Estimation of sample size and testing power (Part 7)

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ABSTRACT: Two-factor factorial design refers to the research involving two experimental factors and the number of the experimental groups equals to the product of the levels of the two experimental factors. In other words, it is the complete combination of the levels of the two experimental factors. The research subjects are randomly divided into the experimental groups. The two experimental factors are performed on the subjects at the same time, meaning that there is no order. The two experimental factors are equal during statistical analysis, that is to say, there is no primary or secondary distinction, nor nested relation. This article introduces estimation of sample size and testing power of quantitative data with two-factor factorial design.

KEYWORDS: statistics, medical; research design; sample size; testing power; parametric estimation

Two-factor factorial design is frequently adopted when the research involves two experimental factors, which are performed on the research subjects at the same time and are equal, and have level 1 interaction which is required to be examined. When there is only one quantitative index, it is called univariate quantitative data with two-factor factorial design, which can be analyzed by the analysis of variance of univariate quantitative data with two-factor factorial design (1). This article introduces the estimation of sample size and testing power of univariate quantitative data with two-factor factorial design. Since the estimation of sample size and testing power of two-factor factorial design is complicated, this article will introduce it through examples when performing the analysis of variance by SAS.

1 Estimation of sample size

1.1 Example 1 A researcher planned to conduct an animal experiment to study the influence of copper (Cu) and vitamin E (VE) on liver injury caused by carbon tetrachloride (CCl4). Thirty healthy male Wistar rats were chosen for pilot test and were equally divided into 10 groups. How many rats were needed to perform the analysis of variance of two-factor factorial design in the formal experiment if the testing power was required to be 80% (1)?

1.2 Analysis Step 1: Based on the information of the pilot test, estimate the population means and the population standard deviations of superoxide
dismutase (SOD) under all the experimental conditions. The corresponding SAS program named exam 1_2.sas is as follows:

```
data step1;
do Cu=0.00,0.05,0.10,0.20,0.40;
do VE=0,150;
do i=1 to 3;
input SOD@@; output;
end;end;end;
cards;
343.8 331.6 318.1 465.2 457.7 449.5 390.0 437.0 419.3 584.5 570.0 576.7 448.1 413.1 454.4 604.3 531.4 605.0 443.7 422.3 468.7 485.6 516.4 485.9 474.6 455.5 464.4 509.8 552.0 498.6
;
run;
ods html;
proc means mean std;
var SOD;
class Cu VE;
run;
ods html close;
```

Program explanation: First, create a new data set named step1; then invoke the MEANS procedure to estimate the means and the standard deviations. Below is the main output:

**Response Variable: SOD**

<table>
<thead>
<tr>
<th>Cu</th>
<th>VE</th>
<th>Sample size</th>
<th>Mean</th>
<th>Standard deviation</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>0</td>
<td>3</td>
<td>331.166 666 7</td>
<td>12.855 478 7</td>
</tr>
<tr>
<td>150</td>
<td>3</td>
<td>457.466 666 7</td>
<td>7.852 600 4</td>
<td></td>
</tr>
<tr>
<td>0.05</td>
<td>0</td>
<td>3</td>
<td>422.100 000 0</td>
<td>6.937 578 8</td>
</tr>
<tr>
<td>150</td>
<td>3</td>
<td>577.066 666 7</td>
<td>7.256 950 7</td>
<td></td>
</tr>
<tr>
<td>0.1</td>
<td>0</td>
<td>3</td>
<td>438.533 333 3</td>
<td>22.250 018 7</td>
</tr>
<tr>
<td>150</td>
<td>3</td>
<td>580.233 333 3</td>
<td>42.292 355 5</td>
<td></td>
</tr>
<tr>
<td>0.2</td>
<td>0</td>
<td>3</td>
<td>441.566 666 7</td>
<td>28.260 455 3</td>
</tr>
<tr>
<td>150</td>
<td>3</td>
<td>495.966 666 7</td>
<td>17.696 421 5</td>
<td></td>
</tr>
<tr>
<td>0.4</td>
<td>0</td>
<td>3</td>
<td>464.833 333 3</td>
<td>9.557 370 6</td>
</tr>
<tr>
<td>150</td>
<td>3</td>
<td>520.133 333 3</td>
<td>28.159 782 2</td>
<td></td>
</tr>
</tbody>
</table>

Step 2: Estimate the sample size based on the result of step 1. The needed SAS program named exam 1_2.sas is as follows:

```
data step2;
do Cu=0.00,0.05,0.10,0.20,0.40;
do VE=0,150;
input mean @ @ output;
end;end;
cards;
331.166667 457.466667 422.100000 577.066667 438.533333 380.233333 441.566667 495.966667 464.833333 520.133333
;
run;
ods html;
proc glmpower data=step2;
class Cu VE;
model mean=Cu|VE;
power stddev=6 to 42 by 12 ntotal=.
power=0.80;
r;quit;
ods html close;
```

Program explanation: First, specify the population means under all the experimental conditions. The "glmpower" procedure performs prospective power analysis. The statement "model mean=Cu|VE" can also be written as "model mean=Cu VE Cu * VE". The option "stddev=6 to 42 by 12" specifies 4 population standard deviations, namely, 6, 18, 30 and 42 since step 1 has already computed the minimum standard deviation as 6.9, and the maximum as 42.3. The option "power=0.80" designates 0.8 as the testing power. The above parameter values can be altered under specific situations.

The main output is as follows:

**Fixed Scenario Elements**

<table>
<thead>
<tr>
<th>Dependent variable</th>
<th>Mean</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nominal Power</td>
<td>0.8</td>
</tr>
<tr>
<td>Alpha</td>
<td>0.05</td>
</tr>
</tbody>
</table>

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The above result presents the needed sample sizes with different standard deviations. When the population standard deviation is 42, the testing power to infer that Cu has a protective effect on liver injury caused by CCl₄ is 81.2%, and the testing power to infer that VE has a protective effect on liver injury caused by CCl₄ is 99.9% as long as the sample size reaches 20. When the sample size is 60, the testing power to infer that there is interaction between Cu and VE is 87.7%. Therefore, when the population standard deviation is 42, 60 rats are needed; that is to say, each group needs 6 rats at least since there are 10 groups.

2 Estimation of testing power

2.1 Example 2 In example 1, suppose that the researcher increased the sample size to 60 and conducted the additional experiments. The researcher wondered whether the testing power was sufficient if adopting the analysis of variance of univariate quantitative data with two-factor factorial design[1].

2.2 Analysis Step 1, based on the information of the pilot test, estimate the population means and the population standard deviations of SOD under all the experimental conditions. See program exam 1.1. sas for reference. The main output is as follows:

The above result shows that when the sample size is 60, the testing power to infer that Cu has a protective effect on liver injury caused by CCl₄ reaches 99.9% and 99.9% (S = 9.6 and S = 45.7), and the testing power to infer that VE has a protective effect on liver injury caused by CCl₄ reaches 99.9% and 99.9% (S = 9.6 and S = 45.7). When the population standard deviation is 9.6, the testing power to infer that there is interaction between Cu and VE reaches 99.9%. When the population standard deviation is 45.7, the testing power reaches 93.65%.
样本量估计与检验效能分析（七）

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摘要：所谓两因素析因设计，是指试验中涉及两个试验因素，不同的试验条件由与两个试验因素的水平数的乘积，即由试验因素的水平全面组合而成。全部受试对象被完全随机地分配到各试验条件组中去。试验时，全部试验因素同时施加（即无先后顺序之分）；对资料进行统计分析时，试验因素位置平等（即不存在主次之分，也不存在嵌套关系）。本文向读者介绍采用两因素析因设计一元定量资料方差分析处理定量资料时，在试验之前如何估计样本含量与检验效能。

关键词：统计学，医学，研究设计；样本大小；检验效能；参数估计