Combinatorial Designs to Explore Large Experimental Search Spaces *

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Abstract

Genomic and proteomic studies take advantage of omic level techniques such as microarrays to achieve species-wide scale. Obtaining an operational model (e.g., a virtual animal) however requires a search in a space consisting of many factors and many values of each factor, all of which may interact. This suggests the need to perform a disciplined search in that space. We explain the use of and some new results in covering arrays, a technique from the theory of combinatorial designs, and explain how to use covering arrays at various stages of experiment and analysis.

1 Introduction:

Your favorite organism can be grown under several conditions by varying light, food, water, and various nutrients. Several natural questions suggest themselves:

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• Which conditions give rise to the best growth, either of the organism as a whole or of some protein of interest?

• Which factors are the most critical?

• Which genes or proteins react most strongly to some factor or collection of factors?

Given unlimited resources and time, you would want to test all possible conditions by testing all values of every factor. Unfortunately, this very soon becomes a daunting task. For example, 10 inputs each having four possible values generates somewhat more than one million experiments. This may be more than most labs can or want to do. Further many of these conditions may yield very similar results to one another because they may differ only in unimportant factors.

Covering arrays, a technique from the theory of combinatorial designs, constitute a disciplined sampling method that give certain coverage guarantees on the search space of conditions while generating few experiments. Before defining the concept formally, let us give an example.

Suppose that we have six factors \( A, B, C, D, E, F \) that we can manipulate, each having three values 0, 1, 2. Covering the entire search space requires \( 3^6 = 729 \) tests. In that case, each of the possible 6-tuples based upon these three values would appear exactly once in the array.

What would a disciplined sampling approach require? First every possible value of every input factor should be present in some experiment. If this were the only condition, then it could be satisfied with the following three experiments:

\[
F : \quad A \quad B \quad C \quad D \quad E \quad F \\
1 \quad 0 \quad 0 \quad 0 \quad 0 \quad 0 \quad 0 \\
2 \quad 1 \quad 1 \quad 1 \quad 1 \quad 1 \quad 1 \\
3 \quad 2 \quad 2 \quad 2 \quad 2 \quad 2 \quad 2
\]
Intuitively, this is unsatisfactory, because it tests no interactions at all. So, let us raise the bar a little. We want to cover every value of every factor as above, but also for every pair of factors, we want to test each possible pair of values. In this way, if two values of two factors entirely dominate the situation, an experiment will discover that condition. Because every pair of factors has 9 possible pairs of values, this gives a lower bound of 9 experiments altogether, but it might seem to be difficult to come close to this few.

Remarkably, one can. A mere 13 experiments is enough:

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</table>

In what sense does this cover every pair of values for each pair of factors? Consider B and E for example. Let us project onto their two columns:
\[ \begin{array}{cc}
B & E \\
0 & 0 \\
1 & 1 \\
2 & 2 \\
0 & 0 \\
0 & 1 \\
1 & 0 \\
2 & 2 \\
1 & 1 \\
0 & 2 \\
1 & 2 \\
2 & 0 \\
2 & 1 \\
0 & 0 \\
\end{array} \]

The first three rows cover the pairs \((0, 0), (1, 1)\) and \((2, 2)\). The fourth row repeats \((0, 0)\) for that pair, but then we get, in succession, \((0, 1), (1, 0)\). After more repeats, we get the rest: \((0, 2), (1, 2), (2, 0),\) and \((2, 1)\). What is clever about this construction is that while taking care of factors B and E we are also taking care of all other pairs of factors.

Let’s review what this 13 experiment design accomplishes. Every value of every factor is tested. Every pair of values of every pair of factors is also tested. Of course, most three and four way interactions are not tested. There is no free lunch. But if you have limited resources, such coverage reveals a lot in few experiments [9]. If you have more resources, then you can do more experiments and cover more interactions. We sketch some of the technicalities below.

Before continuing with the mathematical development, we discuss the variety of ways in which this might apply to biology. First, the working biologist may want to know which factors are likely to be most important in their regulation of some target such as organism growth. Analyzing the results of an experimental set like the one above may uncover strong correlations with one factor or another. Those
correlations are not definitive (there are many confounding factors), but they are suggestive of which factors are most important.

Suppose, in fact, you discovered some factor that seems important. Or alternatively, suppose that you want to focus on that factor in an undirected mode, e.g., to discover which genes or proteins were sensitive to that factor. In either case you could take each value of that factor (or factors) and append a covering array of the other factors. In this example, suppose that we wanted to check the sensitivity to factor $A$, then the following 36 element array will do the job:
<table>
<thead>
<tr>
<th>Exp</th>
<th>A</th>
<th>B</th>
<th>C</th>
<th>D</th>
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<td>36</td>
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</table>
You will notice that the three groups 1-12, 13-24, and 25-36 are the same except in their A value. Thus for example 3, 15 (3+12), and 27 (3+24) have the same values of BCDE, but differ in their A values. So for each A value, all pairwise interactions among BCDE are tested (by 12 tests for these five factors rather than the 13 required for six). Therefore, all three-way interactions involving A are tested overall. If for each experiment triple i, i + 12, i + 24, the value of some target increases, then it is likely that A is inductive for that target. Admittedly, this notion of likelihood is difficult to quantify, because we don’t know the underlying distribution of data, but the construction gives such a large variety of values of BCDE and F that this statement is still an excellent hypothesis.

But what if A has no such consistent pattern. This brings us to a third use of combinatorial designs. Suppose for example that in this last array, experiment 28 gave a strong inductive effect relative to 16 (28-12) and 4 (28-24), whereas 25 did not have a strong inductive effect relative to 13 and 1. Presumably the interaction with BCDE is responsible. If we look at those values in the two cases, we see that 25 and 28 (and therefore pairs (1, 3) and (13, 16)) share the same value for BC. On the other hand, they differ in D and E. So one could establish a context that mixes the two, e.g. D = 0 and E = 2, and test the three experiments:

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<tr>
<th>Exp</th>
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If the effect is strongly inductive, then factor E is critically important in the influence of A. This ability to identify “intermediate” experiments and to zero in on critical factors can make experimental practice far more efficient. We have used this successfully in the study of nitrogen pathways in Arabidopsis [12, 9].
2 Mathematical Definitions and Results

The purpose of this section is to help you understand basic definitions and the number of experiments you would need depending on the number of factors you have and the number of values per factor. In most of the development, we assume the same number of values per factor, but if you have fewer than the maximum for some factor, the software [14] may be able to take advantage of that to reduce the number of needed experiments.

2.1 Covering Arrays

Let us say there are $c$ factors, $v$ values and we want to cover all possible $t$-way interactions. In the 13 experiment example above, there were $c = 6$ factors, each having $v = 3$ values, and we were interested in $t = 2$ way interactions. Here is the general definition.

A $t$-covering array with alphabet size $v$, length $c$, and size $r$ consists of $r$ vectors of length $c$ with entries from $\{0, 1, \ldots, v-1\}$ with the property that in any $t$ columns of the array each of the $v^t$ ordered $t$-tuples occurs at least once [13]. In this terminology, our earlier examples are 2-covering arrays.

The main problem in designing these covering arrays is to minimize $r$ (the number of rows or experiments) for given values of $v, c, t$.

In the case when $v = t = 2$ (two values per factor and two-way interactions), the problem has been completely solved [13]. In particular, for any fixed value of $r$, the maximal number $c$ of factors is determined by

$$c = \binom{r - 1}{\lceil \frac{r}{2} \rceil}.$$

At first glance, this may not seem helpful. Working biologists will often be given a number of factors and want to find out how many experiments are needed, rather
than being given a number of experiments and finding out how many factors can be tested. The formula lets us figure out the number of experiments from the number of factors however. The following table shows how many experiments are needed for each number of conditions for this (two value, two-way interaction) case:\footnote{Such a covering array can be constructed by assuming the first row consists of all zeros, and the remaining $r-1$ rows are taken to be the characteristic vectors of all subsets of weight $\left\lceil \frac{r}{2} \right\rceil$ of a set containing $r-1$ elements.}

<table>
<thead>
<tr>
<th>$c$</th>
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<th>7</th>
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For large $c$, using logs base 2, the number of rows is $r = \log c + \frac{1}{2} \log \log c \ldots$.

For $t = 2, v = 3$ (two-way interactions but three values per factor), a small table of the best known values taken from [13] follows:

<table>
<thead>
<tr>
<th>$c$</th>
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For $t = 2, v > 2$, it is known that for large $c$ the minimal number of rows satisfies $r = \frac{v}{2} \log(c(1 + o(1)))$. Thus it rises as the log of the number of conditions rather than exponentially (which would be the case for a complete search of the entire space).

In [11] the authors study covering arrays and their constructions in more detail. In particular, using sets of mutually orthogonal latin squares, they provide constructions of covering arrays which improve a number of the currently best known parameters for covering arrays; including three cases ($c = 13, r = 15; c = 14, r = 15; \text{ and } c = 15, r = 15$) in the above table for $v = 3$. See [3] or [8] for a discussion of latin squares and sets of mutually orthogonal latin squares.

Using our software [14], for example, computing 2-way interactions for 10 factors each with 3 values requires 15 experiments (as opposed to 59,000 to explore the

1Such a covering array can be constructed by assuming the first row consists of all zeros, and the remaining $r-1$ rows are taken to be the characteristic vectors of all subsets of weight $\left\lceil \frac{r}{2} \right\rceil$ of a set containing $r-1$ elements.
complete space). Computing 2-way interactions for 10 factors each with 4 values requires 52 experiments (as compared with over 1,000,000 to explore the complete space).

Dennis,

Here I think it would be helpful to write out the formulas (in terms of the values $t=2, v=3, c=10, r=15$) that yield the 59,000 and 1,000,000 values.

2.2 Orthogonal Arrays

Whereas we think that covering arrays provide the most parsimonious approach to testing interactions, reasonable scientists might desire a property known as balance. For this we need orthogonal arrays.

An orthogonal array [7] has, like a covering array, $r$ rows (experiments) and $c$ (factors) columns. Again, assuming that each factor has $v$ values, the array has “strength” $t$ (number of interacting factors) and index $\lambda$ if every $r \times t$ subarray (projection of every $t$ columns) of $A$ contains each $t$-tuple in exactly $\lambda$ rows.

For example, for an orthogonal array of strength $t = 2$, each of the $v$ elements occur the same number of times in each column, but each of the $v^2$ possible ordered pairs also occurs the same number of times in any two columns.

Here is the orthogonal array for two-way interactions on six factors ($c = 6$), three values ($v = 3$) per factor. (Recall that we had only 13 experiments before. Now we have ???.)
Chapter 12 of [7] provides an excellent summary of orthogonal arrays and how to construct them when they are possible to construct. When they aren’t, main-effects plans may be used. A main-effects plan is intermediate between a covering array which guarantees nothing about balance and an orthogonal array. In a main-effects plan, the number of occurrences of each ordered is proportional to the number of times that each element appears in each column. For a discussion of main-effect plans, see Section 11.7 of [7]. The website www.research.att.com/~njas/ contains even more recent updates on orthogonal arrays and their constructions.

3 Summary:

Combinatorial designs in general and covering arrays in particular can vastly reduce the number of experiments needed to explore a search space. While they don’t cover every possible combination of the input factors, they sample the search space in a well-separated manner and guarantee that every combination of values in every $t$ factors are tested. If a researcher prefers certain balance properties, then he or she
can use orthogonal arrays or main-effects orthogonal arrays.

Combinatorial designs can be used in an iterative and adaptive fashion.

1. One can start by using it to find those factors that might be important.

2. Given a possibly important factor, one can use combinatorial design to test the consistency of its importance.

3. Given one set of conditions where a factor is important and another set where the factor isn’t, one can create intermediates that will isolate what it is about a context that determines the importance of the factor.

Of course, \( t \)-way covering arrays don’t test \((t + 1)\)-way interactions. Thus, to extend one’s interactivity coverage, one must do more work. But given a certain desired coverage, covering arrays give an inexpensive way to uncover them.

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**References**


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