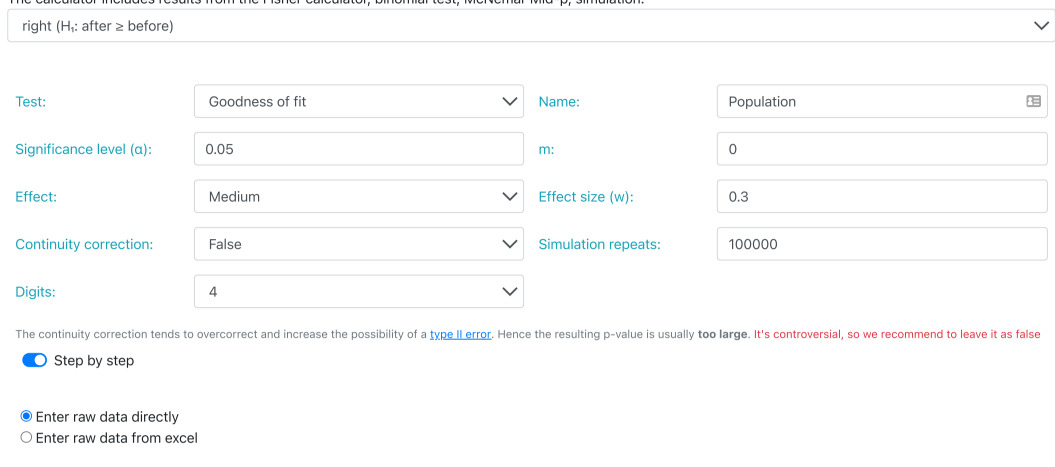
Chi-Square test calculator

Goodness of fit test, Test of independence, McNemar test

Video Information Chi-squared test for variance Chi-Square Calculator

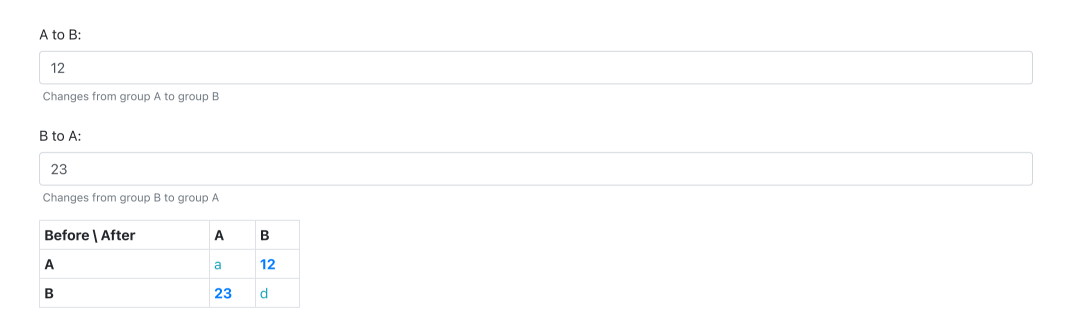
Test calculation

Right-tailed - for the goodness of fit test, the test of independence / the test for association, or the McNemar test, you can use only the right tail test. The calculator includes results from the Fisher calculator, binomial test, McNemar Mid-p, simulation.



McNemar chi-square test

Matched pairs



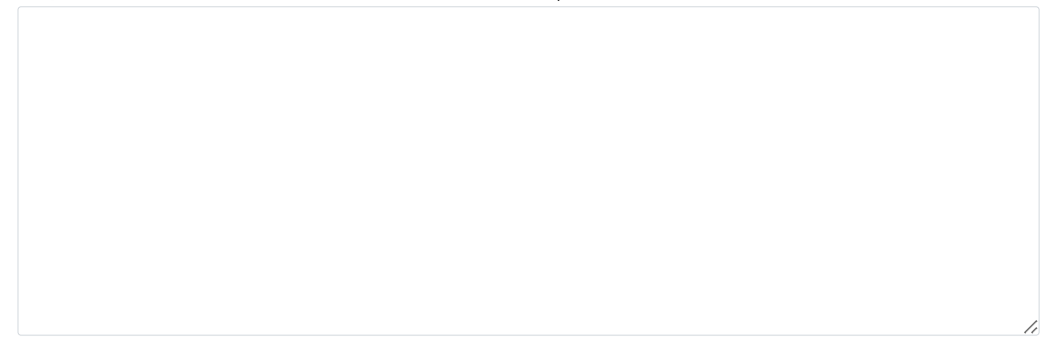
Enter sample data

Categories //	Assertive //	Non-Assertive
Novice	322155531513415:	415212122524252:
Expert	135225524535243	513533324121351:

Enter sample data

You may copy data from Excel, Google sheets or any tool that separate data with **Tab** and **Line Feed**.

Copy the data, **one block** of **3** consecutive columns includes the top **header row** and left **header column**, and paste below. <u>example</u>



It is okay to leave empty cells, empty cells or non numeric cells won't be counted

*Choose Expected Frequencies or Expected Probabilities.

The sum of the **expected values** must be equal to the sum of the observed frequencies or to one.



How to do with R?

Results

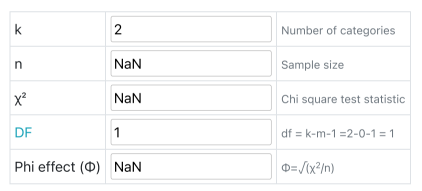
19/07/2022, 20:22

The expected values should be at least 1 for all the values, and at least 5 for 80% of the values Blue - less than 5, but still okay.

Red - potential issue.

$$\chi^2 = \frac{(NaN-NaN)^2}{NaN} + \frac{(NaN-NaN)^2}{NaN} = NaN$$

P-value = 1 - p($\chi^2(1) \le NaN$).









300dness of fit, using χ² distribution (right-tailed) (validation)

<u>. H₀ hypothesis</u>

Since p-value $< \alpha$, H_0 is rejected.

he statistical model does not fit the observations

<u>!. P-value</u>

The p-value equals NaN, ($p(x \le \chi^2) = NaN$). It means that the chance of type I error (rejecting a correct H₀) is small: NaN (NaN%). The smaller the p-value the more it supports H₁.

3. The statistics

The test statistic χ^2 equals NaN, which is not in the 95% region of acceptance: [- ∞ : 3.8415].

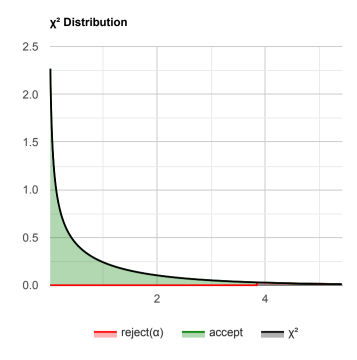
I. Effect size

The observed effect size phi is large, NaN. This indicates that the magnitude of the difference between the observed data and the expected data is large.

f you like the page, please share or like. Questions, comments and suggestions are appreciated. (statskingdom@gmail.com)

Like 322

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More tests

What test should I choose?

Simulation test

The simulation run 100,000 times, the duration is 0.05 seconds. Running more times will have better accuracy, but the default should be sufficient. Since the simulation is random, each run will have a slightly different result.

P-value

P-value equals 0. It means that the chance of type1 error (rejecting a correct H₀) is small: 0 (0%)

The smaller the p-value, the more it supports H₁.

Binomial exact test

The tool calculates the binomial exact test only in the case of two categories.

The two-tailed binomial exact test is equivalent to the one-tailed chi-squared test.

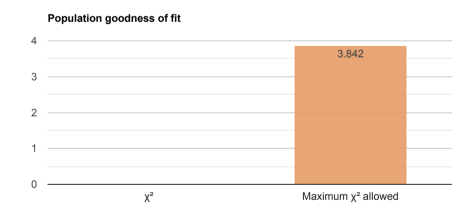
x = 3 2 2 1 5 5 5 3 1 5 1 3 4 1 5 2 5 1 4 2 4 2 1 4 5 2 5 2 5 1 2 1 5 5 2 5, using Binomial(NaN,NaN), two-tailed.

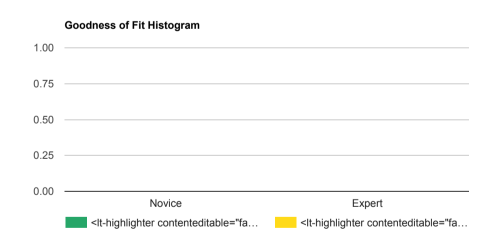
P-value

P-value equals 0. It means that the chance of type1 error (rejecting a correct H_0) is small: 0 (0%)

The smaller the p-value, the more it supports H_1 .

The right-tailed goodness of fit test is equivalent to the two-tailed binomial exact test.





Test validation

The requested test was calculated, it is likely you chose the right test.

• Test power +

Although the priori power is low (NaN), the H_0 is rejected.

Information

Hypotheses

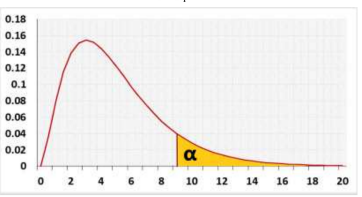
Test statistic

 χ^2 distribution

H₀: Model Fits

 $\chi^{2} = \sum_{i=1}^{n} \frac{(o_{i} - e_{i})^{2}}{e_{i}}$

H₁: Model Doesn't Fit



Target: Check if the statistical model fits the observations.

The test uses Chi-square distribution.

McNemar chi-square test

The test checks only the cases when the status of the dichotomous variable was changed.

The null assumption is that the probability to switch from A to B equals the probability to switch from B to A, equals 0.5.

Before \ After	A	В
Α	No change	A to B
В	B to A	No change

The number of cases with no change, A to A or B to B, doesn't change the result of the McNemar test.

Chi-square test for independence

The null assumption is that the two categorical variables are independent.

R Code

The following R code should produce the same results:

obs<-c(3 2 2 1 5 5 5 3 1 5 1 3 4 1 5 2 5 1 4 2 4 2 1 4 5 2 5 2 5 1 2 1 5 5 2 5,1 3 5 2 2 5 5 2 4 5 3 5 2 4 3 2 1 5 2 3 1 3 4 3 5 3 2 4 3 4 3 2 3 2 2 1 2 5 2)

prob<-c(4 1 5 2 1 2 1 2 2 5 2 4 2 5 2 2 3 1 2 5 2 1 2 2 3 2 1 2 5 2 1 2 1,5 1 3 5 3 3 3 2 4 1 2 1 3 5 1 2 1 4 1 5 2 5 1 2 5 3 5 1 2 1 5 3 2 1 2 4 1 5 1 5 2 5 2 5 1 2 2 5)

prob = prob / sum(prob)

chisq.test(x=obs, y=NULL, correct = FALSE, p=prob)

#Compute p-values by Monte Carlo simulation

chisq.test(x=obs, y=NULL, correct=FALSE, p=prob, simulate.p.value = TRUE, B = 100000)

binom.test(x=3 2 2 1 5 5 5 3 1 5 1 3 4 1 5 2 5 1 4 2 4 2 1 4 5 2 5 2 5 1 2 1 5 5 2 5, n=NaN, p=NaN, alternative='two.sided')

Goodness of fit example: checking a fair dice.

Model: the probability of each side is equal - 1/6.

H₀: fair dice.

 H_1 : unfair dice.

The **groups** are the dice's numbers (1,2,3,4,5,6).

In this example, you throw the dice n times.

Expected frequencies - for each group are n/6.

Observed frequencies - the actual times each number appears.