity conditions for empirical likelihood based methods are weak and natural. However, the empirical likelihood method has not been used widely in the study of accuracy of diagnostic tests. Qin (2007) proposed empirical likelihood based confidence intervals for the sensitivity of a single test at a fixed level of specificity. In this thesis, we are going to expand Qin’s finding (2007) in one continuous-scale test to construct EL-based confidence intervals for the difference between the sensitivities of two continuous-scale tests at a fixed level of specificity.

The thesis is organized as follow. In Chapter 2, we review some existing methods for the interval estimation of the difference between two sensitivities. In Chapter 3, we propose new hybrid empirical likelihood and bootstrap confidence intervals for the difference between two sensitivities at a pre-determined specificity, by using the asymptotic scaled chi-square distribution of the empirical likelihood ratio statistic. In Chapter 4, simulation studies are
conducted to compare the relative performance of the proposed empirical likelihood based intervals with the existing bootstrap intervals (BCa, BTI, and BTII). In Chapter 5, the new empirical likelihood based confidence intervals for the difference between two sensitivities are applied to a real example. A discussion is given in Chapter 6, and simulation tables and S-plus code are provided in the Appendix I and II.
Chapter 2 Existing methods

For two continuous-scale diagnostic tests, it is of interest to compare their sensitivities at a predetermined level of specificity. In this chapter, we give a review of the existing normal-approximation based interval proposed by Wieand (1989) and three bootstrap based intervals recently proposed by Qin et al. (2006) for the difference between two sensitivities at a fixed level of specificity.

2.1 Normal-approximation-based interval

Greenhouse and Mantel (1950) and Linnet (1987) proposed non-parametric procedures for the comparison of two sensitivities at a fixed level of specificity. Wieand et al. (1989) studied asymptotic behaviors of these non-parametric procedures and generalized them to a comparison of two weighted average of sensitivities.

Let \( T_1 \) and \( T_2 \) be two diagnostic tests that yield continuous measurements. It is assumed that both tests are performed on the same \( m \) controls (non-diseased) and \( n \) cases (diseased). \((X_{i1}, X_{2i}), i = 1, 2, \ldots, m\) are i.i.d. bivariate outcomes from the population with a joint distribution \( F(x_1, x_2) \) that represents the non-diseased group, \((Y_{1j}, Y_{2j}), j = 1, 2, \ldots, n\) are i.i.d. bivariate outcomes from the population with a joint distribution \( G(y_1, y_2) \) that represents the diseased group. The marginal distribution functions of \( X_k \) and \( Y_k \) are denoted by \( F_k(x_i) \) and \( G_k(y_j) \) respectively, \( k = 1, 2 \). For a given cut-off point \( \gamma \), the sensitivity and specificity of the test \( T_k, k = 1, 2 \) are defined by

\[
R_k = P(Y_k \geq \gamma) = 1 - G_k(\gamma), Sp_k = P(X_k < \gamma) = F_k(\gamma),
\]
respectively. Thus, the sensitivity of test $T_k$ at a fixed value of specificity $p$, is

$$R_k(p) = 1 - G_k(F_k^{-1}(p)),$$

where $F_k^{-1}(p) = \inf\{t : F_k(t) \geq p\}, k = 1, 2$. The parameter of interest is the difference between two sensitivities at the same fixed value of specificity $p_0$,

$$D(p_0) = R_1(p_0) - R_2(p_0). \quad (2-2)$$

Let $\hat{G}_k$ be the empirical distribution of $G_k$, based on the sample $X_{k1},...,X_{km}$, and $\hat{F}_k^{-1}(p)$ be the empirical estimate for the $p$-th quantile of $F_k$, $k = 1, 2$, based on the sample $Y_{k1},...,Y_{kn}$. The non-parametric estimator for $D(p_0)$ proposed by Linnet (1987) and Wieand et al. (1989) is given as follows:

$$\hat{D}(p_0) = \hat{R}_1(p_0) - \hat{R}_2(p_0), \quad (2-3)$$

where $\hat{R}_k(p_0) = 1 - \hat{G}_k(\hat{F}_k^{-1}(p_0))$.

Let $N=m+n$. Wieand et al. (1989) showed that

$$N^{1/2}(\hat{D}(p_0) - D(p_0)) \xrightarrow{d} N(0, \sigma^2), \quad (2-4)$$

where

$$\sigma^2 = \sigma_1^2 + \sigma_2^2 - 2\sigma_{12},$$

$$\sigma_k^2 = (1 - \lambda)^{-1} R_k(p_0)(1 - R_k(p_0)) + \lambda^{-1} (1 - p_0) p_0 \frac{g_k^2(F_k^{-1}(p_0))}{f_k^2(F_k^{-1}(p_0))} \quad (k = 1, 2),$$

$$\sigma_{12} = (1 - \lambda)^{-1} \{G(F_1^{-1}(p_0), F_2^{-1}(p_0)) - G_1(F_1^{-1}(p_0))G_2(F_2^{-1}(p_0))\} +$$

$$\lambda^{-1} [F(F_1^{-1}(p_0), F_2^{-1}(p_0)) - p_0^2] \frac{g_1(F_1^{-1}(p_0))g_2(F_2^{-1}(p_0))}{f_1(F_1^{-1}(p_0))f_2(F_2^{-1}(p_0))},$$

$$\lambda = m/(m+n),$$
where $f_k$ and $g_k$ are the density functions of $F_k$ and $G_k$ respectively.

If a good estimate for $\sigma^2$ is available, the normal approximation equation (2-4) can be used to construct a confidence interval for the difference between two sensitivities at the same fixed level of specificity. However, the estimation of $\sigma^2$ requires the estimation of density functions $f_k$ and $g_k$, the estimation of bivariate distribution functions $F(x_1,x_2)$ and $G(y_1,y_2)$, and the estimation of quantiles $F_k^{-1}(p)$. Therefore, the performance of the normal approximation based interval is very sensitive to the choice of the smoothing parameters in density and distribution estimations. Selection of satisfactory smoothing parameters in this context is problematic.

### 2.2 Bootstrap based intervals

Qin et al. (2006) proposed three intervals called BCa, BTI and BTII intervals for the difference between sensitivities of two diagnostic tests at a fixed value of specificity by using bootstrap method. The major advantage of these intervals over the normal approximation based interval is that no density and distribution estimation is needed. And the new intervals are computationally easy to implement in practice.

The difference between two sensitivities at the same fixed value of specificity $p_0$ is the difference between two proportions:

$$D(p_0) = R_1(p_0) - R_2(p_0) = P(Y_{ik} \geq F_k^{-1}(p_0)) - P(Y_{2k} \geq F_2^{-1}(p_0)).$$

If $F_k$ were known, an obvious estimator of $D(p_0)$ would be the difference between the observed sensitivities at $p_0$-th quantiles $F_k^{-1}(p_0)$ and $F_2^{-1}(p_0)$, which would be defined as
\[ \hat{D}(p_0) = \frac{1}{n} \sum_{j=1}^{n} I_{[Y_j > F_k^{-1}(p_0)]} - \frac{1}{n} \sum_{j=1}^{n} I_{[Y_j \geq F_k^{-1}(p_0)]}, \quad (2-5) \]

where \( I_A \) is the indicator function of \( A \). We can also regard \( \hat{D}(p_0) \) as the difference between two sample proportions of binomial distributions with proportions \( R_k(p_0) \), \( k = 1, 2 \). However, \( F_k \)'s are unknown, by replacing \( F_k^{-1}(p_0) \) by \( \hat{F}_k^{-1}(p_0) \) in equation (2-5), we acquire an estimator \( \hat{D}(p_0) \) for \( D(p_0) \).

\[ \hat{D}(p_0) = \frac{1}{n} \sum_{j=1}^{n} I_{[Y_j > \hat{F}_k^{-1}(p_0)]} - \frac{1}{n} \sum_{j=1}^{n} I_{[Y_j \geq \hat{F}_k^{-1}(p_0)]} \quad (2-6) \]

Because the indicator variables \( I_{[Y_i > \hat{F}_k^{-1}(p_0)]}, I_{[Y_i \geq \hat{F}_k^{-1}(p_0)]}, \ldots, I_{[Y_n > \hat{F}_k^{-1}(p_0)]} \) are not independent, \( \hat{D}(p_0) \) is no longer the difference between two simple binomial proportions. Depending on whether there is a correlation between the test results from two diagnostic tests, Qin et al. (2006) proposed the following different procedures for the confidence intervals of \( D(p_0) \) by combining bootstrap method with the technique provided by Agresti and Caffo (2000).

2.2.1 Paired uncorrelated samples

If the test results from two diagnostic tests are conditionally uncorrelated, \( \hat{D}(p_0) \) can be considered as the difference between two independent sample proportions. Qin et al. (2006) proposed the following estimator for \( D(p_0) \) instead of \( \hat{D}(p_0) \):

\[ \hat{D}(p_0) = \hat{R}_1(p_0) - \hat{R}_2(p_0), \quad (2-7) \]

where

\[ \hat{R}_k(p_0) = \frac{\sum_{j=1}^{n} I_{[Y_j > \hat{F}_k^{-1}(p_0)]} + Z_{1-\alpha/2}^2 / 2}{n + Z_{1-\alpha/2}^2}, k = 1, 2 \quad (2-8) \]
The above procedure can also be used to the case of two independent samples with different sample size.
2.2.2 Paired dependent samples

When two diagnostic tests are applied to the same patients, the test results from two diagnostic tests are most likely correlated. Qin et al. (2006) proposed to use the following estimates for the sensitivities:

\[
\hat{R}_k(p_0) = \frac{\sum_{j=1}^{n} I_{y_j > \hat{R}_k^{-1}(p_0)} + 1}{n + 2}, k = 1, 2.
\]

The bootstrap estimate \( V^* \) for the variance of \( \hat{D}(p_0) \) is defined as follows:

\[
V^* = V_1^* + V_2^* - 2V_{12}^*,
\]

where \( V_k^* \ (k = 1, 2) \) are defined as before, and

\[
V_{12}^* = \frac{1}{B-1} \sum_{b=1}^{B} (\hat{R}_{1b}^*(p_0) - \bar{R}_1^*(p_0))(\hat{R}_{2b}^*(p_0) - \bar{R}_2^*(p_0))
\]

2.2.3 New bootstrap intervals for \( D(p_0) \)

Qin et al. (2006) proposed three new intervals for \( D(p_0) \). The first two \((1 - \alpha)100\) per cent confidence intervals for \( D(p_0) \) are bootstrap intervals based on the bootstrap variance estimate \( V^* \). They are defined as follows:

(i) The first one, called BTI interval, is

\[
(\hat{D}(p_0) - z_{1-\alpha/2} \sqrt{V^*}, \hat{D}(p_0) + z_{1-\alpha/2} \sqrt{V^*})
\]

where \( \hat{D}(p_0) \) is defined by equation (2-7)

(ii) The second one, called BTII interval, is

\[
(\bar{D}^*(p_0) - z_{1-\alpha/2} \sqrt{V^*}, \bar{D}^*(p_0) + z_{1-\alpha/2} \sqrt{V^*})
\]
Where \( \hat{D}^*(p_0) = \frac{1}{B} \sum_{b=1}^{B} \hat{D}_b^*(p_0) \)

The above two intervals require variance estimation of \( \hat{D}(p_0) \). The third interval for \( D(p_0) \) proposed by Qin et al. (2006) is a BCa-type bootstrap interval in which the direct variance estimation is not needed:

\[
(\hat{D}_{(B\hat{c}/2)}^*(p_0), \hat{D}_{(B(1-\hat{c}/2))}^*(p_0)),
\]

where

\[
\hat{c} = \Phi(w + \frac{w + z_\alpha}{1 - \alpha(w + z_\alpha)})
\]

\[
w = \Phi^{-1}(\frac{1}{B} \sum_{b=1}^{B} I_{[\hat{D}_b^*(p_0) \leq \hat{D}(p_0)]})
\]

\[
\hat{\alpha} = \frac{1}{6} \frac{\sum_{k=1}^{b} l_k^3}{(\sum_{k=1}^{b} l_k^2)^{3/2}}
\]

\[
l_k = (I_{[Y_i \geq \hat{F}_i^{-1}(p_0)]} - I_{[Y_i \geq \hat{F}_i^{-1}(p_0)]}) - (\hat{R}_1(p_0) - \hat{R}_2(p_0))
\]

and \( \Phi \) is the standard normal distribution function, and \( \hat{D}_{(b)}^*(p_0) \) is the \( b \)-th ordered value among \( \{\hat{D}_b^*(p_0), b = 1, 2, ..., B\} \).

Through simulation study, Qin et al. (2006) showed that BTI and BTII intervals perform better than the normal approximation based interval for independent samples, and BCa interval performs better than the normal approximation based interval for paired dependent samples. In addition, BTI and BTII intervals are computationally simpler than the normal approximation based interval. Therefore, we only use BCa, BTI and BTII intervals as a comparison in this thesis.
Chapter 3 Hybrid empirical likelihood based intervals for the difference between two sensitivities

We recently developed an empirical likelihood based method to construct the confidence interval for the difference between two sensitivities from two diagnostic tests at a fixed level of specificity. An introduction of this method is given in this chapter.

Pepe (2003) defined a placement value for a given test value $Y$ from a diseased subject as

$$U = 1 - F(Y).$$

This value is the proportion of the non-diseased population with a test value greater than $Y$. It marks the placement of $Y$ within non-diseased distribution.

It is evident that

$$E(I(U \leq 1 - p)) = P(F(Y) \geq p) = P(Y \geq F^{-1}(p)) = R(p).$$

For two diagnostic tests $T_1$ and $T_2$ that yield continuous measurements, we have

$$U_k = 1 - F_k(y_k), k = 1, 2;$$

$$E[I(U_k \leq 1 - p)] = P(Y_k \geq F_k^{-1}(p)) = 1 - G_k(F_k^{-1}(p)) = R_k(p).$$

Therefore,

$$D(p) = R_1(p) - R_2(p) = E[I(U_1 \leq 1 - p)] - E[I(U_2 \leq 1 - p)]$$

Based on this relationship between $D(p)$ and the placement value $U_k$’s, an empirical likelihood procedure is derived for the difference between two sensitivities. Let
$P_k = (p_{k1}, p_{k2}, \ldots, p_{kn}), k = 1, 2$ be two probability vectors, i.e., \( \sum_{j=1}^{n} p_{kj} = 1 \) and \( p_{kj} \geq 0 \) for all \( j \). The profile EL for \( D(p) \) can be defined as

\[
L(D(p)) = \sup \left\{ \prod_{k=1}^{2} \prod_{j=1}^{n} p_{kj} : \sum_{j=1}^{n} p_{kj} W_{kj}(p) = 1, \sum_{j=1}^{n} p_{kj} V_{kj}(p) = 0, \sum_{j=1}^{n} p_{1j} \hat{V}_{1j} = \sum_{j=1}^{n} p_{2j} \hat{V}_{2j} = D(p), k = 1, 2 \right\},
\]

(3-1)

where

\[
W_{kj}(p) = I(U_{kj} \leq 1 - p) - R_k(p) \equiv V_{kj}(p) - R_k(p),
\]

\[
V_{kj}(p) = I(U_{kj} \leq 1 - p), k = 1, 2.
\]

The placement values, \( U_{kj} \)'s (\( k = 1, 2 \)), depend on the unknown distribution functions \( F_k \)'s (\( k = 1, 2 \)) of the non-diseased populations. Therefore, by replacing \( F_k \) by its empirical distribution \( \hat{F}_k \), we get an adjusted empirical likelihood for \( D(p) \):

\[
\hat{L}(D(p)) = \sup \left\{ \prod_{k=1}^{2} \prod_{j=1}^{n} p_{kj} : \sum_{j=1}^{n} p_{kj} \hat{W}_{kj}(p) = 1, \sum_{j=1}^{n} p_{kj} \hat{V}_{kj}(p) = 0, \sum_{j=1}^{n} p_{1j} \hat{V}_{1j} = \sum_{j=1}^{n} p_{2j} \hat{V}_{2j} = D(p), k = 1, 2 \right\}
\]

where

\[
\hat{W}_{kj}(p) = I(\hat{U}_{kj} \leq 1 - p) - R_k(p) \equiv \hat{V}_{kj}(p) - R_k(p),
\]

\[
\hat{V}_{kj}(p) = I(\hat{U}_{kj} \leq 1 - p), k = 1, 2.
\]

By using the Lagrange multiplier method, we get the corresponding log-EL ratio statistic:

\[
l(D(p)) = 2(\sum_{j=1}^{n} \log(1 + 2t\hat{w}_{1j}(p))) + \sum_{j=1}^{n} \log(1 - 2t\hat{w}_{2j}(p))), \quad (3-2)
\]

where \( t, R_1(p), R_2(p) \) are determined by
Qin (2007) established the following theorem for the asymptotic distribution of the log-EL likelihood ratio statistic.

**Theorem 3.1.** If $D_0(p)$ is the true value of $D(p) = R_1(p) - R_2(p)$ at a fixed level $p$ of specificity, then the limiting distribution of $l(D(p))$, defined by equation (3.2), is a scale chi-square distribution with one degree of freedom. That is,

$$r(p)l(D_0(p)) \xrightarrow{d} \chi^2_1,$$

where the scale constant $r(p)$ is

$$r(p) = \frac{R_1(p)(1 - R_1(p)) + R_2(p)(1 - R_2(p))}{(1 - \lambda)\sigma^2}.$$

The scale constant $r(p)$ in Theorem 3.1 is still unknown. In order to construct confidence intervals for $D(p)$, we propose to use bootstrap method to estimate $r(p)$. The procedure is as follows:

**Step 1:** Draw resample of size $m$, $X_{m_i}^*$'s, with replacement from the non-diseased sample $X_{ki}'s$ and a separate resample of size $n$, $Y_{yi}^*$'s, with replacement from the diseased sample $Y_{yi}'s$. 

\[
\begin{align*}
\frac{1}{n} \sum_{j=1}^{n} \frac{\hat{V}_{ij} - R_i(p)}{1 + 2t(\hat{V}_{ij} - R_i(p))} &= 0 \\
\frac{1}{n} \sum_{j=1}^{n} \frac{\hat{V}_{2j} - R_2(p)}{1 - 2t(\hat{V}_{2j} - R_2(p))} &= 0 \\
\frac{1}{n} \sum_{j=1}^{n} \frac{\hat{V}_{ij} - R_i(p)}{1 + 2t(\hat{V}_{ij} - R_i(p))} - \frac{1}{n} \sum_{j=1}^{n} \frac{\hat{V}_{2j} - R_2(p)}{1 - 2t(\hat{V}_{2j} - R_2(p))} &= D(p)
\end{align*}
\]
Step 2: Calculate the bootstrap versions $\hat{R}_k^* (p)$ of $R_k (p), k = 1, 2$.

$$\hat{R}_k^* (p) = \sum_{i=1}^{n} I[Y_{ki} \geq \hat{R}_{k-1}^* (p) + Z_{1,1/2}^2 / n + Z_{1,1/2}^2 ]/ k = 1, 2.$$  

Step 3: Repeat Steps 1-2 $B$ ($B \geq 150$) times, we get $\{\hat{R}_{1b}^* (p), \hat{R}_{2b}^* (p) : b = 1 ... B\}$ and

$$V_k^* = \frac{1}{B-1} \sum_{b=1}^{B} (\hat{R}_{1b}^* (p) - \bar{R}_k^* (p))^2 , k = 1, 2,$$

$$V_{12}^* = \frac{1}{B-1} \sum_{b=1}^{B} ((\hat{R}_{1b}^* (p) - \bar{R}_1^* (p))(\hat{R}_{2b}^* (p) - \bar{R}_2^* (p)),$$

$$V^* = V_1^* + V_2^* - 2V_{12}^*,$$

where $\bar{R}_k^* (p_0) = \frac{1}{B} \sum_{b=1}^{B} \hat{R}_{xb}^* (p_0), k = 1, 2.$

Hence, the scale constant $r(p)$ can be consistently estimated by

$$r_1^* (p) = \frac{\bar{R}_1^* (p)(1 - \bar{R}_1^* (p)) + \bar{R}_2^* (p)(1 - \bar{R}_2^* (p))}{n*V^*},$$

or

$$r_2^* (p) = \frac{\hat{R}_1 (p)(1 - \hat{R}_1 (p)) + \hat{R}_2 (p)(1 - \hat{R}_2 (p))}{n*V^*}.$$

By using these estimates for $r(p)$, we propose two hybrid bootstrap and empirical likelihood based confidence intervals for $D(p)$.

The first one, called HBELI interval, is defined by

$$\{D(p) : r_1^* (p) I(D(p)) \leq \chi_i^2 (1 - \alpha) \},$$

(3-5)

where $\chi_i^2 (1 - \alpha)$ is the $(1 - \alpha)$-th quantile of $\chi_i^2$. 
The second one, called HBELII interval, is defined by

\[
\left\{ D(p): r_2^*(p) I(D(p)) \leq \chi^2_1(1 - \alpha) \right\}.
\] (3-6)
Chapter 4 Simulation

In this chapter, we conduct two simulation studies using bivariate normal distribution and exponential distribution to evaluate coverage accuracy and interval length of the newly proposed intervals for $D(p)$, the difference of the two sensitivities, when the specificity $p$ is taken to be 0.70, 0.80 or 0.90 in finite-sample sizes. In both studies, we generated 1000 random samples of size $n$ from $G(y_1, y_2)$ for test responses of diseased patients, and another set of independent random samples of size $m$ from $F(x_1, x_2)$ for test responses of non-diseased patients. In this thesis, we didn’t use the normal approximation based interval as a comparison because Qin et al. (2006) have already shown that BTI and BTII intervals perform better than the normal approximation based interval for independent samples, and BCa performs better than the normal approximation based interval for paired dependent samples, and these three intervals are computationally much simpler than the normal approximation based interval.

In the first study, $G(y_1, y_2)$ is chosen to be a bivariate normal distribution having mean $E(Y_1) = \mu_1$, $E(Y_2) = \mu_2$ and with a common standard deviation 2 and correlation $\rho$; $F(x_1, x_2)$ is chosen to be a bivariate normal distribution having means $E(X_1) = 0$, $E(X_2) = 0$ and with a common standard deviation 1 and correlation $\rho$. $\rho$ is chosen as 0 and 0.5 respectively. Thus,

$$R_k(p) = 1 - \Phi\left\{\Phi^{-1}(p) - \mu_k\right\}/2$$

for $k = 1, 2$.

For $D(p) = 0$, we choose $\mu_1 = \mu_2$ such that the sensitivity $R_k(p)$ of the test $T_k(k = 1, 2)$ varies over the points 0.95, 0.90, 0.80, 0.70, 0.60, 0.50, 0.40, 0.30, 0.20, 0.10, respectively.

In the second study, the distributions $G(y_1, y_2)$, $F(x_1, x_2)$ are chosen to be different bivariate exponential distributions that have exponential distributions as their marginal
distributions. Depending on the possible correlation between the test results from two diagnostic
tests, we use two different procedures to generate the random samples of test response.

First we choose the correlation as zero \((\rho = 0)\), and then we generate two independent
samples, \(X_{11}, X_{12}, ..., X_{1m}\) and \(X_{21}, X_{22}, ..., X_{2n}\), from standard exponential distribution; and two
independent samples, \(Y_{11}, Y_{12}, ..., Y_{1\alpha}\) and \(Y_{21}, Y_{22}, ..., Y_{2\beta}\) from exponential distributions with rates
\(\lambda_1, \lambda_2\), respectively. Therefore,

\[
R_k(p) = \exp(\lambda_k \log(1 - p)), \quad \text{for } k = 1, 2.
\]

Similar to the first simulation study, we choose \(\lambda_k, l_k\) \((k = 1, 2)\) such that \(D(p) = 0\) as the
sensitivity \(R_i(p)\) of the test \(T_i\) \((i = 1, 2)\) varies over the points 0.95, 0.90, 0.80, 0.70, 0.60, 0.50,
0.40, 0.30, 0.20, 0.10 respectively.

Secondly, we choose a positive correlation for the bivariate exponential distribution
\((\rho > 0)\). We first generate random sample, \(U_{k1}, U_{k2}, ..., U_{km}\), from an exponential distribution with
rate 0.5, for \(k = 1, 2, 3\); and random samples, \(V_{k1}, V_{k2}, ..., V_{k\alpha}\), from an exponential distribution with
rate \(l_i\), for \(i = 1, 2, \ldots\); and a random sample, \(V_{31}, V_{32}, ..., V_{3\beta}\) from an exponential distributions with
rate 0.02. Then the simulated test responses for a non-diseased patients are
\(X_{ki} = \min(U_{ki}, V_{k3}), k = 1, 2, i = 1, 2, \ldots, m\), which are random samples from two standard exponential
distributions with correlation \(\rho\); and those for diseased patients are
\(Y_{kj} = \min(V_{kj}, V_{3j}), k = 1, 2, j = 1, 2, \ldots, n\), which are random samples from two exponential
distributions with correlation \(\rho\) and rates \(l_1, 0.02, l_2 + 0.02\), respectively. Under this setting,

\[
R_k(p) = \exp[(l_k + 0.02)\lambda_k \log(1 - p)], \quad \text{for } k = 1, 2.
\]
We choose $\lambda_k, l_k \ (k = 1, 2)$ such that $D(p) = 0$ as the sensitivity $R_k(p)$ of the test $T_k (k = 1, 2)$ varies over the points 0.95, 0.90, 0.80, 0.70 respectively.

In the bootstrap step, we draw $B=150$ bootstrap re-samples from the original samples. We construct 95% confidence intervals for $D(p)$. The results of the simulation study are shown in Table I to Table VI in Appendix I. From these tables, the following observations are made.

1. When the correlation $\rho = 0$ and $D(p) = 0$, the BCa, BTI and BTII intervals have better coverage probability, but HBELI and HBELII intervals have shorter interval length.

2. When the correlation $\rho > 0$ and $D(p) = 0$, the five intervals have similar coverage probability, but HBELI and HBELII intervals have shorter interval length.

3. When the correlation $\rho$ is positive, bigger sample sizes $(m,n \geq 150)$ are needed to get better coverage accuracy for all the intervals.

In summary, when correlation exists, the hybrid empirical likelihood and bootstrap based intervals HBELI and HBELII perform better than the bootstrap intervals due to the shorter interval length. When there is no correlation, the bootstrap based intervals BCa, BTI, BTII perform better than the HBELI and HBELII intervals due to better coverage probability.
Chapter 5 Dermatoscope example

Melanoma is a malignant tumor of melanocytes which are found predominantly in skin but also in the bowel and the eye. It is one of the rarer types of skin cancer but causes the majority of skin cancer related deaths. Around 160,000 new cases of melanoma are diagnosed worldwide each year, and it is more frequent in males and Caucasians, especially in Caucasian populations living in sunny climates than other groups. According to the WHO Report about 48,000 melanoma related deaths occur worldwide per annum. Despite many years of intensive laboratory and clinical research, the sole effective cure is surgical resection of the primary tumor before it achieves a thickness greater than 1mm (Wikipedia 2007). Therefore, early diagnose of Melanoma is critical to increase the change to cure the disease.

Dermatoscopy is a hand-held instrument with a dermoscope, a magnifier with a light and a liquid medium between the instrument and the skin, thus illuminating the skin without reflected light. Dermatoscopy is a noninvasive diagnostic technique for the early diagnosis of melanoma and the evaluation of other pigmented and non-pigmented lesions on the skin that are not as well seen with the unaided eye. Stolz et al. (1994) studied the accuracy of clinical evaluations with or without the aid of Dermatoscopy in detecting malignant Melanoma (MM) by using the ABCD rule (Asymmetry, irregular border, different colors, and Diameter larger than 6mm). In this study, two tests were used for detecting MM on the same subjects. The first test is the clinical assessment without the aid of dermoscopy, and the second test is the clinical assessment with the aid of dermoscopy. The data set we used here includes 21 patients with MM and 51 patients with benign melanocytic lesions. The goal is to find out whether the use of dermoscopy can improve for detecting MM. We estimate the difference between two sensitivities of the two tests and construct confidence intervals for the difference by using BCa,
BTI, BTII, HBELI and HBELII methods. The 95% confidence intervals for the difference between two sensitivities when the specificity is fixed at 0.9 or 0.95 respectively are shown in Appendix I Table V.

All the confidence intervals from above five methods contain zero. In summary, we conclude that there is no significant advantage in adopting the clinical assessment with the aid of dermatoscopy in detecting MM. The same conclusion has been obtained in Qin et al. (2006).
Chapter 6 Discussion

When a new method for continuous-scale tests is developed, comparing its effectiveness with existing methods is necessary. Using the confidence intervals for the difference between two sensitivities of two tests is straightforward. In many cases, only a particular value of specificity is relevant (e.g., 70%, 80%, 90%). Therefore, it is of interest to construct a confidence interval for the sensitivity of the test at a fixed level of specificity.

Qin et al. (2006) proposed three bootstrap-based intervals (BCa, BTI and BTII) for the difference between two sensitivities and showed that these intervals outperform the normal-approximation-based interval. In this thesis, we have proposed another two hybrid empirical likelihood and bootstrap confidence intervals (HBELI and HBELII) for the difference between two sensitivities. Simulation studies show that when correlation exists, HBELI and HBELII intervals perform better than the existing bootstrap based intervals (BCa, BTI and BTII) due to shorter interval length. However, when there is no correlation, BCa, BTI and BTII intervals outperform HBELI and HBELII intervals due to better coverage probability in most simulation settings.
References


Qin GS, Hsu YS, and Zhou XH (2006). New confidence intervals for the difference between two sensitivities at a fixed level of specificity. Statistics in Medicine, **25**: 3487-3502

Shapiro DE (1999). The interpretation of diagnostic tests. Statistical Methods in Medical Research, **8**: 113-134


Zhou XH, and Qin GS (2005). Improved confidence intervals for the sensitivity at a fixed level of specificity of a continuous-scale diagnostic test. Statistical in Medicine, **24**: 465-477.
Appendix I: Simulation tables

Table a-1 Level of 95 per cent confidence interval for $D(p)=0$. Bivariate normal distribution with $\rho = 0$

<table>
<thead>
<tr>
<th>Sample size</th>
<th>Method</th>
<th>Specificity=0.7</th>
<th>Specificity=0.8</th>
<th>Specificity=0.9</th>
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</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Coverage probability</td>
<td>Length</td>
<td>Coverage probability</td>
</tr>
<tr>
<td>(20,20)</td>
<td>BCa</td>
<td>0.8200</td>
<td>0.3864</td>
<td>0.9312</td>
</tr>
<tr>
<td></td>
<td>BTI</td>
<td>0.9030</td>
<td>0.4534</td>
<td>0.9557</td>
</tr>
<tr>
<td></td>
<td>BTII</td>
<td>0.9080</td>
<td>0.4534</td>
<td>0.9645</td>
</tr>
<tr>
<td></td>
<td>HBELI</td>
<td>0.8877</td>
<td>0.2591</td>
<td>0.8714</td>
</tr>
<tr>
<td></td>
<td>HBELII</td>
<td>0.8896</td>
<td>0.2591</td>
<td>0.8729</td>
</tr>
<tr>
<td>(50,50)</td>
<td>BCa</td>
<td>0.8905</td>
<td>0.2892</td>
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</tr>
<tr>
<td></td>
<td>BTI</td>
<td>0.9330</td>
<td>0.3191</td>
<td>0.9555</td>
</tr>
<tr>
<td></td>
<td>BTII</td>
<td>0.9380</td>
<td>0.3191</td>
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</tr>
<tr>
<td></td>
<td>HBELI</td>
<td>0.9191</td>
<td>0.2810</td>
<td>0.8714</td>
</tr>
<tr>
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<td>HBELII</td>
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<td>0.8729</td>
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<tr>
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<td>0.2434</td>
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</tr>
<tr>
<td></td>
<td>BTI</td>
<td>0.9430</td>
<td>0.2590</td>
<td>0.9280</td>
</tr>
<tr>
<td></td>
<td>BTII</td>
<td>0.9460</td>
<td>0.2590</td>
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</tr>
<tr>
<td></td>
<td>HBELI</td>
<td>0.9341</td>
<td>0.2490</td>
<td>0.9225</td>
</tr>
<tr>
<td></td>
<td>HBELII</td>
<td>0.9343</td>
<td>0.2490</td>
<td>0.9222</td>
</tr>
<tr>
<td>(150,150)</td>
<td>BCa</td>
<td>0.9505</td>
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</tr>
<tr>
<td></td>
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</tr>
<tr>
<td></td>
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<tr>
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<td>0.9312</td>
</tr>
<tr>
<td></td>
<td>HBELII</td>
<td>0.9413</td>
<td>0.1912</td>
<td>0.9308</td>
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</table>
Table a-1 Level of 95 per cent confidence interval for $D(p)=0$. Bivariate normal distribution with $\rho = 0$ (continued)

<table>
<thead>
<tr>
<th>Sample size</th>
<th>Method</th>
<th>Specificity=0.7</th>
<th></th>
<th>Specificity=0.8</th>
<th></th>
<th>Specificity=0.9</th>
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<td>Length</td>
<td>Coverage probability</td>
<td>Length</td>
<td>Coverage probability</td>
<td>Length</td>
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<td>(50,30)</td>
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<td>0.8635</td>
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<td>0.9360</td>
<td>0.3308</td>
<td>0.9418</td>
<td>0.3511</td>
</tr>
<tr>
<td></td>
<td>BTI</td>
<td>0.9110</td>
<td>0.3753</td>
<td>0.9561</td>
<td>0.3614</td>
<td>0.9603</td>
<td>0.3842</td>
</tr>
<tr>
<td></td>
<td>BTII</td>
<td>0.9220</td>
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<td>0.9667</td>
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<tr>
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<td>0.2866</td>
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<td>0.2590</td>
<td>0.8896</td>
<td>0.2035</td>
</tr>
<tr>
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<td>HBELII</td>
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<td>0.2866</td>
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<td>0.8899</td>
<td>0.2035</td>
</tr>
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<td>BCa</td>
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<td>0.2496</td>
<td>0.9371</td>
<td>0.2530</td>
<td>0.9425</td>
<td>0.2627</td>
</tr>
<tr>
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<td>BTI</td>
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<td>0.2636</td>
<td>0.9519</td>
<td>0.2675</td>
<td>0.9540</td>
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<tr>
<td></td>
<td>BTII</td>
<td>0.9594</td>
<td>0.2636</td>
<td>0.957</td>
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<td>0.9621</td>
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<td>HBELI</td>
<td>0.9329</td>
<td>0.2451</td>
<td>0.9273</td>
<td>0.2201</td>
<td>0.9366</td>
<td>0.1813</td>
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<tr>
<td></td>
<td>HBELII</td>
<td>0.9330</td>
<td>0.2451</td>
<td>0.928</td>
<td>0.2201</td>
<td>0.9365</td>
<td>0.1813</td>
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</tbody>
</table>
Table a-2 Level of 95 per cent confidence interval for D(p)=0. Bivariate normal distribution with $\rho = 0.5$

| Sample size | Method | Specificity=0.7 | | Specificity=0.8 | | Specificity=0.9 | |
|-------------|--------|-----------------|-----------------|-----------------|-----------------|-----------------|
|             |        | Coverage probability | Length | Coverage probability | Length | Coverage probability | Length |
| (20,20)     | BCa    | 0.9667           | 0.4437     | 0.9623           | 0.4493     | 0.9646           | 0.4649     |
|             | BTI    | 0.9180           | 0.3705     | 0.9193           | 0.3760     | 0.9212           | 0.3917     |
|             | BTII   | 0.9407           | 0.3705     | 0.9448           | 0.3760     | 0.9470           | 0.3917     |
|             | HBELI  | 0.9263           | 0.2848     | 0.9099           | 0.2554     | 0.9100           | 0.1909     |
|             | HBELII | 0.9271           | 0.2848     | 0.9094           | 0.2554     | 0.9113           | 0.1910     |
| (50,50)     | BCa    | 0.9687           | 0.3183     | 0.9687           | 0.3225     | 0.9688           | 0.3347     |
|             | BTI    | 0.9111           | 0.2479     | 0.9155           | 0.2525     | 0.9241           | 0.2647     |
|             | BTII   | 0.9294           | 0.2479     | 0.9336           | 0.2525     | 0.9332           | 0.2647     |
|             | HBELI  | 0.9538           | 0.2910     | 0.9425           | 0.2516     | 0.9413           | 0.1945     |
|             | HBELII | 0.9534           | 0.2910     | 0.9425           | 0.2516     | 0.9414           | 0.1946     |
| (80,80)     | BCa    | 0.9719           | 0.2598     | 0.9752           | 0.2647     | 0.9696           | 0.2743     |
|             | BTI    | 0.9114           | 0.1985     | 0.9122           | 0.2016     | 0.9115           | 0.2115     |
|             | BTII   | 0.9264           | 0.1985     | 0.9247           | 0.2016     | 0.9315           | 0.2115     |
|             | HBELI  | 0.9563           | 0.2556     | 0.9477           | 0.2266     | 0.9532           | 0.1789     |
|             | HBELII | 0.9564           | 0.2556     | 0.9481           | 0.2266     | 0.9535           | 0.1789     |
| (150,150)   | BCa    | 0.9733           | 0.1958     | 0.9738           | 0.1991     | 0.9714           | 0.2066     |
|             | BTI    | 0.9141           | 0.1460     | 0.9175           | 0.1485     | 0.9112           | 0.1558     |
|             | BTII   | 0.9247           | 0.1460     | 0.9267           | 0.1485     | 0.9254           | 0.1558     |
|             | HBELI  | 0.9714           | 0.1931     | 0.9563           | 0.1783     | 0.9534           | 0.1470     |
|             | HBELII | 0.9716           | 0.1931     | 0.9561           | 0.1783     | 0.9541           | 0.1470     |
Table a-2 Level of 95 per cent confidence interval for D(p)=0. Bivariate normal distribution with $\rho = 0.5$ (continued)

<table>
<thead>
<tr>
<th>Sample size</th>
<th>Method</th>
<th>Specificity=0.7</th>
<th>Specificity=0.8</th>
<th>Specificity=0.9</th>
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<tbody>
<tr>
<td></td>
<td></td>
<td>Coverage probability</td>
<td>Length</td>
<td>Coverage probability</td>
</tr>
<tr>
<td>(50,30)</td>
<td>BCa</td>
<td>0.9565</td>
<td>0.3589</td>
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<tr>
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<tr>
<td></td>
<td>HBELII</td>
<td>0.9393</td>
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<td>0.9345</td>
</tr>
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<td>(100,80)</td>
<td>BCa</td>
<td>0.9701</td>
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<td>0.9701</td>
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<td>HBELII</td>
<td>0.9606</td>
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</table>
Table a-3  Level of 95 per cent confidence interval for \(D(p)=0\). Bivariate exponential distribution with \(\rho = 0\)

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<th>Sample size</th>
<th>Method</th>
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<td></td>
<td>Coverage probability</td>
<td>Length</td>
<td>Coverage probability</td>
<td>Length</td>
</tr>
<tr>
<td>(20,20)</td>
<td>BCa</td>
<td>0.9323</td>
<td>0.4553</td>
<td>0.9311</td>
</tr>
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<td></td>
<td>BTI</td>
<td>0.9593</td>
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<td>0.9644</td>
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<td>BTII</td>
<td>0.9695</td>
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<td>0.9730</td>
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<td>0.8729</td>
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<td>HBELI</td>
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<td>HBELII</td>
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Table a- 3 Level of 95 per cent confidence interval for $D(p)=0$. Bivariate exponential distribution with $\rho = 0$ (continued)

<table>
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<tr>
<th>Sample size</th>
<th>Method</th>
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<th>Specificity=0.8</th>
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<th></th>
<th>Specificity=0.9</th>
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<td>Coverage probability</td>
<td>Length</td>
<td>Coverage probability</td>
<td>Length</td>
<td>Coverage probability</td>
<td>Length</td>
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</tr>
<tr>
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<td>0.2690</td>
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<td>0.2990</td>
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<td>HBELI</td>
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<td>0.2383</td>
<td>0.9182</td>
<td>0.1450</td>
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Table a- 4 Level of 95 per cent confidence interval for $D(p)=0$. Bivariate exponential distribution with $\rho \neq 0$ (Using 0.02 to generate diseased random sample)

<table>
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<tr>
<th>Sample size</th>
<th>Method</th>
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<th>Specificity=0.8</th>
<th>Specificity=0.9</th>
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<td>Coverage probability</td>
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<td>HBELII</td>
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<tr>
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<tr>
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Table a- 4 Level of 95 per cent confidence interval for $D(p)=0$. Bivariate exponential distribution with $\rho \neq 0$ (Using 0.02 to generate diseased random sample) (continued)

<table>
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<tr>
<th>Sample size</th>
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<th>Specificity=0.8</th>
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<td>Length</td>
<td>Coverage probability</td>
<td>Length</td>
<td>Coverage probability</td>
<td>Length</td>
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Table a- 5 95 per cent confidence interval for the difference of sensitivities between the two clinical assessments with and without the use of dermatoscopy

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<th>BTII</th>
<th>HBELI</th>
<th>HBELII</th>
</tr>
</thead>
<tbody>
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<td>0.90</td>
<td>(-0.261,0.261)</td>
<td>(-0.220,0.394)</td>
<td>(-0.302,0.312)</td>
<td>(-0.183,0.183)</td>
<td>(-0.183,0.183)</td>
</tr>
<tr>
<td>0.95</td>
<td>(-0.609,0.479)</td>
<td>(-0.346,0.346)</td>
<td>(-0.336,0.357)</td>
<td>(-0.052,0.052)</td>
<td>(-0.052,0.052)</td>
</tr>
</tbody>
</table>
Appendix II S-plus code for Simulation

1. Normal distribution

```r
# Functions
# Get sensitivity from abnorm and norm samples at fixed specificity p
# m: number of bootstrap
sensb<-function(abnorm, norm, p, m)
{
  result <- rep(NA, m)
  if(m > 1) {
    for(i in 1:m) {
      t <- sample(abnorm, length(abnorm), replace = T)
      u <- sample(norm, length(norm), replace = T)
      if(max(t) < min(u)) {
        result[i] <- 0
      } else {
        result[i] <- sum(t > quantile(u, p))/length(t)
      }
    }
  } else result[1] <- sum(abnorm > quantile(norm, p))/length(abnorm)
  return(result)
}
solveNonlinear<-function(f,y0,x)
{
  g<-function(x,y0,f) sum((f(x)-y0)^2)
  g$y0<-y0
  g$f<-f
  nlmin(g,x,max.fcal=100,max.iter=100)
}

# Main Program
mm<-1000   # number of repetition
m<-80    # sample sizes of non-diseased samples
n<-80

tt<-0.7   # Specificity level  tt
#tt<-0.8
#tt<-0.7
rho=0
alpha<-0.05
```
Cvalue<-qchisq(1-alpha,1)  #chi-sq(1-alpha,1)

# sensitivity 1 -sensitivity 2 =0
ss1<-c(0.95,0.90,0.80,0.70,0.60,0.50,0.40,0.30,0.20,0.10)  #R1(t) sensitivity 1
ss2<-c(0.95,0.90,0.80,0.70,0.60,0.50,0.40,0.30,0.20,0.10)  #R2(t) sensitivity 2

inrange1<-0  
logltzero<-0  #record the number which less than zero in LDp

Dp.low<-0
Dp.up<-0
nlow<-0  #if the function is converge, nlow+1
nup<-0

Dp2.low<-0
Dp2.up<-0
nlow2<-0  #indicator whether the function is converge, nlow2+1
nup2<-0

inrange2<-0

for (i in 1:length(ss1))
{
  cov1<-cov2<-0        # coverage
  for (j in 1:mm)
  {
    mud1<-qnorm(tt,0,1)-2*qnorm(1-ss1[i],0,1)   # mean of the first
    diseased population
    mud2<-qnorm(tt,0,1)-2*qnorm(1-ss2[i],0,1)   # mean of the second
    diseased population
    Rtt1<-1-pnorm(qnorm(tt,0,1),mud1,2)         #the first true sensitivity
    Rtt2<-1-pnorm(qnorm(tt,0,1),mud2,2)
    Rtt<-Rtt1-Rtt2                #the difference of two true sensitivities
  
  
  
  
  }  
}

# Generate diseased and non-diseased distribution
# Generate two samples from the nondiseased populations:
xx<-rmvnorm(m, mean=c(0,0), cov=matrix(c(1,rho,rho,1),2))
x10<-xx[,1] # the sample from the first nondiseased population
x20<-xx[,2] # the sample from the second nondiseased population

# generate two samples from the diseased populations:
yy<-rmvnorm(n, mean=c(mud1, mud2), ov=matrix(c(4,rho*2*2,rho*2*2,4),2))
# the sample from 2-dimensinal multinomial distribution with mean=c(mud1, mud2),sd=2, and correlation=0.5
y11<-yy[,1] # the sample from the first diseased population
y21<-yy[,2] # the sample from the second diseased population

# Two estimated sensitivities at specificity (tt):
sens1<-sum((yy[,1] >=quantile(xx[,1],tt)))/n    # estimated sensitivity
from the first sample
sens2<-sum(yy[,2] >= quantile(xx[,2],tt))/n  # estimated sensitivity
from the second sample

################################# step 2 #################################
# Bootstrap
# Generate diseased and non-diseased distribution
#################################

B=150

# get sensitivity from bootstrap samples
Rb1<-sensb(y11, x10, tt, B)
Rb2<-sensb(y21, x20, tt, B)

vb1<-sum((Rb1-mean(Rb1))^2)/(B-1)  # V_1^*(t)
vb2<-sum((Rb2-mean(Rb2))^2)/(B-1)

vb2<0
if (rho!=0)
  vb2<-sum( (Rb1-mean(Rb1))*(Rb2-mean(Rb2)) )/(B-1)

vb<-vb1+vb2-2*vb2  # Bootstrap variance estimate

################################# step 3 #################################
R1<0
R2<0
if (vb!=0)
  {  
    R1<-(mean(Rb1)*(1-mean(Rb1))+mean(Rb2)*(1-mean(Rb2)))/(n*vb)  # estimate
     for the scale constant
    R2<-( sens1*(1-sens1) + sens2*(1-sens2) )/(n*vb)

################################# Calculate L(D(p)) #################################
f1<-2
f2<-2

u11hat<-rep(100,n)
u22hat<-rep(100,n)

for(ii in 1:n)  # hat Uk=1-F(Yk)
  {
    u11hat[ii]<-1-mean(x10<=y11[ii])
    u22hat[ii]<-1-mean(x20<=y21[ii])
  }

v11hat<-(u11hat<=tt)*1  # indicator function of U:I(U_j<=p)
v22hat<-(u22hat<=tt)*1

############################ solve R_1(p), R_2(p) and lambda  ####################
g<function(x,v1h=v11hat, v2h=v22hat)
  {
    y_numeric(3)
    y[1].mean( (v11hat-x[1])/(1-2*x[3]*(v11hat-x[1])) )  
  }
\[ y[2]_\text{mean} \left( \frac{(v22hat-x[2])}{(1+2*x[3]*(v22hat-x[2]))} \right) \]
\[ y[3]_\text{mean} \left( \frac{v22hat/(1+2*x[3]*(v22hat-x[2])) - \text{mean}(v11hat/(1-2*x[3]*(v11hat-x[1])))}{v22hat/(1+2*x[3]*(v22hat-x[2]))) - \text{mean}(v11hat/(1-2*x[3]*(v11hat-x[1])))} \right) \]

```r
sol <- solveNonlinear(g, c(0, 0, Rtt), c(Rtt1, Rtt2, 0))
```

# c(Rtt1, Rtt2, 0) are initial values of c(R_1(p), R_2(p), lambda)

newr1 <- sol$x[1]
newr2 <- sol$x[2]
lambda <- sol$x[3]

w11hat <- v11hat - newr1
w22hat <- v22hat - newr2

###### test the number when (1-2*lambda*w11hat or 1+2*lambda*w22hat <0
flag <- 0
for(ii in 1:n) {
  if ((1-2*lambda*w11hat[ii]<0 || (1+2*lambda*w22hat[ii]<0)
    flag <- 1
}
if(flag = 1) logltzero <- logltzero + 1

LDp <- -2 * (sum(log(abs(1-2*lambda*w11hat))) + sum(log(abs(1+2*lambda*w22hat))))

###using abs here
# LDp <- -2 * (sum(log(1-2*lambda*w11hat)) + sum(log(1+2*lambda*w22hat)))

inrange1 <- inrange1 + (R1*LDp < qchisq(1-alpha,1))*1
inrange2 <- inrange2 + (R2*LDp < qchisq(1-alpha,1))*1

####### solove R_1(p), R_2(p), lambda, D(p) to find confidence interval of D(p) 04/24/2007#######

```r
f <- function(x, v1h = v11hat, v2h = v22hat, c = Cvalue) {
  y_numeric(4)
  y[1]_\text{mean} \left( \frac{(v11hat-x[1])}{(1-2*x[3]*(v11hat-x[1]))} \right) \)
  y[2]_\text{mean} \left( \frac{(v22hat-x[2])}{(1+2*x[3]*(v22hat-x[2]))} \right)
  y[3]_\text{mean} \left( \frac{v22hat/(1+2*x[3]*(v22hat-x[2])) - \text{mean}(v11hat/(1-2*x[3]*(v11hat-x[1])))}{v22hat/(1+2*x[3]*(v22hat-x[2]))) - \text{mean}(v11hat/(1-2*x[3]*(v11hat-x[1])))} \right) \)
  y[4]_R1*2*\left( \frac{\sum(\log(\text{abs}(1-2*x[3]*(v11hat-x[1])))\} \right) + \sum(\log(\text{abs}(1+2*x[3]*(v22hat-x[2])))\} \right) - Cvalue
  y
}
```}
solf1 <- solveNonlinear(f, c(0, 0, 0, 0), c((Rtt1+0.1), (Rtt2-0.1), 0, 0.2)) # initial values
if(solf1$converged = T) {
  nlow <- nlow + 1
  Dp.low <- Dp.low + solf1$x[4]
}
solf2 <- solveNonlinear(f, c(0, 0, 0, 0), c((Rtt1-0.1), (Rtt2+0.1), 0, -0.2))
if(solf2$converged = T) {
nup<-nup+1
Dp.up<-Dp.up+solf2$x[4]}

#######solve R_1(p), R_2(p), lambda, D(p) to find confidence interval of D(p) by using R2
04/24/2007###########

f<-function(x,v1h=v11hat, v2h=v22hat c=Cvalue)
{
y_numeric(4)
y[1]_mean( (v11hat-x[1])/(1-2*x[3]*(v11hat-x[1])) )
y[2]_mean( (v22hat-x[2])/(1+2*x[3]*(v22hat-x[2])) )
y[3]_mean( v22hat/(1+2*x[3]*(v22hat-x[2]))) - mean(v11hat/(1-2*x[3]*(v11hat-x[1])))-x[4]
y[4]_R2*2*( sum(log(abs(1-2*x[3]*(v11hat-x[1]))))+sum(log(abs(1+2*x[3]*(v22hat-x[2])))))-Cvalue
}
solf3<-solveNonlinear(f,c(0,0,0,0),c((Rtt1+0.1),(Rtt2-0.1),0,0.2)) #initial values
if(solf3$converged = T)
{
  nlow2<-nlow2+1
  Dp2.low<-Dp.low+solf1$x[4]
}solf4<-solveNonlinear(f,c(0,0,0,0),c((Rtt1-0.1),(Rtt2+0.1),0,-0.2))
if(solf4$converged = T)
{
nup2<-nup2+1
  Dp2.up<-Dp.up+solf2$x[4]
}

} #end of if(vb!=0)

} #end of loop for (j in 1:mm)
} #end of loop for (i in 1:length(ss1))

newcov1<-inrange1/(10*mm)
newcov2<-inrange2/(10*mm)
if(nlow & nup)
{
  Dplow<-min(Dp.low/nlow,Dp.up/nup)
  Dpup<-max(Dp.low/nlow,Dp.up/nup)
}
Dplength<-max(Dpup,Dplow)-min(Dpup,Dplow)
if(nlow2 & nup2)
{
  Dp2low<-min(Dp2.low/nlow2, Dp2.up/nup2)
  Dp2up<-max(Dp2.low/nlow2, Dp2.up/nup2)
}
Dplength2<-max(Dp2up,Dp2low)-min(Dp2up, Dp2low)

#Result Output
sink("D:\Suqin\normalresult1.txt",append = T)

cat("########################################################################");
cat(" specificity=",tt, ",n")
cat(" rho=",rho, ",n")
cat(" Non-disease sample m=", m, " disease sample n=", n, "iteration mm=", mm, ",n")
cat(" Number of log <0 ",logltzero,"n\n");
cat(" Coverage1=", newcov1,"\n");
cat(" Dp  Lower bound 1=", Dplow, " Up bound 1=", Dpup, ",n")
cat(" Coverage length 1 =", Dplength,"n")
cat(" Number of converge nlow1=", nlow, " nup1=", nup,"n\n")

cat(" Coverage2=", newcov2,"\n");
cat(" Dp  Lower bound 2=", Dp2low, " Up bound 2=", Dp2up, ",n")
cat(" Coverage length 2 =", Dplength2,"n")
cat(" Number of converge nlow2=", nlow2, " nup2=", nup2,"n")

cat("########################################################################");
sink();
2. Exponential distribution – no correlation

####### Main Program #######

mm<-1000   # number of repetition
m<-150    # sample sizes of non-diseased samples
n<-80        # sample sizes of diseased samples

## Specificity level tt
#tt<-0.8
#tt<-0.7

rho=0

alpha<-0.05
Cvalue<-qchisq(1-alpha,1)  #chi-sq(1-alpha,1)

# sensitivity 1 - sensitivity 2 =0
ss1<-c(0.95,0.90,0.80,0.70,0.60,0.50,0.40,0.30,0.20,0.10)  #R1(t) sensitivity 1
ss2<-c(0.95,0.90,0.80,0.70,0.60,0.50,0.40,0.30,0.20,0.10)  #R2(t) sensitivity 2

inrange1<-0
logltzero<-0  #record the number which less than zero in LDp

Dp.low<-0
Dp.up<-0
nlow<-0   #if the function is converage, nlow+1
nup<-0

Dp2.low<-0
Dp2.up<-0
nlow2<-0   #idicator whether the function is converage, nlow+1
nup2<-0

inrange2<-0

for (i in 1:length(ss1))
{
        cov1<-cov2<-0        # coverage
        l1<-log(ss1[i])/log(1-tt) # rate of the Exp(l1) (first diseased
                                # population) (rate=1/expectation)
        l2<-log(ss2[i])/log(1-tt) # rate of the Exp(l2) (second diseased
                                # population)
        Rtt1<-exp(l1*log(1-tt))
        Rtt2<-exp(l2*log(1-tt))
        Rtt<-Rtt1-Rtt2  # the difference of two true sensitivities

        for (j in 1:mm)
        {
                # mud1<-qnorm(tt,0,1)-2*qnorm(1-ss1[i],0,1)  # mean of the first diseased
                # population
                # mud2<-qnorm(tt,0,1)-2*qnorm(1-ss2[i],0,1)  # mean of the second diseased
                # population
        }
\# Rtt1<-1-pnorm(qnorm(tt,0,1),mud1,2)          \# the first true sensitivity
\# Rtt2<-1-pnorm(qnorm(tt,0,1),mud2,2)
\# Rtt<-Rtt1-Rtt2                             \# the difference of two true sensitivities

# Exponential distribution
# two dependent samples from the nondiseased populations:
x10<-rexp(m,1)   \# Exp(1): the sample from the first nondiseased population
x20<-rexp(m,1)   \# Exp(1): the sample from the second nondiseased population

# two dependent samples from the diseased populations:
y11<-rexp(n,l1)   \# Exp(l1): the sample from the first diseased population
y21<-rexp(n,l2)   \# Exp(l2): the sample from the second diseased population

sens1<-sum((y11 >=quantile(x10,tt)))/n    \# estimated sensitivity from the first sample
sens2<-sum((y21 >=quantile(x20,tt)))/n    \# estimated sensitivity from the second sample

# Bootstrap
# Generate diseased and non-diseased distribution
B=150
# get sensitivity from bootstrap samples
Rb1<-sensb(y11, x10, tt, B)
Rb2<-sensb(y21, x20, tt, B)

vb1<-sum((Rb1-mean(Rb1))^2)/(B-1)                              \# V_1^*(t)
vb2<-sum((Rb2-mean(Rb2))^2)/(B-1)
vb12<-0
if (rho!=0)
  vb12<-sum( (Rb1-mean(Rb1))*(Rb2-mean(Rb2) ) )/(B-1)
vb<-vb1+vb2-2*vb12     \# Bootstrap variance estimate

# Calculate L(D(p))
f1<-2
f2<-2

ullhat<-rep(100,n)
u22hat<-rep(100,n)
for(ii in 1:n)    # hat Uk=1-F(Yk)
{
  u11hat[ii]<-1-mean(x10<=y11[ii])
  u22hat[ii]<-1-mean(x20<=y21[ii])
}

v11hat<-u11hat<=tt)*1    # indicator function of U:I(U_j<=p)
v22hat<-u22hat<=tt)*1

########solve R_1(p), R_2(p) and lambda ###########

g<-function(x,v1h=v11hat, v2h=v22hat)
{
  y_numeric(3)
  y[1]_mean( (v11hat-x[1])/(1-2*x[3]*(v11hat-x[1])) )
  y[2]_mean( (v22hat-x[2])/(1+2*x[3]*(v22hat-x[2])) )
  y[3]_mean( v22hat/(1+2*x[3]*(v22hat-x[2]))) - mean( v11hat/(1-2*x[3]*(v11hat-x[1])))
  y
}
sol<-solveNonlinear(g,c(0,0,Rtt),c(Rtt1,Rtt2,0))  
  # c(Rtt1,Rtt2,0) are initial values of c(R_1(p), R_2(p), lambda)

newr1<-sol$x[1]
newr2<-sol$x[2]
lambda<-sol$x[3]

w11hat<-v11hat-newr1
w22hat<-v22hat-newr2

###### test the number when (1-2*lambda*w11hat or 1+2*lambda*w22hat < 0
flag<0
for(ii in 1:n)   {
  if ((1-2*lambda*w11hat[ii])<0 || (1+2*lambda*w22hat[ii])<0)
    flag<-1
}
if(flag=1) logltzero<-logltzero+1

LDp<-2*( sum(log(abs(1-2*lambda*w11hat))) + sum(log(abs(1+2*lambda*w22hat))) )
  # using abs here
  # LDp<-2*( sum(log(1-2*lambda*w11hat)) + sum(log(1+2*lambda*w22hat)))
inrange1<-inrange1 + (R1*LDp<qchisq(1-alpha,1))*1
inrange2<-inrange2 + (R2*LDp<qchisq(1-alpha,1))*1

#######solve R_1(p), R_2(p), lambda, D(p) to find confidence interval of D(p)
  04/24/2007###########

f<-function(x,v1h=v11hat, v2h=v22hat c=Cvalue)
{
  y_numeric(4)
  y[1]_mean( (v11hat-x[1])/(1-2*x[3]*(v11hat-x[1])) )
  y[2]_mean( (v22hat-x[2])/(1+2*x[3]*(v22hat-x[2])) )
  y[3]_mean( v22hat/(1+2*x[3]*(v22hat-x[2]))) - mean(v11hat/(1-2*x[3]*(v11hat-x[1])))
    -x[4]
  y[4]_R1*2*( sum(log(abs(1-2*x[3]*(v11hat-x[1]))) + sum(log(abs(1+2*x[3]*(v22hat-x[2])))
    -Cvalue
\[ y \]

solf1 <- solveNonlinear(f, c(0, 0, 0, 0), c((Rtt1+0.1), (Rtt2-0.1), 0, 0.2)) # initial values
if(solf1$converged = T)
{
  nlow <- nlow + 1
  Dp.low <- Dp.low + solf1$x[4]
}
solf2 <- solveNonlinear(f, c(0, 0, 0, 0), c((Rtt1-0.1), (Rtt2+0.1), 0, -0.2))
if(solf2$converged = T)
{
  nup <- nup + 1
  Dp.up <- Dp.up + solf2$x[4]
}

######## solve R_1(p), R_2(p), lambda, D(p) to find confidence interval of D(p) by using R2 04/24/2007###########

f <- function(x, v1h = v11hat, v2h = v22hat, c = Cvalue)
{
  y_numeric(4)
  y[1]_mean((v11hat - x[1]) / (1 - 2 * x[3] * (v11hat - x[1])))
  y[2]_mean((v22hat - x[2]) / (1 + 2 * x[3] * (v22hat - x[2])))

  y
}

solf3 <- solveNonlinear(f, c(0, 0, 0, 0), c((Rtt1+0.1), (Rtt2-0.1), 0, 0.2)) # initial values
if(solf3$converged = T)
{
  nlow2 <- nlow2 + 1
  Dp2.low <- Dp2.low + solf3$x[4]
}
solf4 <- solveNonlinear(f, c(0, 0, 0, 0), c((Rtt1-0.1), (Rtt2+0.1), 0, -0.2))
if(solf4$converged = T)
{
  nup2 <- nup2 + 1
  Dp2.up <- Dp2.up + solf4$x[4]
}

} # end of if(vb!=0)

} # end of loop for (j in 1:mm)

} # end of loop for (i in 1:length(ss1))

newcov1 <- inrange1/(10*mm)
newcov2 <- inrange2/(10*mm)
if(nlow & nup)
{
  Dp.low <- min(Dp.low/nlow, Dp.up/nup)
Dpup<-max(Dp.low/nlow, Dp.up/nup)
}

#Dplow<-Dp.low/(10*mm)
#Dpup<-Dp.up/(10*mm)
Dlength<-max(Dpup, Dplow) - min(Dpup, Dplow)

if(nlow2 & nup2)
{
  Dp2low<-min(Dp2.low/nlow2, Dp2.up/nup2)
  Dp2up<-max(Dp2.low/nlow2, Dp2.up/nup2)
}

Dlength2<-max(Dp2up, Dp2low) - min(Dp2up, Dp2low)

#Result Output
sink("D:\Suqin\Expind_rhoeq0.txt", append = T)

cat("#######Exponential Distribution

specification=", tt, "
"");
cat(" rho=", rho, "
"");
cat(" Non-disease sample m=", m, " disease sample n=", n, "iteration mm=", mm, "
"");
cat(" Number of log <0 ", logltzero, "
"");
cat(" Coverage1=", newcov1, "
"");
cat(" Coverage length 1 =", Dlength, "
"");
cat(" Number of converge nlow1=", nlow, " nup1=", nup, "
"");
cat(" Coverage2=", newcov2, "
"");
cat(" Coverage length 2 =", Dlength2, "
"");
cat(" Number of converge nlow2=", nlow2, " nup2=", nup2, "
"");
cat("#####################################");
sink();
3. Exponential distribution – correlation

...(Specificity and sample size setting are the same as exponential distribution without correlation)...

expcov<-0.02

l1<-log(ss1[i])/log(1-tt)-expcov  # rate of the Exp(l1) (first diseased population) (rate=1/expectation)
l2<-log(ss2[i])/log(1-tt)-expcov  # rate of the Exp(l2) (second diseased population)

Rtt1<-exp((l1+expcov)*log(1-tt))  # the difference of two true sensitivities
Rtt2<-exp((l2+expcov)*log(1-tt))
Rtt<-Rtt1-Rtt2

mnb1<-mnb2<-mnb3<-0
LUb1<-LUb3<-0

for (j in 1:mm)
{

explambda<-0.5
u1<-rexp(m,explambda)
u2<-rexp(m,explambda)
u3<-rexp(m,explambda)

# two dependent samples from the nondiseased populations:
x10<-x20<-0
for (k in 1:m)
{
x10[k]<-min(u1[k],u3[k])  # Exp(1): the sample from the first nondiseased population
x20[k]<-min(u2[k],u3[k])  # Exp(1): the sample from the second nondiseased population
}

# two dependent samples from the diseased populations:
v1<-rexp(n,l1)
v2<-rexp(n,l2)
v3<-rexp(n,expcov)
y11<-y21<-0
for (k in 1:n)
{
y11[k]<-min(v1[k],v3[k])  # Exp(l1+0.01): the sample from the first diseased population
y21[k]<-min(v2[k],v3[k])  # Exp(l2+0.01): the sample from the second diseased population
}

# Two estimated sensitivities at specificity (tt):

X1.hat<-sum((y11 >=quantile(x10,tt)))  # estimated sensitivity from the first sample
sens1<-X1.hat/n

X2.hat<-sum((y21 >=quantile(x20,tt)))  # estimated sensitivity from the second sample
sens2<-X2.hat/n
…(Same as exponential distribution without correlation)...

# Result Output

sink("D:\Suqin\Exponential_Rhogt0_result.txt", append = T)

cat("#####Exponential Distribution \\

" specificity="tt", "\n")
cat("Exponential lambda="explambda," expcov="expcov," \n")
cat(" rho>0","inlambda="inlambda, "\n")
cat(" Non-disease sample m="m, " disease sample n="n, " iteration mm="mm, "\n")
cat(" Number of log <0 ", logltzero, "\n")


cat(" Coverage1=" newcov1,"\n")
cat(" Dp  Lower bound 1=" Dplow, " Up bound 1=" Dpup, "\n")
cat(" Coverage length 1 =", Dplength,"\n")
cat(" Number of converge nlow1=" nlow," nup1=" nup, "\n")

cat(" Coverage2=" newcov2,"\n")
cat(" Dp  Lower bound 2=" Dp2low, " Up bound 2=" Dp2up, "\n")
cat(" Coverage length 2 =", Dplength2,"\n")
cat(" Number of converge nlow2=" nlow2," nup2=" nup2,"\n")

cat("####################################");

sink();
4. Dermatoscope example

mm<-1000   # number of repetition

tt<-0.9
#tt<-0.95

realdataload.table("D:\Suqin\exam4.SSC",header=F,skip=3)
##get non-disease nx and disease number ny;
nx<-0;
for(i in 1:72) if(realdata[i,4]==0) nx<-nx+1;
y<-0;
for(i in 1:72) if(realdata[i,4]==1) ny<-ny+1;
m<-nx; #assign to non-disease group
n<--ny; #assign to disease group

xx<-matrix(0,nx,2)
yy<-matrix(0,ny,2)

xi<-1;
yi<-1;
for (r in 1:72)
{
    if (realdata[r,4]==0) { xx[xi,1]<-realdataloaldata[r,2];xx[xi,2]<-realdataloaldata[r,3]; xi<-xi+1}
    else { yy[yi,1]<-realdataloaldata[r,2]; yy[yi,2]<-realdataloaldata[r,3];yi<-yi+1;}
}

x10<-xx[,1]  # the sample from the first nondisease population
x20<-xx[,2]  # the sample from the second nondisease population

y11<-yy[,1]  # the sample from the first diseased population
y21<-yy[,2]

{The following part is the same as normal distribution}