SWEETPEA: TOWARD UNIFORM SAMPLING

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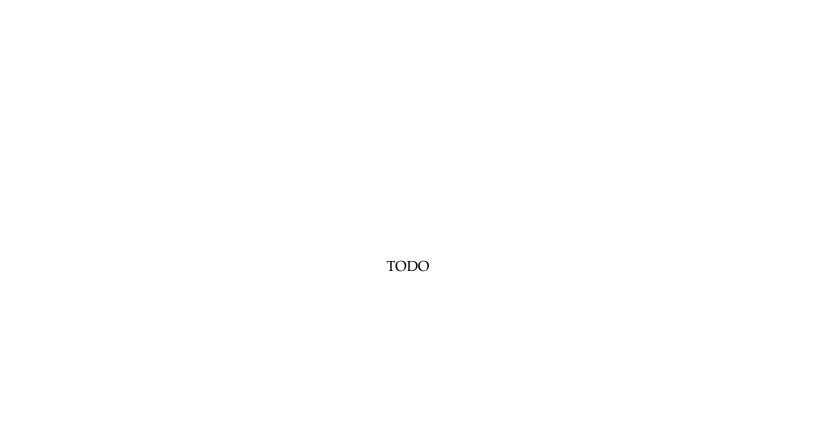
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ABSTRACT

SweetPea is a new language providing a means of declaratively describing experimental designs. Because of their combinatorial nature, realistic experimental designs have too many conforming trial sequences to reasonably enumerate all of them for random sampling. As a result, alternative means of generating uniformly sampled trial sequences must be used. SweetPea currently does this by encoding the design as a SAT formula, to which a SAT sampler is then applied to generate uniformly sampled solutions. Although recent SAT samplers can process formulae with large solution spaces, particular characteristics of experimental design have made this approach unfeasible. We present evidence explaining why SAT samplers in their current state are not viable for this problem and lay the foundation for an alternative encoding, based on a bijection from natural numbers to unique trial sequences. This alternative encoding allows us to generate uniformly-distributed random samples for certain classes of designs in a fraction of the time required by previous methods.



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CHAPTER 1

MOTIVATION

SweetPea [6], a language for experimental design, attempts to generate approximately uniformly sampled trial sequences by relying on the aid of a SAT-sampler to provide uniformity guarantees. In practice, this approach does not scale to support realistic experimental designs. This thesis presents evidence demonstrating the need for an alternative and investigates other approaches to overcome this problem. These include a different encoding scheme as well as techniques for direct construction of a solution based on its numerical index in the sequence of all solutions. These improvements significantly improve SweetPea's capability to guarantee a uniform sampling of trial sequences.

1.1 Introduction

For any valid design configuration, there exists some set of trial sequences that conform to the design. Typically this set is extremely large, as the size grows factorially in the trial sequence length of the design. The magnitude of the solution space for experimental designs has posed a significant challenge for scientists desiring to conduct unbiased resarch.

At present, scientists patch together different scripts and tools in various ways to generate experiment trial sequences. This method, in addition to being brittle, does not facilitate sharing, nor does it produce unbiased sequences, making it nearly impossible for peers to replicate results. SweetPea provides a high-level declarative language for describing all aspects of an experimental design, providing standard tooling where before there was none. In addition, with the aid of a SAT-sampler, it generates approximately uniformly sampled trial sequences from the space of all sequences conforming to the design. Sweet-Pea is composed of a Python library, which provides the language constructs, and a server that generates and processes the design using a SAT-sampler.

At the highest level, an experimental design is composed of a *block*, whose properties may vary depending on its type. At present, only a single block type is provided by

SweetPea, which is the *FullyCrossBlock*. A FullyCrossBlock is composed of a set of *factors*, a set of *constraints*, and a *crossing*. A *factor* is a collection of *levels*, or values to which the factor may be set. In a given trial, each factor takes the value of one of its levels. Levels may be values determined by the designer, or they may be derived from the values of other factors in the design. A *constraint* constrains the experiment in some regard. For example, the designer may wish to prohibit a particular level from being repeated in a trial sequence. Lastly, the *crossing* identifies the factors whose levels should be combined (crossed) in all possible ways in each trial sequence. The output of a design is a set of trial sequences. A single level value from each factor comprises a *trial*.

1.2 Current Implementation

SweetPea works by encoding the experimental design in a boolean satisfiability (also known as SAT) formula. such that any satisfying assignment constitutes a trial sequence that conforms to the design. Currently, a simple positional encoding is used, in which every level of every factor of every trial is represented as a distinct variable.

1.2.1 Boolean Satisfiability

The Boolean satisfiability problem (abbreviated SAT) is the problem of determining whether or not a Boolean formula can be satisfied, or be caused to evaluate to true, by a particular true or false assignment to each of its variables. If a problem can be expressed as a Boolean formula, then techniques from satisfiability theory may be applied to determine whether or not a solution exists to the problem, or to locate a particular solution. Many different SAT solvers exist which accept a Boolean formula as input, and output either that the formula is *unsatisfiable*, or a solution to the formula.

Tools for interacting with SAT formulae typically require the input formula to be in *conjunctive normal form* (abbreviated CNF), which is a form of a Boolean formula with specific properties. A Boolean formula is in conjunctive normal form only when three conditions are met. First, the entire formula must be a conjunction of clauses. Second, each clause consists of an individual variable, or a disjunction of variables. Third, negation may only be applied to individual variables. Any Boolean formula may be expressed in conjunctive normal form, though it usually requires growth in some dimension. SweetPea

uses the Tseitin Transformation [15] to do this conversion, which has the tightest known bounds on formula length, at the expense of allocating additional variables. A SAT formula contains some number of boolean variables and propositional logic clauses. A selection of true or false for each variable in the formula constitutes an *assignment*. If a given assignment causes the entire formula to be true, then the formula is said to be *satisfied* by that assignment, or more generally, the formula is *satisfiable*. If no such assignment exists, the formula is said to be *unsatisfiable*.

1.2.2 Example: Stroop Experiment

The Stroop test [14] is a well-known design among experimental psychologists in which a subject's reaction time is influenced by perceived conflicts in the trial sequence. For example, a subject may be shown printed words representing colors and asked to read the word. However, some of the time the font color of the word will be different than the printed word itself, leading to a delayed reaction due to the conflict. This design is easily represented in SweetPea as follows:

```
color = Factor("color", ["red", "green", "blue"])
text = Factor("text", ["red", "green", "blue"])
```

A FullyCrossBlock with these two factors will yield a sequence of $3 \cdot 3 = 9$ trials. A research may want to constrain the design to ensure that there are never two consective trials in which the color and text are the same, or congruent. Such a constraint requires the addition of another factor:

```
congruent = Factor("congruent?", [
    DerivedLevel("yes", WithinTrial(operator.eq, [color, text])),
    DerivedLevel("no", WithinTrial(operator.ne, [color, text]))
])
```

With the factor now in place, a constraint may be specified:

```
AtMostKInARow(1, ("congruent?", "yes"))
```

The current encoding scheme will allocate 72 variables to represent each level of each

factor in each trial. Additional variables will be allocated to enforce other constraints, such as ensuring that each factor takes only one level in each trial, and requiring that each crossing be present in the sequence. The first step of the encoding can be visualized easily, as demonstrated in Table 1.1.

Table 1.1. Example of an Encoding Diagram

	Color			Text			Color Text Congruen			gruent?
Trial Number	Red	Green	Blue	Red	Green	Blue	Yes	No		
1	1	2	3	4	5	6	7	8		
2	9	10	11	12	13	14	15	16		
3	17	18	19	20	21	22	23	24		
4	25	26	27	28	29	30	31	32		
5	33	34	35	36	37	38	39	40		
6	41	42	43	44	45	46	47	48		
7	49	50	51	52	53	54	55	56		
8	57	58	59	60	61	62	63	64		
9	65	66	67	68	69	70	71	72		

For example, if variable 68 is true, that corresponds to the level red being selected for the factor text in trial number 9. The encoding will also require variables 69 and 70 to be false in this case, as well as require variable 71 to be equivalent to 65. Once a solution is found, it is then decoded back into the original language of the design as a trial sequence. For more details concerning SweetPea's syntax or the current encoding scheme, the reader is referred to the original paper.

1.2.3 One-to-One Correspondence

In order for the uniformity guarantees of sampling tools to tranlate back into the problem space of the original experiment design, it is critical that the encoding scheme maintain a one-to-one relationship between formula solutions and distinct trial sequences. Without this relationship, it would be possible to uniformly sample solutions to the SAT formula while still introducing bias in the generated trial sequences. For example, if three SAT solutions yielded trial sequence S_1 , while only one solution yielded trial sequence S_2 , uniformly distributed solutions to the SAT formula would produce S_1 more frequently than S_2 . A few observations build our confidence that this correspondance is upheld. First, all factor and level selections, including associated constraints for derived levels, in the original problem space are represented directly in the SAT encoding. Each level of each factor is represented by a unique variable for every trial in the sequence, and clauses are added to the CNF to ensure that variable assignments are consistent with the actual constraints of the problem. Referring back to the example in Table 1.1, the levels for the Color factor for trial number 1 are represented by variables 1, 2, and 3 respectively. A popcount circuit is embedded in the SAT formula that requires that exactly one of those variables is set to true. This same circuit is repeated for the Color factor for each of the 9 trials, as well as for all other factors in all 9 trials. Other constraints are also embedded to represent all other rules of the design, including relationships between basic and derived factors, and constraints that require the full spectrum of crossed factor combinations to be represented in the sequence. Because all data from the original design is directly translated into SAT variables and constraints, we believe there is no risk of generating multiple solutions to the formula that represent the same trial sequence.

However, some of these constraints are represented in expressions richer than allowed by CNF, including logical implication. Of necessity, these expressions must be rewritten in an equivalent form using only those primitives allowed by CNF. In order to avoid an exponential explosion in the length of the formula, this rewriting process introduces additional variables to represent repeating clauses. The second observation is that SweetPea applies the Tseytin transformation to do this conversion to CNF, which binds every variable that it introduces to the state of a clause in the original formula. As a result, the number of solutions remains the same, and there is no possibility of underconstrained variables skewing the results.

Lastly, we note that for small designs, we can exhaustively enumerate all possible trial sequences. The number of enumerated sequences matches exactly the result obtained when using a SAT model counter to determine the number of solutions to the SAT formula. Were there not a one-to-one mapping, these figures would disagree.

1.3 Problems

SweetPea sounds panacean, but it has yet to fully realize its objectives. It relies *fully* on a SAT-sampler to approximately uniformly sample some number of solutions to the generated formula. In practice, this works wonderfully for *small* experimental designs, where *small* means designs in which the trial sequence length is less than ten. However, realistic experimental designs are significantly larger, often with trial sequence lengths in the tens or hundreds.

A solution to the formula represents a unique trial **sequence** in the design. Therefore every possible sequence permutation (so long as it upholds the design constraints) is also a distinct solution. Consequentially, the number of solutions grows factorially in the sequence length. In the above example, the sequence length, denoted l, was 9. Assuming the congruence constraint were removed, there would be 9! = 362,880 solutions, which is still reasonable. Consider an experiment however in which l = 36. The solution count grows to $36! \approx 2^{138}$, which is intractable. As will be seen in chapter 3, it is common for experimental designs to have sequence lengths even larger than 36.

Table 1.2. Growth of Solution Space

Sequence Length (1)	#SAT (!!)
4	24
6	720
9	362,880
12	479,001,600
16	20,922,789,888,000
20	2,432,902,008,176,640,000
25	15,511,210,043,330,985,984,000,000
30	265,252,859,812,191,058,636,308,480,000,000
36	371,993,326,789,901,217,467,999,448,150,835,200,000,000

In practice, designs will be constrained, however the constraints only exclude specific patterns in the permutations, therefore the number of solutions remains dominated by *l*!. Clearly, generating all possible solutions is not feasible. We had hoped that a SAT sampler would be able to sample solutions from this space, but in practice this growth rate very quickly overwhelms all tested SAT-samplers. It has been well established that prior SAT-samplers have traded strong uniformity guarantees for scalability [5], and thus

were not suitable for even moderately large formulae. Unigen [5] contributed two primary advancements to bridge this gap between scalability and uniformity guarantees, but only under certain conditions. Unfortunately, the nature of the formulae generated by SweetPea designs violates these conditions, as the target domain of Unigen, constrained-random verification (CRV), exhibits different characteristics than experimental design.

CHAPTER 2

A COMPLEXITY SPECTRUM FOR SWEETPEA DESIGNS

The complexity of SweetPea designs greatly varies depending on the user's requirements. At one end of the spectrum are simple designs, in which there are no structured relationships between trials in a sequence. At the other end are the most complex designs, in which multiple structured relationships exist between multiple trials in a sequence at varying intervals. This chapter will present a precised gradation of 6 complexity tiers for experimental designs in order to facilitate future discussion. Recall that a complete experimental design is defined by a block, which is composed of a set of factors, a crossing of some subset of the factors, and a set of constraints. This section considers the complexity of a design based solely upon the factors present. Constraints on the overall design are not included in this gradation, as their complexity depends primarily on the complexity of the factors that they constrain.

2.1 Factors and Windows

2.1.1 Factor Types

Only two types of factors exist in SweetPea: *Basic* and *Derived*. The levels of a basic factor are literal values. Basic factors form the foundation of all designs. The following example declares a basic factor named direction, whose levels are the directions of the compass:

```
direction = Factor("Direction", ["North", "South", "East", "West"])
```

On the other hand, the levels of a derived factor depend upon values from other factors in the design. The congruency characteristic in the Stroop experiment is expressed as a derived factor:

```
congruency = Factor("Congruency", [
   DerivedLevel("Congruent", WithinTrial(lambda c, t: c == t, [color, text])),
   DerivedLevel("Incongruent", WithinTrial(lambda c, t: c != t, [color, text]))
])
```

The level of the congruency factor is determined by the current levels of the color and text factors, which will vary for each trial in a generated sequence. WithinTrial denotes the *window* of the derived level, which brings us to the next subject.

2.1.2 Window Construction

The construction of a derived level is based on the concept of a *window*, which slices the trial sequence into segments which are considered to determine the current level of a derived factor. A window's dimensions are defined by a *width* and *stride*. The width indicates how many preceding trials, including the current trial, are to be considered. The *stride* indicates how many trials to skip when moving on to the next trial for this factor.

The simplest type of window, for which WithinTrial is an alias, has a width and stride of one. This type of window does not span multiple trials. It only considers the values of basic factors within the same trial.

A slightly more complicated window has a width of two, but still a stride of one. (Aliased by the Transition window.) This type of window considers two adjacent trials for the purpose of inspecting the transition (or lack thereof) between them. For example, a Transition window would be appropriate when seeking to determine if a particular level was repeated in sequence.

```
direction_repeats = Factor("Direction Repetition", [
   DerivedLevel("Repeat", Transition(lambda directions: directions[0] == directions[1], [
   DerivedLevel("No Repeat", Transition(lambda directions: directions[0] != directions[1], [
])
```

The most complicated windows have arbitrary widths and strides. For example, a window with a width of one and a stride of four would consider the value of factors only at every fourth trial in the sequence. For example:

```
direction_is_north = Factor("Direction is North", [
```

```
DerivedLevel("Direction is North", Window(lambda direction: direction == "North", [di DerivedLevel("Direction is not North", Window(lambda direction: direction != "North", [di ])
```

Notice that windows with non-one width and stride will not have a value for every trial in the sequence. Generally speaking, a window with width w and stride s will first apply to trial w, and to every s^{th} trial thereafter.

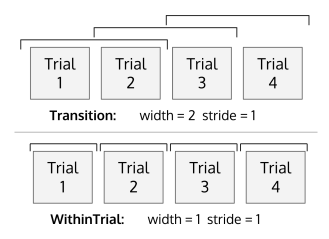


Figure 2.1. Window Properties

2.2 Complexity Gradation

Now understanding the types of factors and windows that can be used to define experimental designs, we can establish the complexity tiers with which to categorize different designs.

2.2.1 Tier 1

Tier 1 designs consist of basic factors only. No derived factors are allowed, and all factors in the design are crossed. Every valid trial sequence is a permutation of the crossed factors.

```
direction = Factor("Direction", ["North", "South", "East", "West"])
time_of_day = Factor("Time of Day", ["Sunrise", "High Noon", "Sunset"])
```

2.2.2 Tier 2

Tier 2 designs still consist of only basic factors, but allow additional basic factors to be present which are uncrossed. Every valid trial sequence is a permutation of the crossed factors and random level selections for the uncrossed factors.

2.2.3 Tier 3

Tier 3 designs add derived factors with windows being limited to width = 1 and stride = 1. (WithinTrial) Every valid trial sequence is a permutation of the crossed factors and random level selections for the uncrossed basic factors. The values for uncrossed derived factors will depend upon the level selections of basic factors.

```
congruency = Factor("Congruency", [
   DerivedLevel("Congruent", WithinTrial(op.eq, [color, text])),
   DerivedLevel("Incongruent", WithinTrial(op.ne, [color, text]))
])
```

2.2.4 Tier 4

Tier 4 designs relax the window limits to width = 2 and stride = 1. (Transition)

```
color_repeats = Factor("Color Repeats", [
   DerivedLevel("Repeat", Transition(lambda c: c[0] == c[1], [color])),
   DerivedLevel("No Repeat", Transition(lambda c: c[0] != c[1], [color]))
])
```

2.2.5 Tier 5

Tier 5 designs allow arbitrary values for the width and stride of all windows.

```
unconnected_color_repeats = Factor("Color Repeats (Unconnected)", [
  DerivedLevel("Repeat", Window(lambda c: c[0] == c[1], [color], 2, 2)),
  DerivedLevel("No Repeat", Window(lambda c: c[0] != c[1], [color], 2, 2))
])
```

CHAPTER 3

CURRENT BENCHMARKS

This chapter introduces several experiment designs and provides benchmarks for their performance as measured using the current approach to sampling, as well as using other available sampling tools. The implementation strategy of each tool will also be discussed lightly, in order to demonstrate shortcomings when sampling solutions to formulae with factorial solution spaces. These results should be convincing of the need for an alternative approach to the problem, as the current method cannot cope with the scale of the solution spaces in question.

3.1 Experiments

Multiple tools will be applied to solve or sample solutions to formulae reprsenting several experimental designs of increasing complexity comparable to realistic experiments written by current researchers. Table 3.1 enumerates the characteristics of each design, including its complexity tier (as defined in chapter 2), properties of its CNF encoding, and a solution count.

For stroop-2 and stroop-3, the solution count is precise, as computed by a SAT model counter. For all other experiments, the solution count is approximated as the factorial of the sequence length, as model counters were unable to provide a count in reasonable amount of time. These approximate solution counts ignore both constraints, which would reduce the solution count, and independent factors, which would increase the solution count. In practice, independent factors contribute more solutions than constraints remove.

3.2 Benchmarks

This section presents the results of testing multiple SAT solving and sampling tools against the experiments presented previously, as well as additional information on the implementation strategy of each tool. All sampling benchmarks were configured to generate

Table 3.1. Benchmark Experiments

Experiment Name	Tier	Length	Variables	Clauses	Total Solutions
stroop-2	3	4	392	1,559	12
stroop-3	3	9	1,965	8,531	151,200
stroop-4	3	16	3,848	16,451	$pprox 2^{44}$
stroop-5	3	25	10,326	45,991	$pprox 2^{83}$
stroop-congruency-balanced	4	17	5,570	24,215	$pprox 2^{48}$
stroop-response	4	33	16,242	72,679	$pprox 2^{122}$
padmala-pessoa	4	37	116,289	559,080	$\approx 2^{143}$
task-switching	4	32	14,246	62,700	$\approx 2^{117}$
task-switching-2	5	146	742,832	3,672,632	$pprox 2^{844}$
task-switching-cue-switching	4	1,025	-	-	$pprox 2^{8778}$

only 10 samples. All benchmarks were run on a PC running Ubuntu 18.04.2 LTS with 16GB of memory and an 8-core Intel Core i7-7700 at 3.60 GHz.

3.2.1 CryptoMiniSat

CryptoMiniSat [10, 13] is the main SAT solver that SweePea relies on for generating non-uniformly sampled trial sequences. SweetPea achieves this with CryptoMiniSat by using the naive approach of solving for any solution, and then constraining the original formula to disallow that particular solution from being produced a second time. Due to the difficulty encountered while attempting to use SAT samplers, it has proven quite valuable to provide researchers with this ability to quickly generate some number of trial sequences conforming to their design, even though they may not be uniformly distributed. CryptoMiniSat is also used internally by Unigen. Lastly, in addition to logical AND, OR, and NOT clauses, CryptoMiniSat also supports XOR claues, although SweetPea has not yet put these to use. This is an opportunity for improvement. Table 3.2 denotes the the time required for CryptoMiniSat to compute a single solution to the SAT formula for each experiment.

The most complicated design for which SweetPea was able to construct a CNF encoding (task-switching-2) took the longest, at 200 seconds. Most other experiments could be solved in less than a second. Since only a single solution is generated, the practical time consideration is of minor importance here.

Table 3.2. Time to Solve for 1 Solution with CryptoMiniSat

Experiment Name	Time to Solve (ms)
stroop-2	4.9
stroop-3	9.1
stroop-4	18.9
stroop-5	64.4
stroop-congruency-balanced	23.9
stroop-response	232
padmala-pessoa	3,380
task-switching	140
task-switching-2	200,000
task-switching-cue-switching	-

3.2.2 Unigen

Unigen [3,5] is a recently developed SAT *sampler*. Given a boolean formula in conjunctive normal form (CNF), Unigen can produce nearly uniformly sampled satisfying assignments. SweetPea relies directly on Unigen for uniformly sampling trial sequnces for a given design. As explained in chapter 1, the SweetPea encoding is designed such that there is a one-to-one relationship between solutions to the formula, and unique trial sequences that conform to the design. Unigen is based on a hashing technique that partitions the solution space and then samples solutions from each partition. Unigen has worked well for some designs, but its capacities are quickly overwhelmed by even moderately sized designs.

Let F denote a boolean formula in CNF, and let R_F denote the set of satisfying assignments, or witnesses, for F. Unigen, as well as several previously developed samplers, are based on a strategy for generating a partition of R_F using a family of 3-independent hash functions which provide strong uniformity guarantees [8]. However, there are two obstacles to using this family of functions: First, it requires an additional parameter m, which influences the size of the subsets of the partition of R_F . Choosing an approparite value for m is non-trivial. Second, these functions rely heavily on the XOR logical operation. Unfortunately, as stated in [5], as the number of variables per XOR clause grows, the difficulty of generating solutions to the formula increases significantly. [7] At the core of Unigen's contributions were techniques for mitigating both of these problems.

3.2.2.1 Model Counting

The best value for m actually depends on $|R_F|$, however in practice this is not known up front. Unigen uses an approximate model counter, ApproxMC [4], to estimate $|R_F|$, from which a narrow range of values for m can be generated and tested for fitness. Unfortunately in practice, ApproxMC has not been able to successfully approximate the count of any of our benchmark experiments beyond stroop-3 in a reasonable amount of time, as shown in Table 3.3. A new version of ApproxMC was recently published [12]; however, it relies on the same family of hash functions, so it is unlikely that it will offer significant improvements.

Table 3.3. Time to Compute Approximate Model Count with ApproxMC

1 1
Time
5.07ms
27.4s
Timed Out
-

In reality, the algorithm implemented by ApproxMC relies on the same family of 3-independent hash functions mentioned previously, therefore it will be subject to the same problem of XOR clause growth, as explained in the next section.

Although available tools struggle to approximate solution counts for SweetPea designs, this is surmountable by estimating $|R_F|$ ourselves using information from the original problem domain. Because each experimental design constructs a sequence of trials based on combinations of factor levels, we can apply techniques for counting sets from combinatorics to estimate $|R_F|$. This will be discussed further in chapter 4.

3.2.2.2 XOR Clause Reduction

Let X be the set of variables in F. Using terminology from [5], X is called the *support* of F. They observed that there is often a subset $S \subseteq X$, referred to as an *independent support*

of X, which fully determines all variable assignments for any satisfying assignment. That is, for all satisfying assignments of F, there are no two assignments whose values differ only in the assignments of variables \overline{S} . In the domain of CRV, |S| is typically significantly smaller than |X|, often by many orders of magnitude. Unigen exploits these observations by only applying variables in S to the hash functions, which yields a significant reduction in the number generated XOR clauses. Therefore it is true that Unigen scales to formulae with hundreds of thousands of variables, and even millions of clauses, but *only* when |S| remains small, typically |S| < 100. In the Unigen benchmarks, over 200 different formulae were sampled. The largest of these, demo3_new, had 865,935 variables and 3,509,158 clauses, but |S| was only 45. Similarly, the largest value of |S| in the benchmarks was 78 in diagStencil_new, with 94,607 variables and 2,838,579 clauses. [2]

How does |S| grow in the formulae generated by SweetPea? SweetPea currently selects all variables that directly represent level values for each factor in each trial as the independent support. While this set does not grow factorially, it still quickly grows into the hundreds and thousands, as shown in table 3.4. While this independent support is certainly not minimal, it is unlikely that improvements to the encoding and the determination of S could scale more than marginally over current performance.

There are a few obvious improvements that could be made. Currently SweetPea is including the variables for *all* factors and levels in the independent support, when in reality only variables for basic factors and derived factors in the crossing need to be included. Additionally, a more efficient encoding could be used based on combinations of variables rather than representing all levels as individual variables. (Although this would increase the complexity of encoding constraints.) Both of these would reduce |S| by some amount, but there is still a theoretical limit by which we are bound. For any formula F with independent support S, there are, by definition, at most $2^{|S|}$ solutions to F. (In practice the number of solutions will be less, as not *all* possible assignments will satisfy the formula.) If |S| < 100 is a practical limit, then even with the most efficient encoding, Unigen would only be able to sample solutions to formulae with at most 2^{100} solutions. As seen in Table 3.1, this would allow for marginal scaling, but not generally for all SweetPea designs.

This shortcoming is proved out in practice, as Unigen fails to count or sample from formulae where |S| > 100. Table 3.5 denotes the time required for Unigen to generate 10

Table 3.4. Independent Support Size for Benchmark Experiments

11	
Experiment Name	S
stroop-2	24
stroop-3	72
stroop-4	160
stroop-5	300
stroop-congruency-balanced	270
stroop-response	526
padmala-pessoa	1,303
task-switching	636
task-switching-2	2,770
task-switching-cue-switching	24,594

Table 3.5. Time to Generate 10 Samples with Unigen

Time to Sample
-
2.18s
Timed Out
-

samples for each experiment when an estimate for $|R_F|$ is provided manually based on the factorial of the sequence length. Nearly all of them fail to complete in the allotted time. Because Unigen can only effectively sample large formulae when |S| is small, in practice |S| < 100, it is insufficient for the needs of SweetPea. Several other samplers were tested as well, but in our experiments, none of them were successful.

3.2.3 KUS

KUS [11] is another SAT sampling tool developed by some of the same team behind Unigen. However, it approaches the problem quite differently. KUS requires a compiled deterministic decomposable negation normal form (d-DNNF) of a formula in conjunctive normal form. There are known polynomial time techniques for many operations on the d-DNNF representation of a boolean formula, including model counting. KUS takes ad-

vantage of these techniques to uniformly sample solutions using this form. This technique offloads much of the complexity to to the d-DNNF compiler, rather than the sampling phase. KUS, while able to sample quickly, is not used in SweetPea due to scaling problems in compiling the formula to d-DNNF. Though the number of variables and clauses in the CNF for a formula is typically modest, compiling the equivalent d-DNNF has proven intractable. Table 3.6 denotes the time required to compile the CNF for each experiment to d-DNNF using the d4 compiler, as well as the resulting d-DNNF file size compared to the original CNF file.

Table 3.6. d-DNNF Compilation for Sampling with KUS

Experiment Name	Time	d-DNNF File Size	CNF File Size
stroop-2	14.4ms	9.7K	24K
stroop-3	1.99s	6.3M	151K
stroop-4	Timed Out	-	303K
stroop-5	Timed Out	-	881K
stroop-congruency-balanced	48.6m	7.3G	447K
stroop-response	Timed Out	-	1.5M
padmala-pessoa	OOM	-	13M
task-switching	Timed Out	-	1.2M
task-switching-2	"Indeterminate"	-	92M
task-switching-cue-switching	-	-	-

For the experiments for which a d-DNNF file could be successfully compiled, table 3.7 denotes the time required to sample solutions using KUS.

Table 3.7. Time to Generate 10 Samples with KUS

Experiment Name	Time
stroop-2	92.7 ms
stroop-3	2.35 s
stroop-4	-
stroop-5	-
stroop-congruency-balanced	OOM
stroop-response	-
padmala-pessoa	-
task-switching	-
task-switching-2	-
task-switching-cue-switching	-

As expected, when a reasonably-sized d-DNNF can be compiled, sampling can be performed quickly. However, the combinatorial nature of experimental designs makes the cost of compiling and interpreting the d-DNNF prohibitive.

3.2.4 Spur

Spur [?] is a more recent SAT sampler that is based on reservoir sampling. In benchmarks, it performs better than Unigen in nearly all cases. However, it does require an exact model count (based on sharpSAT), while an approximation suffices for Unigen. SweetPea does not rely on Spur at present, as it was developed after SweetPea's inception. However it has been considered as it represents the latest research for SAT sampling. Table 3.8 denotes the time required to generate samples for each experiment using Spur, with similar results as previously considered tools.

Table 3.8. Time to Generate 10 Samples with Spur

Experiment Name	Time to Sample
stroop-2	21.9 ms
stroop-3	3.72 s
stroop-4	Timed Out (10 hrs)
stroop-5	Timed Out
stroop-congruency-balanced	OOM
stroop-response	Error
padmala-pessoa	Timed Out
task-switching	Timed Out
task-switching-2	Error
task-switching-cue-switching	-

3.3 Distribution of Samples

As mentioned in chapter 1, there must be a one-to-one relationship between unique trial sequences and solutions to the SAT formula in order for uniformity guarantees to carry over from the SAT sampler results back to the original problem space. In this section we'll examine the distribution of the trial sequences produced by Unigen, KUS, and Spur, providing empirical evidence that the resulting trial sequences are uniformly distributed.

We sampled 10,000 trial sequences for the stroop-3 experiment using Unigen, KUS, and Spur. The resulting samples appeared to be uniformly distributed, each sample only

occurring once for the most part.

3.3.1 Unigen

Unigen sampled 10,010 trial sequences, 9,965 of which were unique. Numbering each unique trial sequence in the order that it appeared and generating a histogram gives a visual confirmation that the sequences are approximately uniform. There were 333 sequences that were selected twice, and only 6 that were selected three times. No sequences were selected more than three times.

Distribution of Unigen Samples

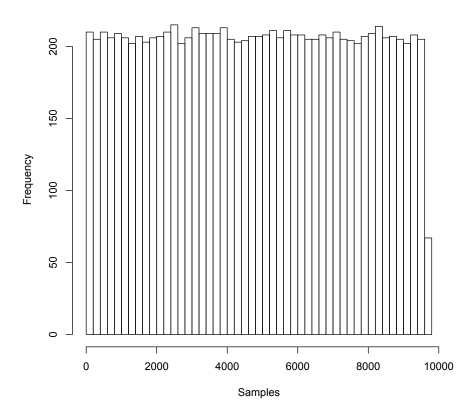


Figure 3.1. Distribution of Sequences Sampled by Unigen

3.3.2 KUS

KUS sampled 10,000 trials sequences, all of which were unique.

Distribution of KUS Samples

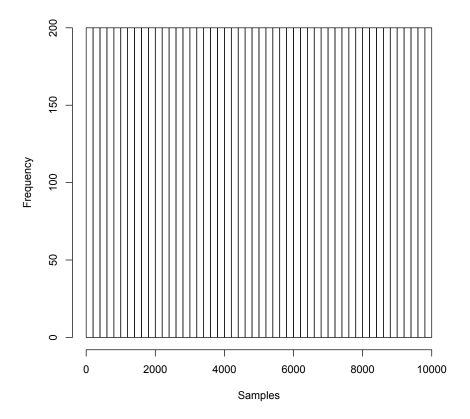


Figure 3.2. Distribution of Sequences Sampled by KUS

3.3.3 Spur

Spur sampled 10,000 trial sequences, 9,664 of which were unique. 332 sequences were selected twice, and 2 were selected three times. No sequences were selected more than three times.

3.3.4 CryptoMiniSat

In contrast to the previous tests, when a SAT solver is used to generate individual samples incrementally by adding negated solutions to the prior formula, the same samples are generated each time. We used CryptoMiniSat to generate 10,000 samples as well, in groups of 100. Exactly the same set of 100 samples was generated each time.

Distribution of Spur Samples

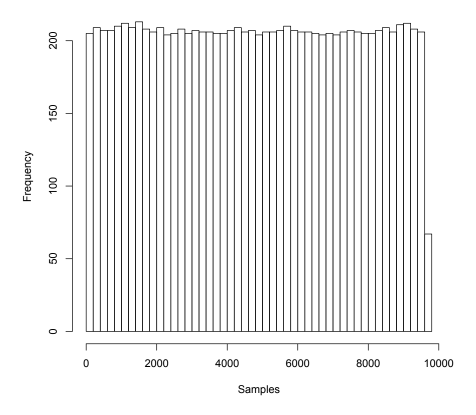


Figure 3.3. Distribution of Sequences Sampled by Spur

3.4 Summary

We had hoped that Unigen would be able to sample from the space of solutions for experimental designs, in particular because the volume of variables and clauses in our generated CNF files fell within the range of example benchmarks provided by Unigen. However, the magnitude of the independent support set plays a critical role that was overlooked. Because the size of the independent support set for experimental designs grows quickly into the hundreds and thousands, a correspondingly large volume of XOR clauses is added to the CNF during sampling, which exceeds the capacities of the SAT solver (CryptoMiniSat). Furthermore, none of the additional tested tools for sampling SAT formulae are capable of handling the scale of practical experimental designs. Although the number of variables and clauses in the CNF for these experiments is reasonable, the factorial explosion in the number of solutions as well as the magnitude of the independent

support set proved too large.

CHAPTER 4

SOLUTION COUNTING

Currently, SweetPea offloads the entire sampling process to an external tool. While an attractive option from an implementation standpoint, this approach does not scale to the sizes needed. As demonstrated in chapter two, the problem spaces that SweetPea is trying to sample from are beyond the capacity of any existing tools. One of the shortcomings of this approach is that external tools cannot take advantage of knowledge of the problem to guide their decisions.

Experimental designs in SweetPea can be viewed as combinatorics problems, each with a countable set of solutions. If we could construct a formula for counting the solutions to an experimental design, we may be able to also discover a bijection between solutions to the design and the natural numbers. This would reduce the burden of guaranteeing uniformity to randomly sampling natural numbers from the uniform distribution.

As a first foray into solution counting for SweetPea designs, this section only considers tier 1, 2, and 3 designs. We also leave the set of design constraints for later consideration.

4.1 Principles of Counting

Before describing a formula for counting solutions to experimental designs, it will be useful to review a few counting principles for future reference, beginning with counting permutations of a set. The standard formula for then number of permutations of r items taken from an n-element set is:

$$P(n,r) = \frac{n!}{(n-r)!}$$

See [1]. When n = r, this becomes simply n!.

When constructing a set S by combining individual elements from mulitple other sets, $P_1, P_2, \cdots P_n$, the size of S is the product of the sizes of sets $P_1, P_2, \cdots P_n$:

$$|S| = \prod_{i=1}^{n} |P_i|$$

This is known as the *Multiplication Principle* [1].

4.2 Partition of an Experimental Design

An experimental design is composed of three elements: a set of factors, a subset of the factors to be combined with each other (crossed), and a set of constraints. The set of crossed factors fundamentally shape the generated trial sequences. The length of a trial sequence is governed by the number of combinations of level values for each crossed factor. For example, consider a design in which there are two crossed factors, each with three levels. By the multiplication principle, there are 3*3=9 unique combinations of level values from each factor, hence trial sequences in this design will be nine trials long.

Let D be the set of all factors in the design. Let C be the set of crossed factors, and \overline{C} be the set of factors not included in the crossing. Therefore $C \subseteq D$ and $\overline{C} \subset D$. C and \overline{C} can both be further divided based on the type of factor: basic or derived. Let C_B be the set of basic factors in C, and C_D be the set of derived factors in C. We will define \overline{C}_B and \overline{C}_D similarly for \overline{C} .

Lastly, we further divide \overline{C}_B into two more sets. Although basic factors in \overline{C} are not included in the set of crossed factors, this does not imply that they are fully independent. It is possible that factors in C_D depend on factors in \overline{C}_B . We'll refer these factors in \overline{C}_B as *source* factors, as they are at least a partial source of the data controlling factors in C_D . We will group such factors in \overline{C}_{B_S} , while \overline{C}_{B_I} represents the truly independent basic factors in \overline{C} .

We could also define C_{B_S} and C_{B_I} , however there will be no benefit to doing so, as the relationship between factors in C_B and dependent derived factors is inverted. Each combination of levels in the crossed set governs the values for C_B , which will then in turn govern dependent derived factors.

The partition of D with which we are concerned is therefore: $\{C_B, C_D, \overline{C}_{B_S}, \overline{C}_{B_I}, \overline{C}_D\}$, as shown in figure 4.1.

Lastly, we define X to be a set of ordered pairs $t_1, t_2, \dots t_n$, representing the crossing of factors in C. Each ordered pair t contains items $i_1, i_2, \dots i_j$, where item i_j is one of the levels

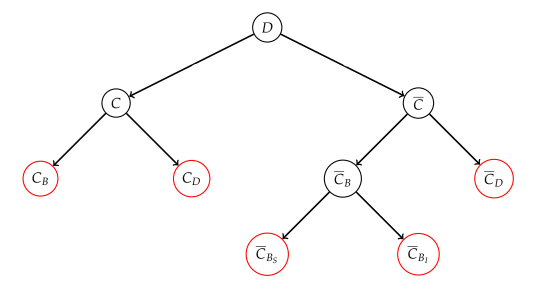


Figure 4.1. Partition of an Experimental Design

from the j_{th} factor in C. Each ordered pair t must be unique, and there is an ordered pair for every possible combination of levels from factors in C. By the multiplication principle, there will be $n = C_1 \cdot C_2 c \cdot C_{|C|}$ such ordered pairs, and there are P(n, n) = n! permutations of this set. We will also use l to refer to the trial sequence length, which is equivalent to |X|.

4.3 Formula

With the partition established, we are now able to develop the formula for counting the number of solutions, s, for an experimental design. For the simplest of designs (tier 1), the ordered pairs in X fully represent all possible trials. $C_B \neq \emptyset$, while every other set in the partition is empty. Every permutaion of X represents a unique trial sequence, therefore there are s = l! solutions to a tier 1 design.

Tier 2 designs introduce basic factors that are not in C, and are thus less constrained. In otherwords, starting with tier 2, $\overline{C}_{B_I} \neq \emptyset$. For a factor f, let |f| denote the number of levels that f has. For each $f \in \overline{C}_{B_I}$, f is fully independent; therefore, any of the |f| levels may be selected for each trial. In a sequence of l trials, we can again apply the multiplication principle to determine that there are $|f|^l$ possible combinations of the levels of f. If $|\overline{C}_{B_I}| > 1$, then we apply the multiplication principle repeatedly to determine the total number of combinations for all factors in \overline{C}_{B_I} . Applying the multiplication principle

one more time, we see that the total number of solutions for a tier 2 design is:

$$s = l! \cdot \prod_{i=1}^{|\overline{C}_{B_I}|} |f_i|^l$$

With tier 3 designs come derived factors. It is now possible that all sets in the partition are non-empty. The only sets in the partition that we have yet to consider are C_D , \overline{C}_{B_S} , and \overline{C}_D . The presence of factors in C_D does not alter the formula, as level selection for factors in C_D is still controlled by X (the crossing). \overline{C}_D also requires no special treatment as the level selection for its factors depends entirely on the levels selected for basic factors. (Whether in C_B , \overline{C}_{B_S} , or \overline{C}_{B_I} .) This leaves only \overline{C}_{B_S} , which will be our next focus.

Derived factors allow the user to provide an arbitrary predicate for each level in the factor. We have not solved the halting problem, so we will need to apply these predicates to different argument combinations in order to determine which inputs are acceptable for each trial. We define X_S to be the set of ordered pairs representing the crossing of factors in \overline{C}_{B_S} , also referred to as the *source crossing*. The next task is to determine, for every element in X, which elements of X_S are compatible using the user-defined predicates.

For every ordered pair $t \in X$, there is some number of items in t corresponding to derived factor levels, labeled $d_1, d_2, \cdots d_{|C_D|}$ comprising the set V_t . Each $d \in V$ is associated with a predicate¹, pred(d). For every $t \in X$, every $b \in X_S$ is consulted to see if the predicates for all $d \in V_t$ are satisfied². If any one of them is not satisfied, then b is not a compatible choice for t, and must be discarded from consideration for t. Once this process is complete, we are left with a list of subsets of X_S , in which the n^{th} subset indicates which elements of X_S are compatible with the n^{th} crossing in X.

More formally, let *S* be a list of items $J_1, J_2, \cdots J_{|X|}$. The following statements are true:

1.
$$\forall J \in S \mid J \subseteq X_S$$

2.
$$\forall t \in X, \forall d \in V_t \mid pred(d) \implies \top$$

¹These predicates, also called derivation functions, are expected to be deterministic. If they are non-deterministic, then inconsistent solutions may be produced as the predicates may be applied more than once to the same arguments. This could also be handled by ensuring that each predicate is only applied once to each argument combination and the results are saved.

 $^{^2}$ If X is very large, then testing all combinations could become intractable. However in practice, X is relatively small, typically |X| < 500. Additionally, SweetPea's current implementation relies on X being small enough for this to work, as a similar process is followed when generating the SAT encoding for derived factors.

Once we have generated *S*, we can apply the multiplication principle one more time to complete the formula for tier 1, 2, and 3 designs:

$$s = l! \cdot \prod_{i=1}^{|\overline{C}_{B_I}|} |f_i|^l \cdot \prod_{k=1}^{|S|} |J_k|$$

The total number of solutions *s* for an experimental design, is the product of the number of permutations of the crossed factors, the number of combinations of each independent factor, and the number of acceptable combinations of all source factors.

It is possible for the user to specify a crossing that is not satisfiable. Referring back to the Stroop example, the user may include all three factors, color, text, and congruent, in the crossing. This is unsatisfiable because some level combinations violate the rules of the derived factor. For example, color=red, text=blue, and congruent=yes is invalid because red and blue are not congruent according to the predicates for congruent. If such a crossing is specified, this will correctly identify the number of potential solutions as zero.

4.4 Example

Consider a design composed of the same factors as the Stroop example:

```
color = Factor("color", ["red", "green", "blue"])
text = Factor("text", ["red", "green", "blue"])

congruent = Factor("congruent?", [
    DerivedLevel("yes", WithinTrial(operator.eq, [color, text])),
    DerivedLevel("no", WithinTrial(operator.ne, [color, text]))
])
```

However, rather than crossing color and text, we will demonstrate an example in which color and congruent? are crossed. For this design, the partition of *D* becomes:

$$C_B = \{ ext{color}\}$$
 $C_D = \{ ext{congruent?}\}$ $\overline{C}_{B_S} = \{ ext{text}\}$ $\overline{C}_{B_I} = \emptyset$ $\overline{C}_D = \emptyset$

The crossing, or X, is the cartesian product of levels of color and congruent?. This is generated exhaustively.

$$X = \{(red, yes), (red, no), (green, yes), (green, no), (blue, yes), (blue, no)\}$$

The size of the crossing determines the sequence length, therefore l = |X| = 6. As a result, the first term, l!, becomes 6! = 720.

In this design, there are no basic independent factors outside the crossing: $\overline{C}_{B_I} = \emptyset$. Therefore there is no value for the second term.

For the third and final term, we need to first determine X_S , the cartesian product of levels of factors in \overline{C}_{B_S} . For this design, the only factor in \overline{C}_{B_S} is text. As a result, X_S is simply the levels in text:

$$X_S = \{red, green, blue\}$$

At this point, each element of the product of X and X_S must be checked for validity under the derived factor definitions in the design to determine the subsets of X_S that are valid for each element in X. In terms of this example, which level selections in X_S do *not* violate the definition of congruent? when paired with each level selection combination in X? These subsets comprise S as defined previously. Table 4.1 gives a visual representation of this process.

Table 4.1. Computing *S* for Solution Counting

		X_S		
X	red	green	blue	J
(red, yes)	√			{(red}
(red, no)		✓	✓	{(green, blue)}
(green, yes)		✓		{(green}
(green, no)	√		✓	{(red, blue)}
(blue, yes)			✓	{(blue}
(blue, no)	✓	✓		{(red, green)}

S is therefore:

$$S = \{(red), (green, blue), (green), (red, blue), (blue), (red, green)\}$$

With S defined, we may now compute the third term of the formula:

$$\prod_{k=1}^{|S|} |J_k| = 1 \cdot 1 \cdot 1 \cdot 2 \cdot 2 \cdot 2 = 8$$

This yields the final solution count of $s = 720 \cdot 8 = 5,760$. This count can also be confirmed by generating the corresponding SAT encoding for this design and applying a model counter to the generated CNF.

CHAPTER 5

SEQUENCE CONSTRUCTION

With an exact count of viable sequences for tier 1, 2, and 3 designs, a mapping from natural numbers to valid trial sequences is now a possibility. Such a mapping would prove valuable for preserving uniformity guarantees, as well as generating solutions more efficiently. This chapter presents an algorithm that guarantees uniformly sampled solutions to tier 1, 2, and 3 designs by randomly sampling natural numbers from the uniform distribution in the range of [0, s). Each sampled number is then used to generate the unique trial sequence associated with that number.

5.1 Enumerating Permutations and Combinations

There are several critical components of constructing a solution based on a natural number. In this section, we will review known methods for generating the n^{th} permutation or combination of a set, as well as mapping individual natural numbers to a unique set of numbers that covers the same space using modular arithmetic. All of these techniques will play a major role in solution construction.

5.1.1 Interval Mapping

Suppose we have n numeric intervals, each from $[0, r_n)$. Revisiting the multiplication principle, we can see that there are $m = \prod_{i=1}^{n} r_i$ unique combinations of individual values from each interval. If j is an number selected from [0, m), we can map j to a unique combination of interval value selections using the modulo and division operators. (This technique will still work even when $j \geq m$, but combinations will begin to repeat.)

We begin by computing $j \mod r_1$, and saving the result as the selection for the first interval. We then recompute j to be $j = \lfloor \frac{j}{r_1} \rfloor$ before moving on to the next range. This process is repeated for $r_2, r_3, ..., r_n$ until the full combination is developed. For example, if we had $U = \{3, 2, 4\}$ as the set of upper range bounds, then there are $3 \cdot 4 \cdot 2 = 24$ possible

combinations of interval values. Suppose we want to compute the interval selections for 17. This would be done as follows:

$$i = 17$$
 $R_1 = 17 \mod 3 = 2$ $i = \left\lfloor \frac{17}{3} \right\rfloor = 5$ $i = 5$ $R_2 = 5 \mod 4 = 1$ $i = \left\lfloor \frac{5}{4} \right\rfloor = 1$ $i = 1$ $R_3 = 1 \mod 2 = 1$

Thus $R = \{2,1,1\}$. Figure 5.1 gives a visual representation of this breakdown, as well as indicates the order in which each interval combination is selected. (TODO)

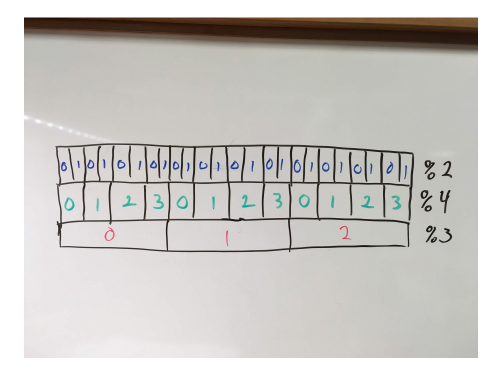


Figure 5.1. Decomposing a Number into Constituent Ranges

This technique is used repeatedly in the construction algorithm.

5.1.2 Permutations

Every permutation of a set can be identified by its *inversion* or *inversion sequence*, which loosely-defined, indicates how out-of-order the permutation is when compared with the

original sequence. The concept of an inversion was first introduced by Hall [9], though we will use on Brualdi's description [1] here.

Let S be the set of numbers $\{1, 2, ..., n\}$ Suppose we have some permutation p of S, $i_1, i_2, ..., i_n$. For each i, there exists some finite number of elements in S whose values are greater than i, yet precede i in p. This is called the inversion for i. An inversion sequence, $a_1, a_2, ...a_n$, is the sequence of inversions for a particular permutation of S.

For example, because the elements $\{1,2,3,4,5\}$ are in order, the corresponding inversion sequence for this arrangement is $\{0,0,0,0,0,0\}$. If we permute the sequence to obtain $\{2,5,4,1,3\}$, the associated inversion sequence is $\{3,0,2,1,0\}$. This can be interpreted to mean that 3 elements larger than 1 precede 1, 0 elements larger than 2 precede 2, 2 elements larger than 3 precede 3, 1 element larger than 4 precedes 4, and 0 elements larger than 5 precede 5. Notice that the final number in an inversion sequence must always be 0.

An inversion sequence can be used to generate the associated permutation by using the inversions to select where to place each element. Given an inversion sequence $a_1, a_2, ... a_n$, begin with n empty locations. Beginning with a_1 , skip a_i empty locations before inserting the i^{th} element of S. Using the previous example, this algorithm is demonstrated in Figure 5.2.

Finally, we can construct j^{th} inversion sequence of a set by applying the interval mapping technique introduced previously. The maximum value of each position in the inversion sequence is used to construct the intervals. To construct an inversion sequence of n inversions, the limits for the intervals are $\{n, n-1, n-2, ..., 2, 1\}$.

5.1.3 Combinations

We can enumerate combinations of a set of elements by again applying the interval mapping technique. For a combination of r elements, taken from a set of k elements, we know that there are k^r possible combinations by the multiplication principle. Applying interval mapping with r intervals, each from [0, k), suffices to enumerate each combination.

5.2 Construction Algorithm

With techniques established for counting solutions and constructing specifc permtutations and combinations, we are now ready to develop the full construction function. At a

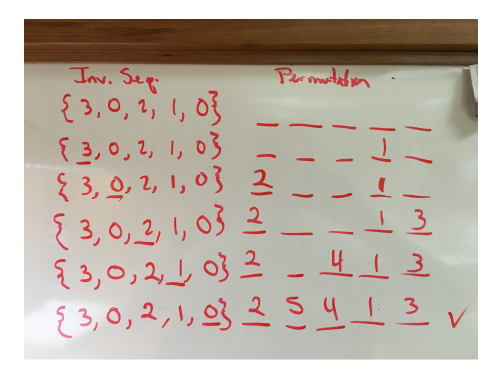


Figure 5.2. Construction of a Permutation from its Inversion Sequence

high level, the construction process has the following steps:

- 1. Determine the total number of solutions, *s*, for the design.
- 2. Randomly sample an integer, i, from the uniform distribution in the range [0, s).
- 3. Use interval mapping to convert i into a unique combination of value selections, R, that identify each component of the final solution.
- 4. Convert each selected value in *R* to the associated factors and levels. Record them as part of the final solution.
- 5. Compute the selected values for factors in \overline{C}_D using the previously selected levels for basic factors.

Accomplishing step 1 was the subject of chapter 4. We will not revisit that here. Step 2 relies on the randrange standard library function in Python 3, which produces uniformly-distributed values from an arbitrarily large range. Step 3 is the complicated step, particularly selecting the ranges for interval mapping.

5.2.1 Interval Mapping for an Experimental Design

For tier 1, 2, and 3 designs, as we saw in chapter 4, there is a partition of every design containing 5 sets: $\{C_B, C_D, \overline{C}_{B_S}, \overline{C}_{B_I}, \overline{C}_D\}$. In this section we will identify the upper bound of each interval to be used for interval mapping that cover all factors in the partition. We will use U to denote the set of these upper bounds. (The lower bounds are uninteresting, as they are always 0.

We first consider C_B and C_D . Recall that together, these sets contain all factors that are crossed with each other to form the crossing, labeled X in chapter 4. Recall also that |X| governs the length of a generated trial sequence: l = |X|, and that there are l! permtutations of the ordered pairs in X. Therefore, the first interval, which will determine which permutation of X to use, is l!. At this point, $U = \{l!\}$.

The next set in the partition is \overline{C}_{B_S} , which is the set of basic factors that are *not* included in C_B , but are sources of data for factors in C_D , and are therefore governed by the values associated with the chosen permutation for a given solution. Recall from chapter 4 that we defined X_S to represent the source crossing, and that there is a distinct subset of X_S , J_n , that represents the source crossings that are compatible with the n^{th} ordered pair in X. We also defined S to be the set of these subsets. We need a distinct interval for each of these subsets, so we add I values to I, one for the size of each subset in I0 will now contain I1 elements: I1 elements: I1 elements: I3 and I4 values to I5 one for the size of each subset in I5.

Lastly, we consider \overline{C}_{B_I} . (\overline{C}_D also remains in the partition, but values for factors in this set are governed by other basic factors, and will be determined in step 5. No additional intervals need be computed.) Recall that \overline{C}_{B_I} represents the set of basic factors that are completely independent, meaning they are not governed by any factors in C_D . Because they are completely independent, every possible l-combination of levels for each factor in \overline{C}_{B_I} is valid. Therefore an interval must also be added for each factor f with upper bound $|f|^l$. (Recall that |f| denotes the number of levels of factor f.) U is now complete:

$$U = \{l!, |J_1|, |J_2|, ..., |J_l|, |f_1|^l, |f_2|^l, ..., |f_i|^l\}$$

Where f_i represents the i^{th} factor in \overline{C}_{B_I} . Interestingly, the upper bounds in U are the same values which were multiplied together in chapter 4 to produce the total solution count.

5.2.2 Mapping Selected Values to Levels

Once a value has been produced for each interval, the next step is to produce the correct levels or level sequences for each factor associated with that interval. The selected permutation value is converted to a unique inversion sequence using interval mapping, which sequence is then used to construct the permutation. This permutation of integers is then converted into a sequence of level values by using each integer as an index into the ordered pairs in *X*.

The selected values for independent basic factors are converted in similar fashion. Each selected value identifies a combination number, which is converted to a set of values using interval mapping. Each value is an index into the list of levels for the associated factor.

Lastly, the selected values for source combinations must be converted. These are slightly more complex, as the viable combinations vary based on the selected levels for each trial. It would be incorrect to select a source combination from the n^{th} set of valid source combinations for the n^{th} trial, because the permutation may have shifted the position of the original n^{th} trial in the sequence. A mult-level indexing scheme is employed to ensure that the selected value is used to map exclusively to valid combinations, in which the permutated values are used to index into the correct set of valid source combinations. This is shown in Figure 5.3.

For example, if the selected permutation places the original crossing value for trial number 2 in the zeroth position in the sequence (trial 0), then the source combination for trial 0 must also be selected from the valid source combinations originally generated for trial number 2.

5.3 Example

Continuing on with the example from Chapter 4, we will construct a single solution to the design for which the total number of solutions was counted. We will choose a solution at random from [1,5,760], 2,590. First, we compute the range of value selections, R, for the $2,590^{th}$ solution. In order to do this, we first compute U, the set of upper-bounds for each range. The upper bound is the number of permutations, l!, for the design. For this example, this was determined to be 720 in chapter 4. Adding the sizes of the subsets in S, we determine $U = \{720, 1, 1, 1, 2, 2, 2\}$. Because $\overline{C}_{B_1} = \emptyset$ and $\overline{C}_D = \emptyset$ in this example, there

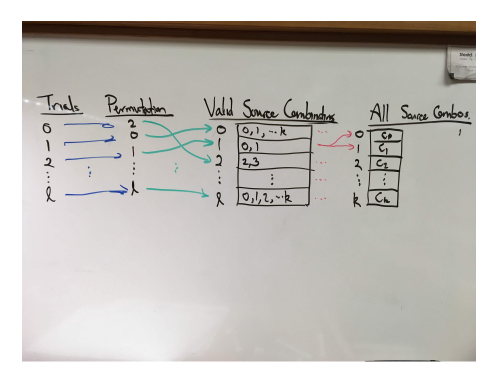


Figure 5.3. Selecting Valid Source Combinations

are no other bounds to compute. When interval mapping is applied to 2,590 to generate values from these ranges, R is determined to be $R = \{430, 0, 0, 0, 1, 1, 0\}$. With R computed, we can now use the selected values to generate the correct permutation, and then select level combinations for the remaining factors.

To generate the 430^{th} permutation, we first construct the 430^{th} inversion sequence by again applying interval mapping with $U = \{6,5,4,3,2,1\}$. This yields $\{4,1,2,0,1,0\}$ as the 430^{th} inversion sequence. Applying the algorithm to convert this sequence to its corresponding permutation, we obtain $\{3,1,5,2,0,4\}$. Recall from chapter 4 that:

$$X = \{(red, yes), (red, no), (green, yes), (green, no), (blue, yes), (blue, no)\}$$

The permutation $\{3, 1, 5, 2, 0, 4\}$ therefore corresponds to:

```
{(green, no), (red, no), (blue, no), (green, yes), (red, yes), (blue, yes)}
```

5.4 Handling Constraints

Recall that the complexity gradation did not account for the external constraints that may be applied to a design. Such constraints typically enforce some kind of rule regarding repetition of particular characteristics. For example, in the design for Stroop experiments, it is common to apply a constraint that requires that the generated sequence never contain two or more congruent trials in sequence. Constraints may be applied to the levels of derived factors as well, therefore the use may impose practically any condition upon the generated sequence.

In the future, the counting and construction algorithms could be enhanced to account for these constraints. This will be discussed further in Chapter 7. In the meantime however, basic rejection sampling is applied ensure that no generated sequences that violate constraints are ever returned to the user. The generated sample is checked against all repetition constraints in the design to ensure that none are violated. More details will be provided in the benchmarks section.

5.5 Benchmarks

Of the ten benchmarks designs that were introduced in chapter 2, 4 can be classified as tier 1, 2, or 3 designs: stroop-2, stroop-3, stroop-4, stroop-5. We will repeat the benchmarks for those designs here using this construction approach to sampling. We will also introduce a few additional designs involving variations on the Stroop design to demonstrate the scalability of this approach, and the empirical performance of the rejection sampling phase. Table 5.1 reviews the basic experiment data for each benchmark design. Table 5.2 summarizes the time required to generate 10,000 samples for each design using a single core, as well as the average number of rejections per sample. (Rejctions caused by constructing a trial sequence that violated external constraints.)

For even large designs, this approach allows a single random sample to be constructed extremely quickly.

Generating a large number of samples can still take a significant amount of time. However, because each sample can be generated fully independently, this algorithm can be easily parallelized for a linear speedup in the number of available cores. This will be discussed further in Chapter 7.

 Table 5.1. Benchmark Experiments for Sequence Construction

Table 511. Benefittativ Experiments for bequence Construction						
Experiment Name	Sequence Length	Unconstrained Solutions	Single Sample			
stroop-2	4	24	0.0009s			
stroop-3	9	362,880	0.003s			
stroop-4	16	20,922,789,888,000	0.007s			
stroop-5	25	$pprox 2^{83}$	0.016s			
stroop-10	100	$pprox 2^{524}$	0.170s			
stroop-20	400	$\approx 2^{2886}$	2.729s			
stroop-20-extra-constraints	400	$pprox 2^{2886}$	3.234s			

 Table 5.2. Sampling Metrics

Experiment Name	10,000 Samples	Avg. Rejections/Sample	
stroop-2	8.763s	0.982	
stroop-3	29.323s	1.393	
stroop-4	72.882s	1.538	
stroop-5	155.699s	1.589	
stroop-10	1908.598s	1.698	
stroop-20	26064.417s	1.684	
stroop-20-extra-constraints	27338.002s	1.898	

CHAPTER 6

FUTURE WORK

This paper has presented an alternative approach to constructing trial sequences for a subset of experimental designs based on a bijection between the natural numbers and viable trial sequences. This approach is beneficial in that sampled sequences are guaranteed to be uniformly distributed, and sequences can be constructed nearly instantaneously. This is an improvement over sampling solutions to the current SAT encoding. However, there are many ways in which it could be improved, and other alternatives that should be researched.

The counting and construction methods presented here may be applied to tier 1, 2, and 3 designs as defined in chapter 3, yet they do not account for external constraints that may be specified as part of the design. They only ensure that no internal constraints (between basic and derived factors) are violated. While rejecting sampling is sufficient in practice for some designs, one can construct design constraints in which valid solutions are so sparse as to render rejection sampling insufficient. Further research to incorporate constraints in the counting and construction process would be worthwhile. Combinatoric techniques for generating permutations with prohibited patterns would be likely be applicable to this process.

Additionally, the existing methods cannot yet construct sequences for designs beyond tier 3 which include derived factors spanning multiple trials. (Complex windows) The approaches presented here could reasonably be extended to apply to tier 4 designs (which allow Transition windows) with small adjustments. Increasing their capability to hand tier 5 designs would also be worthwhile, though significantly more difficult due to potential overlap and staggering between different derived factors. It likely that rejection sampling could continue to handle constraints even for these advanced designs, if the construction methods could be discovered. Lastly, the construction method can be fully

parallelized for a linear speedup in the number of available cores.

There are other approaches that could also be considered which do not rely wholly upon direct construction. Various hybrid approaches have been considered, but not thoroughly researched. For example, a numeric construction approach could be used to construct a partial trial sequence, which could then be completed using a SAT sampler, once the solution space had been sufficiently reduced. Difficulties may encountered in preserving uniforminty guarantees in this approach, as a partially constructed sequence may overprune the remaning search space, leading to biased results.

Lastly, the SAT encoding itself may be altered. The naive encoding used currently is easy to understand and reason about, but is obviously inefficient. If we had more experience with other types of encoding schemes, an alternative may be discovered that could perform better. Such an encoding would still likely need to avoid encoding all permutations as distinct solutions to avoid the explosion. A hybrid approach, the inverse of the previous paragraph, would likely perform well so long as the size of the independent support set were kept small. A SAT encoding could be developed to partially construct a sequence, including sampling the most complex aspects such as derived factors and constraints, while the remaining, simpler, values could be populated using a construction approach, thus reducing the burden on the SAT tooling. In this same realm, SweetPea could also begin utilizing the native XOR support offered by CryptoMiniSat, which may offer performance improvements.

CHAPTER 7

CONCLUSION

The current implementation of SweetPea, a language for experimental design, provides a useful standard for defining experimental designs, but has as yet failed to achieve its objective of providing uniformly sampled trial sequences for realistic experimental designs. This failure is due primarily to the inability of current SAT sampling tools to cope with massive solution spaces. While SAT solvers are still useful for generating non-uniform solutions to SweetPea design's, we have not yet been able to achieve the uniformity guarantees with the current SAT encoding.

This paper has presented the background information and benchmarks demonstrating these problems, and has introduced a complexity gradation for discussing experimental designs of varying complexity. It also introduced methods for enumerating arbitrary solutions to designs that can be classified below tier 4. This allows efficient generation of samples which are guaranteed to be uniformly distributed for these classes of designs. Rejection sampling is used to ensure that no generated sequences that violate design constraints are returned to the user.

While not yet able to solve the problem for all experimental designs, this approach is extremely beneficial for simple designs. It also provides a framework and pattern upon which to build for more complicated designs. We expect that these methods can be extended to generate uniformly distributed samples for all experimental designs in efficient time and space.

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