

Symmetric chain decompositions of partially ordered sets

A THESIS

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Abstract

A partially ordered set, or poset, is a set of elements and a binary relation which determines an order within elements. Various combinatorial properties of finite and ordered posets have been extensively studied during the last 4 decades. The Sperner property states that the size of the largest subset of pairwise incomparable elements does not exceed the size of the largest level set in an ordered poset. Since a symmetric chain decomposition is a sufficient condition for the Sperner property, we may prove the Sperner property by finding a symmetric chain decomposition for a poset.

In this paper we focus on three types of posets: the Boolean algebra, inversion poset and the Young's lattice. An explicit construction for a symmetric chain decomposition is known only for Boolean algebras. No explicit construction has been found for inversion posets and Young's lattices, a symmetric chain decomposition was found only for a small subset of these posets. Using a maximal flow, we introduce an algorithm for finding this decomposition. We present our results and discuss two implementations of this algorithm.

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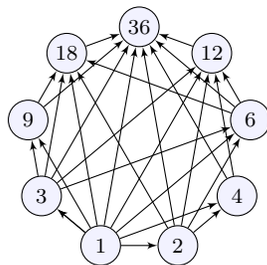
Chapter 1

Introduction

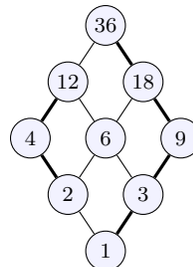
A partially ordered set (poset) is an ordered pair of a set with an order relation, e.g., $(\mathcal{P}, \mathcal{R})$. This relation \mathcal{R} is

- *reflexive*: each element is related to itself, i.e., $\forall a \in \mathcal{P}, (a, a) \in \mathcal{R}$,
- *transitive*: if $(a, b) \in \mathcal{R}$ and $(b, c) \in \mathcal{R}$, then $(a, c) \in \mathcal{R}$,
- *antisymmetric*: if $(a, b) \in \mathcal{R}$ and $a \neq b$, then $(b, a) \notin \mathcal{R}$.

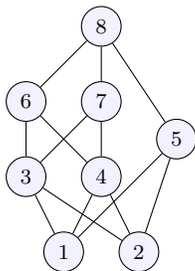
If two elements are related, $(a, b) \in \mathcal{R}$, we use $a \leq b$ to denote this relation. If we would like to emphasize a specific poset \mathcal{P} we use $a \leq_{\mathcal{P}} b$ instead. The set \mathcal{P} with the relation \mathcal{R} can be considered as a directed graph where \mathcal{P} is the vertex set and \mathcal{R} is the edge set. The vertex set \mathcal{P} can be finite or infinite. A simple example is when $\mathcal{P} = \mathbb{Z}^+$ and the relation is defined as $a \leq b$ if $b - a \in \mathbb{N}$. This poset is infinite and since we can compare each pair of positive integers, one integer is always less or equal than another, we say that this order is *total*. A *partial* order on the same set $\mathcal{P} = \mathbb{Z}^+$ is given by divisibility. That is, $a \leq b$ when b is divisible by a . For this order relation, not every pair of positive numbers is comparable. For example if we consider 3 and 5, neither 3 divides 5 nor 3 is divisible by 5. We say 3 and 5 are incomparable. If we restrict \mathcal{P} to all divisors of 36 we get a finite poset. These types of posets are called *division posets*, since the relation is



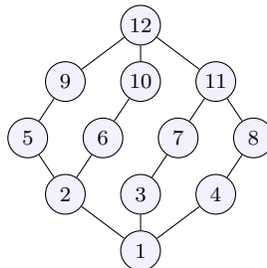
(a) Directed graph of a division poset D_{36} .



(b) Hasse diagram of a division poset D_{36} with a SSCD.



(c) Hasse diagram of a poset without a rank function.



(d) Hasse diagram of a ranked poset.

Figure 1.1: Examples of posets.

defined by division. Finite posets will be our main focus throughout this paper. More types of posets and precise definitions will be introduced in Chapter 2.

The directed graph of the finite division poset of divisors of 36 is shown in Figure 1.1(a). This is the only time we show the entire directed graph of a poset. We will use Hasse diagrams for graphical representations of posets because they contain the same amount of information but have fewer edges than directed graphs. An edge (a, b) is in a Hasse diagram if and only if there is no c such that oriented paths from a to c and from c to b would exist. Additionally, Hasse diagrams do not have oriented edges but we assume that every edge is oriented in upward direction. We can see the difference between a directed graph and a Hasse diagram of a division poset in Figures 1.1(a) and

1.1(b).

Posets have a variety of possible properties. A poset is *ranked* if elements of the Hasse diagram can be partitioned into horizontal level sets such that edges are only between the closest sets. Since there are no edges within each layer nor between layers which are separated by at least one other layer, every oriented path between two elements must have the same length. Each Hasse diagram in Figure 1.1 is ranked except (c). This poset is not ranked because there are two paths from 2 to 8 with different lengths. The path $2 \rightarrow 5 \rightarrow 8$ has length 2 but the path $2 \rightarrow 4 \rightarrow 7 \rightarrow 8$ has length 3.

An *upper bound* of a subset A of \mathcal{P} is an element p such that $a \leq p$ for all $a \in A$. Let U be a set of all upper bounds of A . If there exists an upper bound q such that $q \leq p$ for all $p \in U$, we call q the *least upper bound* of A . *Lower bounds* and the *greatest lower bound* are defined similarly. Notice that the definition of the least upper bound and the greatest lower bound imply uniqueness, since the relation is antisymmetric.

A poset is called a *lattice* if the least upper bound and the greatest lower bound exist for every pair of elements. As we could suppose the division poset and the poset in Figure 1.1(d) are lattices, but the poset in Figure 1.1(c) is not a lattice. If we look at the subset $\{3, 4, 6, 7\}$ in Figure 1.1(c), we can find that $\{3, 4\}$ does not have the least upper bound and $\{6, 7\}$ does not have the greatest lower bound.

A *decomposition* of a poset is a partition of \mathcal{P} into disjoint subsets $\mathcal{P}_0, \mathcal{P}_1, \dots, \mathcal{P}_k$ such that $\cup_{i=0}^k \mathcal{P}_i = \mathcal{P}$. There are many types of decomposition but we will focus on symmetric saturated chain decompositions (SSCD). An ordered n -tuple (c_1, c_2, \dots, c_n) is called a *chain*, with length $n-1$, of the poset \mathcal{P} if $c_i \neq c_{i+1}$ and $c_i \leq c_{i+1}$ for $1 \leq i < n$. A chain (c_1, c_2, \dots, c_n) is *saturated* if it cannot be internally extended, i.e.; (c_i, c_{i+1}) is an edge in the Hasse diagram of the poset for $1 \leq i < n$. A saturated chain is *symmetric* if it starts and ends at levels whose distance from the middle level(s) of the poset is the same. Figure 1.1(b) contains a diagram with two highlighted symmetric saturated chains. A poset has a SSCD if \mathcal{P} can be decomposed into symmetric saturated chains.

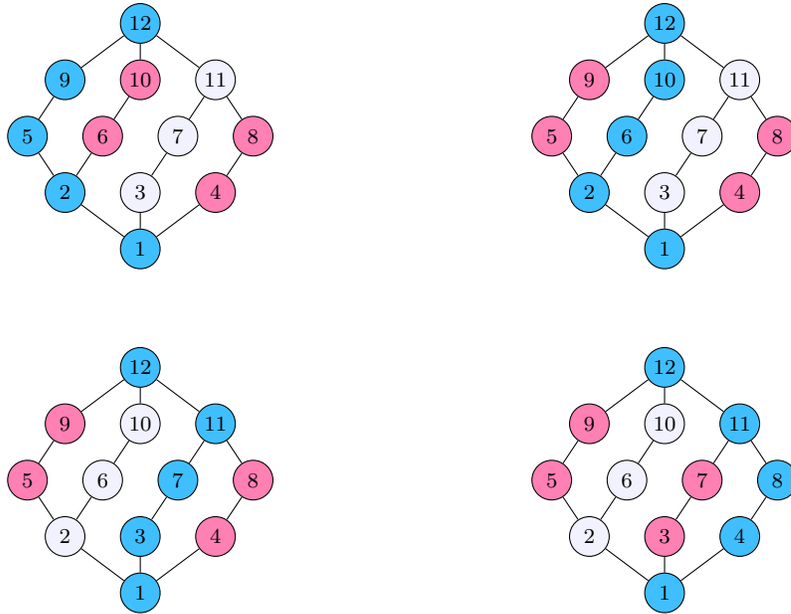


Figure 1.2: Different saturated chain decompositions of a poset depicted in figure 1.1(d).

The division poset (lattice) in Figure 1.1(b) is decomposable into three symmetric saturated chains. The chains, as highlighted in the Figure, are $(1, 3, 9, 18, 36)$, $(2, 4, 12)$ and (6) . The lattice in Figure 1.1(d) cannot be decomposed into symmetric saturated chains. If this lattice were decomposable, there would have to be a longest chain from 1 to 12. We have exactly 4 choices for this chain, the choice depends on which element from the middle layer is included. Each choice will leave two non symmetric chains as we can see in Figure 1.2.

Before we introduce an algorithm for finding a symmetric saturated chain decomposition for a poset; fundamental terminology, a few properties of posets and essential theorems will be introduced in chapter 2. We also explain a connection between Sperner's property and the existence of a SSCD in this chapter.

The rest of the proofs are placed in chapter 3. In this chapter we describe isomorphisms between division and boolean posets and within inversion posets.

Chapter 4 introduces an algorithm for finding a SSCD for posets. This algorithm uses a maximum flow in a graph to construct a SSCD. Two different algorithms for finding a maximum flow in a graph are discussed.

Chapter 5 contains a brief description of two implementations of the algorithm from chapter 4. We provide a table with all posets for which our implementation finds a SSCD and discuss suggestions for a future work.

Both of our implementations have similar interface and a description of these interfaces is given in appendix A. A performance comparison between the two implementations, examples of a SSCD for selected posets and source codes of our implementations are placed in appendixes, also.

Chapter 2

Formal definitions and fundamental properties of posets

The theory of partially ordered sets investigates a poset as a graph or as a set of elements. This set can be infinite but we will focus on finite sets only. At the beginning of this chapter the most common notation and definitions from Graph Theory are introduced. Concrete examples of posets with known attributes will be introduced after definitions. Most of these definitions can be found in [1] or in [10].

Let \mathcal{P} be a poset, \mathcal{R} its relation and p, p', q, q' elements of \mathcal{P} .

We say that p and q are *comparable* if $(p, q) \in \mathcal{R}$ or $(q, p) \in \mathcal{R}$, and we typically write this as $p \leq q$ or $q \leq p$. In the case when p and q are comparable and $p \neq q$ either $p < q$ or $q < p$ is valid. An element p *covers* q , denoted as $q \triangleleft p$, if $q < p$ and if $q < p' \leq p$ implies $p = p'$. An element p is a *minimal element* of \mathcal{P} if $q \leq p$ implies $q = p$; a *maximal element* is defined similarly.

A ranked poset is equipped with a *rank function* ρ . This is a function from \mathcal{P} to \mathbb{N} satisfying $\rho(p) = \rho(q) + 1$ if p covers q . The rank of a poset is defined as the largest rank over all elements from the poset. Since every finite lattice has only one minimal element, the rank of this minimal element is zero.

from the definition of $N_j(\mathcal{P})$, since each layer $N_j(\mathcal{P})$ is empty. A *generating function* of a ranked poset \mathcal{P} is given by

$$GF(\mathcal{P}) = \sum_{w \in \mathcal{P}} q^{\rho(w)} = \sum_{i=0}^{\rho(\mathcal{P})} W_i q^i.$$

We stated that posets in Figures 1.1(b) and 1.1(d) are ranked. Hence, there is a generating function for each poset. The generating function for the poset in Figure 1.1(d) is $1 + 3q + 4q^2 + 3q^3 + q^4$, $GF(D_{36}) = 1 + 2q + 3q^2 + 2q^3 + q^4$.

A poset is *rank unimodal* if there exist j such that $W_0 \leq W_1 \leq \dots \leq W_{j-1} \leq W_j \geq W_{j+1} \geq \dots \geq W_{\rho(\mathcal{P})}$. A *rank symmetric* poset is a ranked poset whose Whitney numbers are symmetric with respect to the middle layer(s), i.e.; $W_i = W_{\rho(\mathcal{P})-i}$ for $0 \leq i < \rho(\mathcal{P})/2$. Again, if we look at Figures 1.1(b) and 1.1(d), sequences of Whitney numbers are $(1, 2, 3, 2, 1)$ and $(1, 3, 4, 3, 1)$, respectively. Both sequences are symmetric and unimodal but only the division poset has a SSCD.

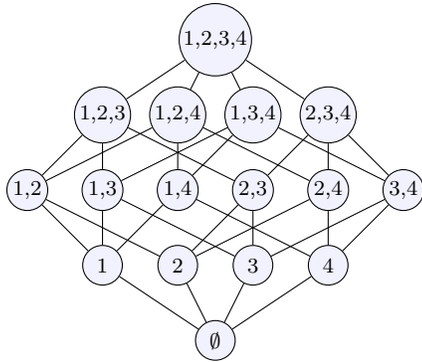
Posets \mathcal{P} and \mathcal{Q} are *isomorphic*, $\mathcal{P} \cong \mathcal{Q}$, if there is a bijective mapping φ from \mathcal{P} onto \mathcal{Q} such that $p \leq_{\mathcal{P}} q$ if and only if $\varphi(p) \leq_{\mathcal{Q}} \varphi(q)$.

Theorem 2.1 (Sperner's Theorem). *Let n be a positive integer and \mathcal{F} be a family of subsets of $[n] := \{1, 2, \dots, n\}$ such that no member of \mathcal{F} is included in another member of \mathcal{F} , that is, for all $X, Y \in \mathcal{F}$ we have $X \not\subset Y$. Then*

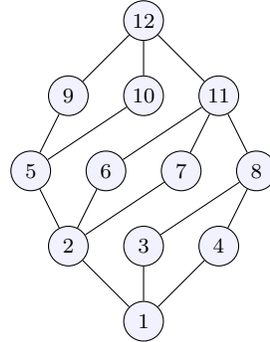
$$|\mathcal{F}| \leq \begin{cases} \binom{n}{n/2} & \text{if } n \text{ is even,} \\ \binom{n}{(n-1)/2} & \text{if } n \text{ is odd.} \end{cases}$$

Sperner's Theorem was at the beginning of the Poset Theory and many important results were obtained by using this theorem. Three different proofs of this theorem can be found in [4], as well as other important results. We will prove this theorem in the next chapter using a symmetric chain decomposition of the family of subsets of $\{1, 2, \dots, n\}$.

Since each level set of a poset contains only incomparable elements, a lower bound



(a) B_4



(b) A lattice without the Sperner property.

Figure 2.2: Examples of two different lattices.

for the size of the largest antichain is given by the largest Whitney number. Sperner's Theorem was stated for the family of subsets of a set ordered by inclusion and it says that this lower bound is tight, the size of the biggest antichain is equal to the largest Whitney number. A generalization of this theorem is available for other types of posets; a ranked poset \mathcal{P} has the *Sperner property* if the size of any antichain is less than or equal to the biggest Whitney number.

The two lattices in Figure 1.1 both have the Sperner property. However, the lattice in Figure 2.2(b) is rank symmetric but it does not have the Sperner property nor a SSCD, the antichain $\{3, 4, 5, 6, 7\}$ is larger than the size of the middle layer.

Unfortunately, it is not true that every poset with the Sperner property has a symmetric saturated chain decomposition. An example of such a poset is in Figure 1.1(d). This poset satisfies the Sperner property, the size of each antichain is not bigger than 4 which is the size of the biggest antichain formed by the middle layer. However, there is no SSCD of this poset as we discussed in Chapter 1.

We will find a use of q -functions in examining rank functions. A q -analog of n is defined as $\{n\}_q = \frac{1-q^n}{1-q}$ and we often want to evaluate the q -analog of n for $q = 1$. This

value does not exist, but we can find a limit as q approaches 1,

$$\lim_{q \rightarrow 1} \frac{1 - q^n}{1 - q} = \lim_{q \rightarrow 1} \frac{(1 - q)(1 + q + q^2 + \cdots + q^{n-1})}{1 - q} = \lim_{q \rightarrow 1} 1 + q + q^2 + \cdots + q^{n-1} = n.$$

Additionally, we will use a continuous extension of q -analog of n ,

$$\frac{1 - q^n}{1 - q} \equiv 1 + q + q^2 + \cdots + q^{n-1},$$

because $\frac{1 - q^n}{1 - q} = 1 + q + q^2 + \cdots + q^{n-1}$ almost everywhere.

A q -factorial of n , $\{n\}_q!$, is defined as a product of q -analogs of i , where $0 < i \leq n$.

Hence,

$$\begin{aligned} \{n\}_q! &= \{n\}_q \cdot \{n-1\}_q \cdots \{1\}_q \\ &= \frac{1 - q^n}{1 - q} \cdot \frac{1 - q^{n-1}}{1 - q} \cdots \frac{1 - q}{1 - q} \\ &\equiv (1 + q + q^2 + \cdots + q^{n-1})(1 + q + q^2 + \cdots + q^{n-2}) \cdots (1 + q)(1). \end{aligned}$$

Similar as the binomial coefficient, the q -binomial coefficient is given by

$$\binom{n}{m}_q = \frac{\{n\}_q!}{\{m\}_q! \{n-m\}_q!}$$

The *Boolean algebra* B_n is the poset of all subsets of a set with n elements, ordered by an inclusion. The size of B_n is 2^n and its Whitney numbers correspond to a row in the Pascal's triangle. The rank of an element is its cardinality.

The *division* poset D_n is a poset of all divisors of n , ordered by divisibility. Let $n = p_1^{a_1} p_2^{a_2} \cdots p_k^{a_k}$ be the prime factorization of n . Each element of the division poset can be written as $p_1^{b_1} p_2^{b_2} \cdots p_k^{b_k}$, where $0 \leq b_i \leq a_i$ for each i . The rank of $p_1^{b_1} p_2^{b_2} \cdots p_k^{b_k}$ is $\sum_{i=1}^k b_i$.

The *Symmetric group* S_n is the group of all permutations of $(1, 2, \dots, n)$. The sum of all Whitney numbers of any partial ordering on S_n has to be $n!$ since there are $n!$ possible arrangements of n elements in a sequence. Suppose $a = (a_1, a_2, \dots, a_n)$ and $b = (b_1, b_2, \dots, b_n)$ are elements of S_n . A pair (i, j) is called an *inversion* of a if $i < j$

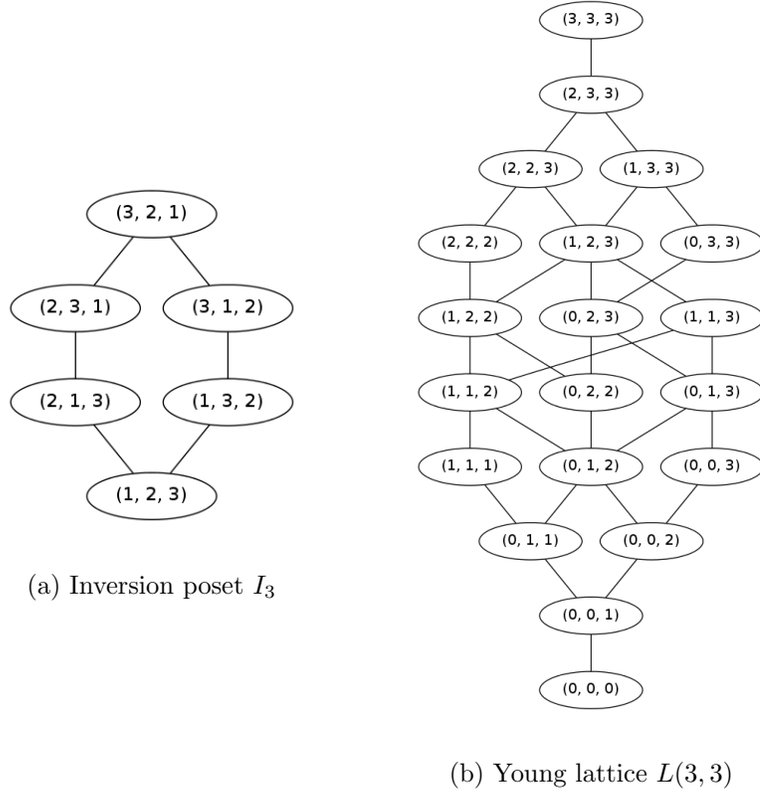


Figure 2.3: Diagrams of an inversion poset and a Young's lattice.

and $a_i > a_j$. The *inversion sequence* of a is a sequence (i_1, i_2, \dots, i_n) where each i_k represents how many elements in a on the left of a_k are greater than a_k [7]. For example, consider a sequence $(7, 5, 8, 3, 2, 6, 1, 4)$. A pair $(2, 7)$ is an inversion since $5 > 1$ and the inversion sequence is $(0, 1, 0, 3, 4, 2, 6, 4)$. The *inversion number* of a permutation is the sum of its inversion sequence, i.e., $\text{inv}(a) = \sum_{j=1}^n i_j$ [11]. The set S_n with a relation given by $a \leq b$, if b is created from a by interchanging a_i with a_{i+1} when $a_i < a_{i+1}$, is called the *inversion poset*, I_n . This binary relation is called the (weak) *Bruhat order*. The generating function is given by

$$GF(I_n) = \{n\}_q! = \prod_{i=0}^{n-1} (1 + q + q^2 + \dots + q^i)$$

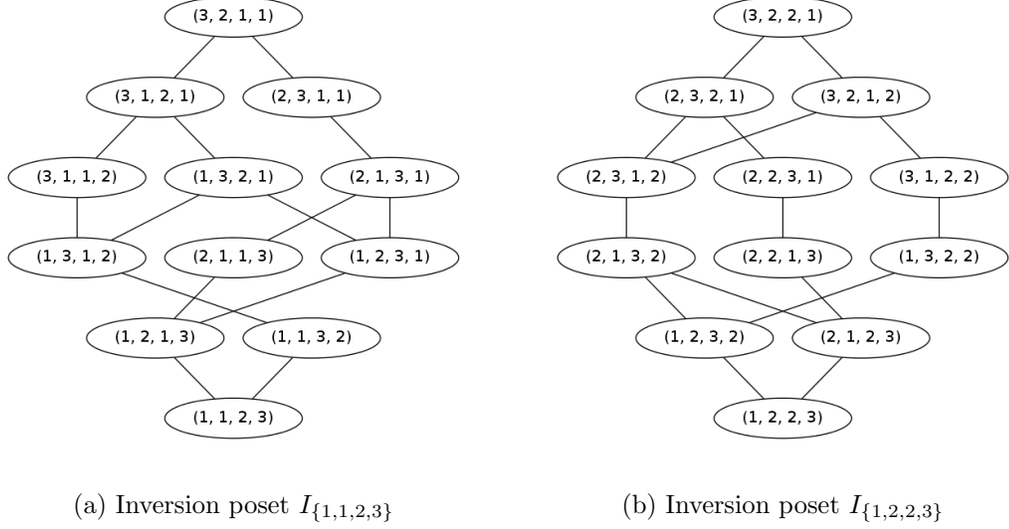


Figure 2.4: Diagrams of inversion posets for multisets.

and the rank of an element is its inversion number.

Similar to inversion posets we define an inversion poset for multisets ordered by inversions. Inversion posets for $\{1, 1, 2, 3\}$ and $\{1, 2, 2, 3\}$, $I_{\{1,1,2,3\}}$ and $I_{\{1,2,2,3\}}$, respectively, are shown in Figure 2.4. The generating function for I_N , where $N = \underbrace{\{n_1, n_1, \dots, n_1\}}_{l_1}, \underbrace{\{n_2, n_2, \dots, n_2\}}_{l_2}, \dots, \underbrace{\{n_k, n_k, \dots, n_k\}}_{l_k}$ is given by a q -multinomial coefficient,

$$GF(I_N) = \binom{\sum_{i=1}^k l_i}{l_1, l_2, \dots, l_k}_q = \frac{\left\{ \sum_{i=1}^k l_i \right\}_q!}{\{l_1\}_q! \cdot \{l_2\}_q! \cdots \{l_k\}_q!}.$$

We will prove this result in Chapter 3. For example, for $I_{\{1,1,2,3\}}$ we have

$$\begin{aligned} GF(I_{\{1,1,2,3\}}) &= \frac{\{2+1+1\}_q!}{\{2\}_q! \{1\}_q! \{1\}_q!} = \frac{(1+q+q^2+q^3)(1+q+q^2)(1+q)(1)}{(1+q)(1)(1)} \\ &= (1+q+q^2+q^3)(1+q+q^2) \\ &= 1+2q+3q^2+3q^3+2q^4+q^5. \end{aligned}$$

Note that the generating function for $I_{\{1,2,2,3\}}$ is the same as the generating function for $I_{\{1,1,2,3\}}$ even though these posets are not isomorphic.

The *Young's* lattice, $L(m, n)$, is a poset of n -tuples (a_1, a_2, \dots, a_n) , where $0 \leq a_1 \leq a_2 \leq \dots \leq a_n \leq m$ with order relation $(a_1, a_2, \dots, a_n) \leq (b_1, b_2, \dots, b_n)$ if $a_i \leq b_i$ for all i . The rank function is given by a sum of all elements in an n -tuple, hence $\rho((a_1, a_2, \dots, a_n)) = a_1 + a_2 + \dots + a_n$. We can find in [11, p. 72] that the generating function for a Young's lattice is

$$GF(L(m, n)) = \binom{m+n}{m}_q = \frac{(1-q^{n+1})(1-q^{n+2}) \dots (1-q^{n+m})}{(1-q)(1-q^2) \dots (1-q^m)}.$$

Theorem 2.2 ([2]). *If \mathcal{P} and \mathcal{Q} are posets with a symmetric saturated chain decomposition, then $\mathcal{P} \times \mathcal{Q}$ has a symmetric saturated chain decomposition.*

Proof. Let P_0, P_1, \dots, P_m and Q_0, Q_1, \dots, Q_n be SSCD's for \mathcal{P} and \mathcal{Q} , respectively. Consider a pair of chains (P_i, Q_j) , say

$$\mathcal{P}_i = p_0 \triangleleft p_1 \triangleleft \dots \triangleleft p_k \text{ and } \mathcal{Q}_j = q_0 \triangleleft q_1 \triangleleft \dots \triangleleft q_h.$$

We can create a new chain E_l , where

$$E_l = \left((p_0, q_l), (p_1, q_l), \dots, (p_{k-l}, q_l), (p_{k-l}, q_{l+1}), \dots, (p_{k-l}, q_h) \right),$$

for $0 \leq l \leq \min\{k, h\}$. Each E_l is evidently a saturated chain in $\mathcal{P} \times \mathcal{Q}$, since the change is only in the first or the second term and both P_i and Q_j are saturated. \mathcal{P}_i and \mathcal{Q}_j are

symmetric, so $\rho(\mathcal{P}) = \rho(p_0) + \rho(p_k)$ and $\rho(\mathcal{Q}) = \rho(q_0) + \rho(q_h)$. Additionally, since

$$\begin{aligned}
\rho((p_0, q_l)) + \rho((p_{k-l}, q_h)) &= \rho(p_0) + \rho(q_l) + \rho(p_{k-l}) + \rho(q_h) \\
&= \rho(p_0) + \rho(q_k) - (k-l) + \rho(p_0) + (k-l) + \rho(q_h) \\
&= \rho(p_0) + \rho(q_k) + \rho(p_0) + \rho(q_h) \\
&= \rho(\mathcal{P}) + \rho(\mathcal{Q}) \\
&= \rho(\mathcal{P} \times \mathcal{Q}),
\end{aligned}$$

E_l is also symmetric. If $k \leq h$, the number of elements in all E_l 's for a pair (P_i, Q_j) is

$$\begin{aligned}
\sum_{l=0}^k |E_l| &= \sum_{l=0}^k (k+h+1-2l) = (k+1)(k+h+1) - 2 \sum_{l=0}^k l \\
&= k^2 + kh + 2k + h + 1 - 2k \frac{k+1}{2} \\
&= k^2 + kh + 2k + h + 1 - k(k+1) = kh + k + h + 1 \\
&= (k+1)(h+1) = |P_i| \cdot |Q_j|.
\end{aligned}$$

If $h < k$, the same result can be obtained by the same procedure. Since we can do this procedure for each pair of chains from the decomposition, the total number of elements in all E_l 's is

$$\sum_{i=0}^m \sum_{j=0}^n \sum_{l=0}^{\min\{k,h\}} |E_l| = \sum_{i=0}^m \sum_{j=0}^n |P_i| \cdot |Q_j| = \sum_{i=0}^m |P_i| \cdot \sum_{j=0}^n |Q_j| = |\mathcal{P}| \cdot |\mathcal{Q}|.$$

Hence we will obtain new symmetric chain decomposition of $\mathcal{P} \times \mathcal{Q}$ by this procedure. \square

For example, consider a direct product of I_3 and a chain of three elements, $C = (c_1, c_2, c_3)$. A diagram of this direct product is shown in Figure 2.5. To follow the proof of Theorem 2.2 we pick a symmetric saturated chain in each poset. These chains are highlighted by thick solid line. These chains will yield, by Theorem 2.2, three symmetric saturated chains in the direct product, since the shortest chain contains three elements. If we follow the same notation, we have chains E_0, E_1 and E_2 . The chain E_0 is the

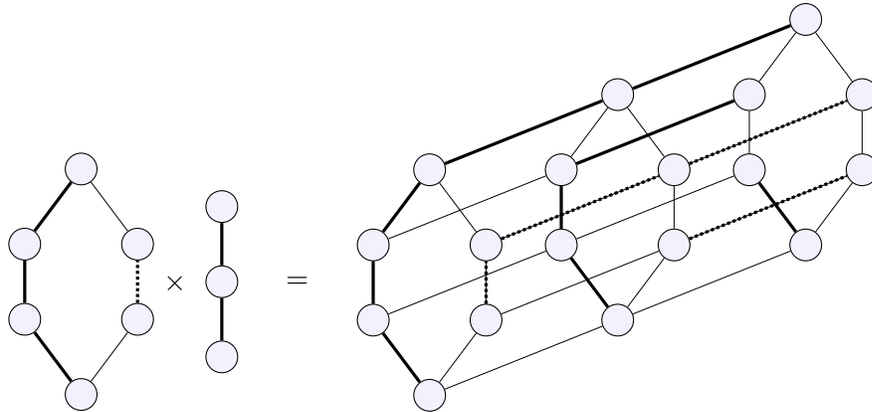


Figure 2.5: Example of theorem 2.2.

longest chain and E_2 is the shortest. Additionally, these chains cover all elements in the Cartesian product of the original highlighted chain in I_3 and C . We can describe this construction easier if a diagram of the direct product is available. Given two saturated symmetric chains \mathcal{P} and \mathcal{Q} , each from a different poset, a SSCD of the direct product $\mathcal{P} \times \mathcal{Q}$ can be found by following an algorithm:

1. Let $S = \mathcal{P} \times \mathcal{Q}$.
2. Pick an element p with the smallest rank in S .
3. Construct a new symmetric saturated chain E_i which starts at p , follows an upward path given by \mathcal{P} as far as it can and then continues to the right following a path given by \mathcal{Q} .
4. If we have not used all elements in $\mathcal{P} \times \mathcal{Q}$ we have to shrink S and find a next chain. In order to do that we have to remove E_i from S and go back to 2.

We will investigate a few types of posets which are created as a direct product of two or more ranked posets. To analyze their properties we prove the following theorems.

Theorem 2.3 ([4]). *The generating function for the direct product of two ranked posets*

\mathcal{P} and \mathcal{Q} is given by

$$GF(\mathcal{P} \times \mathcal{Q}) = GF(\mathcal{P}) \cdot GF(\mathcal{Q}).$$

Proof. Since the generating function of a poset \mathcal{P} is given by $GF(\mathcal{P}) = \sum_{p \in \mathcal{P}} q^{\rho(p)}$, we can use this formula on $\mathcal{P} \times \mathcal{Q}$

$$\begin{aligned} GF(\mathcal{P} \times \mathcal{Q}) &= \sum_{(a,b) \in \mathcal{P} \times \mathcal{Q}} q^{\rho((a,b))} = \sum_{(a,b) \in \mathcal{P} \times \mathcal{Q}} q^{\rho(a) + \rho(b)} = \sum_{a \in \mathcal{P}} \sum_{q \in \mathcal{Q}} q^{\rho(a) + \rho(b)} \\ &= \sum_{a \in \mathcal{P}} q^{\rho(a)} \sum_{b \in \mathcal{Q}} q^{\rho(b)} = GF(\mathcal{P}) \cdot GF(\mathcal{Q}). \end{aligned}$$

□

Theorem 2.4. *A poset with a symmetric saturated chain decomposition has the Sperner property.*

Proof. If a poset \mathcal{P} has a SSCD then the largest Whitney number is $W_{\lfloor \frac{\rho(\mathcal{P})}{2} \rfloor}(\mathcal{P})$. We need to prove that every antichain has size less than or equal to $W_{\lfloor \frac{\rho(\mathcal{P})}{2} \rfloor}(\mathcal{P})$.

Let $A = \{a_1, a_2, \dots, a_l\}$ be an antichain and $C = \{C_1, C_2, \dots, C_k\}$ be a SSCD of \mathcal{P} . Since C contains all elements of \mathcal{P} , suppose, without loss of generality that $a_1 \in C_1$. All elements in C_1 are comparable with a_1 . Hence, without loss of generality, say that $a_2 \in C_2$. Since a_3 is not comparable with neither a_1 nor a_2 it cannot be in C_1 or C_2 , say a_3 is in C_3 . We can conclude, by reordering the C 's as necessary, that $a_i \in C_i$ for each $i \leq l$, therefore $l \leq k$.

It is clear from the definition of SSCD that the biggest Whitney number, the size of the middle layer(s), is equal to the number of chains in a SSCD. Therefore,

$$l \leq k = W_{\lfloor \frac{\rho(\mathcal{P})}{2} \rfloor}(\mathcal{P}) \implies l \leq W_{\lfloor \frac{\rho(\mathcal{P})}{2} \rfloor}(\mathcal{P}),$$

which completes the proof. □

Chapter 3

Additional properties of posets

3.1 Division and Boolean posets

Theorem 3.1. *A division poset D_n is isomorphic to the direct product of chains $C = (1 \triangleleft p_1 \triangleleft p_1^2 \triangleleft \dots \triangleleft p_1^{a_1}) \times (1 \triangleleft p_2 \triangleleft p_2^2 \triangleleft \dots \triangleleft p_2^{a_2}) \times \dots \times (1 \triangleleft p_k \triangleleft p_k^2 \triangleleft \dots \triangleleft p_k^{a_k})$, where $n = p_1^{a_1} p_2^{a_2} \dots p_k^{a_k}$ is the prime factorization of n .*

Proof. Let I be a function from D_n to C given by

$$I(p_1^{b_1} p_2^{b_2} \dots p_k^{b_k}) = (p_1^{b_1}, p_2^{b_2}, \dots, p_k^{b_k}).$$

It is clear that the function I is one-to-one. Let $p = p_1^{b_1} p_2^{b_2} \dots p_k^{b_k}$ and $q = p_1^{c_1} p_2^{c_2} \dots p_k^{c_k}$, where $b_i \leq c_i$ for each i . Apparently, q is divisible by p , $q/p = p_1^{c_1 - b_1} \cdot p_2^{c_2 - b_2} \dots p_k^{c_k - b_k}$ where each $c_i - b_i \geq 0$ and therefore $p \leq_{D_n} q$.

Since $I(p) = (p_1^{b_1}, p_2^{b_2}, \dots, p_k^{b_k})$ we can construct a chain $((p_1^{b_1}, p_2^{b_2}, \dots, p_k^{b_k}) \leq (p_1^{c_1}, p_2^{b_2}, \dots, p_k^{b_k}) \leq (p_1^{c_1}, p_2^{c_2}, \dots, p_k^{b_k}) \leq \dots \leq (p_1^{c_1}, p_2^{c_2}, \dots, p_k^{c_k}) = I(q)$ in C , therefore $I(p) \leq_C I(q)$ from the transitive property of the relation.

To finish the proof we need to show that $I(p) \leq_C I(q)$ implies $p \leq_{D_n} q$. To show this, again suppose that $I(p) = (p_1^{b_1}, p_2^{b_2}, \dots, p_k^{b_k})$ and $I(q) = (p_1^{c_1}, p_2^{c_2}, \dots, p_k^{c_k})$, where $b_i \leq c_i$ for each i . It is clear, since I is one-to-one, that p divides q . \square

We can find the generating function for a division poset by combining Theorems 3.1 and 2.3.

Corollary 3.2. *The generating function for a division poset D_n , where $n = p_1^{a_1} p_2^{a_2} \cdots p_k^{a_k}$ is the prime factorization of n , is given by*

$$GF(D_n) = \prod_{i=1}^k (1 + q + q^2 + \cdots + q^{a_i}).$$

For example the generating function for D_{36} is

$$GF(D_{36}) = GF(D_{2^2 \cdot 3^2}) = (1 + q + q^2) \cdot (1 + q + q^2) = 1 + 2q + 3q^2 + 2q^3 + q^4.$$

We can compare this rank function with a diagram of the division poset D_{36} in Figure 1.1(b), there is one element with rank zero, two with rank one and three with rank two.

Now, we may focus on Boolean posets and show that a Boolean poset has the same properties as a division poset. In fact, a Boolean poset is a special case of a division poset D_n .

Theorem 3.3. *A Boolean poset B_k is isomorphic to D_n , where $n = p_1 p_2 \cdots p_k$ and each p_i is a distinct prime number.*

Proof. Since elements of B_k are subsets of $\{1, 2, \dots, k\}$, we can introduce an one-to-one function I from B_k into D_n . Say

$$I(\{a_1, a_2, \dots, a_i\}) = p_{a_1} p_{a_2} \cdots p_{a_i},$$

where $I(\emptyset) = 1$. Consider two subsets of B_k , A and B , such that $A \subseteq B$. Suppose that $A = \{a_1, a_2, \dots, a_i\}$ and $B = \{a_1, a_2, \dots, a_i, b_1, b_2, \dots, b_j\}$ since A is a subset of B . We have to show that $I(A) \leq_{D_n} I(B)$, i.e., that the function I respects the order. Observe that

$$I(A) = p_{a_1} p_{a_2} \cdots p_{a_i} = \frac{p_{a_1} p_{a_2} \cdots p_{a_i} p_{b_1} p_{b_2} \cdots p_{b_j}}{p_{b_1} p_{b_2} \cdots p_{b_j}} = \frac{I(B)}{p_{b_1} p_{b_2} \cdots p_{b_j}}.$$

Thus

$$\frac{I(B)}{I(A)} = p_{b_1} p_{b_2} \cdots p_{b_j},$$

which indicates that $I(A)$ divides $I(B)$, therefore $I(A) \leq_{D_n} I(B)$. \square

Now we know that a Boolean poset is a ranked poset with a SSCD. Using Corollary 3.2, the generating function for B_n is given by

$$GF(B_n) = \prod_{i=1}^n (1 + q) = \sum_{i=0}^n \binom{n}{i} q^i,$$

which explains why a sequence of Whitney numbers corresponds to a row in the Pascal's triangle.

Theorem 3.4. *Given a division poset D_n , where $n = p_1^{a_1} p_2^{a_2} \cdots p_k^{a_k}$ is the prime factorization of n , we can find an isomorphism I between D_n with a standard order and D_n with a reversed order.*

Proof. We will prove that this isomorphism is given by

$$I(p_1^{b_1} p_2^{b_2} \cdots p_k^{b_k}) = \frac{n}{p_1^{b_1} p_2^{b_2} \cdots p_k^{b_k}}.$$

The function I is one-to-one, therefore we have to check only that if $b \leq_{D_n} c$ then $I(b) \geq_{D_n} I(c)$. Since c is divisible by b we have

$$\frac{c}{b} = \frac{\frac{1}{b}}{\frac{1}{c}} = \frac{1}{b} \cdot \frac{n}{\frac{1}{c}} = \frac{n}{b} = \frac{I(b)}{I(c)},$$

as required. \square

Given a poset P , the poset with the same underlying set but reversed order is called the *dual* of P , P^* [4]. If a poset and its dual are isomorphic, then this poset is called *self-dual* [10].

Theorems 2.2 and 3.1 imply that each division poset has a SSCD, therefore each division poset has the Sperner property by Theorem 2.4. In particular, the Boolean poset B_n satisfies the Sperner property, proving Theorem 2.1.

3.2 Inversion posets and Young's lattices

Let $S(m+n, m)$ be a poset of all bit strings of length $m+n$ with m ones. We define a partial order which is the same as for an inversion poset I_n . That is, $p \leq q$ if q is created by some number of adjacent transpositions of elements in p , where the second element is larger than the first. The smallest and the greatest elements are $\underbrace{0000 \cdots 000}_n \underbrace{11 \cdots 111}_m$ and $\underbrace{11 \cdots 111}_m \underbrace{0000 \cdots 000}_n$, respectively.

Theorem 3.5. $S(m+n, m)$ and $L(m, n)$ are isomorphic.

Proof. Let I be a function from $L(m, n)$ to $S(m+n, m)$ given by

$$I\left((a_1, a_2, \dots, a_n)\right) = \underbrace{1 \cdots 1}_a 0 \underbrace{1 \cdots 1}_{a_2-a_1} 0 \underbrace{1 \cdots 1}_{a_3-a_2} 0 \cdots \underbrace{1 \cdots 1}_{a_n-a_{n-1}} 0 \underbrace{1 \cdots 1}_{m-a_n}$$

To show that I is one-to-one suppose that $a = (a_1, a_2, \dots, a_n)$, $b = (b_1, b_2, \dots, b_n)$ and $I(a) = I(b)$. Then

$$I(a) = I(b)$$

$$\underbrace{1 \cdots 1}_a 0 \underbrace{1 \cdots 1}_{a_2-a_1} 0 \underbrace{1 \cdots 1}_{a_3-a_2} 0 \cdots \underbrace{1 \cdots 1}_{a_n-a_{n-1}} 0 \underbrace{1 \cdots 1}_{m-a_n} = \underbrace{1 \cdots 1}_{b_1} 0 \underbrace{1 \cdots 1}_{b_2-b_1} 0 \underbrace{1 \cdots 1}_{b_3-b_2} 0 \cdots \underbrace{1 \cdots 1}_{b_n-b_{n-1}} 0 \underbrace{1 \cdots 1}_{m-a_n}.$$

If the two strings are the same, then the number of ones separated by a zero has to be

the same. Hence, we get this system of equations

$$\begin{aligned}
a_1 &= b_1 \\
a_2 - a_1 &= b_2 - b_1 \\
a_3 - a_2 &= b_3 - b_2 \\
&\vdots \\
a_n - a_{n-1} &= b_n - b_{n-1} \\
m - a_n &= m - b_n.
\end{aligned}$$

We can use the solution of the first equation, $a_1 = b_1$, to solve the second equation and get $a_2 = b_2$. It is easy to see that the solution of this system is $a_i = b_i$ for $1 \leq i \leq n$ and therefore $a = b$.

To show that the function I preserves order, let $a \leq_L b$. $I(a)$ is less than or equal to $I(b)$ if the number of ones on the left from each zero in $I(a)$ is less than or equal to the number of ones on the left from each zero in $I(b)$. For $I(a)$, the number of ones before the first zero is a_1 , before the second zero it is $a_1 + a_2 - a_1 = a_2$, there are a_3 ones before the third zero, etc., up to m . Similarly, the sequence for b is b_1, b_2, \dots, m . Each a_i is less than or equal to b_i since $a \leq_L b$ and therefore $I(a) \leq I(b)$.

The opposite, $I(a) \leq I(b)$ implies $a \leq_L b$ comes from the same effort of comparing number of ones which precede a zero. □

The function I from the proof above is one-to-one and therefore it has an inverse. The function I^{-1} takes a bit string s and returns a vector containing only those elements from the inversion sequence of s which correspond to zeros in s . An example of an isomorphism between $L(2, 3)$ and $S(5, 2)$ can be seen on Figure 3.1.

Kyle Krueger in his Master's project[8] proved that $\binom{m+n}{m}_q$ is the generating function of $S(m+n, m)$. This result together with the previous theorem shows that the generation function for $L(m, n)$ is

$$GF(L(m, n)) = \binom{m+n}{m}_q.$$

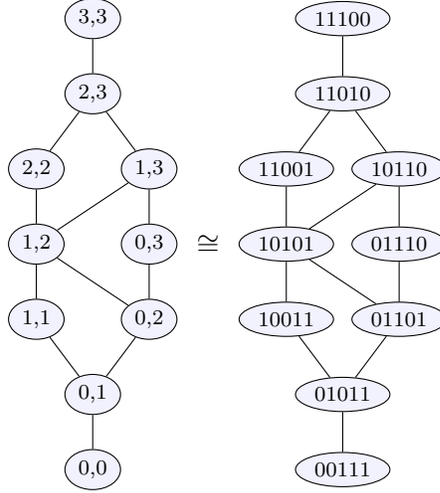


Figure 3.1: An isomorphism between $L(3, 2)$ and $S(5, 3)$.

Theorem 3.6. *The generating function for an inversion poset for a multiset I_N , where*

$$N = \underbrace{\{n_1, n_1, \dots, n_1\}}_{l_1} \underbrace{\{n_2, n_2, \dots, n_2\}}_{l_2} \dots \underbrace{\{n_k, n_k, \dots, n_k\}}_{l_k} \text{ is}$$

$$\left(\begin{array}{c} \sum_{i=1}^k l_i \\ l_1, l_2, \dots, l_k \end{array} \right)_q.$$

Proof. We prove this theorem using strong induction on k , the number of distinct elements in N . Let $k = 2$ be our base case. Krueger in [8] provides the formula for the generating function in this base case as a q -binomial coefficient,

$$\binom{l_1 + l_2}{l_1}_q = \frac{\{l_1 + l_2\}_q!}{\{l_1\}_q! \cdot \{l_2\}_q!}.$$

This is the same formula that we get from our hypothesis, since

$$\left(\begin{array}{c} \sum_{i=1}^2 l_i \\ l_1, l_2 \end{array} \right)_q = \frac{\{l_1 + l_2\}_q!}{\{l_1\}_q! \cdot \{l_2\}_q!}.$$

Assume that the formula holds for each k between 1 and m . The generating function for I_N , where $N = \underbrace{\{n_1, n_1, \dots, n_1\}}_{l_1}, \underbrace{\{n_2, n_2, \dots, n_2\}}_{l_2}, \dots, \underbrace{\{n_m, n_m, \dots, n_m\}}_{l_m}$, can be found in a few steps. First, let us recall a definition of the generating function for the inversion poset

$$GF(I_N) = \sum_{w \in I_N} q^{\text{inv}(w)},$$

where the inversion number of w is a sum of its inversion sequence. Additionally, we introduce a bijection $\varphi : I_N \rightarrow W' \times W''$, where $\varphi(w) = (w', w'')$. The permutation w' is a copy of w without any n_m 's and w'' is a copy of w where each element smaller than n_m is replaced by zero. For example if $N = \{1, 1, 2, 2, 2, 3, 3, 4, 4\}$ and $w = (3, 1, 2, 4, 1, 2, 4, 3, 2)$ then $w' = (3, 1, 2, 1, 2, 3, 2)$, $w'' = (0, 0, 0, 4, 0, 0, 4, 0, 0)$, $\text{inv}(w) = 13$, $\text{inv}(w') = 7$ and $\text{inv}(w'') = 6$.

This bijection splits the inversion number of w into two parts. The inversion number of w'' expresses the contribution of the n_m 's to the inversion number of w and the inversion number of w' indicates the contribution of all other elements. We can represent this relation by the following equation,

$$\text{inv}(w) = \text{inv}(w') + \text{inv}(w'').$$

It is apparent that φ is an injective function. To show that it is also surjective, given a pair w', w'' we can find w by replacing all zeros in w'' with elements from w' in a given order.

This bijection allows to rewrite the generating function for I_N ,

$$\begin{aligned} \sum_{w \in I_N} q^{\text{inv}(w)} &= \sum_{\substack{w \in I_N \\ (w', w'') = \varphi(w)}} q^{\text{inv}(w') + \text{inv}(w'')} = \sum_{(w', w'') \in W' \times W''} q^{\text{inv}(w')} \cdot q^{\text{inv}(w'')} \\ &= \sum_{w' \in W'} q^{\text{inv}(w')} \cdot \sum_{w'' \in W''} q^{\text{inv}(w'')}. \end{aligned}$$

Since W' contains all permutations of $\{\underbrace{n_1, \dots, n_1}_{l_1}, \underbrace{n_2, \dots, n_2}_{l_2}, \dots, \underbrace{n_{m-1}, \dots, n_{m-1}}_{l_{m-1}}\}$, the generating function will be the same as for $I_{\{\underbrace{n_1, \dots, n_1}_{l_1}, \underbrace{n_2, \dots, n_2}_{l_2}, \dots, \underbrace{n_{m-1}, \dots, n_{m-1}}_{l_{m-1}}\}}$ and therefore by the induction assumption,

$$\sum_{w' \in W'} q^{\text{inv}(w')} = \binom{\sum_{i=1}^{m-1} l_i}{l_1, l_2, \dots, l_{m-1}}_q = \frac{\left\{ \sum_{i=1}^{m-1} l_i \right\}_q!}{\{l_1\}_q! \cdot \{l_2\}_q! \cdots \{l_{m-1}\}_q!}.$$

Similarly, W'' contains all permutations of $\{\underbrace{0, 0, \dots, 0, 0}_{l_1+l_2+\dots+l_{m-1}}, \underbrace{n_m, \dots, n_m}_{l_m}\}$ and therefore the generating function will be the same as the generating function for $I_{\{\underbrace{0, 0, \dots, 0, 0}_{l_1+l_2+\dots+l_{m-1}}, \underbrace{n_m, \dots, n_m}_{l_m}\}}$.

This function is our base case, thus

$$\sum_{w'' \in W''} q^{\text{inv}(w'')} = \binom{\sum_{i=1}^{m-1} l_i + l_m}{\sum_{i=1}^{m-1} l_i}_q = \frac{\left\{ \sum_{i=1}^m l_i \right\}_q!}{\left\{ \sum_{i=1}^{m-1} l_i \right\}_q! \cdot \{l_m\}_q!}.$$

Using the general definition of a generating function for an inversion poset we can describe the generating function for $I_{\{\underbrace{n_1, n_1, \dots, n_1}_{l_1}, \underbrace{n_2, n_2, \dots, n_2}_{l_2}, \dots, \underbrace{n_m, n_m, \dots, n_m}_{l_m}\}}$ as a product of the q-multinomial coefficient and the q-binomial coefficient,

$$\begin{aligned} \frac{\left\{ \sum_{i=1}^{m-1} l_i \right\}_q!}{\{l_1\}_q! \cdot \{l_2\}_q! \cdots \{l_{m-1}\}_q!} \cdot \frac{\left\{ \sum_{i=1}^m l_i \right\}_q!}{\left\{ \sum_{i=1}^{m-1} l_i \right\}_q! \cdot \{l_m\}_q!} &= \frac{\left\{ \sum_{i=1}^m l_i \right\}_q!}{\{l_1\}_q! \cdot \{l_2\}_q! \cdots \{l_m\}_q!} \\ &= \binom{\sum_{i=1}^m l_i}{l_1, l_2, \dots, l_m}_q, \end{aligned}$$

as required. \square

Theorem 3.7. *An inversion poset for a multiset I_N is self-dual.*

Proof. Let $F : I_N \rightarrow I_N^*$ be given by

$$F\left((p_1, p_2, \dots, p_m)\right) = (p_m, p_{m-1}, \dots, p_2, p_1).$$

Let a and b be two permutations of I_N such that $F(a) = F(b)$. Elements a and b must be the same, since their reverses are the same and therefore F is one-to-one.

Let $a \leq b$, thus b can be constructed from a by some number of swaps of consecutive elements where the element on the right is greater than the element on the left. Let $C = (a, c_1, c_2, \dots, c_m, b)$ be a saturated chain between a and b . Since $c_m \leq b$, there has to be a pair of consecutive elements in b where the element on the right is less than the element on the left. Thus, we know that $F(b) \leq F(c_m)$ since F reverses the order of b and c_m . Similarly, we can show that $F(c_m) \leq F(c_{m-1}) \leq \dots \leq F(c_1) \leq F(a)$ and therefore $F(b) \leq F(a)$. \square

The next theorem introduces another isomorphism between inversion posets which is convenient when a SSCD is being found.

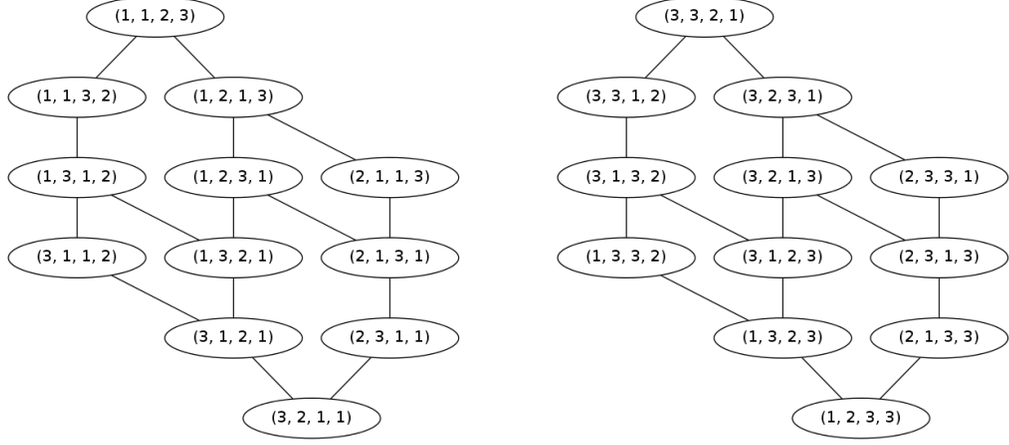
Theorem 3.8. *The inversion poset $I_{\{\underbrace{n_1, \dots, n_1}_{l_1}, \underbrace{n_2, \dots, n_2}_{l_2}, \dots, \underbrace{n_k, \dots, n_k}_{l_k}\}}$ is isomorphic to the dual of the inversion poset $I_{\{\underbrace{n_1, \dots, n_1}_{l_k}, \underbrace{n_2, \dots, n_2}_{l_{k-1}}, \dots, \underbrace{n_k, \dots, n_k}_{l_1}\}}$.*

Proof. For the sake of simplicity let $\bar{n}_i = n_{k-i+1}$ and $m = \sum_{i=1}^k l_i$. Let $F : I_{N_1} \rightarrow I_{N_2}^*$ be a function given by

$$F\left((p_1, p_2, \dots, p_m)\right) = (\bar{p}_1, \bar{p}_2, \dots, \bar{p}_m),$$

where $N_1 = \{\underbrace{n_1, \dots, n_1}_{l_1}, \underbrace{n_2, \dots, n_2}_{l_2}, \dots, \underbrace{n_k, \dots, n_k}_{l_k}\}$ and $N_2 = \{\underbrace{n_k, \dots, n_k}_{l_1}, \underbrace{n_{k-1}, \dots, n_{k-1}}_{l_2}, \dots, \underbrace{n_1, \dots, n_1}_{l_k}\}$.

Consider two permutations a and b of I_{N_1} such that $F(a) = F(b)$. It is clear from the definition of F that since $F(a) = F(b)$ then $a = b$ and therefore F is one-to-one.



(a) The dual of the inversion poset $I_{\{1,1,2,3\}}$. (b) The inversion poset $I_{\{1,2,3,3\}}$.

Figure 3.2: An illustration of an isomorphism between $I_{\{1,1,2,3\}}$ and $I_{\{1,2,3,3\}}$.

To prove the isomorphism we have to show that $a \leq b$ if and only if $F(a) \geq F(b)$. Since we showed that F is one-to-one it is enough to show that $a < b$ if and only if $F(a) > F(b)$. To show this, we will prove that $a < b$ if and only if $F(a) > F(b)$ because these statements are equivalent since there is always a saturated chain between a and b if $a < b$.

Let $a = (a_1, a_2, \dots, a_i, a_{i+1}, \dots, a_m)$, where $a_i < a_{i+1}$ and suppose that $a < b$, with $b = (a_1, a_2, \dots, a_{i+1}, a_i, \dots, a_m)$.

$$F(b) = (\overline{a_1}, \overline{a_2}, \dots, \overline{a_{i+1}}, \overline{a_i}, \dots, \overline{a_m})$$

Let's focus on the pair $(\overline{a_{i+1}}, \overline{a_i})$. Since $a_i < a_{i+1}$ we can say that $n_{j_1} = a_i < a_{i+1} = n_{j_2}$, where $j_1 < j_2$. Then $\overline{a_i} = \overline{n_{j_1}} = n_{k-j_1+1}$ and $\overline{a_{i+1}} = \overline{n_{j_2}} = n_{k-j_2+1}$. It follows from the inequality $j_1 < j_2$ that $k - j_2 + 1 < k - j_1 + 1$, $n_{k-j_2+1} < n_{k-j_1+1}$ and hence $\overline{a_{i+1}} < \overline{a_i}$. Therefore, we can conclude that

$$F(b) = (\overline{a_1}, \overline{a_2}, \dots, \overline{a_{i+1}}, \overline{a_i}, \dots, \overline{a_m}) < (\overline{a_1}, \overline{a_2}, \dots, \overline{a_i}, \overline{a_{i+1}}, \dots, \overline{a_m}) = F(a),$$

as required.

If we follow previous steps in the opposite direction we get that $F(a) \succ F(b)$ implies $a < b$, which completes the proof. \square

The previous theorem implies that $I_{\{1,1,2,3\}}$ is isomorphic to the dual of $I_{\{1,2,3,3\}}$. Additionally, the dual of the inversion poset $I_{\{1,2,3,3\}}$ is isomorphic to $I_{\{1,2,3,3\}}$, because the inversion poset is self-dual. Therefore, Theorems 3.7 and 3.8 yield the following corollary.

Corollary 3.9. *The inversion poset $I_{\{\underbrace{n_1, \dots, n_1}_{l_1}, \underbrace{n_2, \dots, n_2}_{l_2}, \dots, \underbrace{n_k, \dots, n_k}_{l_k}\}}$ is isomorphic to the inversion poset $I_{\{\underbrace{n_1, \dots, n_1}_{l_k}, \underbrace{n_2, \dots, n_2}_{l_{k-1}}, \dots, \underbrace{n_k, \dots, n_k}_{l_1}\}}$.*

Chapter 4

The algorithm

We described, with a proof, an algorithm for finding a SSCD for a division poset in Chapter 3. We introduced Inverse posets and Young's lattices but we didn't indicate any decomposition for these posets. The reason for this is that there is no general proof that these posets have a SSCD. Stanley gave a combinatorial proof that a Young's lattice, $L(m, n)$, is rank unimodal and has the Sperner property in [9]. Although, it was proven by Sylvester in 1878 that $L(m, n)$ is rank unimodal, Stanley gave the first combinatorial proof. He also conjectured that $L(m, n)$ has a SSCD for every positive m and n . This conjecture has not yet been proven for a general case, only cases where $\min(m, n) \leq 4$ were proven [3].

We will introduce an enhanced algorithm for finding a SSCD for a general ranked poset \mathcal{P} based on the previous work of Katsumata in [7]. The algorithm starts with $W_{\lfloor \rho(\mathcal{P})/2 \rfloor}$ chains located in the middle of \mathcal{P} since the center of every symmetric saturated chain is located in the middle layer(s). It will create chains of length 0 or 1 based on the number of layers and then it will augment some of these chains in a way that they will remain symmetric and saturated.

Our algorithm differs significantly from the algorithm provided by Katsumata, since it uses a flow algorithm for the augmentation of chains and it can work with any poset

given by its adjacency list, not just with an inversion poset I_n . If the program finds a SSCD of the poset, it saves this decomposition into a text file and creates four pictures of Hasse diagrams with highlighted chains. Pictures for selected posets are displayed in appendix B.

4.1 Finding a flow in a graph

A directed graph can represent many structures from everyday life, like public transportation, supply chains, road maps, computer networks or simplified water supply system. Each of these structures can be represented as a system of pipes, connected with joints, with a flow of an incompressible fluid. We will focus on a system with only one point of entry and only one exit point for the fluid and we will try to investigate what the maximal flow through the system is.

This abstraction gives rise to three constraints on the flow based on physics. There is a limitation on how fast we can transport the fluid through a pipe based on the material of the pipe, its radius and a viscosity of the fluid and other factors. This limitation is called a capacity and each pipe can have a different capacity. The next constraint is called antisymmetry and it means that a flow in a pipe is uniform – the flow is always in only one direction. The last constraint is that fluid can leave the system only at one joint and it can enter only at one joint and these joints have to be different. The exact definitions follow.

Let a directed graph $G = (V, E)$ with a capacity function $c : V^2 \rightarrow \mathbb{R}_0^+$, where $c(e) = 0$ if $e \notin E$, is given. Furthermore, denote sizes of V and E by n and m , respectively. A *flow* between two distinct vertices, source s and sink t , is given by a function $f : V^2 \rightarrow \mathbb{R}_0^+$ such that $\forall e \in E$:

$$f(e) \leq c(e) \quad (\text{capacity constraint}), \quad (4.1)$$

$$f((u, v)) = -f((v, u)) \text{ where } (u, v) = e \quad (\text{antisymmetry constraint}), \quad (4.2)$$

$$\sum_{(u,v) \in V \setminus \{s,t\}} f((u,v)) = 0 \quad (\text{flow conservation constraint}). \quad (4.3)$$

The *value* $|f|$ is defined as the total flow into the sink, i.e. $|f| = \sum_{u \in V} f((u,t))$. A *maximum flow* is a flow of maximum value. We define a *residual capacity* $c_r(e)$ to be a difference between a capacity of an edge and a flow through an edge, $c_r(e) = c(e) - f(e)$. An edge is *saturated* if its residual capacity is zero.

Many algorithms are available for finding a maximum flow in a graph [5]. We will focus on two types of flow algorithms, on the Ford–Fulkerson algorithm and the Push–Relabel algorithm.

The Ford–Fulkerson algorithm

This algorithm was first published in 1956 and improved versions of this algorithm were published by Dinic in 1970 and by Edmonds and Karp in 1972 [5]. This algorithm works as follows:

1. $f((u,v)) \leftarrow 0$ for every pair of vertices in V
2. while there exist a path p between s and t such that $c_r(e) > 0$ for every edge in this path p :
 - (a) denote the smallest residual capacity in p by r
 - (b) for each edge (u,v) in p :
 - i. $f((u,v)) \leftarrow f((u,v)) + r$
 - ii. $f((v,u)) \leftarrow f((v,u)) - r$

The Ford–Fulkerson algorithm finds a path through unsaturated edges in each iteration and it adds this path to the flow. This path is frequently called an augmenting path and the performance of this algorithm depends on an algorithm used for finding this path. Finding a path in a graph is generally implemented as a depth–first or breadth–first search on vertices. Both of these methods have their advantages. The Ford–Fulkerson

algorithm has a running time of $O(n^2m)$ or $O(nm^2)$ depending on whether a breadth-first or a depth-first search is used, respectively [6].

Using the Ford–Fulkerson algorithm, we have to find a new path between s and t in every cycle. This disadvantage is addressed in the next maximum flow algorithm.

The Push–Relabel algorithm

This algorithm does not find only one path from the source to the sink as the Ford–Fulkerson algorithm, but it pushes through the graph as much flow as is possible. For this purpose it manages a preflow and a vertex labeling d . *Preflow* is a function from V^2 to \mathbb{R} which satisfies equations (4.1) and (4.2) for every edge in the graph. Preflow does not have to conserve the flow, it allows an incoming flow into a vertex to be bigger than an outgoing flow from the vertex. The algorithm starts with a preflow and it improves it into a maximal flow. Any valid preflow can be used, but commonly a preflow f_p where $f_p((s,p)) = c((s,p))$ for every $(s,p) \in E$, where s is the source, and $f_p(e) = 0$ for other edges is used.

After an initialization the algorithm either pushes a flow through a vertex or it updates the labeling. This vertex labeling provides estimates for a distance from the source and a distance to the sink for each vertex. The first implementation of this algorithm, by Goldberg in 1985, had a running time of $O(n^3)$ [6]. One year later, Goldberg and Tarjan published an implementation with a running time of $O(nm \log(n^2/m))$ [5].

4.2 Algorithm description

Our algorithm, similar to an algorithm of Katsumata, is implemented in three stages. These stages are making a poset, finding a SSCD and arrangement of chains.

1. Firstly, we create the Hasse diagram of a given poset or we read the poset structure from files. We represent the poset as a directed graph, e.g. as a list of vertices and an adjacency list, V and E , respectively. This graph can be partitioned into

levels since we are representing the Hasse diagram and we store these levels in another list to improve the performance. Denote the rank of the poset by k and let $N = \{N_0, N_1, \dots, N_k\}$ be the collection of level sets. Additionally, we will create an empty list C for storing a SSCD.

When the first stage is finished, the list N has $k+1$ elements in it: N_0, N_1, \dots, N_k . Each of these lists contains vertices with the same rank.

2. If k is odd, there are two middle levels in the poset, we have to find a perfect matching between them. This matching yields the two middle elements from each chain and it can be found by making a graph H . This graph contains vertices from $N_{(k-1)/2}$, $N_{(k+1)/2}$ and two other vertices s and t . We add all edges between $N_{(k-1)/2}$ and $N_{(k+1)/2}$ from E to H , connect s with every vertex from $N_{(k-1)/2}$ and connect each vertex from $N_{(k+1)/2}$ to t . Finding a maximum flow in H between s and t , where the capacity of every edge is 1, yields the matching. If the flow value is equal to $|N_{(k-1)/2}|$, we erase edges with no flow in H , all incidence edges of s , t and the vertices s and t . Edges which are left are saved into C . If the value is smaller, the poset does not have a SSCD.

If k is even, we fill C with vertices from $N_{k/2}$.

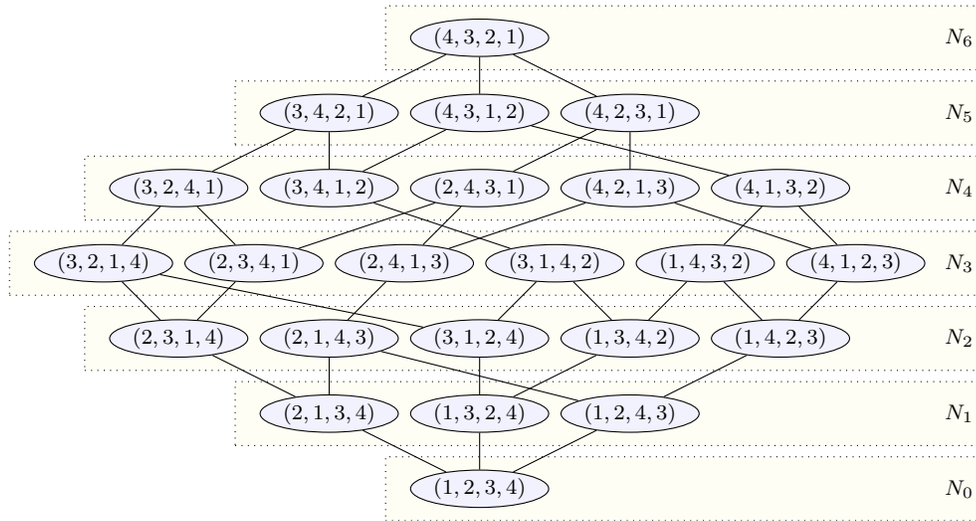
For $i = \lfloor k/2 \rfloor, \lfloor k/2 \rfloor - 1, \lfloor k/2 \rfloor - 2, \dots, 1$ perform following:

- (a) Make a graph H which contains vertices from N_{i-1} , N_i , N_{k-i} , $N_{k-(i-1)}$ and two extra vertices s and t . Connect s with each vertex in N_{i-1} and each vertex in $N_{k-(i-1)}$ with t . If there is an edge in E between vertices from N_{i-1} and N_i , connect the corresponding vertices in H . Similarly, for layers N_{k-i} and $N_{k-(i-1)}$. Layers N_i and N_{k-i} are connected using chains in C , each chain which starts in N_i and ends in N_{k-i} will be represented as an edge in H .
- (b) Find a maximum flow between s and t , where the capacity of every edge

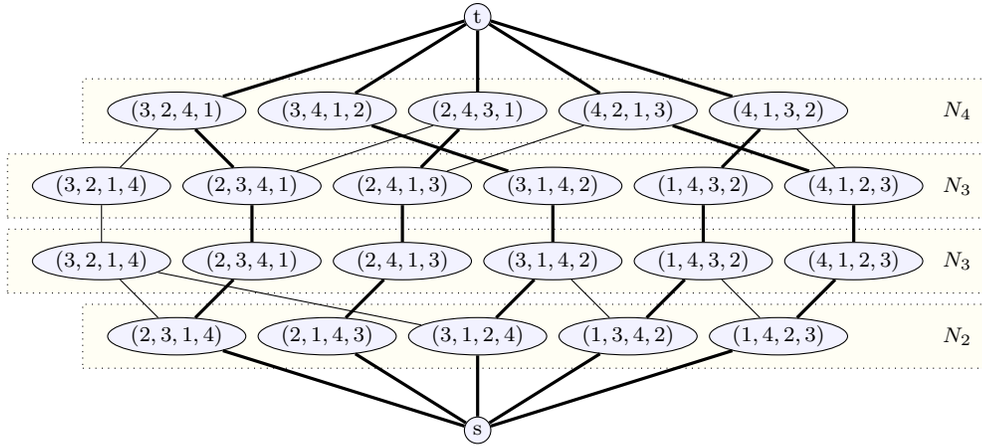
is one. If the value of the flow is equal to $|N_i|$, we remove all edges without any flow, vertices s and t and their adjacent edges. The next step is extending the chains in C , which correspond to the edges between N_i and N_{k-i} , with the flow. The list C contains now a SSCD for subgraph induced by $N_{i-1}, N_i, N_{i+1}, \dots, N_{k-(i-1)}$ levels of G . If the value is smaller, the poset does not have to have a SSCD.

3. A symmetric saturated chain decomposition is stored in C .

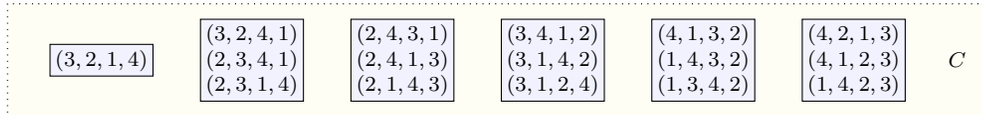
Let us follow steps of this algorithm on I_4 . We start by creating the Hasse diagram of the poset and dividing its vertices into seven level sets: N_0, N_1, \dots, N_6 as shown below.



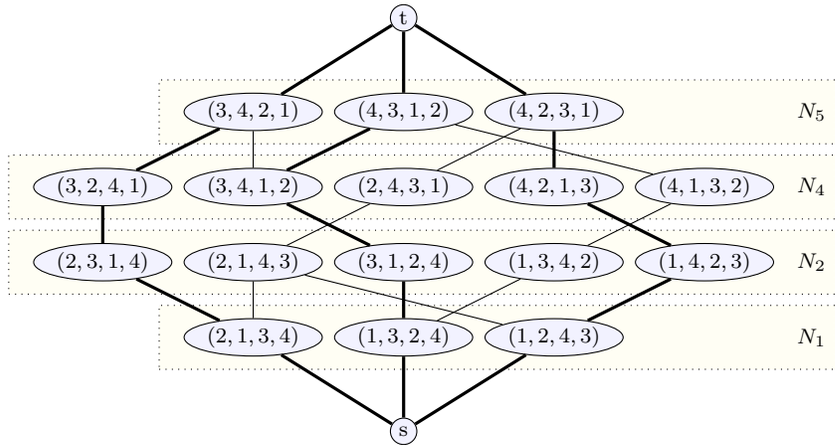
In the next step we save N_3 into C and set $i = 3$ since the rank of I_4 is 6. A graph H contains vertices from levels N_2, N_3, N_3 and N_4 of the original graph. The edges between N_2 and N_3 are determined by the original graph, likewise the edges between N_3 and N_4 . Additionally, corresponding vertices in N_3 and N_3 are connected by an edge. The next picture contains a diagram of H with a maximum flow for $i = 3$. The edges with a nonzero flow are denoted by a thicker line.



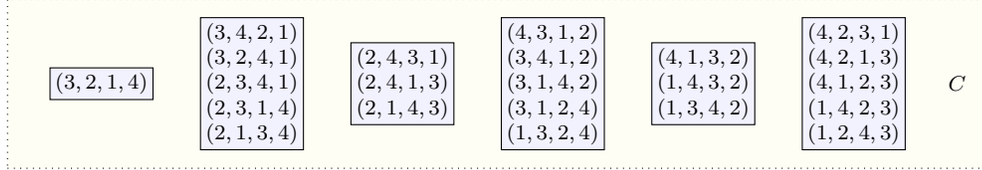
When we find a maximum flow, we remove all edges without any flow and vertices s and t . Remaining chains are used for extending chains in C . The list C is shown beneath.



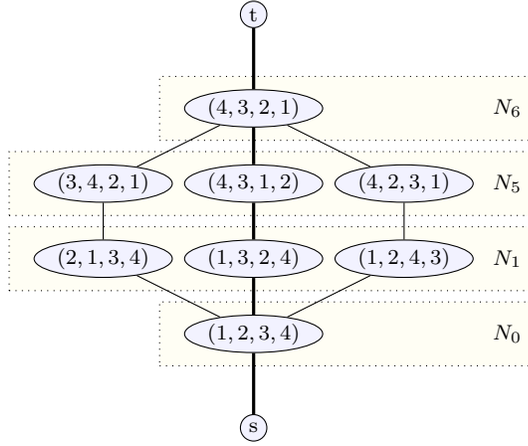
In the next iteration, when $i = 2$, the graph H will contain levels N_1, N_2, N_4 and N_5 . Additional vertices s and t are connected to N_1 and N_5 , respectively. The diagram of H with a maximum flow is shown below.



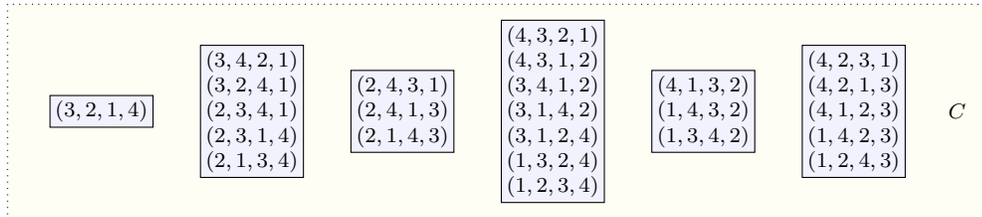
Removing unsaturated edges and vertices s and t from H yields three saturated chains of length 3. These three chains are used to extend some of the existing chains in C . The list C after the extension is illustrated below.



In the last iteration, the graph H contains levels N_0, N_1, N_5 and N_6 since $i = 1$.



A maximum flow in G yields only one chain because N_0 and N_6 contain only one vertex. Therefore, we expand only one chain in C , this is the longest chain in I_4 .



When the last chain is extended, C will contain a SSCD of I_4 .

The running time of this algorithm is polynomial, since it does not include backtracking. For this reason, the algorithm might not find a SSCD even if a SSCD exists.

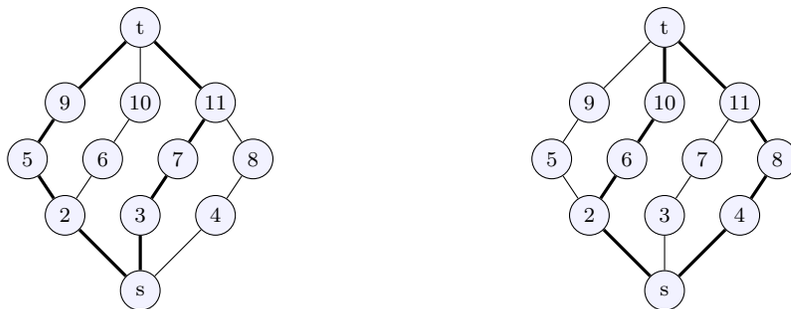


Figure 4.1: Two different flows in a graph.

If the algorithm finds a maximal flow, with its value smaller than the number of neighbors of s or t , it fails to find a SSCD. The algorithm has a running time of $O(n^3m)$ or $O(n^2m^2)$, using the Ford–Fulkerson algorithm, depending on whether a breadth–first or depth–first search is used. The running time is $O(n^2m \log(n^2/m))$ if the Push–Relabel algorithm is used.

Anomalies

Some posets do not have a SSCD. If we run the described algorithm on these posets we should be able to recognize this. One of the first steps of the algorithm, if we run it on the poset shown in Figure 1.1(d), is constructing the graph in Figure 4.1 and finding a maximal flow. Two maximal flows are highlighted in this picture and as we can see, there is always one edge adjacent to s or t with no flow. The algorithm continues with removing s, t and edges without any flow, this step will continue without any problems. But contracting all remaining edges will not create symmetric saturated chains and the algorithm fails. Actually, the algorithm fails on this poset every time; there is at least one edge adjacent to s or t with no flow and our implementations of this algorithm contains such a check. This behavior is needed, since there is no SSCD of this poset.

As mentioned before, the described algorithm does not include backtracking. The algorithm processes a poset from middle layer(s) outward and it does not return to already processed layers. This is the main reason why the running time is polynomial,

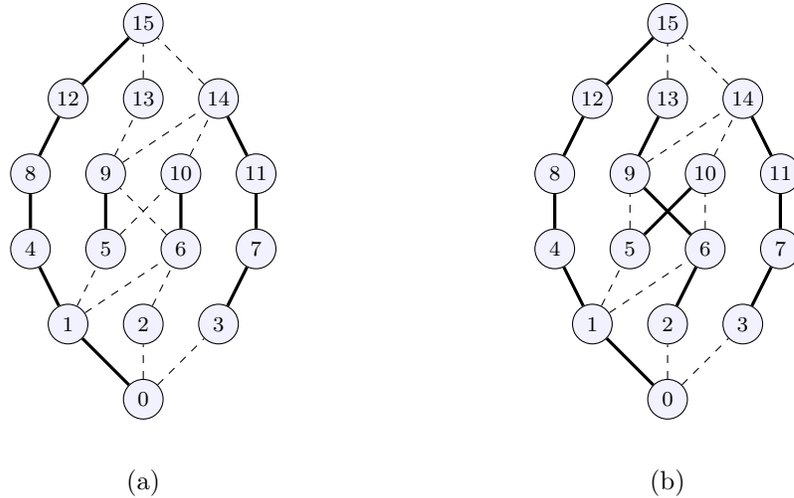


Figure 4.2: An example of two different maximal flows between middle layers and how they can affect the poset decomposition.

but on the other hand this may cause additional problems. While a SSCD contains a maximal flow between layers which are symmetric to the middle layer(s), not every maximal flow between these layers is extendable to a SSCD. There are two choices for a maximal flow between the middle layers for the poset shown in Figure 4.2. If the algorithm picks the maximal flow which contains the edge $(5, 9)$, then it fails to find a SSCD because there will be always two unconnected vertices as in Figure 4.2(a). On the other hand if we choose the second maximal flow between middle layers, we would be able to extend this chains into a SSCD, this is shown in Figure 4.2(b).

There is no simple solution for this problem, the algorithm would have to contain backtracking to solve this and the total running time of such an algorithm would be significantly larger. Fortunately, we were able to find a SSCD for all important posets we ran this algorithm on, even with this imperfection. This was surprising and led us to additional statistics described in the next chapter focused on maximal chains in a poset.

Chapter 5

Results

Two computer programs, which are listed in appendix C, were written using Python and C++. Both of these programs use advanced graph libraries – NetworkX and PyGraphviz in Python and Boost libraries in C++. Both programs can find a SSCD for an inversion poset given its multiset or for a Young’s lattice given its two constants m and n . The program written in C++ can additionally find a SSCD for a Boolean algebra and it can read a poset from files. To make sure that our programs provide correct results we checked some decompositions by another program. This additional program checked that the decomposition contained each element of a posed exactly once and that the chains were valid and symmetric.

While the Boost libraries implement the Push–Relabel algorithm, the NetworkX library in Python does not implement this algorithm. Unfortunately, NetworkX implements only the Ford–Fulkerson algorithm with a depth–first search approach. The running time of this algorithm allowed us to find a SSCD, in a reasonable time, for all inversion posets where the multiset contained less than 11 elements. This was a big improvement from the work of Katsumata but we were able to successfully run this program on an inversion poset with 11 distinct elements after improving the Ford–Fulkerson algorithm.

We observed that the performance of the Ford–Fulkerson algorithm with the depth–first search varies significantly. The algorithm is fast at the beginning but it slows down rapidly. On the other hand, the breadth–first search is slower than the depth–first search for paths at the beginning, but its performance does not change over time. It seems that the best solution can be achieved by combining these two approaches. Our implementation of the Ford–Fulkerson algorithm uses the depth–first search first and it switches to the breadth–first search when the breadth–first search is faster.

Additionally, we managed to improve the breadth–first search as well. The regular breadth–first search from a source in a graph would visit all neighbors of the source first, then it would visit neighbors of those neighbors, etc. Our improved implementation of the breadth–first search randomly chooses some of source’s neighbors and performs full breadth–first search on them. If it will not find a path from the source to the sink it continues with the next part of nonvisited neighbors of the source until it finds a path from the source to the sink. We achieved the best performance when the algorithm processed 2% of source’s neighbors at a time.

The implementation in C++ does not need this improvement because Boost libraries contain an implementation of the Push–Relabel algorithm. This flow algorithm is faster than our improved Ford–Fulkerson algorithm. Since this program is faster than implementation in Python, we extended the input options for it. It can find a SSCD for inversion posets, Young’s lattices, Boolean algebras and it can read a poset from files with its structure.

As mentioned in Chapter 3, we can use Corollary 3.9 when testing all inversion posets. Suppose that we would like to find a SSCD for all inversion posets where the multiset contains 4 distinct elements and its length is 6. Let us denote the elements as 1, 2, 3 and 4, where the order is naturally $1 < 2 < 3 < 4$. There are 10 different possible multisets. This means that we would have to run the program ten times, but using Corollary 3.9 we can reduce the number of instances to 6 because some of these multisets will create isomorphic posets. A list with these combinations of multisets follows.

Poset name	Size
Boolean algebra B_n	$n \leq 25$
Inversion poset for a multiset I_N	$ N \leq 11$
Young's lattice $L(m, n)$	$m + n \leq 30$

Table 5.1: Summary of posets with the maximal size for which a symmetric saturated chain decomposition was found by our program.

1. $\{1, 1, 1, 2, 3, 4\}, \{1, 2, 3, 4, 4, 4\}$
2. $\{1, 2, 2, 2, 3, 4\}, \{1, 2, 3, 3, 3, 4\}$
3. $\{1, 1, 2, 2, 3, 4\}, \{1, 2, 3, 3, 4, 4\}$
4. $\{1, 1, 2, 3, 3, 4\}, \{1, 2, 2, 3, 4, 4\}$
5. $\{1, 1, 2, 3, 4, 4\}$
6. $\{1, 2, 2, 3, 3, 4\}$

Each multiset in one row will yield the same poset, up to isomorphism. There are 2047 multisets with less than 12 elements, but only 1086 of them are different up to isomorphism.

As mentioned before, the program written in C++ is faster than the program in Python and therefore all our tests were performed with this program. Table 5.1 contains all posets we tested and for which we found a SSCD. It is surprising that the program never failed and always found a SSCD for all tested inversion posets, Young's lattices and Boolean algebras since we provided a poset for which our program fails. It may be possible to extend these boundaries a little bit, but this depends on the computer which runs the program.

n	Number of different longest chains in I_n	Bad chains
4	$16 = \frac{6!}{5 \cdot 3 \cdot 3}$	4
5	$768 = \frac{10!}{7 \cdot 5 \cdot 5 \cdot 3 \cdot 3 \cdot 3}$	147
6	$292,864 = \frac{15!}{9 \cdot 7 \cdot 7 \cdot 5 \cdot 5 \cdot 5 \cdot 3 \cdot 3 \cdot 3}$	24,363
7	$1,100,742,656 = \frac{21!}{11 \cdot 9^2 \cdot 7^3 \cdot 5^4 \cdot 3^5}$	Out of reach
8	$48,608,795,688,960 = \frac{28!}{13 \cdot 11^2 \cdot 9^3 \cdot 7^4 \cdot 5^5 \cdot 3^6}$	Out of reach
9	$29,258,366,996,258,488,320 = \frac{36!}{15 \cdot 13^2 \cdot 11^3 \cdot 9^4 \cdot 7^5 \cdot 5^6 \cdot 3^7}$	Out of reach
10	$273,035,280,663,535,522,487,992,320 = \frac{45!}{17 \cdot 15^2 \cdot 13^3 \cdot 11^4 \cdot 9^5 \cdot 7^6 \cdot 5^7 \cdot 3^8}$	Out of reach

Table 5.2: Number of different longest chains in inversion posets I_n , where $4 \leq n \leq 10$.

5.1 Future work

There are several ways this work might be extended. From a computational point of view, the main reason we could not run our programs on bigger posets was the amount of memory needed. Our implementations construct a poset and keep it as list of vertices and list of edges in the memory. However, because we have to keep only three layers of the poset in memory, the whole poset can be saved in external memory and loaded into internal memory only when needed. This approach, when using a compression, should be investigated in the future.

The memory issue may have an another solution. Distributing the vertex and edge lists over more computers should allow to improve the upper bounds we set in this paper. Distributed flow algorithms are available, although the required communication can be a new limiting factor.

We modified our program to count the number of maximal chains in inversion posets I_n and to find a SSCD given a maximal chain. The number of maximal chains corresponds to the number of different standard Young tableaux of shape $(n-1, n-2, \dots, 2, 1)$ and it can be found by the Hook formula [11]. This is another piece of evidence that

our program performs correctly. The program did not find a SSCD for some maximal chains; we call these chains bad chains. Table 5.2 shows how many maximal chains I_n has and for small n , how many bad chains I_n has. For example, in I_4 , 4 of the 16 maximal chains cannot be extended to a SSCD by our algorithm. Since our algorithm can fail to find a SSCD, table 5.2 provides only upper bounds on the number of bad chains. However, we checked that the exact number of bad chains for I_4 is 4.

Another way to find a SSCD can be by finding symmetric chains from the longest to the shortest. This approach itself may not perform well, but it may be combined with the approach used in our algorithm, since it seems that the majority of the longest chains will not make the rest of the poset undecomposable into symmetric saturated chains.

From a mathematical point of view, Stanley's conjecture that there is a SSCD for any Young's lattice as well as for any inversion poset is still open. However, some progress has been made in determining the Sperner property and other combinatorial properties of these and similar posets [3].

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Appendix A

Program interface

We explain how to compile and run programs we developed and what each input parameter means in this chapter. In our description we will assume that we work on a Debian-based Linux machine.

A.1 Python

To successfully run this program we need to have files *diagram.py* and *main.py* inside a folder and install two additional Python libraries, NetworkX and PyGraphviz. The program can be called from a terminal by using this command

```
|| $ python main.py [-h] [--inversion 1 2 3 4 ...] [--young m n] [--no_pict]
```

The optional arguments are

- `--h` or `--help`
 - This argument prints a help message and exits the program.
- `--inversion`
 - This argument starts the program for an inversion poset and a name of the minimal vertex with the smallest rank follows as a list of numbers, each

separated by a space. For example a SSCD for I_5 can be found by this command

```
|| $ python main.py --inversion 1 2 3 4 5
```

- `--young`

- Two numbers, m and n , separated by a space follow this argument. The program tries to find a SSCD for a Young’s lattice $L(m, n)$. For example a SSCD for $L(5, 5)$ can be found by this command

```
|| $ python main.py --young 5 5
```

- `--no_pict`

- The program automatically stores the decomposition in a text file and it creates pictures of the Hasse diagram of the poset with the decomposition highlighted. When using this argument, no pictures will be created. Generating pictures for big posets like I_{10} is time consuming and the usability of such pictures is questionable.

If the program finds a SSCD it saves it together with its output into a text file and creates four pictures, depending on the `--no_pict` argument.

“..._FULL.png” Contains all vertices with labels. Each level is ordered exactly as it is saved in the memory.

“..._FULL_NO_LABELS.png” Contains all vertices without labels. Each level is ordered exactly as it is saved in the memory.

“..._SPARSE.png” Contains all vertices with labels. Each level is ordered such that the final graph has less intersecting edges.

“..._SPARCE_NO_LABELS.png” Contains all vertices without labels. Each level is ordered such that the final graph has less intersecting edges.

A.2 C++

To be able to compile the program we need to install Boost libraries. Optionally, we can install the OpenMP library if we have more CPUs available. The program can be compiled by this command

```
$ g++ -o main main.cpp node.cpp node.h -lboost_program_options
-lboost_serialization -lboost_system -lboost_filesystem
-I/usr/include/python2.7 -lpython2.7 -fopenmp -O3
```

assuming that Python 2.7 is installed on the system. This will generate an executable file named “main” which contains our program. The program can be called by using this command

```
$ ./main [--bool n] [--file name] [--inversion 1 2 3 4 ...] [--young m n]
[--no_pict] [--help]
```

The program in C++ shares some arguments with the program in Python and their effects are the same. These arguments are `--inversion`, `--young`, `--no_pict` and `--help`. The rest of arguments are described below.

- `--bool`

- This argument together with a number n will find a SSCD for a Boolean algebra B_n . The program call for B_6 is

```
$ ./main --bool 6
```

- `--file`

- A string (name) follows this argument. The current folder has to contain two files, “name_levels” and “name_nodes”. As their names may suggest,

“name_levels” contains nodes separated into layers and “name_nodes” contains neighbors of each vertex corresponding to the line number. Vertices have to be identified by sequential numbers starting from zero. For example the poset, for which our algorithm fails, from Figure 4.2 can be saved into two files:

my_levels:

```
0 0
1 1,2,3
2 4,5,6,7
3 8,9,10,11
4 12,13,14
5 15
```

my_nodes:

```
0 1,2,3
1 4,5,6
2 6
3 7
4 8
5 9,10
6 10,9
7 11
8 12
9 13,14
10 14
11 14
12 15
13 15
14 15
15 15
```

As we see, the last vertex has to be connected to itself. The algorithm will fail to find a SSCD for this configuration, but the poset has a SSCD.

If we change the sixth line of *my_nodes* to “9,10” instead of “10,9” the algorithm will find the SSCD because it will find a maximal flow which is shown in Figure 4.2(b). The program can be called by

```
|| $ ./main --file my
```

Additionally, all pictures are generated by the Python class *diagram.py* and therefore this class has to be in the same folder as the executed *main* file if the argument `--no_pict` is not used. If the argument `--no_pict` is used, then program generate the same pictures as the program written in Python.

Appendix B

Performance of programs and generated figures

We provide running times for our programs in selected posets in Table B.1 and Figures B.1 and B.2. Note that these times do not include the time needed to generate the poset in memory.

Program version		Time needed to find a SSCD					
Inversion poset, I_n							
$n =$	6	7	8	9	10	11	
C++	1 s	1 s	1 s	15 s	175 s	38.5 min	
Python, Ford–Fulkerson from NetworkX	1 s	5.1 s	278.7 s	11.72 h	n/a	n/a	
Python, improved Ford–Fulkerson	1 s	2.5 s	75.2 s	2.86 h	n/a	n/a	
Young’s lattice, $L(m, n)$							
$(m, n) =$	(7,7)	(8,8)	(9,9)	(10,10)	(11,11)	(12,12)	
C++	1 s	1 s	2 s	8 s	35 s	170 s	
Python, Ford–Fulkerson from NetworkX	2.1 s	25.6 s	453.4 s	148.9 min	56.3 h	n/a	
Python, improved Ford–Fulkerson	1.3 s	12.4 s	141.0 s	37.1 min	12.4 h	n/a	

Table B.1: Performance of each program on selected posets.

When we focus on the graphs which correspond to the program written in C++, it appears that the last three points of each graph form a line. Since the axes of both graphs have a logarithmic scale, we can estimate the program complexity using the

slopes of these lines. The slope for the line which approximates the performance for inversion posets is 1.07; this implies that the running time is $O(n^{1.07})$. Similarly, the slope of the line which approximates the last three points for Young's lattices is 1.14 which corresponds to $O\left(\binom{m+n}{n}^{1.14}\right)$. Using these estimations we can conclude that if we had a computer with enough memory, if the program did not fail, we would need 9.2 and 22.2 hours to find a SSCD for I_{12} and $L(16, 16)$, respectively.

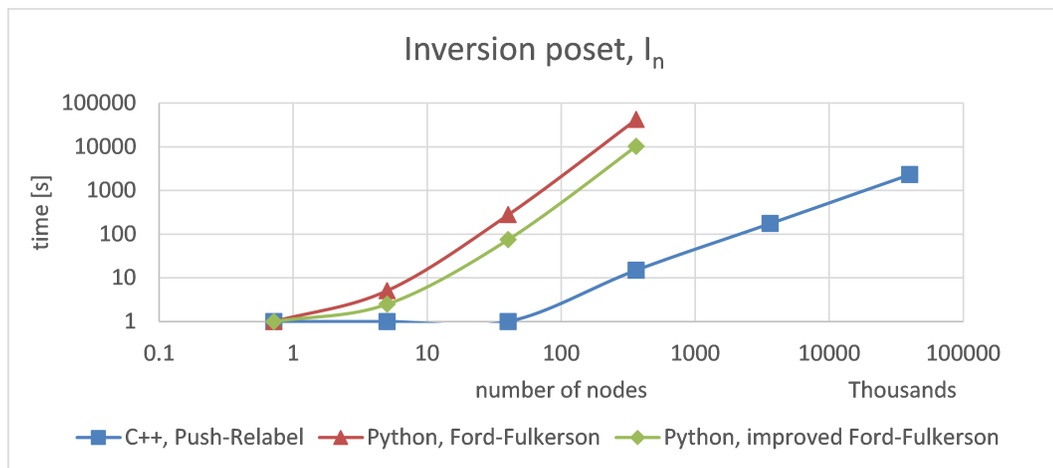


Figure B.1: Performance of programs on inversion posets.

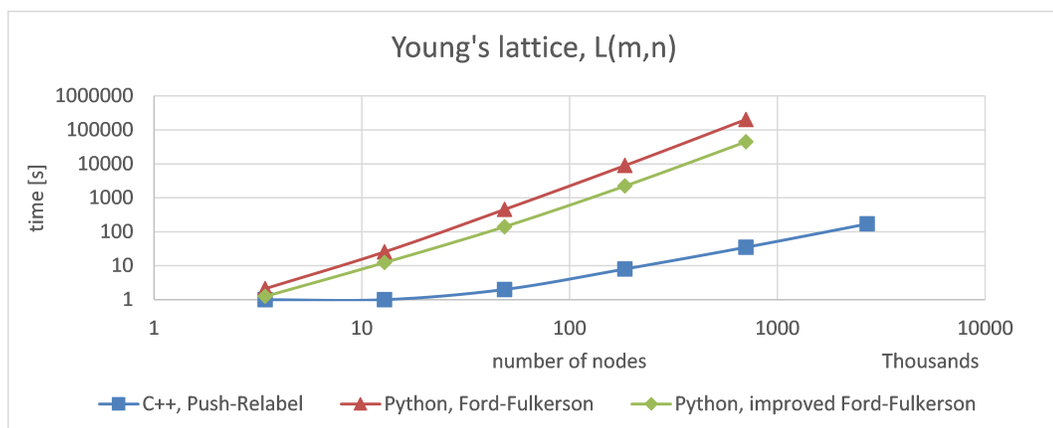
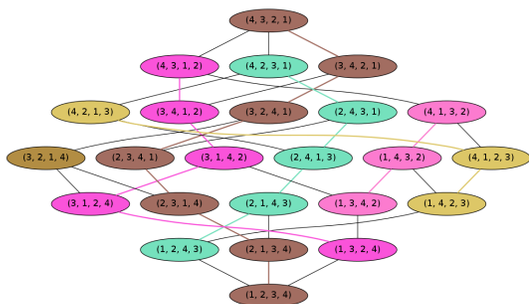
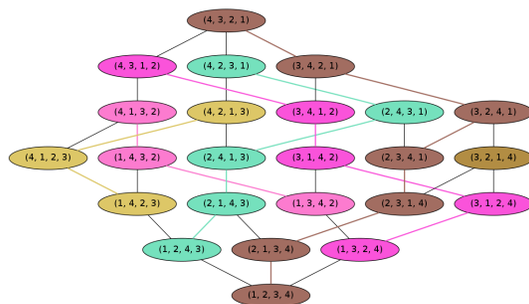


Figure B.2: Performance of programs on Young's lattices.

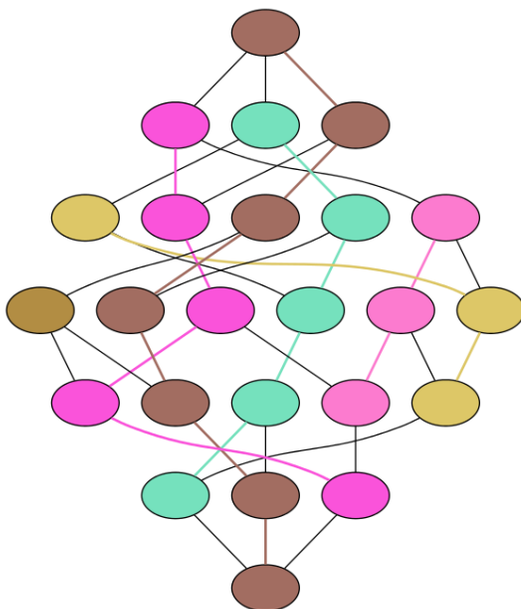
Selected figures generated by the program are displayed below and on the following pages.



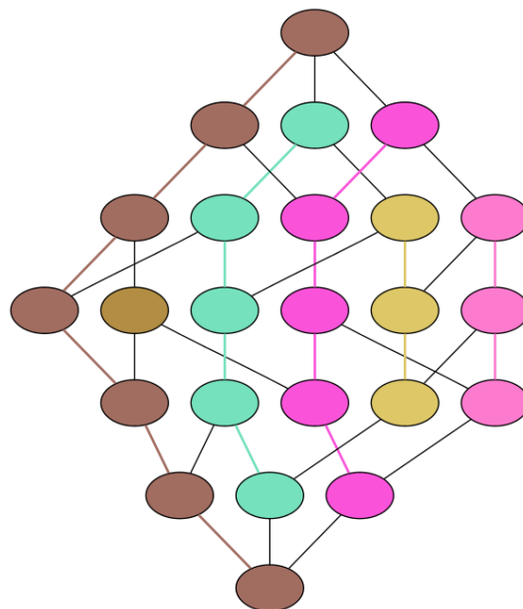
(a) 1,2,3,4_FULL.png



(b) 1,2,3,4_SPARSE.png



(c) 1,2,3,4_FULL_NO_LABELS.png



(d) 1,2,3,4_SPARSE_NO_LABELS.png

Figure B.3: Inversion poset, I_4

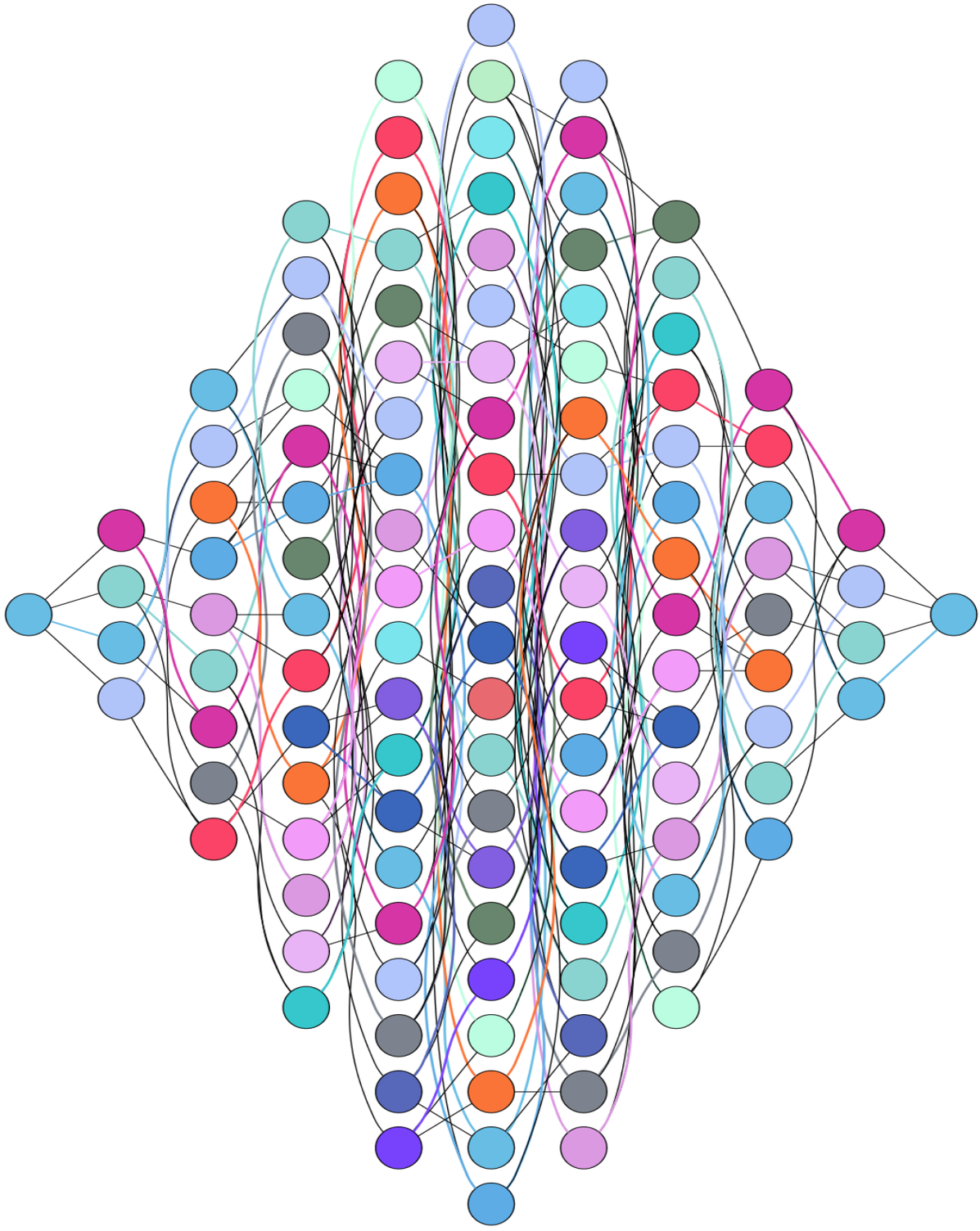


Figure B.4: I_5 : 1,2,3,4,5_FULL_NO_LABELS.png

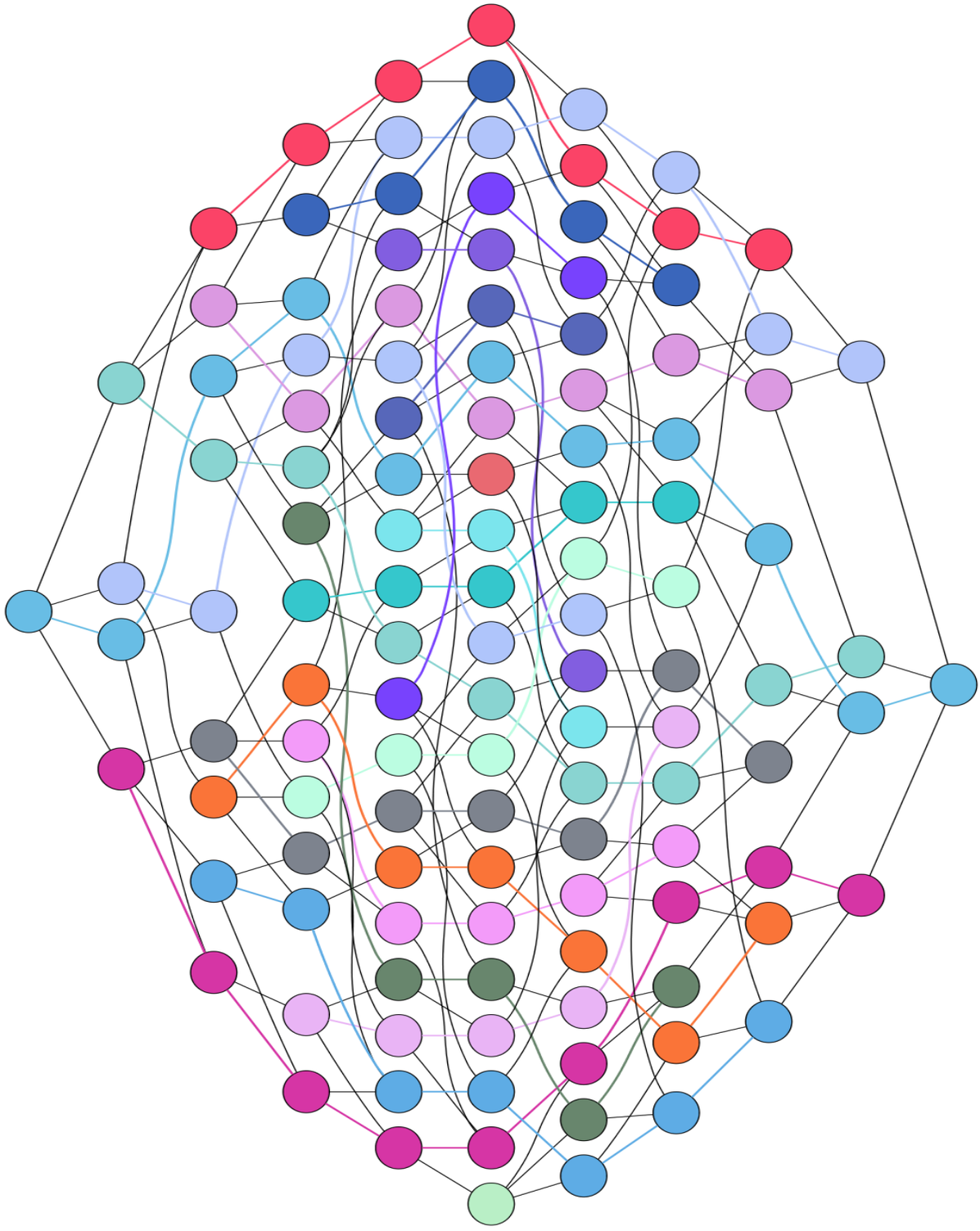


Figure B.5: I_5 : 1,2,3,4,5_SPARSE_NO_LABELS.png

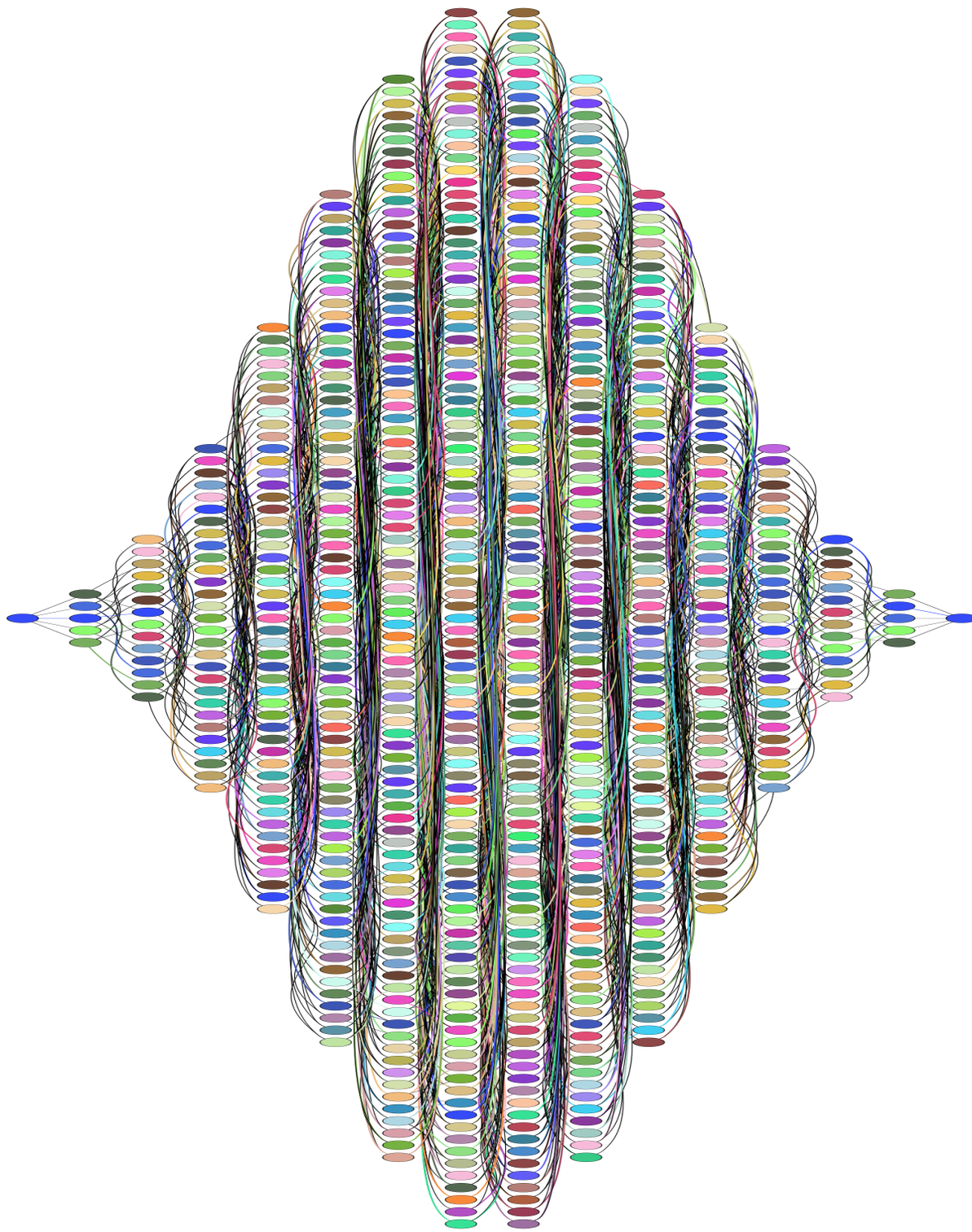


Figure B.6: I_6 : 1,2,3,4,5,6_FULL_NO_LABELS.png

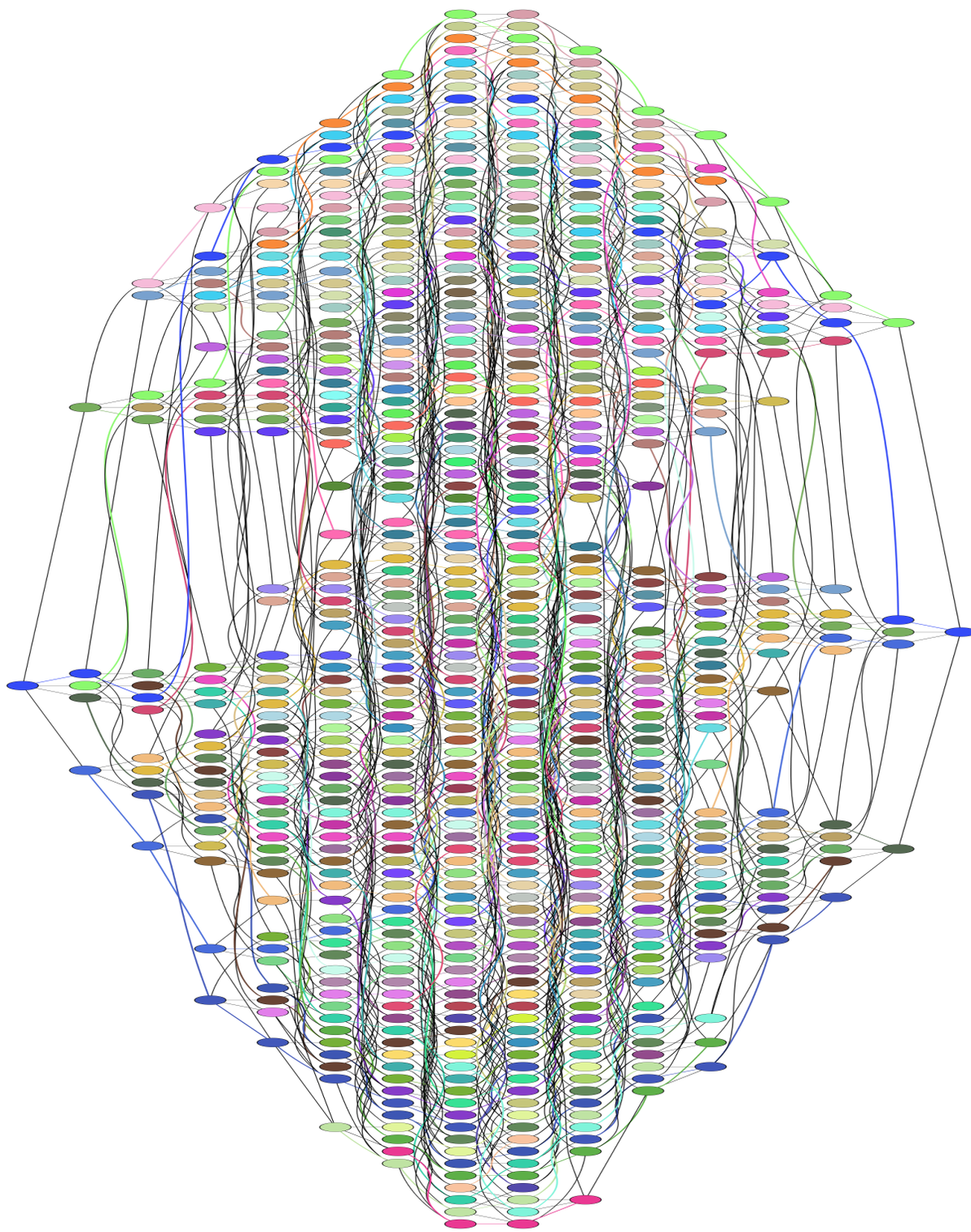
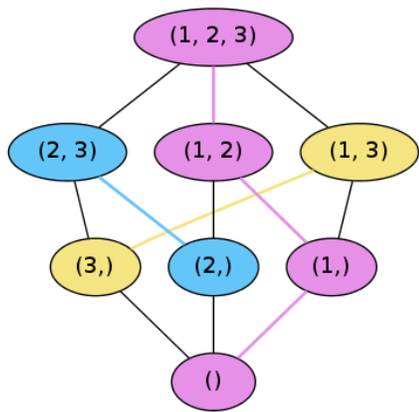
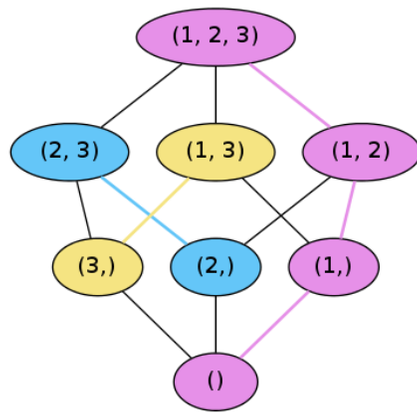


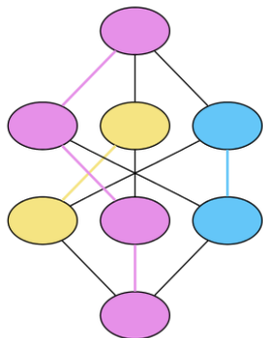
Figure B. 7: I_6 : 1,2,3,4,5,6_SPARSE_NO_LABELS.png



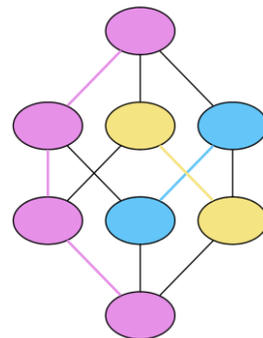
(a) 3_FULL.png



(b) 3_SPARSE.png

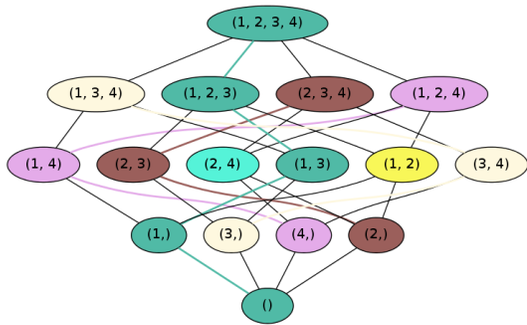


(c) 3_FULL_NO_LABELS.png

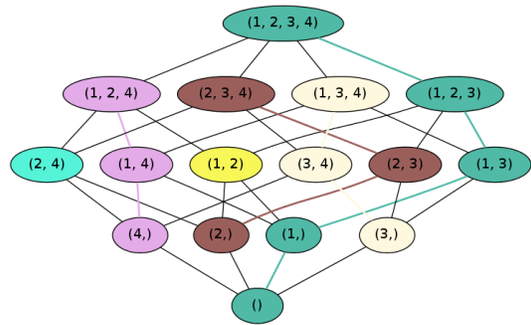


(d) 3_SPARSE_NO_LABELS.png

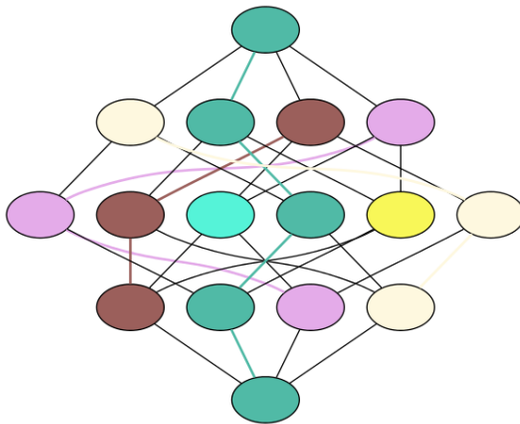
Figure B.8: Boolean algebra, B_3



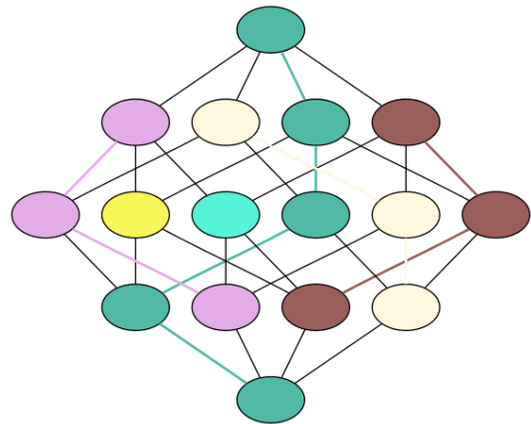
(a) 4_FULL.png



(b) 4_SPARSE.png



(c) 4_FULL_NO_LABELS.png



(d) 4_SPARSE_NO_LABELS.png

Figure B.9: Boolean algebra, B_3

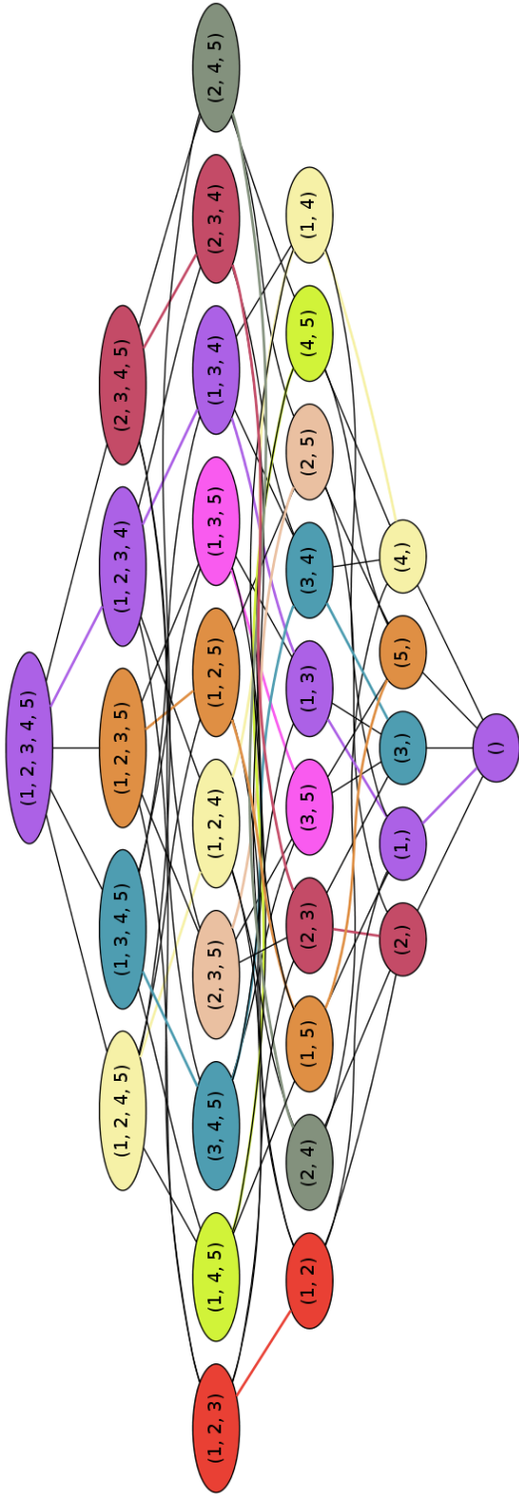


Figure B.10: B_5 : 5_FULLL.png

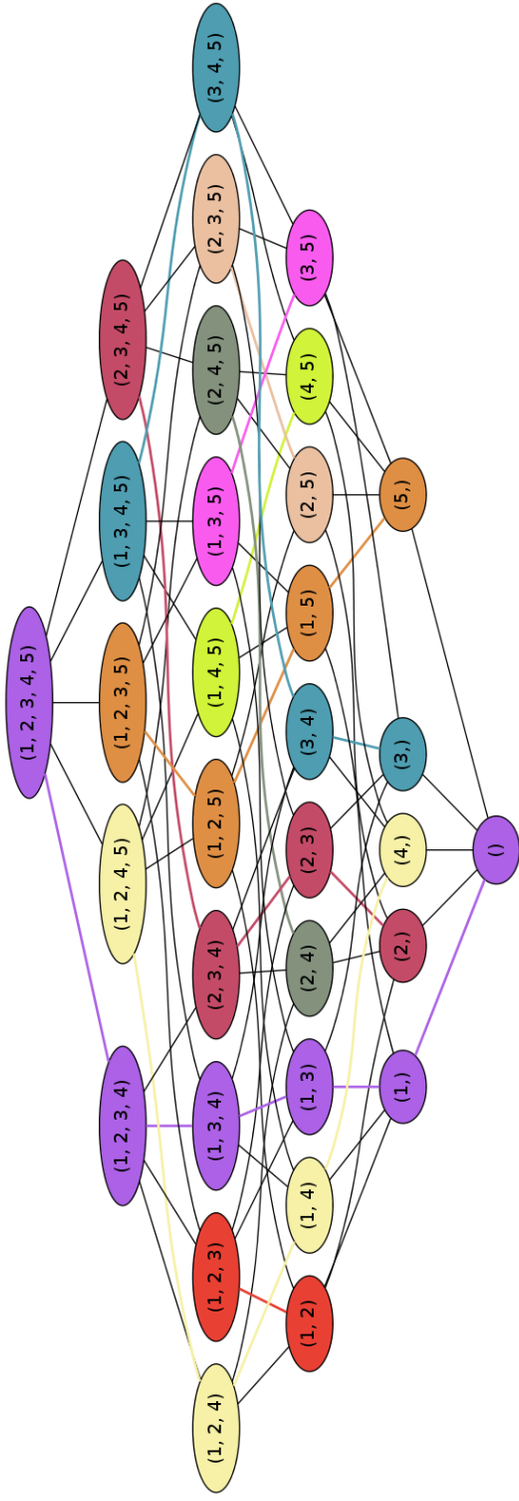
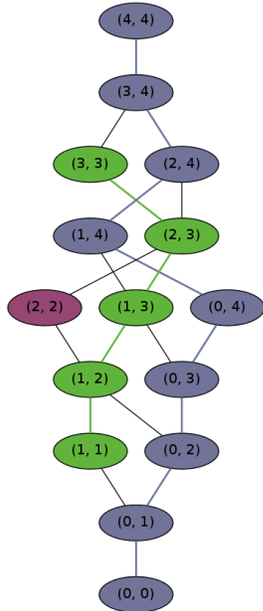
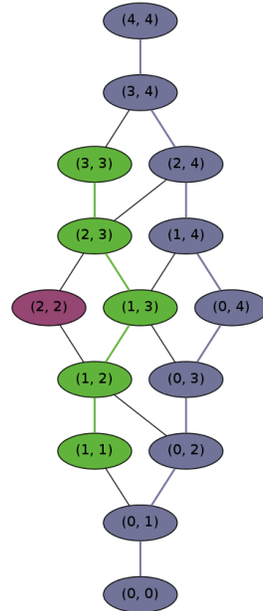


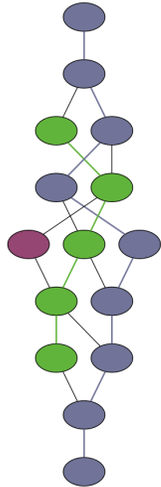
Figure B.11: B_5 : 5-SPARSE.png



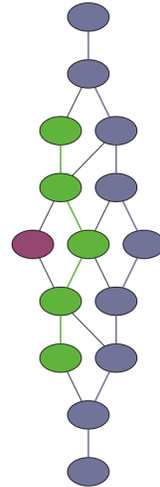
(a) 4,4.FULL.png



(b) 4,4.SPARSE.png



(c) 4,4.FULL_NO_LABELS.png



(d) 4,4.SPARSE_NO_LABELS.png

Figure B.12: Young's lattice, $L(4,2)$

Appendix C

Source codes

The purpose of this section is to provide two implementations, in Python and in C++, of the algorithm given in the text.

C.1 Python

To run the Python implementation we need two external libraries NetworkX and PyGraphviz. The Ford–Fulkerson algorithm is implemented in the NetworkX library and the PyGraphviz library creates figures of the poset and its symmetric saturated chain decomposition.

The first script tries to find a SSCD. If this script finds a SSCD, it saves the decomposition into a text file and calls the second script (`diagram.py`) to generate figures of the poset and its decomposition. The first script accompanied with comments is listed below.

main.py:

```
1 #!/usr/bin/python
2
3 ## list of libraries
4 from array import array
5 import networkx as nx
6 import itertools
```

```

7 import time
8 import collections
9 import sys
10 import math
11 import random
12 import argparse
13 import diagram
14
15 ## global variable for the number of already found paths
16 global gCount
17
18 class Node:
19     ## each node has 4 properties
20     ## name: name of the node
21     ## level: the number of the level
22     ## neighbors: list of neighbors in the next level
23     ## nameOfMatchingNode: name of the matched node
24     def __init__(self, name, level):
25         self.name = name
26         self.level = level
27         self.neighbors = list()
28         self.nameOfMatchingNode = name
29     ## checks if there is possible inversion of its elements
30     def isLast(self):
31         for i in xrange(len(self.name)-1):
32             if self.name[i] < self.name[i+1]:
33                 return False
34         return True
35     ## generate next inversion
36     def generateNext(self, dictOfNextLevel):
37         for i in xrange(len(self.name)-1):
38             if self.name[i] < self.name[i+1]:
39                 tmpName = list(self.name)
40                 tmp = self.name[i]
41                 tmpName[i] = tmpName[i+1]
42                 tmpName[i+1] = tmp
43                 self.neighbors.append(tuple(tmpName))
44                 if ~dictOfNextLevel.has_key(tuple(tmpName)):
45                     dictOfNextLevel[tuple(tmpName)] = Node(tmpName, self.level+1)
46
47     @property
48     def Name(self):
49         return tuple(self.name)
50
51     @property
52     def NameOfMatchingNode(self):
53         return tuple(self.nameOfMatchingNode)
54
55 ## logger for easier handling with output
56 def loggerOut(*message):
57     msg = ""
58     for m in message:
59         msg = ''.join([msg, str(m)])
60     print msg
61     f.write(''.join([msg, '\n']))
62     sys.stdout.flush()
63
64 ## constructing an Inverse poset given a multiset
65 def makeGraphOfInversePartiallyOrderedMultiSet(rootList):
66     listOfLevels = [dict(), dict()]
67     root = Node(rootList, 0)

```

```

66     listOfLevels[0][tuple(root.name)] = root
67     root.generateNext(listOfLevels[1])
68     level = 1
69     while (len(listOfLevels[level]) > 1) or (not listOfLevels[level].values()[0].
70         isLast()):
71         listOfLevels.append(dict())
72         d = listOfLevels[level]
73         for node in d.itervalues():
74             node.generateNext(listOfLevels[level+1])
75         level += 1
76     return listOfLevels
77
78     ## constructing a Young lattice given its size
79     def makeGraphOfYoungsLattice(m,n):
80         listOfLevels = [dict() for x in xrange(m*n+1)]
81         tmp = [0 for i in xrange(n)]
82         root = Node(tmp,0)
83         listOfLevels[0][root.Name] = root
84         for i in xrange(len(listOfLevels)-1):
85             for node in listOfLevels[i].values():
86                 if sum(node.Name) < m*n:
87                     for ii in xrange(n):
88                         if (node.Name[ii] < m) and ( (ii == n-1) or (node.Name[ii] <
89                             node.Name[ii+1]) ):
90                             new = list(node.name)
91                             new[ii] += 1
92                             newNode = Node(new, i+1)
93                             if listOfLevels[i+1].has_key(newNode.Name) == True:
94                                 node.neighbors.append(listOfLevels[i+1][newNode.Name
95                                     ].Name)
96                             else:
97                                 node.neighbors.append(newNode.Name)
98                                 listOfLevels[i+1][newNode.Name] = newNode
99         return listOfLevels
100
101     ## Expanding the graphs for additional layers.
102     def expandGraph(graph, middleLayer, listOfLevels, intActualLayer):
103         if (len(graph) == 1) and (intActualLayer == -1): return
104         graph.clear()
105         for node in listOfLevels[intActualLayer].values():
106             graph.add_edges_from([(node.Name, listOfLevels[intActualLayer+1][neighbor
107                 ].NameOfMatchingNode) for neighbor in node.neighbors], capacity=1.0)
108         for node in listOfLevels[len(listOfLevels)-intActualLayer-2].values():
109             graph.add_edges_from([(node.NameOfMatchingNode, listOfLevels[len(
110                 listOfLevels)-intActualLayer-1][neighbor].Name) for neighbor in node.
111                 neighbors], capacity=1.0)
112
113     ## Adding 'source' and 'sink' nodes for a flow algorithm.
114     def addEndNodes(graph, bottomLevel, upperLevel):
115         graph.add_edges_from([( 'rootDown', node) for node in bottomLevel], capacity
116             =1.0)
117         graph.add_edges_from([(node, 'rootUp') for node in upperLevel], capacity=1.0)
118
119     ## Removing edges without any flow.
120     def removeExtraEdges(graph, flow, middleLayer):
121         graph.remove_nodes_from(('rootUp', 'rootDown'))
122         for node in middleLayer.values():
123             for nodeTmp in graph.successors((node.NameOfMatchingNode, 'u')):
124                 if flow[(node.NameOfMatchingNode, 'u')][nodeTmp] == 0:

```

```

118         graph.remove_edge((node.NameOfMatchingNode, 'u'), nodeTmp)
119     for nodeTmp in graph.predecessors((node.NameOfMatchingNode, 'd')):
120         if flow[nodeTmp][(node.NameOfMatchingNode, 'd')] == 0:
121             graph.remove_edge(nodeTmp, (node.NameOfMatchingNode, 'd'))
122
123     ## Contracting edges with a flow – exanding a list with the SSCD.
124     def contractEdges(graph, bottomLayer, middleLayer, upperLayer):
125         graphNew = nx.DiGraph()
126         for node in middleLayer.values():
127             if len(graph.successors((node.NameOfMatchingNode, 'u'))) > 0:
128                 graph.add_path((graph.predecessors((node.NameOfMatchingNode, 'd'))[0],
129                                 node.NameOfMatchingNode, graph.successors((node.NameOfMatchingNode, 'u'))[0]))
129             else:
130                 graph.add_node(node.NameOfMatchingNode)
131                 graph.remove_nodes_from(((node.NameOfMatchingNode, 'u'), (node.
132                                         NameOfMatchingNode, 'd')))
132         for node in bottomLayer.keys():
133             neigh = graph.neighbors(node)[0]
134             graph.remove_node(node)
135             nodeNew = (node, neigh, graph.neighbors(neigh)[0])
136             graph.remove_nodes_from(nodeNew[1:3])
137             graphNew.add_node(nodeNew)
138             bottomLayer[nodeNew[0]].nameOfMatchingNode = nodeNew
139             upperLayer[nodeNew[2]].nameOfMatchingNode = nodeNew
140         return graphNew, graph.nodes()
141
142     result = None
143
144     ## Finding a path between source and sink using depth-first approach.
145     def find_pathDepthFirst(graph, visitedNodes, source, sink, path):
146         result = None
147         visitedNodes.append(source)
148         if source == sink:
149             return path
150         for successor in graph.successors(source):
151             residual = graph.edge[source][successor]['capacity'] - graph.edge[source][
152                 successor]['flow']
153             if residual > 0 and not successor in visitedNodes:
154                 if not successor in path:
155                     result = find_pathDepthFirst( graph, visitedNodes, successor, sink
156                                                     , path + [successor] )
157                 if result != None:
158                     return result
159         for predecessor in graph.predecessors(source):
160             residual = graph.edge[predecessor][source]['flow']
161             if residual > 0 and not predecessor in visitedNodes:
162                 if not predecessor in path:
163                     result = find_pathDepthFirst( graph, visitedNodes, predecessor,
164                                                     sink, path + [predecessor] )
165                 if result != None:
166                     return result
167
168     ## Finding a path between source and sink using enhanced breadth-first approach.
169     This function process nodes attached to the source in blocks. Each block
170     contain certain percentage of neighbors to source. The breadt-first search
171     then continue with those nodes. If no path to the sink is found, then the
172     function investigate next block of neighbors.
173     def find_pathBreadthFirst(graph, sink, actualLayer, layers, percentage):

```

```

167     if percentage == 0:
168         raise Exception('Percentage_cannot_be_0!')
169     visitedNodes = dict()
170     ## erase visited tokens in graph at the beginning
171     if actualLayer == 0:
172         if not layers.has_key(1):
173             layers[1] = list()
174         for source, path in layers[0]:
175             visitedNodes[source] = True
176             for successor in graph.successors(source):
177                 residual = graph.edge[source][successor]['capacity'] - graph.edge[
178                     source][successor]['flow']
179                 if (residual > 0) and (not successor in path) and (not
180                     visitedNodes.has_key(successor)):
181                     visitedNodes[successor] = True
182                     layers[1].append([successor, path+[successor]])
183             actualLayer += 1
184     ## change from percentage to a number of nodes
185     if (actualLayer == 1) and (percentage <= 1):
186         percentage = math.ceil(len(layers[1]) * percentage)
187
188     while actualLayer > 0:
189         newNodes = False
190         if not layers.has_key(actualLayer+1):
191             layers[actualLayer+1] = list()
192             ## neighbors of the source node
193             if actualLayer == 1:
194                 for i in xrange(int(percentage)):
195                     if len(layers[actualLayer]) == 0:
196                         break
197                     item = layers[actualLayer].pop(random.choice(xrange(len(layers[
198                         actualLayer]))))
199                     source = item[0]
200                     path = item[1]
201                     if source == sink:
202                         return path
203                     for successor in graph.successors(source):
204                         residual = graph.edge[source][successor]['capacity'] - graph.
205                             edge[source][successor]['flow']
206                         if residual > 0 and (not successor in path) and (not
207                             visitedNodes.has_key(successor)):
208                             layers[actualLayer+1].append([successor, path+[successor]])
209                             visitedNodes[successor] = True
210                             newNodes = True
211                     for predecessor in graph.predecessors(source):
212                         residual = graph.edge[predecessor][source]['flow']
213                         if residual > 0 and (not predecessor in path) and (not
214                             visitedNodes.has_key(predecessor)):
215                             layers[actualLayer+1].append([predecessor, path+[
216                                 predecessor]])
217                             visitedNodes[predecessor] = True
218                             newNodes = True
219             ## Investigate the whole path for nodes not attached to source.
220             else:
221                 for source, path in layers[actualLayer]:
222                     if source == sink:
223                         return path
224                     for successor in graph.successors(source):

```

```

218         residual = graph.edge[source][successor]['capacity'] - graph.
219             edge[source][successor]['flow']
220         if residual > 0 and (not successor in path) and (not
221             visitedNodes.has_key(successor)):
222             layers[actualLayer+1].append([successor, path+[successor]])
223             visitedNodes[successor] = True
224             newNodes = True
225         for predecessor in graph.predecessors(source):
226             residual = graph.edge[predecessor][source]['flow']
227             if residual > 0 and (not predecessor in path) and (not
228                 visitedNodes.has_key(predecessor)):
229                 layers[actualLayer+1].append([predecessor, path+[
230                     predecessor]])
231                 visitedNodes[predecessor] = True
232                 newNodes = True
233     if newNodes:
234         actualLayer += 1
235     else:
236         actualLayer -= 1
237
238     return None
239
240 ## Ford-Fulkerson algorithm with adaptive switching between depth and breadth
241 search
242 def myFord_fulkerson(graph, source, sink):
243     for x in graph.edge:
244         for y in graph.edge[x]:
245             graph.edge[x][y]['flow'] = 0
246     path = find_pathDepthFirst(graph, [], source, sink, [source])
247     tmpTime = time.time()
248     tmp = find_pathBreadthFirst(graph, sink, 0, {0:[source,[source]]}, .02)
249     timeBreadth = time.time() - tmpTime
250     value = 0
251     breadth = False
252     while path != None:
253         value += 1
254         for i in xrange(1, len(path)):
255             if graph.edge[path[i-1]].has_key(path[i]):
256                 graph.edge[path[i-1]][path[i]]['flow'] = 1
257             else:
258                 graph.edge[path[i]][path[i-1]]['flow'] = 0
259         if breadth:
260             path = find_pathBreadthFirst(graph, sink, 0, {0:[source,[source
261                 ]]]}, .02)
262         else:
263             start = time.time()
264             path = find_pathDepthFirst(graph, [], source, sink, [source])
265             if time.time() - start > timeBreadth*2:
266                 breadth = True
267                 loggerOut('---_Breadth-search_activated!_---')
268     flow = dict()
269     for x,y in graph.edges():
270         f = graph.edge[x][y].pop('flow')
271         if flow.has_key(x):
272             flow[x][y] = f
273         else:
274             flow[x] = {y: f}
275     return value, flow

```

```

271  ## Extending the middle layer and finding a maximal flow through a graph.
272  def extendAndFlow(graph, middleLayer, bottomLayer):
273      for node in middleLayer.values():
274          neighDown = graph.predecessors(node.NameOfMatchingNode)
275          neighUp = graph.successors(node.NameOfMatchingNode)
276          graph.remove_node(node.NameOfMatchingNode)
277          graph.add_edge((node.NameOfMatchingNode, 'd'), (node.NameOfMatchingNode, 'u'
278                        ), capacity=1.0)
279          graph.add_edges_from([( (node.NameOfMatchingNode, 'u'), x) for x in neighUp
280                               ], capacity=1.0)
281          graph.add_edges_from([(x, (node.NameOfMatchingNode, 'd')) for x in
282                               neighDown], capacity=1.0)
283      value, flow = myFord_fulkerson(graph, 'rootDown', 'rootUp')
284      return value, flow
285
286  ## Main body of the program.
287  def main(f, listOfLevels):
288      gCount = 0
289      start = time.time()
290      numberOfLayers = len(listOfLevels)-1
291      decomposition = list()
292      graph = nx.DiGraph()
293      middleLayer = list()
294      if numberOfLayers%2 == 0:
295          ## there is one layer in the middle
296          loggerOut('—_One_layer_in_the_middle_—')
297          intActualLayer = numberOfLayers/2-1
298          ## Make a graph with three layers.
299          loggerOut('—_Middle_part_start_—')
300          for i in xrange(numberOfLayers/2-1, numberOfLayers/2+1):
301              for item in listOfLevels[i].values():
302                  for iitem in item.neighbors:
303                      graph.add_edge(item.Name, iitem, capacity=1.0)
304          for item in listOfLevels[numberOfLayers/2].values():
305              middleLayer.append(item.Name)
306          loggerOut('—_Middle_part_done_—', (time.time() - start), "seconds.")
307      else:
308          ## There are two middle layers -> find matching, create a graph with 3 layers
309          loggerOut('—_Two_layers_in_the_middle_—')
310          intActualLayer = numberOfLayers/2
311          for item in listOfLevels[intActualLayer].values():
312              for itemNeighbor in item.neighbors:
313                  tmpNode = listOfLevels[intActualLayer+1].get(itemNeighbor)
314                  graph.add_edge(item.Name, tmpNode.Name)
315          tmp = nx.DiGraph()
316          loggerOut('—_matching_start_—', (time.time() - start), "seconds.")
317          matching = nx.max_weight_matching(graph)
318          loggerOut('—_matching_end_—', (time.time() - start), "seconds.")
319          ## Contract the matching into symmetric chains.
320          for item in matching.keys():
321              if listOfLevels[intActualLayer].has_key(item):
322                  newNode = (item, matching[item])
323                  listOfLevels[intActualLayer].get(item).nameOfMatchingNode = newNode
324                  listOfLevels[intActualLayer+1].get(matching[item]).nameOfMatchingNode =
325                      newNode
326                  middleLayer.append(newNode)
327          intActualLayer -= 1
328          tmp.add_nodes_from(middleLayer)
329          graph = tmp

```

```

326  ## Add bottom neighbors for a matching.
327  for item in listOfLevels[intActualLayer].values():
328      for itemNeighbor in item.neighbors:
329          tmpNode = listOfLevels[intActualLayer+1].get(itemNeighbor).NameOfMatchingNode
330          graph.add_edge(item.Name, tmpNode)
331  ## Add upper neighbors for a matching.
332  for item in listOfLevels[numberOfLayers-intActualLayer-1].values():
333      for itemNeighbor in item.neighbors:
334          tmpNode = listOfLevels[numberOfLayers-intActualLayer].get(itemNeighbor).Name
335          graph.add_edge(item.NameOfMatchingNode, tmpNode)
336
337      while (intActualLayer >= 0):
338  loggerOut('—_ActualLayer_is_—', intActualLayer, '/', numberOfLayers, '—', (time.
339          time() - start), "seconds.")
340  addEndNodes(graph, listOfLevels[intActualLayer].keys(), listOfLevels[
341          numberOfLayers-intActualLayer].keys())
342  loggerOut('—_flow_start_—', (time.time() - start), "seconds.")
343  value, flow = extendAndFlow(graph, listOfLevels[intActualLayer+1], listOfLevels[
344          intActualLayer].keys())
345  if value < len(listOfLevels[intActualLayer]):
346      raise Exception('NO_matching')
347  loggerOut('—_flow_end_—', (time.time() - start), "seconds.")
348  removeExtraEdges(graph, flow, listOfLevels[intActualLayer+1])
349  loggerOut('—_removeExtraEdges_end_—', (time.time() - start), "seconds.")
350  graph, disconnectedNodes = contractEdges(graph, listOfLevels[intActualLayer],
351          listOfLevels[intActualLayer+1], listOfLevels[numberOfLayers-intActualLayer])
352  decomposition.extend(disconnectedNodes)
353  intActualLayer -= 1
354  loggerOut('—_expandGraph_start_—', (time.time() - start), "seconds.")
355  expandGraph(graph, middleLayer, listOfLevels, intActualLayer)
356  loggerOut('—_expandGraph_done_—', (time.time() - start), "seconds.")
357  decomposition.extend(graph.nodes())
358  end = time.time()
359  elapsed = end - start
360  loggerOut("Time: _", (time.time() - start), "seconds.")
361  loggerOut('\nDecomposition:')
362  for i in decomposition:
363  if type(i[0]) == tuple:
364      msg = ""
365      for m in i:
366          msg = ''.join([msg, str(m)])
367  else:
368      msg = str(i)
369  f.write(''.join([msg, '\n']))
370  f.close()
371
372  ## Processing user input, running the main function and generating graphs
373  parser = argparse.ArgumentParser(description='Program_for_finding_a_symmetric_
374  saturated_chain_decomposition, _created_as_a_part_of_Master_Thesis_by_Ondrej_
375  Zjevik_at_University_of_Minnesota.', usage='%(prog)s [-h] [--inversion 1 2 3 4
376  ...] [--young m.n]')
377  parser.add_argument('--inversion', type=int, nargs='+', help='element_with_rank_0
378  ', metavar='1 2 3 4 ...', dest='root')
379  parser.add_argument('--young', type=int, nargs=2, help="constants_of_a_Young's_
380  lattice", metavar=('m', 'n'))
381  parser.add_argument('--no_pict', dest='picture', default=True, const=False,
382          action='store_const', metavar='', help="will_not_generate_pictures")
383  args = parser.parse_args()
384  if (args.root == None and args.young == None):

```

```

375     parser.print_help()
376     exit()
377 sys.setrecursionlimit(10000)
378 if(args.root != None):
379     listOfLevels = makeGraphOfInversePartiallyOrderedMultiSet(args.root)
380     f = open(''.join(['', '.join(map(str, args.root)), '.txt']), 'w')
381     main(f, listOfLevels)
382     if args.picture:
383         diagram.makeGraphOfInversePartiallyOrderedMultiSet(f, args.root)
384 else:
385     listOfLevels = makeGraphOfYoungsLattice(args.young[0], args.young[1])
386     f = open(''.join(['', '.join(map(str, [args.young[0] for i in xrange(args.young
387     [1]))]), '.txt']), 'w')
387     main(f, listOfLevels)
388     if args.picture:
389         diagram.makeGraphOfYoungsLattice(f, args.young[0], args.young[1])

```

The second script written in Python generates pictures from an output from the first script or from the script written in C++. The content of the second script follows.

diagram.py:

```

1  #!/usr/bin/python
2
3  from array import array
4  import pygraphviz as pgv
5  from random import randrange
6  color = True;
7
8  class Node:
9      def __init__(self, name, level):
10         self.name = name
11         self.level = level
12         self.neighbors = list()
13         self.nameOfMatchingNode = name
14     def isLast(self):
15         for i in xrange(len(self.name)-1):
16             if self.name[i] < self.name[i+1]:
17                 return False
18         return True
19     def generateNext(self, dictOfNextLevel, listOfNextLevel, graph):
20         graph.add_node(self.Name)
21         for i in xrange(len(self.name)-1):
22             if self.name[i] < self.name[i+1]:
23                 tmpName = list(self.name)
24                 tmp = self.name[i]
25                 tmpName[i] = tmpName[i+1]
26                 tmpName[i+1] = tmp
27                 self.neighbors.append(tuple(tmpName))
28                 graph.add_edge(self.Name, tuple(tmpName), weight=1, style='solid')
29                 if dictOfNextLevel.has_key(tuple(tmpName)):
30                     tmpNode = Node(tmpName, self.level+1)
31                     dictOfNextLevel[tuple(tmpName)] = tmpNode
32                     listOfNextLevel.append(tmpNode)
33     def generateNextBool(self, dictOfNextLevel, listOfNextLevel, graph, size):
34         graph.add_node(self.Name)

```

```

35     for i in xrange(1, size+1):
36         if self.name.count(i) == 0:
37             tmpName = list(self.name)
38             tmpName.append(i)
39             tmpName.sort()
40             self.neighbors.append(tuple(tmpName))
41             graph.add_edge(self.Name, tuple(tmpName), weight=1, style='solid')
42             if dictOfNextLevel.has_key(tuple(tmpName)):
43                 tmpNode = Node(tmpName, self.level+1)
44                 dictOfNextLevel[tuple(tmpName)] = tmpNode
45                 listOfNextLevel.append(tmpNode)
46
47 @property
48 def Name(self):
49     return tuple(self.name)
50
51 @property
52 def NameOfMatchingNode(self):
53     return tuple(self.nameOfMatchingNode)
54
55 def makeBool(graph, size):
56     dictOfLevels = [dict(), dict()]
57     listOfLevels = [list(), list()]
58     root = Node(tuple(), 0)
59     dictOfLevels[0][tuple(root.name)] = root
60     listOfLevels[0].append(root)
61     root.generateNextBool(dictOfLevels[1], listOfLevels[1], graph, size)
62     level = 1
63     while (len(dictOfLevels[level]) > 1):
64         dictOfLevels.append(dict())
65         listOfLevels.append(list())
66         d = dictOfLevels[level]
67         for node in d.itervalues():
68             node.generateNextBool(dictOfLevels[level+1], listOfLevels[level+1],
69                                 graph, size)
70         level += 1
71     sparseGraph = graph.copy()
72     for level in xrange(1, len(listOfLevels)):
73         for node in dictOfLevels[level-1].keys():
74             for node2 in dictOfLevels[level].keys():
75                 if graph.has_edge(node2, node) == False:
76                     graph.add_edge(node, node2, weight=1, style='invis')
77     return sparseGraph
78
79 def makeInversePartiallyOrderedMultiSet(graph, root):
80     dictOfLevels = [dict(), dict()]
81     listOfLevels = [list(), list()]
82     root = Node(root, 0)
83     dictOfLevels[0][tuple(root.name)] = root
84     listOfLevels[0].append(root)
85     root.generateNext(dictOfLevels[1], listOfLevels[1], graph)
86     level = 1
87     while (len(dictOfLevels[level]) > 1) or (not dictOfLevels[level].values()[0].
88         isLast()):
89         dictOfLevels.append(dict())
90         listOfLevels.append(list())
91         d = dictOfLevels[level]
92         for node in d.itervalues():
93             node.generateNext(dictOfLevels[level+1], listOfLevels[level+1], graph)
94         level += 1
95     sparseGraph = graph.copy()

```

```

92     for level in xrange(1, len(listOfLevels)):
93         for node in dictOfLevels[level-1].keys():
94             for node2 in dictOfLevels[level].keys():
95                 if graph.has_edge(node2, node) == False:
96                     graph.add_edge(node, node2, weight=1, style='invis')
97     return sparseGraph
98
99 def makeYoungsLattice(graph, m, n):
100     dictOfLevels = [dict() for x in xrange(m*n+1)]
101     tmp = [0 for i in xrange(n)]
102     root = Node(tmp, 0)
103     dictOfLevels[0][root.Name] = root
104     for i in xrange(len(dictOfLevels)-1):
105         for node in dictOfLevels[i].values():
106             if sum(node.Name) <= m*n:
107                 for ii in xrange(n):
108                     if (node.Name[ii] < m) and ( (ii == n-1) or (node.Name[ii] <
109                                             node.Name[ii+1]) ):
110                         new = list(node.name)
111                         new[ii] += 1
112                         newNode = Node(new, i+1)
113                         graph.add_edge(newNode.Name, node.Name, weight=1, style='
114                                     solid')
115                         if dictOfLevels[i+1].has_key(newNode.Name) == True:
116                             node.neighbors.append(dictOfLevels[i+1][newNode.Name
117                                     ].Name)
118                         else:
119                             node.neighbors.append(newNode.Name)
120                             dictOfLevels[i+1][newNode.Name] = newNode
121     sparseGraph = graph.copy()
122     for level in xrange(1, m*n):
123         for node in dictOfLevels[level-1].keys():
124             for node2 in dictOfLevels[level].keys():
125                 if graph.has_edge(node2, node) == False:
126                     graph.add_edge(node2, node, weight=1, style='invis')
127     return sparseGraph
128
129 def makeFromFile(graph, fileName):
130     listOfLevels = []
131     f = open(fileName+"_levels", 'r')
132     level = -1;
133     for line in f:
134         listOfLevels.append(list())
135         level += 1;
136         line.strip()
137     for node in line.split(','):
138         listOfLevels[level].append(int(node))
139
140     f = open(fileName+"_nodes", 'r')
141     nodeId = -1;
142     for line in f:
143         nodeId += 1;
144         if nodeId > level: continue
145         line.strip()
146     for neighbor in line.split(','):
147         graph.add_edge((int(nodeId),), (int(neighbor),), weight=1, style='solid')

```

```

148         for node in listOfLevels[level-1]:
149             for node2 in listOfLevels[level]:
150                 if graph.has_edge((node2,),(node,)) == False:
151                     graph.add_edge((node,),(node2,),weight=1,style='invis')
152     return sparseGraph
153
154 def makeGraphOfBool(oldFile, size):
155     graph = pgv.AGraph(rankdir='BT')
156     sparseGraph = makeBool(graph, size)
157
158     decomp = False;
159     i = 0
160     d = dict()
161     f = open(oldFile.name, 'r')
162     generatePictures(f, graph, sparseGraph)
163
164 def makeGraphOfInversePartiallyOrderedMultiSet(oldFile, root):
165     graph = pgv.AGraph(rankdir='BT')
166     sparseGraph = makeInversePartiallyOrderedMultiSet(graph, root)
167
168     decomp = False;
169     i = 0
170     d = dict()
171     f = open(oldFile.name, 'r')
172     generatePictures(f, graph, sparseGraph)
173
174 def makeGraphOfYoungsLattice(oldFile, m, n):
175     f = open(oldFile.name, 'r')
176     graph = pgv.AGraph()
177     numberOfLayers = m*n
178     sparseGraph = makeYoungsLattice(graph, m, n)
179     generatePictures(f, graph, sparseGraph)
180
181 def makeGraphFromFile(decomposition, fileName):
182     graph = pgv.AGraph(rankdir='BT')
183     sparseGraph = makeFromFile(graph, fileName)
184     f = open(decomposition.name, 'r')
185     generatePictures(f, graph, sparseGraph)
186
187 def generatePictures(f, graph, sparseGraph):
188     decomp = False;
189     i = 0
190     d = dict()
191     for line in f:
192         i += 1
193         line.strip()
194
195     if decomp:
196         line = str.replace(line, '(', '(')
197         line = str.replace(line, ')', ')')
198         line = str.replace(line, ')', ')')
199         line = str.replace(line, ')', ')')
200         line = str.replace(line, '\n', '\n')
201         if str.count(line, ',') == 0:
202             line = str.replace(line, ',,,-')
203             lineList = list()
204             for part in line.split(',') [1:]:
205                 if len(part) > 0:
206                     lineList.append(tuple(map(int, part.split(','))))

```

```

207     else:
208         lineList.append(tuple())
209         c = "#%s" % "".join([hex(randrange(50, 255))[2:] for i in range(3)])
210         if(color): graph.node_attr['style'] = 'filled'
211         graph.get_node(lineList[0]).attr['fillcolor'] = c
212         if(color): sparseGraph.node_attr['style'] = 'filled'
213         sparseGraph.get_node(lineList[0]).attr['fillcolor'] = c
214         for ii in xrange(1, len(lineList)):
215             graph.get_node(lineList[ii]).attr['fillcolor'] = c
216         if (color):
217             graph.add_edge(lineList[ii-1], lineList[ii], weight=1, color=c, penwidth=2)
218         else:
219             graph.add_edge(lineList[ii-1], lineList[ii], weight=1)
220             sparseGraph.get_node(lineList[ii]).attr['fillcolor'] = c
221         if(color):
222             sparseGraph.add_edge(lineList[ii-1], lineList[ii], weight=1, color=c,
223                                 penwidth=2)
224         else:
225             sparseGraph.add_edge(lineList[ii-1], lineList[ii], weight=1)
226     if (line == 'Decomposition:\n'): decomp = True
227
228     graphNoLabels = graph.copy()
229     graphNoLabels.node_attr['label'] = '_'
230     sparseGraphNoLabels = sparseGraph.copy()
231     sparseGraphNoLabels.node_attr['label'] = '_'
232     graph.layout(prog='dot')
233     graph.draw(str.replace(f.name, '.txt', '')+"_FULL.png")
234     sparseGraph.layout(prog='dot')
235     sparseGraph.draw(str.replace(f.name, '.txt', '')+"_SPARSE.png")
236     graphNoLabels.layout(prog='dot')
237     graphNoLabels.draw(str.replace(f.name, '.txt', '')+"_FULL_NO_LABELS.png")
238     sparseGraphNoLabels.layout(prog='dot')
239     sparseGraphNoLabels.draw(str.replace(f.name, '.txt', '')+"_SPARSE_NO_LABELS.png")

```

C.2 C++

We present the source code for the C++ language in this section. The program contains two classes, *main* and *node*. Each instance of the node class represents one vertex of the poset and each instance stores an id, vertex label, level index and a list of its neighbors. The classes are given below.

node.h:

```

1
2 #ifndef NODE_H
3 #define NODE_H
4

```

```

5 #include <vector>
6 #include <string>
7 #include <map>
8 #include <fstream>
9 #include <boost/serialization/vector.hpp>
10 #include <boost/serialization/set.hpp>
11 #include <boost/serialization/array.hpp>
12 #include <boost/archive/text_oarchive.hpp>
13 #include <boost/archive/text_iarchive.hpp>
14 #include <boost/shared_ptr.hpp>
15 #include <boost/make_shared.hpp>
16
17 using namespace std;
18
19 typedef unsigned int IdType;
20 typedef unsigned char NameElementType;
21
22 class Node
23 {
24 public:
25     unsigned char level; // Good for up to 255 levels
26     IdType id; // Can store up to 12! elements
27     IdType boolSize; // store the number of elements in a set
28
29     vector<NameElementType> name;
30     vector<IdType> neighbors;
31
32
33     Node();
34     Node(std::vector<NameElementType> name, int level, IdType id);
35     ~Node();
36     void generateNextBool( map<vector<NameElementType>, boost::shared_ptr<Node> >*>
37         tmp_level, vector<vector<IdType>*> adjacentMatrix, vector<IdType>*>
38         nextLevel, std::ofstream* fileWithNodeNames);
39     void generateNextInverse( map<vector<NameElementType>, boost::shared_ptr<Node> >*>
40         tmp_level, vector<vector<IdType>*> adjacentMatrix, vector<IdType>*>
41         nextLevel, std::ofstream* fileWithNodeNames);
42     void generateNextYoung( map<vector<NameElementType>, boost::shared_ptr<Node> >*>
43         tmp_level, vector<vector<IdType>*> adjacentMatrix, vector<IdType>*>
44         nextLevel, std::ofstream* fileWithNodeNames, int m);
45     string getName();
46     string getNameYoung();
47 };
48
49 #endif // NODE.H

```

node.cpp:

```
1 // libraries :
2 #include "node.h"
3 #include <vector>
4 #include <string>
5 #include <boost/concept_check.hpp>
6 #include <boost/lexical_cast.hpp>
7 using namespace std;
8
9 Node::Node(vector<NameElementType> name, int level, IdType id){
10     this->name = name;
11     this->level = level;
12     this->id = id;
13 }
14 // Default constructors :
15 Node::Node(){
16 }
17
18 Node::~Node(){
19 }
20
21 // generate next vertices in the boolean lattice
22 void Node::generateNextBool( map<vector<NameElementType>, boost::shared_ptr<Node>
23     >* tmp_level, vector<vector<IdType>*> * adjacentMatrix, vector<IdType>*
24     nextLevel, std::ofstream* fileWithNodeNames)
25 {
26     for (NameElementType i = 1; i <= this->boolSize; i++)
27     {
28         if (name.size() == 0 || find(name.begin(), name.end(), i) == name.end() )
29         {
30             vector<NameElementType> tmp;
31             for(int j = 0; j < name.size(); j++) tmp.push_back(name[j]);
32             tmp.push_back(i);
33             sort(tmp.begin(), tmp.end());
34             if(tmp_level->count(tmp) == 0){
35                 boost::shared_ptr<Node> newNode = boost::make_shared<Node>(tmp, level+1,
36                     adjacentMatrix->size());
37                 newNode->boolSize = this->boolSize;
38                 string name = newNode->getName();
39                 int stringSize = (boolSize <= 9 ? 2+boolSize*2 : 2+9*2+(boolSize-9)*3);
40                 (*fileWithNodeNames) << name;
41                 for(int k = 0; k < stringSize-name.length(); k+)(*fileWithNodeNames) <<
42                     "_";
43                 (*fileWithNodeNames) << endl;
44                 fileWithNodeNames->flush();
45                 adjacentMatrix->at(id)->push_back(adjacentMatrix->size());
46                 nextLevel->push_back(adjacentMatrix->size());
47                 adjacentMatrix->push_back(new vector<IdType>());
48                 (*tmp_level)[tmp] = newNode;
49             } else{
50                 adjacentMatrix->at(id)->push_back(tmp_level->at(tmp)->id);
51             }
52         }
53     }
54 }
55 // generate next vertices using only inversion on an instantiation
```

```

52 void Node::generateNextInverse( map<vector<NameElementType>, boost::shared_ptr<
    Node> >* tmp_level, vector<vector<IdType>*>* adjacentMatrix, vector<IdType>*>*
    nextLevel, std::ofstream* fileWithNodeNames)
53 {
54     for (unsigned int i = 0; i < name.size()-1; i++)
55     {
56         if (name.at(i) < name.at(i+1))
57         {
58             vector<NameElementType> tmp;
59             for(int j = 0; j < name.size(); j++) tmp.push_back(name[j]);
60             NameElementType t = tmp.at(i);
61             tmp.at(i) = tmp.at(i+1);
62             tmp.at(i+1) = t;
63             if(tmp_level->count(tmp) == 0){
64                 boost::shared_ptr<Node> newNode = boost::make_shared<Node>(tmp, level+1,
                    adjacentMatrix->size());
65                 string name = newNode->getName();
66                 (*fileWithNodeNames) << name << endl;
67                 fileWithNodeNames->flush();
68                 adjacentMatrix->at(id)->push_back(adjacentMatrix->size());
69                 nextLevel->push_back(adjacentMatrix->size());
70                 adjacentMatrix->push_back(new vector<IdType>());
71                 (*tmp_level)[tmp] = newNode;
72             } else{
73                 adjacentMatrix->at(id)->push_back(tmp_level->at(tmp)->id);
74             }
75         }
76     }
77 }
78 // generate next vertices in a Young's lattice
79 void Node::generateNextYoung( map<vector<NameElementType>, boost::shared_ptr<Node
    > >* tmp_level, vector<vector<IdType>*>* adjacentMatrix, vector<IdType>*>*
    nextLevel, std::ofstream* fileWithNodeNames, int m)
80 {
81     for (int i = 0; i < name.size(); i++)
82     {
83         if (name.at(i) < m && (i == 0 || name[i-1] > name[i]) )
84         {
85             vector<NameElementType> tmp;
86             for(int j = 0; j < name.size(); j++) tmp.push_back(name[j]);
87             tmp.at(i)++;
88             if(tmp_level->count(tmp) == 0){
89                 boost::shared_ptr<Node> newNode = boost::make_shared<Node>(tmp, level+1,
                    adjacentMatrix->size());
90                 string name = newNode->getNameYoung();
91                 (*fileWithNodeNames) << name << endl;
92                 fileWithNodeNames->flush();
93                 adjacentMatrix->at(id)->push_back(adjacentMatrix->size());
94                 nextLevel->push_back(adjacentMatrix->size());
95                 adjacentMatrix->push_back(new vector<IdType>());
96                 (*tmp_level)[tmp] = newNode;
97             } else{
98                 adjacentMatrix->at(id)->push_back(tmp_level->at(tmp)->id);
99             }
100         }
101     }
102 }
103 // return name as a string
104 string Node::getName()

```

```

105 {
106     string ret = "(";
107     for(int i = 0; i < (int)name.size()-1; i++){
108         ret += boost::lexical_cast<string>((int)name.at(i));
109         ret += ",";
110     }
111     if(name.size() > 0) ret += boost::lexical_cast<string>((int)name.at(name.size()
112         -1));
113     ret += ")";
114     return ret;
115 }
116 // return name as a string
117 string Node::getNameYoung()
118 {
119     string ret = "(";
120     for(unsigned int i = name.size()-1; i > 0; i--){
121         if((int)name.at(i) < 10) ret += "0";
122         ret += boost::lexical_cast<string>((int)name.at(i));
123         ret += ",";
124     }
125     if((int)name.at(0) < 10) ret += "0";
126     ret += boost::lexical_cast<string>((int)name.at(0));
127     ret += ")";
128     return ret;
129 }

```

main.cpp:

```
1 //libraries:
2 #include "algorithm"
3 #include <iostream>
4 #include <fstream>
5 #include <limits>
6 #include <time.h>
7 #include <iterator>
8 #include <vector>
9 #include <map>
10 #include <boost/program_options.hpp>
11 #include <boost/program_options/options_description.hpp>
12 #include <boost/algorithm/string.hpp>
13 #include <boost/graph/adjacency_list.hpp>
14 #include <boost/graph/graphviz.hpp>
15 #include <boost/property_map/property_map.hpp>
16 #include <boost/graph/boykov_kolmogorov_max_flow.hpp>
17 #include <boost/graph/edmonds_karp_max_flow.hpp>
18 #include <boost/graph/push_relabel_max_flow.hpp>
19 #include <boost/graph/properties.hpp>
20 #include <boost/graph/max_cardinality_matching.hpp>
21
22 #include <boost/serialization/map.hpp>
23 #include <boost/serialization/list.hpp>
24 #include <boost/serialization/vector.hpp>
25 #include <boost/serialization/shared_ptr.hpp>
26 #include <boost/archive/text_oarchive.hpp>
27 #include <boost/archive/text_iarchive.hpp>
28 #include <boost/archive/binary_oarchive.hpp>
29 #include <boost/archive/binary_iarchive.hpp>
30 #include "boost/graph/graph_traits.hpp"
31
32 #include "node.h"
33 #include <Python.h>
34
35 using std::vector;
36 using namespace boost;
37 namespace po = boost::program_options;
38
39 typedef adjacency_list_traits < vecS, vecS, directedS > Traits;
40
41 // vertex structure
42 struct Vertex{
43     IdType id;
44     unsigned char level;
45     long distance;
46     default_color_type color;
47     Traits::edge_descriptor predecessor;
48 };
49 // edge structure
50 struct Edge{
51     long capacity;
52     long residual_capacity;
53     Traits::edge_descriptor reverse;
54 };
55
56 typedef adjacency_list < vecS, vecS, directedS, Vertex, Edge> Graph;
57
```

```

58 typedef Graph::vertex_descriptor NodeId;
59 typedef Graph::edge_descriptor EdgeId;
60 Graph graph;
61
62 vector<vector<IdType>* > listOfLevels;
63 time_t start_time;
64 // contains ids of neighbors of a node
65 vector<vector<IdType>* > arrayOfEdges;
66
67 void removeUnusedEdges(Graph *graph, property_map<Graph, long Edge::*>::type*
    m_e_c, property_map<Graph, long Edge::*>::type* m_e_r_c, int bottom_layer,
    int upper_layer);
68 // output file
69 static std::ofstream* output;
70 // logger
71 static void loggerOut(string message){
72     cout << message << "\n";
73     *output << message << "\n";
74 }
75 unsigned int charInLine = 0;
76 // helper class
77 std::istream& GotoLine(std::istream& file, unsigned int num){
78     if(charInLine == 0){
79         file.seekg(std::ios::beg);
80         string line;
81         getline(file, line);
82         charInLine = file.tellg();
83     }
84     file.seekg(charInLine*num);
85     return file;
86 }
87 // add an edge between two vertices and set its parameters
88 EdgeId* AddEdge(Graph::vertex_descriptor &v1, Graph::vertex_descriptor &v2, const
    int capacity, Graph* g)
89 {
90     EdgeId e1 = add_edge(v1, v2, *g).first;
91     EdgeId e2 = add_edge(v2, v1, *g).first;
92     g->operator [] (e1).capacity = 1;
93     g->operator [] (e1).reverse = e2;
94     g->operator [] (e2).capacity = 0;
95     g->operator [] (e2).reverse = e1;
96     g->operator [] (v2).predecessor = e2;
97     return &e1;
98 }
99 // Uses Push-Relabel flow algorithm to find a matching between middle layers.
100 void findMatching(Graph* graph, int layer)
101 {
102     loggerOut("—_Middle_part_start_—");
103     Graph g = *graph;
104     int l = listOfLevels.size()-1 - layer;
105     NodeId sink = add_vertex(*graph);
106     graph->operator [] (sink).id = -1;
107     for (vector<IdType>::const_iterator it = listOfLevels[l]->begin(), e =
        listOfLevels[l]->end(); it != e; ++it)
108     {
109         //add the upper layer
110         NodeId n = add_vertex(*graph); //(*it) - (*listOfLevels[l]->begin()) + 1;
        //id of a node in the graph
111         graph->operator [] (n).id = *it;

```

```

112     AddEdge(n, sink, l, graph);
113 }
114 l--;
115
116 NodeId src = add_vertex(*graph);
117 for (vector<IdType>::const_iterator it = listOfLevels[l]->begin(), e =
118     listOfLevels[l]->end(); it != e; ++it)
119 {
120     //add bottom layer and connect nodes with its neighbors
121     add_vertex(*graph);
122     NodeId n = (*it) - (*listOfLevels[l]->begin()) + 2 + listOfLevels[l+1]->size
123     (); //id of a node in the graph
124     graph->operator [] (n).id = *it;
125     AddEdge(src, n, l, graph); //connect with source
126
127     for (vector<IdType>::const_iterator neighbour = arrayOfEdges[*it]->begin(), ee
128         = arrayOfEdges[*it]->end(); neighbour != ee; ++neighbour)
129     {
130         NodeId nn = (*neighbour) - (*listOfLevels[l+1]->begin()) + 1;
131         AddEdge(n, nn, l, graph);
132     }
133 }
134 property_map<Graph, long Edge::*>::type map_edge_capacity (get(&Edge::capacity, *
135 graph) );
136 property_map<Graph, long Edge::*>::type map_edge_residual_capacity (get(&Edge::
137 residual_capacity, *graph) );
138 property_map<Graph, Graph::edge_descriptor Edge::*>::type map_edge_reverse (get
139 (&Edge::reverse, *graph) );
140 property_map<Graph, vertex_index_t >::type map_vertex_index = get(
141 vertex_index, *graph);
142
143 long flow = 0;
144 time_t time_start, time_end;
145 double time_push;
146 time(&time_start);
147 flow = push_relabel_max_flow(*graph, src, sink, map_edge_capacity,
148     map_edge_residual_capacity, map_edge_reverse, map_vertex_index);
149 time(&time_end);
150 time_push = difftime(time_end, time_start);
151 time(&time_start);
152
153 loggerOut("—_Middle_part_done_—"+lexical_cast<string>(time_push));
154 removeUnusedEdges(graph, &map_edge_capacity, &map_edge_residual_capacity, l, l+1)
155 ;
156 }
157 // removes edges without any flow
158 void removeUnusedEdges(Graph *graph, property_map<Graph, long Edge::*>::type*
159 m_e_c, property_map<Graph, long Edge::*>::type* m_e_r_c, int bottom_layer,
160     int top_layer){
161     property_map<Graph, long Edge::*>::type map_edge_capacity = *m_e_c;
162     property_map<Graph, long Edge::*>::type map_edge_residual_capacity = *m_e_r_c;
163     graph_traits<Graph>::vertex_iterator u_iter, u_end, u_it;
164     graph_traits<Graph>::edge_iterator ei, e_end;
165     int id_bottom_first = listOfLevels[bottom_layer]->at(0), id_bottom_last =
166     listOfLevels[bottom_layer]->at(listOfLevels[bottom_layer]->size()-1);
167     int id_top_first = listOfLevels[top_layer]->at(0), id_top_last = listOfLevels[
168     top_layer]->at(listOfLevels[top_layer]->size()-1);

```

```

157     int id_before_top_first = listOfLevels[top_layer-1]->at(0), id_before_top_last
158         = listOfLevels[top_layer-1]->at(listOfLevels[top_layer-1]->size()-1);
159
160     for(int i = id_bottom_first; i <= id_bottom_last; i++) arrayOfEdges[i]->clear()
161         ; // erase all links for bottom layer
162     for(int i = id_before_top_first; i <= id_before_top_last; i++) arrayOfEdges[i
163         ]->clear(); // erase all links to upper layer
164
165     for(boost::tie(ei, e_end) = edges(*graph); ei != e_end; ei++){
166         if ( map_edge_capacity[*ei] == 1 && (map_edge_capacity[*ei] -
167             map_edge_residual_capacity[*ei]) == 1 ){
168             //keep the edge - save the connected node into arrayOfEdges as the only
169                 neighbor
170             IdType id_source = graph->operator [] (source(*ei, *graph)).id;
171             IdType id_target = graph->operator [] (target(*ei, *graph)).id;
172             if(id_source >= id_bottom_first && id_source <= id_bottom_last && id_target
173                 != -1 && id_target != id_source){
174                 //edge goes from bottom layer
175                 arrayOfEdges[id_source]->push_back(id_target);
176             } else
177             if(id_target >= id_top_first && id_target <= id_top_last && id_target != -1
178                 && id_target != id_source){
179                 //edge goes to the top layer
180                 arrayOfEdges[id_source]->push_back(id_target);
181             }
182         }
183     }
184 }
185
186 // extends the middle layer and run the flow algorithm on the constructed graph
187 int extendAndFlow(Graph* graph, int layer) //layer is the # of actual layer and
188     it's going to zero
189 {
190     loggerOut("_flow_start_");
191     int upperLayer = listOfLevels.size()-1 - layer;
192     NodeId sink = add_vertex(*graph);
193     graph->operator [] (sink).id = -1;
194     NodeId src = add_vertex(*graph); //source node
195     graph->operator [] (src).id = -1;
196
197     //upper layer
198     for (vector<IdType>::const_iterator it = listOfLevels[upperLayer]->begin(), e =
199         listOfLevels[upperLayer]->end(); it != e; ++it)
200     {
201         NodeId n = add_vertex(*graph);
202         graph->operator [] (n).id = *it;
203         AddEdge(n, sink, 1, graph);
204     }
205
206     //middle layer
207     upperLayer--;
208     for (vector<IdType>::const_iterator it = listOfLevels[upperLayer]->begin(), e =
209         listOfLevels[upperLayer]->end(); it != e; ++it)
210     {
211         NodeId n = add_vertex(*graph);
212         graph->operator [] (n).id = *it;
213
214         NodeId nn = add_vertex(*graph);
215         graph->operator [] (nn).id = *it;
216         AddEdge(n, nn, 1, graph); //edge in the middle

```

```

206     for (vector<IdType>::const_iterator neighbour = arrayOfEdges[*it]->begin(), ee
207           = arrayOfEdges[*it]->end(); neighbour != ee; ++neighbour)
208     {
209         NodeId tmp = vertex((*neighbour) - (*listOfLevels[upperLayer+1]->begin()) +
210                             2,*graph);
211         AddEdge(nn,tmp,1,graph);
212     }
213 //bottom layer
214 for (vector<IdType>::const_iterator it = listOfLevels[layer]->begin(), e =
215       listOfLevels[layer]->end(); it != e; ++it)
216 {
217     NodeId n = add_vertex(*graph);
218     graph->operator [] (n).id = *it;
219     AddEdge(src,n,1,graph);
220
221     for (vector<IdType>::const_iterator neighbor = arrayOfEdges[*it]->begin(), ee
222           = arrayOfEdges[*it]->end(); neighbor != ee; ++neighbor){
223         NodeId nn = *neighbor;
224
225         for (int i = layer; i < upperLayer-1; i++) nn = arrayOfEdges[nn]->at(0);
226         nn = vertex(2*(nn - listOfLevels[upperLayer]->at(0))+2+listOfLevels[
227             upperLayer+1]->size(),*graph);
228         AddEdge(n,nn,1,graph);
229         graph->operator [] (nn).id = *neighbor;
230     }
231 }
232
233 property_map<Graph, long Edge::*>::type map_edge_capacity (get(&Edge::capacity,*
234 graph) );
235 property_map<Graph, long Edge::*>::type map_edge_residual_capacity (get(&Edge::
236 residual_capacity,*graph) );
237 property_map<Graph, Graph::edge_descriptor Edge::*>::type map_edge_reverse (get
238 (&Edge::reverse,*graph) );
239 property_map<Graph, vertex_index_t >::type map_vertex_index = get(
240 vertex_index,*graph);
241
242 long flow = 0;
243 time_t time_start, time_end;
244 double time_push;
245 time(&time_start);
246 flow = push_relabel_max_flow(*graph,src,sink,map_edge_capacity,
247 map_edge_residual_capacity,map_edge_reverse,map_vertex_index);
248 time(&time_end);
249 time_push = difftime(time_end,time_start);
250 time(&time_start);
251
252 loggerOut("—_flow_end_—" +lexical_cast<string>(time_push));
253
254 removeUnusedEdges(graph,&map_edge_capacity,&map_edge_residual_capacity,layer,
255 upperLayer+1);
256 return flow;
257 }
258 // the main class, which tries to find a Symmetric Saturated Chain Decomposition
259 of poset saved in listOfLevels and in arrayOfEdges
260 void findSSCD(Graph *g, boost::program_options::variables_map vm){
261     Graph graph = *g;

```

```

253 int layer;
254 string name = "nodes.dat";
255
256 if (listOfLevels.size() % 2 == 0)
257 {
258     //Two layers in the middle
259     loggerOut("—Two_layers_in_the_middle—");
260     //layer is the number of the actual layer; layer goes to zero
261     layer = listOfLevels.size()/2 - 1;
262     graph.clear();
263     findMatching(&graph, layer--);
264
265 }
266 else
267 {
268     //One layer in the middle
269     loggerOut("—One_layer_in_the_middle—");
270     layer = listOfLevels.size()/2 - 1;
271     graph.clear();
272     extendAndFlow(&graph, layer--);
273 }
274 while(layer >= 0){
275     graph.clear();
276     loggerOut(" Actual_layer_is_" + lexical_cast<string>(layer) + "/" + lexical_cast<
277         string>(listOfLevels.size()));
278     if (extendAndFlow(&graph, layer--) != listOfLevels[layer+1]->size()){
279         loggerOut("—Possibly_no_symmetric_chain_decomposition!_Terminating...--");
280         output->flush();
281         exit(EXIT_FAILURE);
282     }
283 }
284
285 //Print founded symmetric chains decomposition
286 std::ifstream fileWithNodeNameRead(name.c_str());
287 std::istream& nodeName = fileWithNodeNameRead;
288 name = "decomposition.dat.tmp";
289 std::ofstream fileWithTmpDecomposition(name.c_str(), ios::trunc);
290 string line;
291
292 time_t tmp;
293 time(&tmp);
294 loggerOut("—Total_time:_"+lexical_cast<string>(tmp-start_time)+"—");
295
296 loggerOut("Decomposition:");
297 if(listOfLevels.size() % 2 == 1){
298     for (vector<IdType>::const_iterator it = listOfLevels[listOfLevels.size()
299         /2]->begin(), e = listOfLevels[listOfLevels.size()/2]->end(); it != e; ++
300         it){
301         if(arrayOfEdges[*it]->size() == 0){
302             GotoLine(nodeNames, *it);
303             getline(nodeNames, line);
304             trim(line);
305             (vm.count("file")) ? (fileWithTmpDecomposition << "(" << *it << ")" <<
306                 endl) : (fileWithTmpDecomposition << line << endl);
307         }
308     }
309 }
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308     for (vector<IdType>::const_iterator it = listOfLevels [i]->begin(), e =
309           listOfLevels [i]->end(); it != e; ++it){
310         IdType tmp = *it, tmp_new;
311         if (arrayOfEdges [tmp]->size () > 0){
312             fileWithTmpDecomposition << endl;
313         }
314         bool print = false;
315         if (arrayOfEdges [tmp]->size () > 0){
316             do
317             {
318                 print = true;
319                 GotoLine (nodeNames, tmp);
320                 getline (nodeNames, line);
321                 trim (line);
322                 (vm.count ("file")) ? (fileWithTmpDecomposition << "(" << tmp << ")") :
323                     (fileWithTmpDecomposition << line);
324                 tmp_new = arrayOfEdges [tmp]->at (0);
325                 arrayOfEdges [tmp]->clear ();
326                 tmp = tmp_new;
327             } while (arrayOfEdges [tmp]->size () > 0);
328             if (print && tmp != arrayOfEdges .size ()-1){
329                 GotoLine (nodeNames, tmp);
330                 getline (nodeNames, line);
331                 trim (line);
332                 (vm.count ("file")) ? (fileWithTmpDecomposition << "(" << tmp << ")") :
333                     (fileWithTmpDecomposition << line);
334             }
335         }
336     }
337     fileWithTmpDecomposition .close ();
338
339     for (int i = 0; i < arrayOfEdges .size (); i++) delete arrayOfEdges [i];
340     for (int i = 0; i < listOfLevels .size (); i++) delete listOfLevels [i];
341
342     //erase empty lines printed to decomposition .dat .tmp
343     name = "decomposition .dat .tmp";
344     std :: ifstream fileWithTmpDecompositionRead (name .c_str ());
345     std :: istream & decomposition = fileWithTmpDecompositionRead;
346     name = "decomposition .dat";
347     std :: ofstream fileWithDecomposition (name .c_str (), ios :: trunc);
348     while (getline (decomposition, line)){
349         if (line .size () > 0){
350             fileWithDecomposition << line << endl;
351             loggerOut (line);
352         }
353     }
354     fileWithTmpDecompositionRead .close ();
355     remove ("decomposition .dat .tmp");
356
357     fileWithNodeNameRead .close ();
358     fileWithDecomposition .close ();
359     output ->flush ();
360     output ->close ();
361     delete output;
362     remove ("nodes .dat");
363     remove ("decomposition .dat");

```

```

364 }
365 // helper to load the Inverse poset into the memory
366 void makeBoolPoset(Graph *g, boost::program_options::variables_map vm){
367     Graph graph = *g;
368     int layer;
369     string name = "nodes.dat";
370     std::ofstream fileWithNodeName(name.c_str(), ios::trunc);
371     boost::shared_ptr<Node> root;
372
373     vector<NameElementType> rootName;
374     root = boost::make_shared<Node>(rootName,0,0);
375     root->boolSize = vm["bool"].as<int>();
376     string fileName = boost::lexical_cast<string>(root->boolSize)+" .txt";
377     output = new std::ofstream(fileName.c_str(), ios::trunc);
378     arrayOfEdges.push_back(new vector<IdType>());
379
380     name = root->getName();
381     int stringSize = (root->boolSize <= 9 ? 2+root->boolSize*2 : 2+9*2+(root->
382         boolSize-9)*3);
383     fileWithNodeName << name;
384     for(int k = 0; k < stringSize-name.length(); k++)fileWithNodeName << "_";
385     fileWithNodeName << endl;
386
387     map<vector<NameElementType>, boost::shared_ptr<Node> >* layerAct = new map<
388         vector<NameElementType>, boost::shared_ptr<Node> >;
389     layerAct->operator [] (root->name) = root;
390
391     vector<IdType>* tmp_level = new vector<IdType>();
392     tmp_level->push_back(root->id);
393     listOfLevels.push_back(tmp_level);
394     map<vector<NameElementType>, boost::shared_ptr<Node> >* layerNext;
395     map<vector<NameElementType>, boost::shared_ptr<Node> >::const_iterator it, e;
396
397     // generating all elements
398     while(true){
399         layerNext = new map<vector<NameElementType>, boost::shared_ptr<Node> >();
400         tmp_level = new vector<IdType>();
401
402         for (it = layerAct->begin(), e = layerAct->end(); it != e; ++it)
403         {
404             it->second->generateNextBool(layerNext, &arrayOfEdges, tmp_level, &
405                 fileWithNodeName);
406         }
407         if(tmp_level->size() == 0) break;
408         listOfLevels.push_back(tmp_level);
409
410         layerAct->clear();
411         delete layerAct;
412
413         layerAct = layerNext;
414     }
415     // include small loop for the top node for printing decomposition
416     arrayOfEdges[arrayOfEdges.size()-1]->push_back(arrayOfEdges.size()-1);
417     delete layerAct;
418     fileWithNodeName.close();
419     // calling Python diagram class to generate pictures
420     if(listOfLevels.size() > 1){

```



```

475     layerAct->clear();
476     delete layerAct;
477
478
479     layerAct = layerNext;
480
481 }
482 // include small loop for the top node for printing decomposition
483 arrayOfEdges[arrayOfEdges.size()-1]->push_back(arrayOfEdges.size()-1);
484 delete layerAct;
485 fileWithNodeName.close();
486 // calling Python diagram class to generate pictures
487 if(listOfLevels.size() > 1){
488     findSSCD(&graph, vm);
489     if(vm.count("no_pict") == 0){
490         Py_Initialize();
491         PyRun_SimpleString("import os, sys");
492         PyRun_SimpleString("sys.path.append(os.getcwd())");
493         PyRun_SimpleString("import diagram");
494         ofstream py("python.in");
495         py << "f_=_open('" << fileName << "', 'r')\n".close()\ndiagram.\n
496             makeGraphOfInversePartiallyOrderedMultiSet(f, ["<< root->getName().substr
497                 (1, root->getName().size()-2) <<"])" ;
498         py.close();
499         FILE *fp = fopen("python.in", "r");
500         PyRun_SimpleFile(fp, "python.in");
501         fclose(fp);
502
503         Py_Finalize();
504         remove("python.in");
505     }
506 }
507 else
508     loggerOut("—The_poset_has_only_one_vertex_The_SSCD_is_obvious.—");
509 }
510 // helper to load the Young's lattice into the memory
511 void makeYoungsLattice(Graph *g, boost::program_options::variables_map vm){
512     Graph graph = *g;
513     int layer;
514     string name = "nodes.dat";
515     std::ofstream fileWithNodeName(name.c_str(), ios::trunc);
516     boost::shared_ptr<Node> root;
517
518     int m = vm["young"].as<vector<int>>()[0], n = vm["young"].as<vector<int>>()[1];
519     vector<NameElementType> rootName;
520     for (int i = 0; i < n; i++){
521         rootName.push_back(0);
522     }
523
524     root = boost::make_shared<Node>(rootName, 0, 0);
525     string fileName = "";
526     for (int i = 1; i < n; i++) fileName += lexical_cast<string>(m) + ",";
527     fileName += lexical_cast<string>(m) + ".txt";
528     output = new std::ofstream(fileName.c_str(), ios::trunc);
529     arrayOfEdges.push_back(new vector<IdType>());
530
531     fileWithNodeName << root->getNameYoung() << endl;

```

```

530 map<vector<NameElementType>, boost::shared_ptr<Node> >* layerAct = new map<
      vector<NameElementType>, boost::shared_ptr<Node> >;
531 layerAct->operator [] (root->name) = root;
532
533 vector<IdType>* tmp_level = new vector<IdType>();
534 tmp_level->push_back(root->id);
535 listOfLevels.push_back(tmp_level);
536 map<vector<NameElementType>, boost::shared_ptr<Node> >* layerNext;
537 map<vector<NameElementType>, boost::shared_ptr<Node> >::const_iterator it, e;
538
539 // generating all elements
540 while(true){
541     layerNext = new map<vector<NameElementType>, boost::shared_ptr<Node> >();
542     tmp_level = new vector<IdType>();
543
544     for (it = layerAct->begin(), e = layerAct->end(); it != e; ++it)
545     {
546         it->second->generateNextYoung(layerNext, &arrayOfEdges, tmp_level, &
            fileWithNodeName,m);
547     }
548     if(tmp_level->size() == 0) break;
549     listOfLevels.push_back(tmp_level);
550
551     layerAct->clear();
552     delete layerAct;
553
554     layerAct = layerNext;
555
556 }
557 // include small loop for the top node for printing decomposition
558 arrayOfEdges[arrayOfEdges.size()-1]->push_back(arrayOfEdges.size()-1);
559 delete layerAct;
560 fileWithNodeName.close();
561 // calling Python diagram class to generate pictures
562 if(listOfLevels.size() > 1){
563     findSSCD(&graph, vm);
564     if(vm.count("no_pict") == 0){
565         Py_Initialize();
566         PyRun_SimpleString("import os, sys");
567         PyRun_SimpleString("sys.path.append(os.getcwd())");
568         PyRun_SimpleString("import diagram");
569         ofstream py("python.in");
570         py << "f_=_open(' " << fileName << " ', 'r')\n f.close()\n diagram .
            makeGraphOfYoungsLattice(f, "+lexical_cast<string>(m)+", "+lexical_cast<
            string>(n)+")";
571         py.close();
572         FILE *fp = fopen("python.in", "r");
573         PyRun_SimpleFile(fp, "python.in");
574         fclose(fp);
575
576         Py_Finalize();
577         remove("python.in");
578     }
579 }
580 else
581     loggerOut ("—The_poset_has_only_one_vertex..The_SSCD_is_obvious.—");
582 }
583 // helper to load a poset from the given file into the memory
584 void readPosetFromFile(Graph *g, boost::program_options::variables_map vm){

```

```

585 // Load all nodes
586 Graph graph = *g;
587 string fileName = vm["file"].as<string>()+".txt";
588 output = new std::ofstream(fileName.c_str(), ios::trunc);
589 string name = vm["file"].as<string>()+ "_nodes";
590 std::ifstream ifFile(name.c_str());
591 std::istream& isFile = ifFile;
592 string line;
593 while (getline(isFile, line)){
594     vector<string> strs;
595     boost::split(strs, line, boost::is_any_of(", "));
596     arrayOfEdges.push_back(new vector<IdType>());
597     for(int i = 0; i < strs.size(); i++){
598         if(strs[0].size() == 0) continue;
599         if(strs.at(i).at(strs.at(i).size()-1) == 13) strs.at(i).resize(strs.at(i).
600             size() - 1);
601         arrayOfEdges[arrayOfEdges.size()-1]->push_back(lexical_cast<IdType>(strs.at
602             (i)));
603     }
604 }
605 ifFile.close();
606
607 // load the level structure
608 name = vm["file"].as<string>()+ "_levels";
609 ifFile.open(name.c_str());
610 isFile.copyfmt(ifFile);
611 while (getline(isFile, line)){
612     vector<string> strs;
613     boost::split(strs, line, boost::is_any_of(", "));
614     listOfLevels.push_back(new vector<IdType>());
615     for(int i = 0; i < strs.size(); i++){
616         if(strs.at(i).at(strs.at(i).size()-1) == 13) strs.at(i).resize(strs.at(i).
617             size() - 1);
618         listOfLevels[listOfLevels.size()-1]->push_back(lexical_cast<IdType>(strs.at
619             (i)));
620     }
621 }
622 // calling Python diagram class to generate pictures
623 if(listOfLevels.size() > 1){
624     findSSCD(&graph, vm);
625     if(vm.count("no_pict") == 0){
626         Py_Initialize();
627         PyRun_SimpleString("import os, sys");
628         PyRun_SimpleString("sys.path.append(os.getcwd())");
629         PyRun_SimpleString("import diagram");
630         ofstream py("python.in");
631         py << "f=_open('" << fileName << "', 'r')\nf.close()\ndiagram."
632             << "makeGraphFromFile(f, '"+vm["file"].as<string>()+ "'");
633         py.close();
634         FILE *fp = fopen("python.in", "r");
635         PyRun_SimpleFile(fp, "python.in");
636         fclose(fp);
637
638         Py_Finalize();
639         remove("python.in");
640     }
641 }
642 else
643     loggerOut("—The_poset_has_only_one_vertex._The_SSCD_is_obvious.—");

```

```

639 }
640
641 int main(int argc, char** argv)
642 {
643     time(&start_time);
644     boost::program_options::options_description desc("Allowed options");
645     desc.add_options()
646         ("bool", po::value<int>(), "set the number of elements in a set; --bool 4")
647         ("file", po::value<string>(), "--file [name]\nRead a poset from files. There
        have to be two files in the current folder, '[name]_nodes' and '[name]
        _levels'. Each line of '[name]_nodes' contains a list of neighbors of
        node with id equal to the line number starting from zero. Each line of '[
        name]_levels' contains a list of nodes in the layer which number
        corresponds to the number of the current line starting from zero.\n[name]
        _levels:\n0\n1,2\n3\n\n[name]_nodes:\n1,2\n3\n3\n3")
648         ("inversion", po::value<vector<int>>()->multitoken(), "element with rank 0;
        --inversion 1 2 3 4 [7...]")
649         ("young", po::value<vector<int>>()->multitoken(), "set m and n values; --
        young 3 4")
650         //("sum", po::value<int>(), "construct a poset using sum decomposition up to
        given number; --sum_decomposition 5")
651         ("no_pict", "will not generate pictures")
652         ("help", "produce help message")
653     ;
654     po::variables_map vm;
655     po::store(po::parse_command_line(argc, argv, desc), vm);
656     po::notify(vm);
657
658     if (vm.count("help") || ( !vm["young"].empty() && vm["young"].as<vector<int>>
        >().size() != 2 ) || ( vm["young"].empty() && vm["inversion"].empty() && vm
        ["sum"].empty() && vm["file"].empty() && vm["bool"].empty() ) ) {
659         cout << desc << "\n";
660         return 1;
661     }
662
663     Graph graph;
664     if (!vm["bool"].empty())
665         makeBoolPoset(&graph, vm);
666     if (!vm["file"].empty())
667         readPosetFromFile(&graph, vm);
668     if (!vm["inversion"].empty())
669         makeInversePoset(&graph, vm);
670     if (!vm["young"].empty())
671         makeYoungsLattice(&graph, vm);
672     return 0;
673 }

```