

DEPARTMENT OF COMPUTER SCIENCE

TDT4900 — MASTER'S THESIS

Exploring Matroids in Fair Allocation: Building the Matroids.jl Library

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 $\mathrm{May}\ 22,\ 2023$

In matroid theory, we explore A structure with properties galore From bases to circuits It never discourages Mathematicians always wanting more

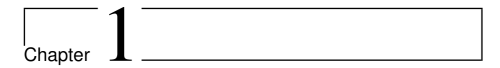
The axioms it holds are so grand And its applications vast and unplanned From optimization to graphs It can solve so many tasks Matroid theory, truly a wonderland

ChatGPT

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Introduction

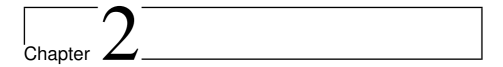
Fair allocation is the problem of *fairly* partitioning a set of resources among individuals with different preferences over these resources. This has been a hot topic of interest since antiquity (the 2000-year old Babylonian Talmud includes a discussion on how to distribute the estate of a deceased debtor among his creditors, for example [3]), and remains so today. As societies are faced with rising economical and environmental pressures have to do more with less, the problem of achieving fair and efficient allocations will remain central.

The mathematical study of fair allocation started with Hugo Steinhaus as late as 1948 [20], and for decades the focus was largely on the *divisible* case, in which the resources can be divided into arbitrary small pieces, and envy-free allocations (in which no agent values another agent's bundle higher than her own) always exist [2]. More recently, fair allocation of *indivisible* goods has garnered the attention of computer scientists, who have brought algorithmic techniques to the field to great effect.

- Computer science offers a fresh angle to further the research agenda for indivisible fair allocation
- 2. EF, EFX, egalitarian welfare, Nash welfare all NP-hard...
- 3. ...in the general, additive case. Reduced problem instances!
- 4. Matroids are great, Yankee Swap is really fair...
 - strong possibility results, polynomial-time computable

5. ...but little tooling exists for empirical working

One goal for this project is to create a library for the Julia programming language [5], supplying functionality for generating and interacting with (random) matroids. Throughout the text I will refer to this library as Matroids.jl. In the preparatory project delivered fall of 2022, I implemented Knuth's 1974 algorithm for the random generation of arbitrary matroids via the erection of closed sets [17]. With this, I was able to randomly generate matroids of universe sizes $n \leq 12$, but for larger values of n my implementation was unbearably slow.



Preliminary matroid theory

For simplicity, we also assume that every point in a geometry is a closed set. Without this additional assumption, the resulting structure is often described by the ineffably cacaphonic term "matroid", which we prefer to avoid in favor of the term "pregeometry".

Gian-Carlo Rota [8]

If a mathematical structure can be defined or axiomatized in multiple different, but not obviously equivalent, ways, the different definitions or axiomatizations of that structure make up a cryptomorphism. The many obtusely equivalent definitions of a matroid are a classic example of cryptomorphism, and belie the fact that the matroid is a generalization of concepts in many, seemingly disparate areas of mathematics.

Matroids were first introduced by Hassler Whitney in 1935 [24], in a seminal paper where he described two axioms for independence in the columns of a matrix, and defined any system obeying these axioms to be a "matroid" (which unfortunately for Rota is the term that has stuck). Whitney's key insight was that this abstraction of "independence" is applicable to both matrices and graphs. As a result of this, the terms used in matroid theory are borrowed from analogous concepts in both graph theory and linear algebra. Matroids have also

received attention from researchers in fair allocation, as their properties make them useful for modeling user preferences; for instance, matroid rank functions are a natural way of formally describing course allocation for students [4].

Independent sets

The most common way to characterize a matroid is as an *independence system*. An independence system is a pair (E, \mathcal{I}) , where E is the ground set of elements, $E \neq \emptyset$, and \mathcal{I} is the set of independent sets, $\mathcal{I} \subseteq 2^E$. The *dependent sets* of a matroid are $2^E \setminus \mathcal{I}$.

In practice, the ground set E represents the universe of elements in play, and the independent sets of typically represent the legal combinations of these items. In the context of fair allocation, the independent sets represent the legal (in the case of matroid constraints) or desired (in the case of matroid utilities) bundles of items.

A matroid is an independence system with the following properties [24]:

- (1) If $A \subseteq B$ and $B \in \mathcal{I}$, then $A \in \mathcal{I}$.
- (2) If $A, B \in \mathcal{I}$ and |A| > |B|, then there exists $e \in A \setminus B$ such that $B \cup \{e\} \in S$.
- (2') If $S \subseteq E$, then the maximal independent subsets of S are equal in size.

Properties (2) and (2') are equivalent. To see that (2) \Longrightarrow (2'), consider two maximal subsets of S. If they differ in size, (2) tells us that there are elements we can add from one to the other until they have equal cardinality. We get (2') \Longrightarrow (2) by considering $S = A \cup B$. Since |A| > |B|, they cannot both be maximal, and some $e \in A \setminus B$ can be added to B to obtain another independent set.

When S=E, (2') gives us a fundamental property of the *bases*, or maximal independent sets of a matroid, namely that all bases are of the same size. This is the rank of the matroid.

Rank

Given a matroid $\mathfrak{M}=(E,\mathcal{I})$, the matroid rank function (MRF) is a function $r:2^E\to\mathbb{Z}^+$ that gives the rank of a set $A\subseteq E$, defined to be the size of the largest independent subset of A. Formally,

$$r(A) = \max\{|X| : X \subseteq A \text{ and } X \in \mathcal{I}\}.$$

Matroid rank functions are binary submodular. Binary because they have binary marginals, that is, $r(A \cup \{e\}) - r(A) \in \{0,1\}$, for all $A \subseteq 2^E$ and $e \in E$. Submodularity refers to rank functions' natural diminishing returns property, namely that for any two sets $X, Y \subseteq E$, we have

$$r(X \cup Y) + r(X \cap Y) \le r(X) + r(Y).$$

Every binary submodular function is the rank function of some matroid [23]. The diminishing returns property makes the rank function useful for modeling user preferences, as we will see in fair allocation with matroidal valuations.

Closed sets

We also need to establish the concept of the *closed sets* of a matroid. A closed set is a set whose cardinality is maximal for its rank. Equivalently to the definition given above, we can define a matroid as $\mathfrak{M} = (E, \mathcal{F})$, where \mathcal{F} is the set of closed sets of \mathfrak{M} , satisfying the following properties [17]:

- 1. The set of all elements is closed: $E \in \mathcal{F}$
- 2. The intersection of two closed sets is a closed set: If $A,B\in\mathcal{F},$ then $A\cap B\in\mathcal{F}$
- 3. If $A \in \mathcal{F}$ and $a, b \in E \setminus A$, then b is a member of all sets in \mathcal{F} containing $A \cup \{a\}$ if and only if a is a member of all sets in \mathcal{F} containing $A \cup \{b\}$

The closure function is the function $cl: 2^E \to 2^E$, such that

$$cl(S) = \{x \in E : r(S) = r(S \cup \{x\})\}.$$

That is to say, the closure function, when given a set $S \subseteq E$, returns the set of elements in $x \in E$ such that x can be added to S with no increase in rank. It returns the closed set of the same rank as S, that contains S.



Generating matroids

The overarching goal for this project is to make Matroids.jl, a proof-of-concept library for working programmatically with matroids, specifically in the context of fair allocation. This chapter covers how Matroids.jl enables the creation of specific matroids and the generation of random ones, as well as how to access important properties such as independent sets, closed sets, circuits, bases, the rank function and the closure function. The first part of the chapter focuses on implementing these features for various types of matroids, including uniform, linear (vector), graphic, and partition matroids. The final part of the chapter is a significant portion of the thesis as a whole, and describes the implementation of Knuth's interesting algorithm [17] for the erection of arbitrary rank-r matroids.

3.1 Uniform matroids and partition matroids

A uniform matroid U_n^r is the matroid over n elements where the independent sets are exactly the sets of cardinality at most r. The free matroid $U_n^n = (E, 2^E)$ is a special case of the uniform matroid and is the simplest, biggest and least interesting type of matroid, being the trivial case in which every subset of E is an independent set. In Matroids.jl, we represent uniform matroids with a simple struct.

struct UniformMatroid
n::Integer

```
r::Integer
end
FreeMatroid(n) = UniformMatroid(n, n)
```

3.2 Linear matroids

3.3 Graphic matroids

We begin with defining the graph theory terms used in this section. An undirected graph G = (V, E) is said to be connected if there exists at least one path between each pair of nodes in the graph; otherwise it is disconnected. A disconnected graph consists of at least two connected subsets of nodes. These connected subgraphs are called components. A tree is a connected acyclic graph, and a forest is a disconnected graph consisting of some number of trees. A spanning tree of G is a subgraph with a unique simple path between all pairs of vertices of G. A spanning forest of G is a collection of spanning trees, one for each component. The degree of a node v is the number of edges for which v is an endpoint. A regular graph is a graph in which all nodes have the same degree. An induced subgraph G[S], where S is either a subset of the nodes of G (in which case G[S] is a node-induced subgraph) or of the edges of G (edge-induced).

Given a graph G = (V, E), let $\mathcal{I} \subseteq 2^E$ be the family of subsets of the edges E such that, for each $I \in \mathcal{I}$, (V, I) is a forest. It is a classic result of matroid theory that $\mathfrak{M} = (E, \mathcal{I})$ is a matroid [19, p. 657]. To understand how, we will show that it adhers to axioms (1) and (2'), as given in Section 2. (1) holds trivially, as all subsets of a forest are forests. To see that (2') holds, consider the bases (maximal independent sets) $\mathcal{B} \subseteq \mathcal{I}$. By definition, each basis $B \in \mathcal{B}$ is a maximal forest over G. Since a spanning tree of a graph with n nodes must needs have n-1 edges, we have |B| = |V| - k, where k is the number of components of G. This is the same for every $B \in \mathcal{B}$, which proves property (2'). Any matroid given by a graph G, denoted by $\mathfrak{M}(G)$, is called a graphic matroid.

3.3.1 Random graphs

Since generating random graphic matroids will require us to generate random graphs, let us take a look at some of the options available to us for this. Luckily for us, random graphs has been an area of extensive study for more than sixty years, and several models with different properties exist.

The Erdős-Rényi (ER) model (also known as Erdős-Rényi-Gilbert [11]) picks uniformly at random a graph from among the $\binom{n}{2}$ possible graphs with n nodes and M edges, or, alternatively, constructs a graph with n nodes where each edge is present with some probability p [9, 13]. This model produces mostly disconnected graphs, and the size distribution of its components with respect to the number of edges has been studied extensively. With n nodes and fewer than $\frac{n}{2}$ edges, the resulting graph will almost always consist of components that are small trees or contain at most one cycle. As the number of edges exceeds $\frac{n}{2}$, however, the so-called "giant" component of size $\mathcal{O}(n)$ emerges, and starts to absorb the smaller components [16]. The ER model is the oldest and most basic random graph model, and is often referred to simply as the random graph, denoted by G(n, p).

Variations of the ER model have been developed by physicists and network scientists to produce phenomena commonly seen in real-world networks [11]. These variations include the Barabási-Albert model, which grows an initial connected graph using preferential attachment (a mechanism colloquially known as "the rich get richer"), in which more connected nodes are more likely to receive new connections. This results in graphs in which a small number of nodes ("hubs") have a significantly higher degree than the rest, creating a power-law distribution of node degrees. This property is known as scale-freeness and is thought to be a characteristic of the Internet [1].

Another approach is the Watts-Strogatz model, which starts with a ring lattice, a regular graph with n nodes, each with degree k, and then rewires each edge with some probability p. By changing p, one is able to 'tune' the graph between regularity (p=0) and disorder (p=1). For intermediate values of p, Watts-Strogatz produces so-called "small-world" graphs, which exhibit both a high degree of clustering (how likely two nodes with a common neighbor are to be adjacent), and short average distance between nodes. This phenomenon is found in many real-world networks, such as social systems or power grids [22].

3.3.2 Properties of random graphic matroids

We will use the Graphs.jl library [10] for handling graphs in Matroids.jl. This library has built-in methods for the random graph models described in the previous chapter¹.

When constructing matroids, we want to be able to specify the size of the ground set, and perhaps also the rank of the matroid. Let us see how we

¹https://docs.juliahub.com/Graphs/VJ6vx/1.4.1/generators/

can achieve this with the random graph models we have discussed. The method barabasi_albert (n, k) generates a Barabási-Albert model random graph with n nodes. It starts with an initial graph of k nodes, and adds the remaining n-knodes one at a time, each new node receiving k edges via preferential attachment. Thus, the final graph has |E| = (n-k)k edges. To specify a matroid with m edges, we pick some k|m and solve for n. Remember that the rank of a graphic matroid is the size of a spanning tree over the graph, which is n-1 when the graph is connected. If we select a smaller k from among the factors of |E|, we get a larger final rank, and vice versa. We can generate a Watts-Strogatz model random graph with the method watts_strogatz(n, k, β), where n is the number of nodes, k the node degree and β the probability of rewiring. The number of edges of a regular graph with n nodes and degree k (and thus the size of the ground set of the induced graphic matroid) is given by $\frac{nk}{2}$, so nk must be even. Erdős-Rényi is the simplest model for our purposes, as the method erdos_renyi (nv, ne) simply takes in the desired number of nodes and edges. However, since the resulting graph has a large number of components for $|E| < \frac{n}{2}$, we have less fine-grained control over the final rank of the induced graphic matroid.

In Matroids.jl, we "generate" a graphic matroid by simply accepting some graph, and figure out the rank of the matroid using Kruskal's algorithm for maximal spanning forests, which runs in $\mathcal{O}(|E|\lg|E|)$ time [6]. Implementing the methods for finding the properties of our graphic matroids is simple, as they reduce to well-known algorithms (implemented by Graphs.jl) for finding the properties of the graphs they are derived from.

```
using Graphs
struct GraphicMatroid
g::Graph
n::Integer
r::Integer
GraphicMatroid(g::Graph) = new(g, ne(g), length(kruskal_mst(g)))
end
```

The rank function returns the size of a spanning forest of the subgraph induced by some subset of the edges. This is the rank of that subset. Thus, the rank of a subset $S \subseteq E$ can be found in $\mathcal{O}(|S| \lg |S|)$ time (when the MST is found using Kruskal's algorithm).

```
function rank(m::GraphicMatroid, S)
```

```
edgelist = [e for (i, e) in enumerate(edges(g)) if i in S]
subgraph, _vmap = induced_subgraph(m.g, edgelist)
return length(kruskal_mst(subgraph))
end
```

The indepence oracle returns whether the subgraph induced by a supplied subset of edges is acyclic. While independence can also be determined with the rank function, by checking whether the cardinality of a set equals its rank, this uses a DFS behind the scenes², which runs in linear time [6].

```
function is_indep(m::GraphicMatroid, S)
  edgelist = [e for (i, e) in enumerate(edges(g)) if i in S]
  subgraph, _vmap = induced_subgraph(m.g, edgelist)
  return !is_cyclic(subgraph)
end
```

The circuit oracle

```
function is_circuit(m::GraphicMatroid, S)
#TODO
end
```

The closure function accepts a set of elements S, and returns the largest set of elements cl(S) such that $S \subseteq cl(S) \subseteq E, r(S) = r(cl(S))$. In a graph context, given a graph G = (V, E) and an edge-induced subgraph $G[S] = (V, S), S \subseteq E$, this is the same as finding the largest edge-induced subgraph $G[T], S \subseteq T \subseteq E$, in which a spanning tree has the same number of edges as one in G[S]. Since the size of a spanning tree in G[S] is given by |V| - 1, G[T] cannot contain any edges to nodes not in V, as this would increase the rank of G[T]. Therefore, we get that the closure of S is the largest set T of edges between nodes that are present in the edge-induced subgraph G[S]. The method closure below returns the set of all edges whose endpoints are both located in the subgraph induced by S.

```
function closure(m::GraphicMatroid, S)
  edgelist = [e for (i, e) in enumerate(edges(m.g)) if i in S]
  _sg, vmap = induced_subgraph(m.g, edgelist)
  return [e for e in edges(m.g) if [e.src, e.dst] ⊆ vmap]
end
```

 $^{^2} https://docs.juliahub.com/Graphs/VJ6vx/1.4.1/pathing/\#Graphs.is_cyclical control of the con$

3.4 Matroid erection

Before we delve into Knuth's general matroid construction, we need to establish a few concepts. The rank-k truncation of a matroid $\mathfrak{M} = (E, \mathcal{I})$, is the matroid $\mathfrak{M}^{(k)} = (E, \mathcal{I}^{(k)})$, where

$$\mathcal{I}^{(k)} = \{ I \in \mathcal{I} : |I| \le k \}.$$

When \mathfrak{M} is of rank r, its truncation is given as $T(\mathfrak{M})=\mathfrak{M}^{(r-1)}$. As a simple example, we have that the uniform matroid $U_n^{n-1}=T(\mathfrak{F}_n)$, where \mathfrak{F}_n is the free matroid with n elements. The erection of the matroid \mathfrak{M} is the matroid \mathfrak{N} such that $\mathfrak{M}=T(\mathfrak{N})$ [7]. A matroid can have many erections. For example, it is easy to see that U_n^4 is an erection of U_n^3 , since U_n^4 by definition has all the same independent sets of the rank-3 matroid U_n^3 , along with all size-4 subsets of E. However, let \mathfrak{U} be the rank-4 matroid over E where every subset of E of rank ≤ 3 is independent, and which has one size-4 rank-3 dependent set. The rank-3 independent sets of \mathfrak{U} are the same as those of U_n^3 , so we have $U_n^3 = T(\mathfrak{U})$, however the introduction of a size-4 dependent set of rank 3 has erected a different rank-4 matroid.

3.5 Knuth's matroid construction

In the preparatory project delivered fall of 2022, I implemented Knuth's 1974 algorithm for the random generation of arbitrary matroids via the erection of closed sets [17]. With this, I was able to randomly generate matroids of universe sizes $n \leq 12$, but for larger values of n my implementation was unbearably slow. In this section, Knuth's method for random matroid construction will be described, along with the steps I have taken to speed up my initial, naïve implementation.

KNUTH-MATROID (given in Algorithm 1) accepts the ground set E and a list X such that $X[i] \subseteq 2^E$, and produces the rank-r matroid \mathfrak{M} such that $\operatorname{rank}(X) = k$ for each $X \in X[k]$. This is done in a bottom-up manner through r sequential erections starting from the empty rank-0 matroid, $\mathfrak{M}^{(0)}$, each iteration i producing the erection $\mathfrak{M}^{(i+1)}$ from $\mathfrak{M}^{(i)}$ and X[i]. The algorithm outputs the tuple (E, F), where $F = [F_0, \ldots, F_r]$, r being the final rank of \mathfrak{M} and F_i the family of closed sets of rank i. In the paper, Knuth shows that $\bigcup_{i=0}^r F[r] = \mathcal{F}$, where \mathcal{F} is the set of closed sets of a matroid, and so the algorithm produces a valid matroid represented by its closed sets.

Algorithm 1 KNUTH-MATROID(E, X)

Input: The ground set of elements E, and a list of enlargements X.

Output: The list of closed sets of the resulting matroid grouped by rank, $F = [F_0, \ldots, F_r]$, where F_i is the set of closed sets of rank i.

```
1 r = 0, F = [\{\emptyset\}]
   while TRUE
2
        Push!(F, Generate-Covers(F, r, E))
3
        F[r+1] = F[r+1] \cup X[r+1]
4
        Superpose!(F[r+1], F[r])
5
        if E \not\in F[r+1]
6
             r \leftarrow r + 1
7
8
        else
9
            return (E, F)
```

To understand the procedure, let us investigate what Algorithm 1 does at iteration 1 < i < r, where r is the final rank \mathfrak{M} , the matroid under construction. At iteration i, we produce a rank-(i+1) erection $\mathfrak{M}^{(i+1)}$ of $\mathfrak{M}^{(i)}$, which is represented by its closed sets $F = [F_0, F_1, ..., F_i]$, where F_i is the set of closed sets of rank i. We want to produce the set F_{i+1} of rank-r closed sets of an erection of $\mathfrak{M}^{(i)}$ such that each $X \in X[i]$ is contained in some rank-r closed set. First we find the "covers" of each closed set in F_i . The covers of a closed set A of rank r are the sets obtained by adding one more element from E to A. The covers are generated with GENERATE-COVERS(F, r, E).

```
GENERATE-COVERS(F, r, E) 
1 return \{A \cup \{a\} : A \in F[r], a \in E \setminus A\}
```

Given no enlargements $(X[i] = \emptyset)$, the resulting matroid $\mathfrak{M}^{(i+1)}$ is the free erection of $\mathfrak{M}^{(i)}$, and there are no essential closed sets in F_{i+1} . Arbitrary matroids can be generated by supplying different lists X. When enlarging, the sets in X[r+1] are simply added to F[r+1], before Superpose! is run to ensure that the newly enlarged family of closed sets of rank r+1 is valid (ie. in accordance with the closed set axioms given in Section 2). If F_{r+1} contains two sets A, B whose intersection $A \cap B \not\subseteq C$ for any $C \in F_r$ (in other words, their intersection

is not a closed set), replace A, B with $A \cup B$. Repeat until no two sets exist in F_{r+1} whose intersection is not contained within some set $C \in F_r$.

```
Superpose!(F_{r+1}, F_r)
     for A \in F_{r+1}
 1
 2
            for B \in F_{r+1}
                   flag \leftarrow TRUE
 3
                   for C \in F_r
 4
                         if A \cap B \subseteq C
 5
                                flag \leftarrow FALSE
 6
 7
 8
                   if flag = TRUE
                         F_{r+1} \leftarrow F_{r+1} \setminus \{A, B\}
 9
                         F_{r+1} \leftarrow F_{r+1} \cup \{A \cup B\}
10
```

3.5.1 Randomized KMC

In the randomized version of KNUTH-MATROID, we generate matroids by applying a supplied number of random coarsening steps, instead of enlarging with supplied sets. This is done by applying SUPERPOSE! immediately after adding the covers, then choosing a random member A of F[r+1] and a random element $a \in E \setminus A$, replacing A with $A \cup \{a\}$ and finally reapplying SUPERPOSE!. The parameter $p = (p_1, p_2, \ldots)$ gives the number of such coarsening steps to be applied at each iteration of the algorithm.

The pseudocode given up to this point corresponds closely to the initial Julia implementation, which can be found in Appendix ??. It should already be clear that this brute force implementation leads to poor performance – for instance, the Superpose! method uses a triply nested for loop, which should is a candidate for significant improvement. Section 3.5.2 describes the engineering work done to create a more performant implementation.

3.5.2 Improving performance

When recreating Knuth's table of observed mean values for the randomly generated matroids, some of the latter configurations of n and $(p_1, p_2, ...)$ was unworkably slow, presumably due to my naïve implementation of the algorithm. Table 3.1 shows the performance of this first implementation.

Algorithm 2 RANDOMIZED-KNUTH-MATROID(E, p)

Input: The ground set of elements E, and a list $p = [p_1, p_2, ...]$, where

 p_r is the number of coarsening steps to apply at rank r in the

construction.

Output: The list of closed sets of the resulting matroid grouped by rank, $F = [F_0, ..., F_r]$, where F_i is the set of closed sets of rank i.

```
1 r = 0, F = [\{\emptyset\}]
    while TRUE
         Push!(F, Generate-Covers(F, r, E))
 3
         Superpose!(F[r+1], F[r])
 4
         if E \in F[r+1] return (E,F)
 5
         while p[r] > 0
 6
 7
              A \leftarrow \text{a random set in } \mathbf{F}[r+1]
              a \leftarrow \text{a random element in } E \setminus A
 8
              replace A with A \cup \{a\} in F[r+1]
 9
              SUPERPOSE!(F[r+1], F[r])
10
              if E \in F[r+1] return (E,F)
11
              p[r] = p[r] - 1
12
13
         r = r + 1
```

The performance was measured using Julia's <code>@timed^3</code> macro, which returns the time it takes to execute a function call, how much of that time was spent in garbage collection and the number of bytes allocated. As is evident from the data, larger matroids are computationally quite demanding to compute with the current approach, and the time and space requirements scales exponentially with n. Can we do better? As it turns out, we can; after the improvements outlined in this section, we will be able to generate matroids over universes as large as n = 128 in a manner of seconds and megabytes.

³https://docs.julialang.org/en/v1/base/base/#Base.@timed

Table 3.1: Performance of random_kmc_v1.

\overline{n}	(p_1,p_2,\ldots)	Trials	Time	GC Time	Bytes allocated
10	(0, 6, 0)	100	0.0689663	0.0106786	147.237 MiB
10	(0, 5, 1)	100	0.1197194	0.0170734	$251.144~\mathrm{MiB}$
10	(0, 5, 2)	100	0.0931822	0.0144022	$203.831~\mathrm{MiB}$
10	(0, 6, 1)	100	0.0597314	0.0094902	$132.460~\mathrm{MiB}$
10	(0, 4, 2)	100	0.1924601	0.0284532	$406.131~\mathrm{MiB}$
10	(0, 3, 3)	100	0.3196838	0.0463972	$678.206~\mathrm{MiB}$
10	(0, 0, 6)	100	1.1420602	0.1671325	$2.356~\mathrm{GiB}$
10	(0, 1, 1, 1)	100	2.9283978	0.3569357	$5.250~\mathrm{GiB}$
13	(0, 6, 0)	10	104.0171128	9.9214449	$161.523~\mathrm{GiB}$
13	(0, 6, 2)	10	11.4881308	1.3777947	$20.888~\mathrm{GiB}$
16	(6, 0, 0)	1	-	-	-

Representing sets as binary numbers

The first improvement we will attempt is to represent our closed sets using one of Julia's Integer types of bit width at least n, instead of as a Set⁴ of elements of E. Appendix ?? contains all the code referenced in this chapter; the Julia implementation at this point can be found in ??.

The idea is to define a family of closed sets of the same rank as Set {UInt16}. Using UInt16 we can support ground sets of size up to 16. Each 16-bit number

 $^{^4} https://docs.julialang.org/en/v1/base/collections/\#Base.Set$

represents a set in the family. For example, the set $\{2, 5, 7\}$ is represented by

$$164 = 0 \times 0004 = 0 \times 00000000010100100 = 2^7 + 2^5 + 2^2.$$

At either end we have $\emptyset \equiv 0$ x0000 and $E \equiv 0$ xffff (if n = 16). The elementary set operations we will need have simple implementations using bitwise operations.

Set operation	Bitwise operation
$A \cap B$	A AND B
$A \cup B$	$A ext{ OR } B$
$A \setminus B$	A AND NOT B
$A \subseteq B$	A AND B = A

We can now describe the bitwise versions of the required methods. The bitwise implementation of Generate-Covers finds all elements in $E \setminus A$ by finding each value $0 \le i < n$ for which A & 1 << i === 0, meaning that the set represented by 1 << i is not a subset of A. The bitwise implementation of Superpose! is unchanged apart from using the bitwise set operations described above.

Table 3.2: Performance of random_kmc_v2.

\overline{n}	(p_1,p_2,\ldots)	Trials	Time	GC Time	Bytes allocated
10	[0, 6, 0]	100	0.0010723	0.0001252	$1.998~\mathrm{MiB}$
10	[0, 5, 1]	100	0.0017543	0.0001431	$3.074~\mathrm{MiB}$
10	[0, 5, 2]	100	0.0008836	0.0001075	$2.072~\mathrm{MiB}$
10	[0, 6, 1]	100	0.0007294	6.73 e-5	$1.700~\mathrm{MiB}$
10	[0, 4, 2]	100	0.0020909	0.0001558	$3.889~\mathrm{MiB}$
10	[0, 3, 3]	100	0.0024636	0.0002139	$4.530~\mathrm{MiB}$
10	[0, 0, 6]	100	0.007082	0.0004801	$9.314~\mathrm{MiB}$
10	[0, 1, 1, 1]	100	0.0132477	0.0008307	$17.806~\mathrm{MiB}$
13	[0, 6, 0]	10	0.042543	0.0014988	$31.964~\mathrm{MiB}$
13	[0, 6, 2]	10	0.0183313	0.0012176	$21.062~\mathrm{MiB}$
16	[0, 6, 0]	10	1.2102877	0.0146129	$450.052~\mathrm{MiB}$

The performance of random_kmc_v2 is shown in Table 3.2. It is clear that representing closed sets using binary numbers represents a substantial improvement – we are looking at performance increases of 100x-1000x across the board. Great stuff!

Sorted superpose

Can we improve the running time of our implementation further? It is clear that SUPERPOSE! takes up a large portion of the compute time. In the worst case, when no enlargements have been made, F_{r+1} is the set of all r+1-sized subsets of E, $|F_{r+1}| = \binom{n}{r+1}$. Comparing each $A, B \in F_{r+1}$ with each $C \in F_r$ in a triply nested for loop requires $\mathcal{O}(\binom{n}{r+1}^2\binom{n}{r})$ operations. In the worst case, no enlargements are made at all, and we build the free matroid in $\mathcal{O}(2^{3n})$ time (considering only the superpose step).

After larger closed sets have been added to F[r+1], SUPERPOSE! will cause sets to merge, so that only maximal dependent sets remain. Some sets will even simply disappear. In the case where $X=\{1,2\}$ was added by GENERATE-COVERS, and the $Y=\{1,2,3\}$ was added manually as an enlargement, the smaller set will be fully subsumed in the bigger set, as $\{1,2\} \cap \{1,2,3\} = \{1,2\}$ (which is not a subset of any set in F[r]) and $\{1,2\} \cup \{1,2,3\} = \{1,2,3\}$. In this situation, Y would "eat" the covers $\{1,3\}$ and $\{2,3\}$ as well. This fact is reflected in the performance data – compare the memory allocation differences between the 10-element matroid with p=[0,0,6] and the one with p=[0,6,0] in any of the performance tables in this section. Making enlargements at earlier ranks result in smaller matroids as more sets get absorbed.

```
function sorted_bitwise_superpose!(F, F_prev)
As = sort!(collect(F), by = s -> length(bits_to_set(s)))
while length(As) !== 0
A = popfirst!(As)

for B in setdiff(F, A)
    if should_merge(A, B, F_prev)
        insert!(As, 1, A | B)
        setdiff!(F, [A, B])
        push!(F, A | B)
        break
    end
end
end
return F
end
```

Since the larger sets will absorb so many of the smaller sets (around $\binom{p}{r+1}$), where p is the size of the larger set and r+1 is the size of the smallest sets allowed to be added in a given iteration), might it be an idea to perform the superpose operation in descending order based on the size of the sets? This should result in fewer calls to Superpose!, as the bigger sets will remove the smaller sets that fully overlap with them in the early iterations, however, the repeated sorting of the sets might negate this performance gain. This is the idea behind sorted_bitwise_superpose!, which was used in random_kmc_v3. The full code can be found in Appendix $\ref{eq:continuous}$?

Unfortunately, as Table 3.3 shows, this implementation is a few times slower and more space demanding than the previous implementation. This is might be due to the fact that an ordered list is more space inefficient than the hashmap-based Set.

\overline{n}	(p_1,p_2,\ldots)	Trials	Time	GC Time	Bytes allocated
10	[0, 6, 0]	100	0.0023382	0.0001494	4.042 MiB
10	[0, 5, 1]	100	0.001853	0.0001433	$4.383~\mathrm{MiB}$
10	[0, 5, 2]	100	0.0017845	0.0001341	$4.043~\mathrm{MiB}$
10	[0, 6, 1]	100	0.0015145	0.0001117	$3.397~\mathrm{MiB}$
10	[0, 4, 2]	100	0.0030704	0.0002125	$6.385~\mathrm{MiB}$
10	[0, 3, 3]	100	0.0037838	0.0002514	$7.018~\mathrm{MiB}$
10	[0, 0, 6]	100	0.008903	0.000557	14.159 MiB
10	[0, 1, 1, 1]	100	0.0142828	0.0008823	$21.838~\mathrm{MiB}$
13	[0, 6, 0]	10	0.0627633	0.002094	$51.492~\mathrm{MiB}$
13	[0, 6, 2]	10	0.0106478	0.0007704	$20.774~\mathrm{MiB}$
16	[0, 6, 0]	10	0.6070136	0.0095656	310.183 MiB

Iterative superpose

The worst-case $\mathcal{O}(\binom{n}{r+1}^2\binom{n}{r})$ runtime of Superpose! at step r is due to the fact that it takes in F after all covers and enlargements have been indiscriminately added to F[r+1] and then loops through to perform the superposition. Might

there be something to gain by inserting new closed sets into the current family one at a time, and superposing on the fly?

```
# Superpose (random_kmc_v4)
push!(F, Set()) # Add F[r+1].
while length(to_insert) > 0
A = pop!(to_insert)
push!(F[r+1], A)

for B in setdiff(F[r+1], A)

if should_merge(A, B, F[r])
    push!(to_insert, A | B)
    setdiff!(F[r+1], [A, B])
    push!(F[r+1], A | B)
    end
end
end
```

In random_kmc_v4, the full code of which can be found in Appendix ??, the covers and enlargements are not added directly to F[r+1], but to a temporary array to_insert. Each set A is then popped from to_insert one at a time, added to F[r+1] and compared with the other sets $B \in F[r+1] \setminus \{A\}$ and $C \in F[r]$ in the usual SUPERPOSE! manner. This results in fewer comparisons, as each set is only compared with the sets added before it; the first set is compared with no other sets, the second set with one other and the sets in F[r], and so on. The number of such comparisons is therefore given by the triangular number $T_{\binom{n}{r+1}}$, and so we should have roughly halved the runtime at step r. It is worth noting that this implementation of SUPERPOSE! uses a subroutine should_merge that returns early when it finds one set $C \in F[r]$ such that $C \supseteq A \cap B$, so in practice it usually does not require $\binom{n}{r}$ comparisons in the innermost loop.

Table 3.4 shows that the iterative superpose was a meaningful improvement. For most input configurations, it is a few times faster and a few times less space demanding than random_kmc_v2.

Rank table

While Superpose! is getting more efficient, it is still performing the same comparisons over and over again. Let's consider what we are really trying to achieve with this function, to see if we can't find a smarter way to go about it.

After adding the closed sets for a rank, Superpose! is run to maintain the closed set properties of the matroid (given in Section 2). These are maintained

Table 3.4: Performance of random_kmc_v4.

\overline{n}	(p_1,p_2,\ldots)	Trials	Time	GC Time	Bytes allocated
10	[0, 6, 0]	100	0.0014585	3.94 e-5	724.635 KiB
10	[0, 5, 1]	100	0.0007192	9.39 e-5	$659.729~\mathrm{KiB}$
10	[0, 5, 2]	100	0.0005943	3.53 e-5	$617.668~\mathrm{KiB}$
10	[0, 6, 1]	100	0.0003502	2.88e-5	$408.666~\mathrm{KiB}$
10	[0, 4, 2]	100	0.001013	5.36e-5	$887.618~\mathrm{KiB}$
10	[0, 3, 3]	100	0.0011847	5.03e-5	$1.003~\mathrm{MiB}$
10	[0, 0, 6]	100	0.0015756	9.7e-5	$1.066~\mathrm{MiB}$
10	[0, 1, 1, 1]	100	0.0046692	0.0001385	$2.455~\mathrm{MiB}$
13	[0, 6, 0]	10	0.0118201	0.0005486	$6.289~\mathrm{MiB}$
13	[0, 6, 2]	10	0.0075668	0.0002458	$4.666~\mathrm{MiB}$
16	[0, 6, 0]	10	0.2819294	0.0040792	$81.317~\mathrm{MiB}$
16	[0, 6, 1]	10	0.8268207	0.0070206	$154.451~\mathrm{MiB}$
16	[0, 0, 6]	10	95.1959596	0.0290183	$553.597~\mathrm{MiB}$

by ensuring that, for any two newly added sets $A, B \in F[r+1]$, there exists $C \in F[r]$ such that $A \cap B \subseteq C$. Up to this point, this has been done by checking if the intersection of each such A, B is contained in a set C of rank r. We remember that one of the properties of the closed sets of a matroid is that the intersection of two closed sets is itself a closed set. Therefore, we do not need to find a closed set C of rank r that contains $A \cap B$, since if A and B are indeed closed sets, their intersection will be equal to some closed set C of any rank C C. This insight leads us to the next improvement: if we keep track of all added closed sets in a rank table, then we can memoize SUPERPOSE! and replace the innermost loop with a constant time dictionary lookup.

```
# The rank table maps from the representation of a set to its assigned rank.
rank = Dict{T, UInt8}(0=>0)

[...]

# Superpose.
push!(F, Set()) # Add F[r+1].
while length(to_insert) > 0
A = pop!(to_insert)
```

```
push!(F[r+1], A)
  rank[A] = r

for B in setdiff(F[r+1], A)
  if !haskey(rank, A&B) || rank[A&B] >= r
    # Update insert queue.
    push!(to_insert, A | B)

# Update F[r+1].
    setdiff!(F[r+1], [A, B])
    push!(F[r+1], A | B)

# Update rank table.
    rank[A|B] = r
    break
  end
end
end
```

Table 3.5: Performance of random_kmc_v5.

\overline{n}	(p_1,p_2,\ldots)	Trials	Time	GC Time	Bytes allocated
10	[0, 6, 0]	100	0.0001335	0.0	138.966 KiB
10	[0, 5, 1]	100	0.0001436	0.0	$158.691~\mathrm{KiB}$
10	[0, 5, 2]	100	0.0001928	0.0	$167.487~\mathrm{KiB}$
10	[0, 6, 1]	100	0.0002204	0.0	$148.812~\mathrm{KiB}$
10	[0, 4, 2]	100	0.0001578	0.0	$173.455~\mathrm{KiB}$
10	[0, 3, 3]	100	0.0001743	0.0	$202.566~\mathrm{KiB}$
10	[0, 0, 6]	100	0.0003433	0.0	$431.089~\mathrm{KiB}$
10	[0, 1, 1, 1]	100	0.0004987	0.0	$439.511~\mathrm{KiB}$
13	[0, 6, 0]	100	0.0004776	0.0	$422.431~\mathrm{KiB}$
13	[0, 6, 2]	100	0.0003469	0.0	$441.621~\mathrm{KiB}$
16	[0, 6, 0]	100	0.0009073	0.0	$1010.452~\mathrm{KiB}$
16	[0, 6, 1]	100	0.0007939	0.0	$997.022~\mathrm{KiB}$
16	[0, 0, 6]	100	0.0066951	0.0	$8.564~\mathrm{MiB}$
20	[0, 6, 0]	100	0.0030797	0.0	$4.042~\mathrm{MiB}$
20	[0, 6, 2]	10	0.0022849	0.0	$4.547~\mathrm{MiB}$
32	[0, 6, 2, 1]	10	0.0269912	0.0	$63.082~\mathrm{MiB}$

The full code for random_kmc_v5 can be found in Appendix ??. Table 3.5 shows that implementing a rank table was an extremely significant improvement. For smaller matroids, it is around 5-10x faster, however it is for larger matroids that it truly outshines its predecessors – random_kmc_v5 is a whopping 13 000 times faster than random_kmc_v4 with n = 16, p = [0, 0, 6] as input.

Non-redundant cover generation

Up to this point, our cover generation routine has not taken into account that any two sets of rank r will have at least one cover in common. To see this, consider a matroid-under-construction with n=10 where $A=\{1,2\}$ and $B=\{1,3\}$ are closed sets of rank 2. Currently, GENERATE-COVERS will happily generate the cover $C=\{1,2,3\}$ twice, once as the cover of A and subsequently as the cover of B. Throughout this analysis, we will assume the worst case scenario of no enlargements, as any enlargements will strictly lower the number of sets in play at a given rank. In this case, $|F[r]| = \binom{n}{r}$, and for each closed set A of rank r we are generating $|E \setminus A| = (n-r)$ covers, giving us a total of $\binom{n}{r}(n-r)$ covers generated at each rank r, including the duplicates. With no enlargements, we know that there are $\binom{n}{r+1}$ covers, and

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

$$= \frac{n!}{r!(n-r-1)!}$$

$$= (r+1)\frac{n!}{(r+1)!(n-r-1)!}$$

$$= (r+1)\binom{n}{r+1}.$$

For each step r, we are generating r+1 times as many covers as we need to. Over the course of all steps $0 \le r \le n$, we are generating

$$\sum_{r=0}^{n} (r+1) = \sum_{r=1}^{n+1} r = T_{n+1}$$

times the actual number of covers, where $T_{n+1} = \frac{(n+1)(n+2)}{2}$ is the triangular number. In other words, if we find a way to generate each cover only once, we will have shaved off an n^2 factor from the asymptotic complexity of our implementation.

When generating covers, random_kmc_v6 improves upon the brute force cover generation described above by only adding the covers

$$\Big\{A \cup \{a\}: A \in \mathcal{F}[r], a \in E \setminus A, a \notin \bigcup \big\{B: B \in \mathcal{F}[r+1], A \subseteq B\big\}\Big\}.$$

In other words, we find the covers of A, that is, the sets obtained by adding one more element a from E to A, but we do not include any a that is to be found in another, already added, cover B that contains A. This solves the problem described above; the cover $\{1,2,3\} = B \cup \{2\}$ will not be generated, as $2 \in C$ and $B \subseteq C$. This is implemented in the following manner:

```
# Generate minimal closed sets for rank r+1 (random_kmc_v6)
for y in F[r] # y is a closed set of rank r.
    t = E - y # The set of elements not in y.
# Find all sets in F[r+1] that already contain y and remove excess elements
        from t.
for x in F[r+1]
    if (x & y == y) t &= ~x end
        if t == 0 break end
end
# Insert y U a for all a \in t.
while t > 0
    x = y|(t&-t)
    add_set!(x, F, r, rank)
    t &= ~x
end
end
```

We have extracted the iterative superpose logic described above into its own function to allow it to be performed on a cover-per-cover basis:

```
function add_set!(x, F, r, rank)
  if x in F[r+1] return end
for y in F[r+1]
  if haskey(rank, x&y) && rank[x&y]<r
    continue
  end

# x \cap y has rank > r, replace with x \cup y.
  setdiff!(F[r+1], y)
  return add_set!(x|y, F, r, rank)
end

push!(F[r+1], x)
  rank[x] = r
end
```

As such, F[r+1] is empty when the first cover $y \in F$ is generated, and all covers $\{y \cup \{a\} : a \in E \setminus y\}$ are added. For later sets y, we are comparing with the previously added covers, and dropping any element to be found in a cover x that fully includes y. This way, we avoid re-generating the cover x.

The full code for random_kmc_v6 can be found in Appendix ??.

Table 3.6: Performance of random_kmc_v6.

\overline{n}	(p_1,p_2,\ldots)	Trials	Time	GC Time	Bytes allocated
10	[0, 6, 0]	100	0.000157	0.0	11.306 KiB
10	[0, 5, 1]	100	0.0001427	0.0	$12.257~\mathrm{KiB}$
10	[0, 5, 2]	100	0.000121	0.0	$11.568~\mathrm{KiB}$
10	[0, 6, 1]	100	8.61e-5	0.0	$10.447~\mathrm{KiB}$
10	[0, 4, 2]	100	0.0001237	0.0	$13.597~{ m KiB}$
10	[0, 3, 3]	100	0.0001233	0.0	$14.029~{ m KiB}$
10	[0, 0, 6]	100	0.0002856	0.0	$15.414~\mathrm{KiB}$
10	[0, 1, 1, 1]	100	0.0001942	0.0	$14.446~\mathrm{KiB}$
13	[0, 6, 0]	100	0.0004483	0.0	$19.117~\mathrm{KiB}$
13	[0, 6, 2]	100	0.0004541	0.0	$18.957~\mathrm{KiB}$
16	[0, 6, 0]	10	0.0014919	0.0	$34.531~\mathrm{KiB}$
16	[0, 6, 1]	10	0.0014731	0.0	$36.016~\mathrm{KiB}$
16	[0, 0, 6]	10	0.0168858	0.0	$127.652~\mathrm{KiB}$
20	[0, 6, 0]	10	0.0061574	0.0	$81.573~\mathrm{KiB}$
20	[0, 6, 2]	10	0.0059717	0.0	$82.323~\mathrm{KiB}$
32	[0, 6, 2, 1]	10	0.1599507	0.0	$279.531~\mathrm{KiB}$
63	[0, 6, 4, 2, 1]	1	11.138914	0.0	$4.912~\mathrm{MiB}$
64	[0, 6, 4, 4, 2, 1]	1	12.508729	0.0	$4.912~\mathrm{MiB}$
128	[0,6,6,4,4,2,1]	1	1232.8570	0.0114583	$102.159~\mathrm{MiB}$

3.5.3 What do the generated matroids look like?

Observations

 The average cardinality of the closed sets of a given rank is usually not very much higher than the rank. If the average cardinality were to stray much, all the sets merge instead and the sole closed set of that rank would become E. This might be a useful heuristic when finding the rank of a set.

2

3.5.4 Finding the properties of arbitrary matroids

To build a general matroid library, we want to be able to access all properties of a generated matroid \mathfrak{M} , viz. its bases \mathcal{B} , independent sets \mathcal{I} , circuits \mathcal{C} , and rank function $rank: 2^E \to \mathbb{Z}^*$.

In this section, I will first describe an extension of KNUTH-MATROID that is also fully enumerates \mathcal{I} and \mathcal{C} for \mathfrak{M} when n is small enough. However, this approach does not scale well for larger values of n. For values of n up to 128, we will therefore restrict our attention to independent sets and the rank function, as these are the matroid properties that are relevant to our usecase of fair allocation.

Up-front enumeration of circuits and independent sets for smaller matroids

In his 1974 paper [17], Knuth includes an ALGOL W [25] implementation that also enumerates all circuits and independent sets for the generated matroid. A later implementation in C called ERECTION.W can be found at Knuth's home page [18]. random_erect is an extension of random_kmc_v6 that finds \mathcal{I} and \mathcal{C} by pre-populating the rank table with all subsets of E. The full source code for random_erect can be found in Appendix XXX.

```
# Populate rank table with 100+cardinality for all subsets of E.
k=1; rank[0]=100;
while (k<=mask)
  for i in 0:k-1 rank[k+i] = rank[i]+1 end
  k=k+k;
end</pre>
```

Covers are generated and sets inserted in the same manner as in $random_kmc_v6$. After all covers and enlargements have been inserted and superposed (meaning F[r+1] contains the closed sets of rank r+1), a new operation, $mark_independent_subsets!$ is called on each closed set.

Determining matroid properties after erection

In his 1989 paper, Greene introduces the concept of descriptive sufficiency. A subcollection of closed sets of a matroid is descriptively sufficient if it can be used to identify the fundamental properties of the matroid using certain easily applied conditions [14].

```
ппп
```

```
function is_indep(M::ClosedSetsMatroid, S::Integer)

Determines whether a given set S is independent in the matroid M, given by the closed sets of M grouped by rank. Uses (I.1) in Greene (1989).

"""
function is_indep(M::ClosedSetsMatroid, S::Integer)
t = Base.count_ones(S)

for F in M.F[t]
    if Base.count_ones(S&F) > t-1 return false end end

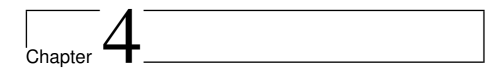
return true
end
```

Indepence oracle.

Circuit oracle.

```
function minimal_spanning_subsets(M::ClosedSetsMatroid, A::Integer)
A modification of Algorithm 3.1 from Greene (1989) that finds all minimal
      spanning subsets of A \subseteq E, given a matroid M = (E, F). If A = E, this finds
       the bases of M.
minimal_spanning_subsets(M::ClosedSetsMatroid, A::Integer) = _mss_all(M, 0, A)
\textbf{function} \ \_\texttt{mss\_all} \ (\texttt{M}:: \texttt{ClosedSetsMatroid}, \ \texttt{j}:: \texttt{Integer}, \ \bar{\texttt{A}}:: \texttt{Integer})
  B = [\bar{A}\&F \text{ for } F \text{ in } M.F[j+1] \text{ if } Base.count\_ones(\bar{A}\&F) > j]
  while length(B) == 0
    if j >= Base.count_ones(A)-1 return A end
     B = [\bar{A}\&F \text{ for } F \text{ in } M.F[j+1] \text{ if } Base.count\_ones}(\bar{A}\&F) > j]
   end
  bases = Set()
   t = reduce(|, B)
   while t > 0
     x = t&-t
     bases = bases \cup _mss_all(M, j, \bar{A}\&\sim x)
     t &= ~x
   end
return bases
```

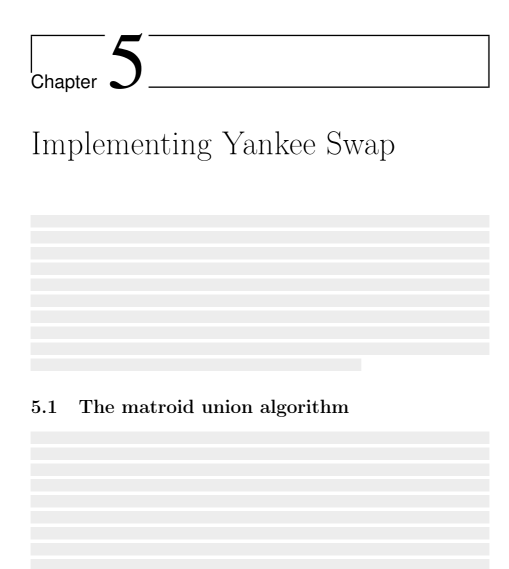
end		
Minimal spanning subsets and l	bases.	



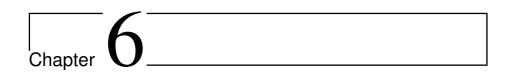
The Matroids.jl API

In the previous chapter, I described how our library generates and represents matroids, as well as how to access the basic properties of our matroids. If all one wanted was to play around with matroids in and of themselves, one might have declared the library finished at this point. However, as mentioned in the introduction, the raison d'être of Matroids.jl is to enable the empirical study of fair allocation algorithms that utilize matroids in some capacity. In this chapter, I consider what fair allocation-specific methods the Matroids.jl API should expose to achieve this goal, and how they might be implemented. The implementation will draw inspiration from, and be designed to interoperate with, Hummel and Hetland's well-organized Allocations.jl library [15].

We will have fair allocation's two main use cases for matroids in mind when deciding what the Matroids.jl API should look like: matroidal rank valuations and matroidal constraints. This chapter will consist of a literature study, the goal of which is to understand what functionality algorithms for these cases will require, after which I discuss the implementation in Matroids.jl. In the next chapter, I will as a proof of concept describe an implementation of the Yankee Swap algorithm for fair allocation with matroidal valuations [21] that make use of the functionality developed in this section, and present some experimental results. Before all this though, we should establish the required fair allocation concepts.



5.2	Some experimental results								



Conclusions

6.1	Limitations		

6.2 Future work

Notes

- 1. Skrive mer om hvordan $\text{Set}\{\text{Set}\{\text{Integer}\}\}$ lagres i minnet og fordelene med å gå over til $\text{Set}\{\text{Integer}\}.$
- $2.\,$ @Benchmarking. Histogrammer. Beskrive variansen i matroide-størrelse ifht input.

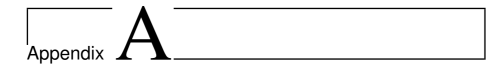
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Appendices



Sets as numbers – some useful tricks

Section* 3.5.2 details a number of steps taken in order to build a performant Julia implementation of Knuth-Matroid. Perhaps chief among these steps in terms of sheer performance gain compared to the initial, naïve implementation, was the transition from representing subsets of E as a Set of integers (or whatever type the elements of E might have), to representing them as a single integer, whose 1-bits denote which elements are in the set. This is possible as long as n is less than the widest available integer type (in off-the-shelf Julia, 128 bits, though one can go wider with the help of libraries [12]). We reiterate the bitwise equivalents of the basic set operations in Table A.1. These bitwise

Table A.1: Set operations and their equivalent bitwise operations

Set operation	Bitwise equivalent
$A \cap B$	$A \wedge B$
$A \cup B$	$A \lor B$
$A \setminus B$	$A \wedge \neg B$
$A \subseteq B$	$A \wedge B = A$

equivalents allow us to perform the set operations in constant time (right???),

resulting in significant performance increases. In the code snippets included throughout Section 3.5.2 and Appendix ??, a number of "tricks" are performed with bitwise operations whose workings and purpose might be a bit obtuse. This appendix came to be as I got to grips with working with sets in this manner.

How do I...

...create a singleton set?

The left-shift operator (<<) can be used to set the *i*th bit to 1 and the others to 0. In general, $\{a\} = 1 << a$. This is used in an early version of GENERATE-COVERS:

...find the smallest element of a set?

Using the two's complement of a set T, denoted by $-T = \neg T + 1$, we can find the smallest element with the operation $T \wedge -T$. This is used in the next trick.

...enumerate all elements of a set one by one?

Using the previous trick, we can repeatedly pop the smallest element in the following manner:

This outputs all numbers from 1 to 0xff with a Hamming weight of 1.

...get a random element from a set?

We find all the positions at which the reversed bitstring of the set has a '1' character, and choose a random one.

```
function rand_el(S::Integer)
  x = rand([2^(i-1) for (i,c) in enumerate(reverse(bitstring(S))) if c == '1'])
  return convert(typeof(S), x)
end
```