# <span id="page-0-1"></span>Evaluating and extending speedup techniques for optimal crossing minimization in layered graph drawings



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<span id="page-0-0"></span>Fig. 1: We aim to create faster and more scalable methods of fnding layered graph layouts with the minimum number of crossings. We characterize nine techniques to improve the performance of an integer linear programming (ILP) formulation and empirically test their improvement. We call these *switches* since they can be toggled and combined. Here, we show the performance of two switches and highlight an optimal control flow graph layout from our case study, with final node placements generated by the bendiness reduction of Di Bartolomeo et al. [\[9\]](#page-9-0) performed in sequence after crossing minimization (full layouts available at <https://osf.io/5vq79>). These control flow graphs can grow very large but benefit from minimal crossing visualizations to aid human task performance.

**Abstract**—A layered graph is an important category of graph in which every node is assigned to a layer, and layers are drawn as parallel or radial lines. They are commonly used to display temporal data or hierarchical graphs. Previous research has demonstrated that minimizing edge crossings is the most important criterion to consider when looking to improve the readability of such graphs. While heuristic approaches exist for crossing minimization, we are interested in optimal approaches to the problem that prioritize human readability over computational scalability. We aim to improve the usefulness and applicability of such optimal methods by understanding and improving their scalability to larger graphs. This paper categorizes and evaluates the state-of-the-art linear programming formulations for exact crossing minimization and describes nine new and existing techniques that could plausibly accelerate the optimization algorithm. Through a computational evaluation, we explore each technique's effect on calculation time and how the techniques assist or inhibit one another, allowing researchers and practitioners to adapt them to the characteristics of their graphs. Our best-performing techniques yielded a median improvement of 2.5–17× depending on the solver used, giving us the capability to create optimal layouts faster and for larger graphs. We provide an open-source implementation of our methodology in Python, where users can pick which combination of techniques to enable according to their use case. A free copy of this paper and all supplemental materials, datasets used, and source code are available at <https://osf.io/5vq79>.

**Index Terms**—Integer linear programming, layered graph drawing, layered network visualization, crossing minimization, edge crossings

# **1 INTRODUCTION**

In a layered graph, every node in the graph is assigned to a layer and edges connect nodes in different layers. Layered graphs (also called layered networks) are commonly used to represent sequential and hierarchical relationships across a wide variety of domains, including machine learning [\[33,](#page-9-1) [54\]](#page-10-0), biology [\[5\]](#page-9-2), the humanities [\[20\]](#page-9-3), and more. Usually, layered graphs are shown using node-link visualizations, which means that human task performance in reading them heavily depends on the spatial layout of nodes and edges [\[2\]](#page-9-4). Computing this layout is most often done using heuristic algorithms which are fast, but produce suboptimal

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*xx xxx. 201x. Date of Publication xx xxx. 201x; date of current version xx xxx. 201x. For information on obtaining reprints of this article, please send e-mail to: [reprints@ieee.org.](mailto:reprints@ieee.org) Digital Object Identifer: xx.xxxx/TVCG.201x.xxxxxxx*  results, particularly for sparsely connected graphs [\[31,](#page-9-5) [46\]](#page-10-1). Conversely, other methods focus on the optimality of the result but require more time and computational resources and are, therefore, less scalable.

Straightline edge crossing minimization is the most important aesthetic criterion for graph readability [\[25,](#page-9-6) [42\]](#page-10-2). The goal is to create a node-link visualization for a given graph with the fewest number of edges that cross over each other. Traditionally, this is done using heuristics which quickly produce layouts with few crossings. Notably, the Sugiyama framework for graph visualization [\[46\]](#page-10-1) breaks this into steps: nodes are frst assigned to layers, and are reordered within layers using a barycentric method to produce a drawing with few crossings. This approach creates very readable graph visualizations, and is widely used in graph drawing libraries [\[4,](#page-9-7) [17\]](#page-9-8).

Typically, methods are implemented with the goal of balancing the runtime and optimality of the layouts generated, but in this paper we focus on the creation of layouts with a provably-optimal number of overlapping edges. Not only does the optimal crossing minimization approach create more readable graphs, but it can also be used to

<span id="page-1-2"></span>benchmark the speed of heuristic layout algorithms and the readability of their layouts. However, optimal layouts can take a long time to compute, making heuristics more suitable for larger graphs.

This paper focuses on speeding up the creation of provably-optimal layouts using Integer Linear Programming (ILP). The advantage of ILP formulations is that they allow us to use powerful existing solvers, many of which are commercially available and very quick in practice. As ILP techniques and computer speeds improve, the computational cost of these exact methods approaches the realm of practicality for larger and larger graphs, leading researchers to speculate that optimal techniques will eventually replace heuristics for layered graph layouts [\[16\]](#page-9-9). Another beneft is that ILP is extremely fexible—we can combine constraints for whichever criteria we desire into a combined ILP model and explicitly input how much we want the result to favor one criterion over the others [\[9,](#page-9-0) [16\]](#page-9-9). Finally, ILP is uniquely positioned to solve specifc problems, such as the layout of sparse graphs, which are often in greatest need of minimal crossing visualization due to the increased importance of individual edges. While heuristics perform worse on sparser graphs, ILP's performance skyrockets [\[31\]](#page-9-5).

In this paper, we present two standard formulations for layered crossing minimization ILP and analyze nine techniques for improving runtime across a range of input sizes. To our knowledge, no prior research has reported on the use of two of these nine techniques, nor has any research empirically validated any of the techniques or transitivity formulations we present. To make optimal layered crossing minimization feasible on larger graphs, we categorize existing approaches and classify the performance of each of nine techniques for each of two transitivity types on a 3,200-graph dataset. We then perform a comprehensive benchmark study of 1,280 different ILP formulations on a dataset of 1,700 graphs, to compare the combined performance benefts of the different techniques. Finally, we illustrate our formulation with a case study on visualizing code control fow and compare it with existing formulations in this application.

Specifcally, this paper contributes to layered graph visualization:

- 1. An empirical comparison of nine new and existing techniques for faster exact crossing minimization and their scalability to larger graphs, using two different base formulations and two ILP solvers,
- 2. A case study demonstrating the practicality of our recommended method applied to control flow graphs, and
- 3. An open-source implementation of our exact crossing minimization algorithm in Python for both Gurobi 10.0 [\[21\]](#page-9-10) (license required) and HiGHS 1.5 [\[26\]](#page-9-11) (fully open source), available on OSF at <https://osf.io/5vq79>.

# **2 BACKGROUND AND RELATED WORK**

In the following section, we discuss relevant extant layered graph drawing algorithms. We also survey important heuristics, and describe notable optimal algorithms, including ILP formulations and optimization approaches. A review of general graph visualization and graph drawing is outside the scope of this discussion. However, for an introduction, see Tamassia's book [\[47\]](#page-10-3), Gibson et al.'s survey [\[19\]](#page-9-12), and the following systematic review of computational evaluations of graph layout algorithms [\[7\]](#page-9-13).

## <span id="page-1-1"></span>**2.1 Layered Graphs**

A *layered graph G*, also referred to as a *layered graph*, is a structure consisting of a set of vertices *V* (also referred to as nodes) and edges *E*, where each edge connects two vertices, and a layering *L*. The *layering*  is a function  $L: V \to \{1,2,3,...,K\}$ , where *K* is the number of layers in *G*. We call a layered graph *proper* if, for all  $(i, j) \in E$ , we have  $|L(i) - L(j)| = 1$ . That is, all edges in a proper layered graph are between consecutive layers, an important property for the use of the [formulations](#page-1-0)  outlined in this paper. When graphs contain edges traversing more than one layer, we split them up into sections using *anchor or dummy nodes*  as per Gansner et al. [\[17\]](#page-9-8). We refer to the number of nodes in the graph after this dummy node insertion as *total nodes*. We then consider a *drawing* to be an assignment of nodes to 2-dimensional coordinates, where nodes in the same layers are drawn aligned, and these layer lines are parallel and ordered consecutively. For the purposes of layouts in this paper, layers are drawn vertically.

Layered node-link graph visualizations are used to represent hierarchical and sequential relationships between entities [\[23\]](#page-9-14). These are useful for visualizing medical time series [\[1\]](#page-9-15), SQL queries [\[9\]](#page-9-0), and navigation techniques [\[24\]](#page-9-16), for example. Layers can be provided as part of the data or assigned as a step in the algorithm. Many criteria infuence the readability of these visualizations. Purchase found edge-crossing reduction to have the most signifcant impact on the readability of relations in a graph [\[42\]](#page-10-2). In later work, she formalized seven metrics to measure the aesthetic criteria that infuence the readability of graph layouts [\[43\]](#page-10-4). Hence, layout algorithms range from optimizing some of these criteria, such as minimizing edge bends [\[9](#page-9-0)[,46\]](#page-10-1), to constraining the design conventions of specifc domains, like calculating readable metro maps [\[37,](#page-10-5) [41\]](#page-10-6). While many of these metrics are studied, we focus on crossing minimization as it is the most important metric for many tasks [\[42,](#page-10-2) [53\]](#page-10-7).

Many fast but suboptimal heuristic algorithms have been proposed to minimize edge crossings for layered graphs. Among the most prominent, the Sugiyama framework [\[23,](#page-9-14) [45,](#page-10-8) [46\]](#page-10-1) consists of assigning layers to a graph and then sweeping through them to permute nodes with a barycenter heuristic. Eades and Wormald proposed a similar method in 1994, with a median heuristic to compute the positions of nodes [\[14\]](#page-9-17). In their paper, they also utilized [dummy nodes](#page-1-1) to ensure multilevel graphs have a [proper layering.](#page-1-1) This technique is still commonly used and allows us to make [formulations](#page-1-0) for layered graphs that would otherwise not be [proper.](#page-1-1) A few years later, Eades, alongside Lin and Tamassia, presented a degree-weighted barycentric method [\[13\]](#page-9-18). All these methods focus on iteratively sweeping through layers and computing "average" positions of nodes, with the main differences among them being how they defne average. The original barycentric heuristic [\[46\]](#page-10-1) has exceptional performance [\[31\]](#page-9-5) and is often implemented in graph visualization libraries [\[27\]](#page-9-19). Matuszewski et al. [\[36\]](#page-10-9) propose a *k*-layer sifting method that often outperforms the barycentric method at the expense of being slower.

## <span id="page-1-0"></span>**2.2 Optimal Algorithms for Layered Graphs**

Alternative to heuristic approaches, graph layouts can be computed by using precise mathematical expressions, which can be solved for provably optimal solutions. However, fnding a minimum crossing drawing for a layered graph is NP-Complete [\[18\]](#page-9-20) due to the combinatorial nature of reordering nodes. Hence, many approaches in the literature have focused on improving the performance and scalability of existing algorithms. In presenting their framework, Suyigama et al. proposed exact algorithms for permuting nodes within layers subject to a prior layer assignment step. For the exact layout, they designed their objective function to minimize the vertical distance between every pair of nodes at the end of each edge, which is a quadratic optimization problem [\[46\]](#page-10-1). Gansner et al. added more constraints to linearize the method proposed by Sugiyama [\[17\]](#page-9-8).

Optimal Formulations using ILP. Jünger and Muntzel proposed a branch-and-bound algorithm to minimize crossings in 2-layer graphs [\[30\]](#page-9-21). That same year Valls et al. also proposed another branch-and-bound method [\[50\]](#page-10-10).

We call a *formulation* a way to defne a desired property as a set of integer variables and linear constraints. An *ILP model* is an encoded formulation (or multiple formulations) with a corresponding optimization goal given to an ILP solver. ILP solvers programmatically fnd an integer assignment to the variables that satisfy all constraints such that the optimization expression is minimized/maximized. ILP solvers leverage years of research [\[28\]](#page-9-22), and commercial solvers such as Gurobi report yearly performance increases [\[21\]](#page-9-10). Therefore, we expect ILP solvers' increased performance to translate to more viable and scalable optimal formulations. This paper aims to compare the performance of existing ILP models by combining techniques that alter the formulations themselves or infuence the solver. Below, we discuss the varied formulations and optimization techniques found in the literature.

Extant ILP Formulations. In 1997, Jünger and Mutzel proposed an ILP formulation to permute the nodes in the 2-layer case to reduce crossing numbers [\[30\]](#page-9-21). They further developed a multilayer approach [\[31\]](#page-9-5) using the [dummy node](#page-1-1) strategy. Their formulation directly ensures the transitivity of nodes by outlining constraints on the vertical position variables for each pair of nodes. Di Bartolomeo et al. presented another notable formulation in Stratisfmal Layout [\[9\]](#page-9-0). While many other formulations

## Table 1: Notation Used in This Paper

<span id="page-2-5"></span><span id="page-2-1"></span>
$$
G = (V, E, L)
$$
  
\nA layered graph *G* with vertex set *V*, edge set *E*, and  
\nlayer assignment on the vertices  $L: V \rightarrow \{1, 2, ..., K\}$ .  
\nTransformed node set of *G* with dummy nodes inserted  
\non long edges and |*V'* | total nodes.  
\nAll  $(u, v) \in E$  such that  $L(u) = r$  and  $L(v) = r + 1$ .  
\nK  
\nBinary variable denoting the relative position of nodes

 $i$  and  $j$  in the layout.  $c_{(i,k),(j,l)}$  Binary variable denoting if edges  $(i,k)$  and  $(j,l)$  cross.

ensured transitivity via direct constraints, Stratisfmal Layout implies this property by assigning indexed positions to nodes within layers when minimizing edge length. Indices are natural numbers, which are transitive on the "greater than" relation (>). Di Bartolomeo et al. formalized vertical position constraints as a set of implications relating direct transitivity constraints to node positions. They proceed to linearize these implications with a conversion that introduces auxiliary variables to provide constraints for the ILP. Stratisfmal Layout still implements the direct transitivity constraints, which creates redundancy in their [formulation.](#page-1-0)

Notable Speedup Approaches. The corpus of [ILP formulations](#page-1-0) also consists of strategies to improve the solver's performance. To our knowledge, we are the frst to empirically evaluate these when combined and compared across different transitivity formulations. Gange et al. provide a comprehensive evaluation of ILP and MIP solutions for layered crossing minimization using a few of the switches outlined in our work [\[16\]](#page-9-9) the closest computational evaluation to ours in scale of anything we could fnd in the literature. They propose symmetry breaking, collapsing leaf nodes, and an improved set of constraints based on the vertex exchange graph introduced by Healy and Kuusik [\[22\]](#page-9-23) as potential improvements.

Mutzel proposed leveraging the planarity of graphs on their formulation by considering constraints based on subgraphs that guarantee the existence of a crossing [\[39\]](#page-10-11). Their work started with the 2-layer case [\[40\]](#page-10-12) and was later extended to *k*-levels by Healy and Kuusik [\[22\]](#page-9-23). Notably, Mutzel described how cycles in 2-layer graphs guarantee the existence of crossings. Their approach involved creating constraints that checked for the existence of such subgraphs. Conversely, we use this insight to remove crossing constraints with known solutions.

Zarate et al. abstracted Sankey Diagrams as layered graphs and propose an ILP approach for their layouts [\[56\]](#page-10-13). They added symmetric variables on crossings to hone in on possible solutions to increase performance. Some ILP literature discusses how adding redundant variables can help the [formulation](#page-1-0) determine solutions faster [\[32\]](#page-9-24). They noticed that if there is a crossing between two vertices, it does not matter which vertex we consider the frst one in our notation the crossing is still there. However, adding redundancy is theoretically justifed but not experimentally verifed. They also proposed enhancing the solver's performance by branching on the binary position variables (*x*-vars). Branch-and-bound techniques are often used in ILP problems to fnd candidate solutions [\[38\]](#page-10-14). Their formulation included a crossing constraint, with  $x, c \in \{0,1\}$ . However, they highlight that since *x*-vars constraints are binary, the crossing variables (*c*-vars) must be integers. Therefore, they mention they do not have to enforce the constraints that c-vars are also binary since the crossing constraints imply this. Nonetheless, they claim that adding branching priorities on x-vars is more effcient, telling the solver these are more important in fnding the solution space.

Some of these optimizations inspire the "switches" used for the different experiments we run to evaluate the performance of different models, which we further discuss in their respective sections. We also compare the [formulations](#page-1-0) of Jünger et al. [\[29\]](#page-9-25), Zarate et al. [\[56\]](#page-10-13), Gange et al. [\[16\]](#page-9-9), and Di Bartolomeo et al. [\[9\]](#page-9-0) to our own in the context of a case study.

# **3 METHODOLOGY**

Modern ILP solvers allow complex problems to be solved much more quickly than brute force, provided the problem can be formulated as a set of linear constraints. Exact layered crossing minimization is one such problem. We frst defne the standard [formulation](#page-1-0) for layered crossing minimization, which is a way to transform any input graph into a set of

variables and linear constraints. This formulation has the property that if we can fnd the assignment to the variables with minimum crossing variable sum, it will correspond to the layout of the graph with the minimum possible edge crossings. We then describe two variations of this formulation which we will later compare empirically. Next, we detail nine "switches", which modify the formulation and ILP solver in ways that could improve the speed of exact crossing minimization while still returning an optimal solution.

# <span id="page-2-3"></span>**3.1 ILP Formulation**

Existing techniques for formulating layered crossing minimization using ILP follow the same standard template. We assume a [proper](#page-1-1)  layered graph *G* is given, with a [layer assignment](#page-1-1)  $L: V \rightarrow \{1,2,...,K\}$ for every node in *V*, such that all edges in *E* connect nodes in different layers. Given *G*, we say the following variables and constraints defne the standard model  $M_G$ :

- *Position variables*, denoted *xi*, *j*, corresponding to two nodes *i* and *j*. These are defned for every pair of nodes in the same layer. If *i* is drawn above *j* in that layer we write  $x_{i,j} = 1$ , and if *i* is below *j* we write  $x_{i,j}=0$ .
- *Crossing variables*, denoted  $c_{(i,k),(j,l)}$ , corresponding to whether or not edges  $(i,k)$  and  $(j,l)$  cross in the drawing of the graph. They are defned when *i*, *j*,*k*, and *l* are all unique, for every pair of edges between the same consecutive layers. If  $(i,k)$  and  $(j,l)$  cross, we write  $c_{(i,k),(j,l)} = 1$ ; if they do not, we write  $c_{(i,k),(j,l)} = 0$ .
- *Crossing constraints*, which are given in [\[9,](#page-9-0) [56\]](#page-10-13), and in [\[29\]](#page-9-25) as a set of inequalities equivalent to:

<span id="page-2-4"></span>
$$
c_{(i,k),(j,l)} + x_{j,i} + x_{k,l} \ge 1
$$
  
\n
$$
c_{(i,k),(j,l)} + x_{i,j} + x_{l,k} \ge 1
$$
 (1)

These are defined for every crossing variable  $c_{(i,k),(j,l)}$ , and enforce  $c_{(i,k),(j,l)} \geq 1$  if  $(i,k)$  and  $(j,l)$  cross. This concept is illustrated here:



The first equation enforces  $c_{(i,k),(j,l)} = 1$  when *i* is above *j* but *k* is below *l* (illustration 2 above); the second enforces  $c_{(i,k),(i,l)} = 1$ when  $i$  is below  $j$  and  $k$  is above  $l$  (illustration 3). Note that these constraints can be added only if *G* is a [proper](#page-1-1) layered graph.

• *Transitivity constraints*, see [Sec. 3.2.](#page-2-0)

When combined into an [ILP model,](#page-1-0) valid assignments to position and crossing variables correspond to a layout of the input graph. The sum of all crossing variables is the number of crossings in this [drawing.](#page-1-1) Therefore, the assignment to these variables with the smallest sum

<span id="page-2-2"></span>
$$
OBJ = \sum_{r=1}^{K-1} \sum_{e_1, e_2 \in E_r} c_{e_1, e_2} \tag{2}
$$

represents a layout with the fewest crossings. This is the *objective function* the ILP solver seeks to minimize. The goal of the ILP solver is to assign binary values to all *x*- and *c*-variables such that the crossing and transitivity constraints are satisfed and the sum [Eq. \(2\)](#page-2-2) is minimal.

So, given some input graph, we encode it using the above constraints to form an [ILP model.](#page-1-0) This is passed to an ILP solver, which works to fnd assignments minimizing the objective function *OBJ*, and we can post-process these assignments to create and visualize the crossing-minimized [drawing.](#page-1-1)

## <span id="page-2-0"></span>**3.2 Transitivity Constraints**

Since node placements are encoded using their position relative to other nodes instead of absolute position, it is necessary to include constraints that prevent non-transitive assignments. In order for the variable assignments produced by the ILP solver to correspond to a valid [layout,](#page-1-1) <span id="page-3-8"></span>there must be additional constraints that ensure the *x*[-variables](#page-2-1) are not assigned values by the ILP solver which disobey transitivity. For instance, if we start with a graph that has three nodes *i*, *j*, and *k* in the same layer, nothing stops our ILP solver from setting  $x_{i,j} = 1$ ,  $x_{j,k} = 1$ , and  $x_{i,k} = 0$ . But if we decode what this is actually saying, we fnd out that the ILP solver has just told us to draw *i* above *j*, *j* above *k*, and *k* above *i*—this is impossible since we cannot draw *i* simultaneously above and below *k*.

There are multiple ways of encoding this transitivity relationship as an ILP constraint, which we will compare empirically.

## <span id="page-3-7"></span>3.2.1 Direct transitivity constraints

The original [formulation](#page-1-0) of Jünger et al. [\[29\]](#page-9-25) enforces transitivity directly on all triples of nodes in the same layer:

<span id="page-3-0"></span>
$$
x_{i,j} + x_{j,k} - x_{i,k} \le 1
$$
  
\n
$$
x_{i,j} + x_{j,k} - x_{i,k} \ge 0
$$
\n(3)

[Equation \(3\)](#page-3-0) enforces the relationships  $x_{i,j} = 1 \wedge x_{j,k} = 1 \Rightarrow x_{i,k} = 1$ and  $x_{i,j} = 0 \land x_{j,k} = 0 \Rightarrow x_{i,k} = 0$ , which comes directly from the definition of a transitive relation. That is, if we have that  $i$  is above  $j$ and *j* is above *k*, we require that *i* is above *k*. Similarly for the converse, if we have *i* below *j* and *j* below *k*, we require that *i* is below *k*. These are added to the ILP model for every combination of nodes *i*, *j*,*k* in the same layer, guaranteeing a transitive relationship across the entire layer. This generates  $O(|V|^3)$  constraints.

## <span id="page-3-6"></span>3.2.2 Vertical position transitivity

Di Bartolomeo et al. [\[9\]](#page-9-0) defne an additional integer-valued variable *y*  for each node representing the vertical position of the node when drawn. For this to work, we require constraints that ensure that assignments to *x*[-variables](#page-2-1) (the relative positions of nodes) are consistent with assignments to *y*-variables (the absolute position of each node). The following inequalities from Di Bartolomeo et al. accomplish this:

$$
z_{i,j} - M \cdot x_{i,j} \le 0
$$
  
\n
$$
z_{i,j} - y_i - M \cdot x_{i,j} \ge -M
$$
  
\n
$$
y_j - z_{i,j} - x_{i,j} \ge 0
$$
  
\n
$$
z_{i,j} - y_i \le 0
$$
  
\n
$$
z_{i,j} \ge 0
$$
\n(4)

<span id="page-3-1"></span>These constraints implicitly enforce transitivity on the *x*-variables, and the authors also use them to relate the *x*-variables to vertical position *y*-variables to better control the placement of nodes in the resultant layout. *M* is a fixed upper bound which is set to the size of the largest layer. Eq. (4) also requires an additional variable  $z_{i,j}$  for each variable  $x_{i,j}$ , and adds  $O(|V|^2)$  constraints to the model in total. For large graphs, this adds far fewer constraints than [Eq. \(3\).](#page-3-0) Advanced solvers such as Gurobi [\[21\]](#page-9-10) can perform this linearization automatically provided  $x_{i,j} = 1 \implies y_i < y_j$ and  $x_{i,j} = 0 \Longrightarrow y_j > y_i$  for each  $x_{i,j}$ , the constraints encoded by [Eq. \(4\).](#page-3-1)

## **3.3 Switches**

Having defned the ILP formulation for crossing minimization, we now detail nine techniques we call "switches". In contrast to [formulations,](#page-1-0) switches can be toggled on and off but never alter the value of the solution found by the model. *Switches* are defned as modifcations to the variables and constraints in an [ILP model,](#page-1-0) or alterations to the behavior of the ILP solver, in a way that does not affect the optimal solution.

The purpose of these switches is to try and improve the runtime of the ILP solver and its ability to scale to larger graphs. This is accomplished by restricting the space of all possible solutions or giving hints to the solver that guide it toward the optimal solution. Due to the nature of exponential growth, [ILP models](#page-1-0) will always reach a point where the inputs get too big, causing runtime to rapidly increase, thereby dramatically decreasing the feasibility of solution-fnding for graphs past a certain size. Our goal in defning these switches is to try to reduce the time complexity of crossing minimization, even if some additional overhead is required. This expands the range of feasible graph sizes for the optimizer. To that end, we frst summarize the action behind

## Table 2: Switch Descriptions

<span id="page-3-5"></span>

each switch in [Tab. 2](#page-3-5) before describing them in detail and explaining our intuition for why it may improve performance.

## <span id="page-3-2"></span>Switch 1: Symmetry breaking

Gange et al. introduce symmetry breaking in their model, which is the process of selecting a single *x*-variable and fxing it to 0 prior to optimization [\[16\]](#page-9-9). This works because there is functionally no difference, most importantly in crossing number, between a layered graph layout and the upside-down version of the same layout. This is visible in [Fig. 1,](#page-0-0) where the two layouts of the graph at top-left are functionally the same. So, we can fix  $x_{u,v} = 0$  before optimizing the model without repercussion. So, fxing one single variable breaks this symmetry and eases the burden on the ILP solver by one decision variable.

Gange et al. implement this by selecting the *x*-variable that corresponds to the frst pair of nodes in the same layer. We extend this technique by selecting not the frst *x*-variable, but the *x*-variable that appears in the greatest number of [crossing constraints.](#page-2-3) This gives the solver a starting point to leverage for fnding the optimal assignment to the other variables and simplifes the model as much as possible.

## <span id="page-3-3"></span>Switch 2: Butterfly reduction

This switch involves crossing variables whose edges are part of a  $2 \times 2$ biclique, or *butterfy* [\[44\]](#page-10-15). This graph motif is 2-level non-planar [\[39\]](#page-10-11). Zarate et al. [\[56\]](#page-10-13) make use of this fact by describing an additional ILP constraint for any butterfly  $\{(i, j), (k, l), (i, l), (k, j)\} \subseteq E$ :

$$
c_{(i,j),(k,l)} + c_{(i,l),(k,j)} = 1
$$
\n(5)

In [Fig. 2a](#page-4-6) we can see that swapping the within-layer positions of nodes *k* and *l* does not change that the graph has exactly one crossing. Likewise for swapping the positions of *i* and *j*.

# <span id="page-3-4"></span>Switch 3: Polyhedral constraints

Jünger et al. study the polytope associated with the solution space of the layered crossing minimization problem [\[29\]](#page-9-25), and derive several classes of constraints which are "facet-defning", meaning they restrict the solution space of the crossing minimization problem as much as possible.

<span id="page-4-7"></span><span id="page-4-6"></span>

Fig. 2: Butterfy reduction pictured in (a), 3-claw motif and dome-path respectively pictured in (b) and (c).

This switch adds all facet-defning constraints described by Jünger et al., with the exception of two sets of constraints which are cycle-based. The fndings detailed in the appendices at<https://osf.io/5vq79> and the recommendation of Gange et al. for cycle-based constraints [\[16\]](#page-9-9) suggest that these constraints do not signifcantly improve solution time. Furthermore,  $\Box$  [switch 2](#page-3-3) already studies constraints regarding the 4-cycle.

The only other minimal non-planar 2-layer motif besides the butterfy is the *3-claw* [\[39\]](#page-10-11), shown in [Fig. 2b.](#page-4-6) Jünger et al. describe additional constraints for any occurrence of this subgraph [\[29\]](#page-9-25).

$$
\sum_{e_1, e_2 \in W} c_{e_1, e_2} \ge 1 \tag{6}
$$

For any pair of edges  $e_1, e_2 \in W$  such that the vertex point *v* of *W* is part of exactly one of the two edges, for all 3-claws *W* in *G*.

The other set of constraints involves *dome-paths* [\(Fig. 2c\)](#page-4-6), which are added for any 2-layer path on four edges and fve nodes *i*, *j*,*k*,*l*,*m*:

$$
x_{k,l} - 2x_{k,m} + x_{l,m} - c_{(i,k),(j,l)} - c_{(i,l),(j,m)} \le 0
$$
  

$$
-x_{k,l} + 2x_{k,m} - x_{l,m} - c_{(i,k),(j,l)} - c_{(i,l),(j,m)} \le 0
$$
 (7)

## <span id="page-4-0"></span>Switch 4: Mirrored variables with symmetry constraints

This switch doubles the number of variables by adding both  $x_{i,j}$  and  $x_{j,i}$ to the model. Without this switch on, we assume only one of the two is added, say  $x_{i,j}$ . If a constraint requires  $x_{j,i}$  we instead substitute  $1 - x_{i,j}$ (the negation of  $x_{i,j}$ ). The statement "*i* is above *j*" being false implies that "*i* is below *j*" is true, hence this substitution is valid. Turning this switch on also implements the following symmetry constraints from Zarate et al. [\[56\]](#page-10-13):  $x_{i,j} = 1 - x_{j,i}$  and  $c_{e_1,e_2} = c_{e_2,e_1}$  for all *x*- and *c*-variables, enforcing the symmetric relationship of the mirrored variables.

This doubles the number of variables used in the model; however, redundancy in ILP models has been shown to sometimes improve solver performance [\[32\]](#page-9-24). Other previous works typically follow the precedent of Jünger et al. [\[29\]](#page-9-25) who do not use mirrored variables, so for the purpose of this paper we use the more minimal model as default.

## <span id="page-4-1"></span>Switch 5: Cycle constraints

Healy and Kuusik describe a structure called the vertex exchange graph which gives rise to additional ILP constraints [\[22\]](#page-9-23). The vertex exchange graph is created from an input graph *G* by defining a node  $\langle u_1, u_2 \rangle$  for every same-layer node pair in *G* (i.e., one node for every *x*-variable in the standard ILP model) and connecting pairs of nodes  $\langle u_1, u_2 \rangle, \langle v_1, v_2 \rangle$ with an edge if  $(u_1, v_1)$  and  $(u_2, v_2)$  are edges in *G* (thus the edges correspond with *c*-variables in the ILP model). Cycles in this new graph are *odd-labeled* if the sum of the *c*-variables of the cycle edges is odd and *even-labeled* otherwise. If this switch is on, add the following constraints for each fundamental cycle *C* in the vertex exchange graph:

For odd-labeled cycles *C*:

$$
\sum_{e \in C} c_e \ge 1\tag{8}
$$

For even-labeled cycles *C*:

$$
\sum_{e \in C} c_e \le |C| - 1 \tag{9}
$$



Fig. 3: Example of  $\Box$  [leaf node collapse.](#page-4-2) The left graph contains a node with four leaves, highlighted in red, which are combined into a single node as shown on the right.

These a reduced set of the original paper's [\[22\]](#page-9-23) constraints per the recommendation of Gange et al. [\[16\]](#page-9-9) (see the appendices at [https:](https://osf.io/5vq79) [//osf.io/5vq79](https://osf.io/5vq79) for an evaluation supporting this recommendation).

## <span id="page-4-2"></span>**O** Switch 6: Collapse leaf nodes

Introduced by Gange et al. [\[16\]](#page-9-9), this switch is unique from the other switches in that it directly modifes the input graph. The principle of the technique is to select a leaf node subgraph—two or more nodes connected only to a single parent node—and remove all the leaf nodes, replacing them with a single "collapsed" node and edge. Once the crossing minimized layout is found, the nodes in this subgraph are re-inserted back into the graph by squeezing them all into the spot left by the collapsed node. To ensure optimality, the objective function in [Eq. \(2\)](#page-2-2) must be updated to

$$
OBJ = \sum_{r=1}^{K-1} \sum_{e_1, e_2 \in E_r} w_{e_1} w_{e_2} c_{e_1, e_2}
$$
 (10)

where  $w_e$  is the weight of the collapsed edge and  $w_e = 1$  if *e* is not a collapsed edge. This procedure works because if it is optimal to place one leaf node in a certain position, then any other leaf nodes must also be optimally placed if they are immediately adjacent—all leaves will incur the same number of crossings.

# <span id="page-4-3"></span>Switch 7: Branching on *x*-variables

This switch sets the branching priority of all *x*-variables to 1 and all other variables to 0. ILP solvers search for potential solutions by *branching*  on a decision variable, which involves looking at the potential solutions when the branch variable is fxed. At each branching point, a variable is selected randomly from the set of variables with the highest branching priority that has not yet been branched on [\[21\]](#page-9-10). Zarate et al. claim this yields up to  $10\times$  performance improvement for large instances [\[56\]](#page-10-13).

#### <span id="page-4-4"></span>Switch 8: Heuristic starting assignments

This switch uses the straightforward iterated degree-weighted barycenter heuristic [\[13\]](#page-9-18) to provide an initial starting assignment for the solver. ILP solvers perform an iterative process of fnding assignments to the variables which progressively get closer to the optimal solution, but modern solvers allow the user to input a valid starting assignment to the variables. This is not the same as fxing the variable, as is done in  $\bullet$  [switch 1,](#page-3-2) because the solver can change the assignments made by the starting value parameter.

The barycenter heuristic is widely used for crossing minimization on layered graphs We use it as a reasonable starting point for the ILP solver. Providing a starting point is suggested to speed up solution time as it potentially skips early iterations of the ILP algorithm [\[21\]](#page-9-10).

## <span id="page-4-5"></span>Switch 9: Continuous variables

Mixed-integer programming (MIP) solvers have traditionally been used to solve layered crossing minimization, utilizing a branch-and-cut approach [\[29\]](#page-9-25). Confusingly, the terms ILP and MIP are sometimes used synonymously. With modern solvers, however, users can defne the integrality of each variable in the model explicitly, and the solver selects the best algorithms to use on the back end. This switch makes sure the

Table 3: Datasets Used for Empirical Study

<span id="page-5-4"></span><span id="page-5-1"></span>

Experiment	Graphs	V'	Per Bin	By Collection	
				Rome-Lib	AT&T
Ind. switches	3170	10-399	100	2959	211
All combos.	1713	10-399	50	1615	98

*x*[-variables](#page-2-1) are kept as binary variables, but the crossing *c*[-variables](#page-2-1)  and vertical position *y*[-variables](#page-3-6) (if [vertical position transitivity](#page-3-6) is being used) are explicitly defned to be real-valued.

This does not affect the optimal solution of the model:

- [Equation \(1\)](#page-2-4) ensures  $c > 0$  if *c*'s edges do not cross and  $c > 1$  if they do. Since our objective is to minimize the sum of all *c*-variables, real-valued *c*-variables will still all converge to either 1 or 0 in the optimal solution.
- $\bullet$  " $\lt$ " is a transitive relation on the real numbers and the integers, so [Eq. \(4\)](#page-3-1) still enforces transitivity on real-valued *y*-variables.

A number of solvers have made advancements in algorithms for mixed-integer programming (MIP)problems, with Gurobi 10.0 claiming  $24\%$  improvement over the previous release for large models.<sup>1</sup> It may improve solution time if we can expressly defne some variables in our model to be real-valued since MIP algorithms may be more efficient than ILP algorithms for the same problem [\[51\]](#page-10-16), or signal to the solver a more efficient way to treat these continuous variables.

# **4 EXPERIMENT**

We will now analyze the effect these nine [switches](#page-3-5) and the [transitivity](#page-2-0)  [formulations](#page-2-0) have on optimization time via a computational evaluation.

# <span id="page-5-3"></span>**4.1 Dataset**

The experiment dataset was composed of graphs from standard benchmark collections AT&T [\[10\]](#page-9-26) and Rome-Lib [\[11\]](#page-9-27), detailed in [Tab. 3.](#page-5-1) Graphs used are available from the [Graph Drawing Benchmark](https://visdunneright.github.io/gd_benchmark_sets/)  [Datasets Repository](https://visdunneright.github.io/gd_benchmark_sets/) [\[8\]](#page-9-28).

For our frst experiment on individual switch evaluation, we combined the entire AT&T collection of 1,276 graphs with the Rome-Lib collection of 11,528 graphs and randomly sampled 100 graphs for every 10-node interval up to 400 [total nodes.](#page-2-1) Past the [270,280) interval there were fewer than 100 graphs per 10-node interval, so no sampling was necessary. In practice, no run of the experiment completed the full dataset without being cutoff due to timing out. Note that graph size |*V* ′ | used for the sampling is in terms of the post-processed number of vertices, see [Sec. 4.2.1.](#page-5-2) This dataset contains 3,170 graphs and is included at<https://osf.io/5vq79>with the code that performed the sampling.

For the second experiment evaluating all combinations of switches, we performed the same steps as for the previous experiment, with the exception that we sampled 50 graphs per 10-node interval up to 400 [total nodes.](#page-2-1) Sampling was necessary up to interval [310,320), and as before, no run of the experiment completed the full dataset. The dataset for the all-combinations experiment contains 1,713 graphs.

# **4.2 Procedure**

We now discuss how we pre-processed and modifed the graphs to create [proper](#page-1-1) layered graphs. We also detail the evaluation conducted for individual switches and all combinations of switches.

## <span id="page-5-2"></span>4.2.1 Graph pre-processing

The Rome-Lib [\[11\]](#page-9-27) and AT&T [\[10\]](#page-9-26) graphs do not have a predetermined [layer assignment,](#page-1-1) hence one must be created for each graph.

First, we interpret the input graph as directed by assuming the frst node in each edge is the source and the second is the target for the Rome-Lib graphs. The AT&T graphs are already both directed and acyclic [\[10\]](#page-9-26). We then applied the greedy cycle removal heuristic of Eades et al. [\[12\]](#page-9-29), followed by [assigning layers](#page-1-1) using the minimum width layering heuristic of Tarassov et al. [\[48\]](#page-10-17). They suggest using the layering with minimum width across all combinations of input parameters  $UBW = 1,2,3,4$  and  $c = 1,2$ , which are tuning parameters specific to the minimum-width algorithm. We use only  $UBW = 4$  and  $c = 2$  as we do not require perfect width minimization, and these parameter choices still consistently produce layered graphs with "rectangular" shapes. I.e., they had a similar number of nodes in each layer. This rectangular shape is desirable as it better approximates real-world layered graphs such as Storyline graphs and time-series data [\[1,](#page-9-15) [20\]](#page-9-3), and gives the drawing a conventional aspect ratio without much unused space. More trivial approaches, such as assigning layers based on each node's level in the tree created by a breadth-frst search, tend to have a few layers with many more nodes than the others. Finally, we added back the edges removed by the cycle removal step to re-create the original graph topology.

After pre-processing to ensure we had [layered graphs,](#page-1-1) we ensured each graph was [proper](#page-1-1) by replacing edges that skip layers by adding [dummy nodes,](#page-1-1) following the procedure described by Eades and Wormald [\[14\]](#page-9-17). That is, if  $(u, v)$  is a long edge, where  $L(u) + 1 < L(v)$  $L(u) + 1 < L(v)$ , we remove edge  $(u, v)$  and add nodes  $d_1, d_2, \ldots, d_r$  where:

$$
L(d_1) = L(u) + 1
$$
  
\n
$$
L(d_2) = L(u) + 2
$$
  
\n
$$
\vdots
$$
  
\n
$$
L(d_r) = L(u) + r = L(v) - 1
$$

We then add dummy edges  $(u,d_1), (d_1,d_2),..., (d_r,v)$ .

This process transforms the graph into a [proper](#page-1-1) graph so that crossing constraints can be imposed correctly. We refer to the number of nodes in this processed graph (including [dummy nodes\)](#page-1-1) as [total nodes,](#page-2-1) which may be larger than the number of nodes in the original graph. We henceforth report data using total nodes as the independent variable, since we fnd that it more closely correlates with solver runtime than the number of original nodes.

The resultant graph is transformed into its corresponding set of linear constraints as described in [Sec. 3.1,](#page-2-3) and they are modifed according to the choice of switches. These are provided to Gurobi 10.0, a state-of-the-art linear programming solver, which computes the optimal layered graph drawing.

#### 4.2.2 Individual switch evaluation

For both [transitivity formulations,](#page-2-0) we ran a baseline experiment with all switches disabled. We also evaluated the effect of turning on only one of the switches at a time. Therefore, we performed 18 runs of the experiment for each pair of the two transitivity formulations and nine switches. With the baseline runs, this left us with 20 sets of results.

Each run comprised solving every graph in the experiment dataset [\(Secs. 4.1](#page-5-3) and [4.2.1\)](#page-5-2) and recording the solver runtime. We separated graphs into bins by their [total nodes](#page-2-1) in 10-node intervals. When less than 75% of the graphs in a bin completed optimization within the cutoff time of 5 minutes, the run was halted. The two baseline runs were continued for every graph completed in the nine individual-switch runs for that formulation. This ensured all data points in the experiment had a corresponding baseline data point for comparison.

#### 4.2.3 Evaluation of all switch combinations

To fnd the best-performing sets of [switches,](#page-3-5) and to make comparisons between the different [transitivity formulations,](#page-2-0) we ran an experiment for all 1024 combinations of switches  $(2^9)$  and formulations  $(\times 2)$ . We used a smaller set of graphs to run the experiment within the time available. All 1024 formulation-switch combinations were run on the all-combinations dataset of 1,713 graphs [\(Tab. 3\)](#page-5-1), with the experiment halted once fewer than half of the graphs in any 10-node bin were solved within the cutoff time. The 5-minute cutoff time per graph was changed to a shorter 1-minute cutoff for this experiment to reduce the number of compute-hours to perform the experiment. Additionally, the halting condition was relaxed for this experiment to better compare switch combinations. Instead, all combinations of switches were evaluated on all graphs solved by the best-performing combination.

<span id="page-5-0"></span><sup>1</sup><https://www.gurobi.com/whats-new-gurobi-10-0/>

<span id="page-6-2"></span><span id="page-6-0"></span>

Fig. 4: The left fgure shows performance improvement over the baseline for individual switches. All fgures plot median runtime as a natural log ratio over the respective baseline, where line color corresponds to the color of the switch. The two center figures show more detailed information for the symmetry breaking (1) and mirrored variables (4). The lines are 25% and 75% quartiles, and the darker center line is the median. Data is truncated once fewer than 75% of the graphs per 10-node interval are solved within the 5-minute cutoff by both the switch technique and the baseline. [Symmetry breaking](#page-3-2) gives a large performance improvement which increases as input sizes grows, while **CO** [mirrored variables](#page-4-0) also improved with input size but hurt performance on average for most sizes tested. On the right, individual switch results for the HiGHS solver are shown, where **O** [symmetry breaking,](#page-3-2) [continuous variables,](#page-4-5) commirrored variables, and constraints all perform well. All figures shown combine the direct transitivity and vertical position transitivity results. See appendices at <https://osf.io/5vq79> for more in-depth results figures.

## 4.2.4 Evaluation using different solver

We repeated both of the experiments described above using HiGHS, an open-source linear optimization solver  $[26]$ .  $\Box$  Heuristic start and  $\bullet x$ [-var branch priority](#page-4-3) are unavailable due to the required functionality being not yet available for HiGHS, and so were left out of both experiments. All experiment parameters were otherwise kept the same.

## 4.2.5 Analysis

We report results for these experiments by calculating, for each 10-node bin, the natural logarithm of the ratio of mean experimental runtime to the mean baseline runtime. Natural log ratios are additive, symmetric, normed indicators of relative change [\[49\]](#page-10-18), allowing us to more easily compare the impact of a switch on different [formulations.](#page-1-0)

Units. Each data point is reported in *optimization time*, the time in seconds it took the ILP solver to determine the optimal solution, with independent variable [total nodes.](#page-2-1) This ignores setup and pre-processing time, which took at most a few seconds for the largest graphs.

Hardware/software. We ran the experiments in CentOS Linux with 8 GB RAM & Gurobi 10.0 [\[21\]](#page-9-10). For the secondary solver, we used HiGHS 1.5 [\[26\]](#page-9-11) available through SciPy 1.10's linprog.

## <span id="page-6-1"></span>**4.3 Discussion and Results**

In the following section we discuss key takeaways and ramifcations of our experimental results. We also outline ways practitioners can change their models to improve performance, and the degree of speedup that can be expected.

# 4.3.1 Switch performance

Use  $\Box$  [symmetry breaking](#page-3-2) to reliably reduce optimization time. Compared to the baseline, only one [switch](#page-3-5) had a large runtime impact for both solvers when used in isolation. The overall results are shown in [Fig. 4.](#page-6-0) Deleting the symmetry in the model by fxing one variable makes fnding a solution approximately twice as fast (in general), with even more speedup for larger graphs. For both [transitivity formulations,](#page-2-0) every combination of switches that included fxing one variable greatly outperformed the combinations that did not [\(Fig. 6\)](#page-7-0). It is incredibly helpful as a universal technique and shows that even advanced ILP solvers do not always have the capability to recognize the inherent symmetry of the problem. Adopting this approach may improve runtime performance for other problems in visualization and beyond. We recommend that anyone implementing an ILP/MIP model examine the problem for symmetries that allow you to fx even a single variable.

Combining switches is very successful, but only when using HiGHS. With the open-source solver HiGHS [\[26\]](#page-9-11), all

three of  $\Box$  [symmetry breaking,](#page-3-2)  $\Box$  [mirrored variables,](#page-4-0) and [continuous variables](#page-4-5) contributed signifcant performance improvements when using direct transitivity—see [Fig. 5.](#page-7-1) Moreover, the formulation using these three switches was the best-performing combination of the experiment, solving an impressive 88% of the 600 graphs tested within one minute. The baseline, meanwhile, solved only 60%. This corresponds to a more than  $17\times$  median speedup across all graphs tested—the combination had a median runtime of 1.9 seconds while the baseline took a median of 32.5 seconds to complete.

A number of other switches also contributed to strong performance results when using HiGHS with direct transitivity, namely  $\Box$  butterfly [reduction,](#page-3-3)  $\bullet$  [cycle constraints,](#page-4-1) and  $\bullet$  [leaf node collapse.](#page-4-2) Generally speaking, combining switches together greatly improved the performance of the HiGHS solver.

This contrasts with the Gurobi results, for which  $\bigcirc$  symmetry [breaking](#page-3-2) drastically improved runtime while the inclusion of additional switches often slowed the solver down. For instance, using symmetry breaking with no other switches was a top-performing combination. Solving the 1150 graphs took a median runtime of 1.7 seconds while the baseline took 4.2 seconds—a more modest  $2.5 \times$  speedup. We suspect that since Gurobi is a very efficient solver optimized for finding high-quality potential solutions, the addition of some of our described techniques is more of a distraction. That is, the solver wastes time each iteration verifying the additional constraints added by many of the techniques we describe are upheld when that time would often be better spent applying solution-fnding and pruning heuristics to advance towards the optimal solution.

The remaining switches give little performance increase. By comparison, the remaining six switches do not have much, if any, positive impact when used in isolation, suggesting their usefulness is more restricted to specifc graphs, or larger input sizes and cutoff times. Results for all individual switches are included in the supplemental materials at <https://osf.io/5vq79>.

However, switch performance dependends on more than node count. We were concerned with the variability exhibited in the ILP solver times and performed additional studies on layer counts and edge density with a more controlled dataset. Procedures and results are described in the appendices at <https://osf.io/5vq79>. We fnd that when using Gurobi with direct transitivity, both  $\bigcirc$  [symmetry breaking](#page-3-2) and  $\bullet$  *x*[-var branch priority](#page-4-3) delay the exponential spike in runtime that happens when increasing the edge density and number of layers—for the 50-node graphs studied,  $\Box$  [symmetry breaking](#page-3-2) in particular was able to solve graphs at up to 35% edge density within 5 minutes, while the baseline was unable to solve the graphs with 25% edge density.

<span id="page-7-3"></span><span id="page-7-1"></span>

Fig. 5: Median *improvement* provided by each switch for direct and vertical transitivity formulations, provided as an increase in the number of graphs solved in the all-combinations experiment dataset (1,150 graphs for Gurobi and 600 for HiGHS). Average improvements over 1%—12 graphs for the Gurobi experiment and 6 for HiGHS—are bolded. We define improvement for a switch as the difference between the number of graphs successfully solved for a given formulation and the same formulation without that switch, presenting the median improvement over all formulations including the given switch. We see that including **O** [symmetry breaking](#page-3-2) resulted in being able to solve an average of 160 (13.9%) additional graphs when using Gurobi and direct transitivity. This is a huge improvement, seeing as each 10-node interval included 50 graphs: including this switch allowed us to routinely solve graphs 30 nodes larger than without the switch, within the same amount of time.

Guidelines for using switches. We provide the following summary of the improvement provided by each switch technique.



For the best guarantee of fast optimal layouts, it is recommended to select a number of combinations of switches shown to perform well on average for your choice of solver and run them in parallel. This also helps offset some of the variability inherent to the ILP solving process. For an example of this, refer to [Sec. 5.](#page-7-2)

## 4.3.2 Transitivity constraint performance

Direct transitivity generally outperforms vertical position transitivity. Each combination of switches with direct transitivity solved, on average, 21 more graphs (1.9% of the dataset) than the same combination with vertical position transitivity. It is worth noting that direct transitivity combinations took more setup time than vertical position transitivity combinations, which never took more than 1 second. This is due to the direct transitivity requiring more constraints than vertical position transitivity—for large inputs, this  $O(|V|^3)$  versus  $O(|V|^2)$ difference starts to become noticeable. Additionally, vertical position transitivity performs better than direct transitivity for specifc graphs, oftentimes large graphs with many layers but fewer nodes per layer, as in [Sec. 5.](#page-7-2) However, for the HiGHS solver all-combinations experiment, direct transitivity formulations solved 7.0% more graphs on average, a much more substantial increase than Gurobi. Direct transitivity formulations also responded better to the addition of switches, seen in [Fig. 5.](#page-7-1) Therefore, we recommend using direct transitivity constraints, although experimentation with vertical position transitivity is benefcial for large many-layered graphs when using effcient solvers.

<span id="page-7-0"></span>

Fig. 6: All 512 combinations for each transitivity formulation plotted against their baseline using the natural log ratio of median runtimes. In the top row, all combinations using  $\Box$  [symmetry breaking](#page-3-2) are highlighted in red, showing that the use of the switch completely partitions the space of all combinations beyond 50 total nodes. <br> **O** [Polyhedral constraints,](#page-3-4) meanwhile, tend to be on the bottom half of the two partitions performance-wise.

Some types of redundancy are helpful, but others are not. Lalla-Ruiz et al. claim that redundancy can both help and hinder the solver performance [\[32\]](#page-9-24). We can corroborate this since some of our techniques introduce redundancy into the model. Specifcally, **O** [mirrored variables](#page-4-0) doubles the variables and constraints in the model, but does not typically slow down solution time, even improving it substantially for some solvers and formulations. However, a study we conducted found that including both direct transitivity and vertical position transitivity simultaneously had a large negative impact on solution time—see appendices at <https://osf.io/5vq79>.

## <span id="page-7-2"></span>**5 CASE STUDY: VISUALIZING SOFTWARE CONTROL FLOW**

Control Flow Graph (CFG) readability is crucial to help reverse engineers [\[35,](#page-9-30)[52\]](#page-10-19) and malware researchers [\[55\]](#page-10-20) extract knowledge from decompiled binary fles, including malware. Therefore minimizing edge crossing should be prioritized to facilitate tasks like tracking variables and tracing activation code in the diagram.

Reverse engineering control flow graphs can be represented as layered [graphs,](#page-1-1) where nodes correspond to blocks of decompiled assembly code, and edges represent jumps taken between code blocks [\[35\]](#page-9-30). Code blocks are often ended by JMP or JNZ instructions, followed by an edge connecting to the next code block. Specifc structures can appear in CFG visualizations which tell engineers about the nature of the code. For example, switch instructions generate a large number of outgoing edges from a single node. The [layer assignment](#page-1-1) for these [drawings](#page-1-1) is typically done by trying to maximize the number of call arrows pointing in the same direction before assigning positions within each layer, using a Sugiyamastyle approach [\[6\]](#page-9-31). The layouts of these drawings are conventionally computed with heuristic-based algorithms (Radare2, for instance, cites Buchheim et al. [\[3\]](#page-9-32)). The optimal approach we propose can be used to produce more readable visualizations for reverse engineers. For example, the readability and utility of control fow graphs generated by tools like the recent work by Devkota et al. [\[6\]](#page-9-31) could be further improved by using layouts with the optimally fewest crossings. While optimal approaches have rarely been applied in this context in the past because of their high

<span id="page-8-1"></span><span id="page-8-0"></span>

Table 4: Control Flow Graph optimization results, reported as the median of fve trials on each graph. All formulations were tested using Gurobi. The formulations for comparison are, in order, a baseline with direct transitivity and no switches, the same baseline with  $\Box$  [symmetry breaking,](#page-3-2) Di Bartolomeo et al.'s Stratisfmal Layout (SL) [\[9\]](#page-9-0), Zarate et al.'s Optimal Sankey Diagrams (OSD) [\[56\]](#page-10-13), Jünger et al.'s Polyhedral Approach to Crossing Minimization (PA) [\[29\]](#page-9-25), and Gange et al.'s Optimal *k*-Level Planarization and Crossing Minimization (OkLCM) [\[16\]](#page-9-9). Our ensemble method, taking advantage of running multiple formulations in parallel, is quickest for all three graphs.

computational requirements, we demonstrate that our approach is viable for use with control flow graphs with very large numbers of nodes.

We compare an ensemble method of our best switch combinations to formulations from the literature, which we can also represent as switch combinations. Di Bartolomeo et al.'s Stratisfmal Layout [\[9\]](#page-9-0) uses vertical position transitivity with no other switches. Zarate et al.'s Optimal Sankey Diagrams [\[56\]](#page-10-13) use direct transitivity with  $\Box$  mirrored [variables,](#page-4-0)  $\bullet$  butterfly reduction, and  $\bullet$  *x*[-var branch priority.](#page-4-3) Jünger et al.'s Polyhedral Approach [\[29\]](#page-9-25) uses direct transitivity with **D** [polyhedral constraints.](#page-3-4) Gange et al.'s Optimal *k*-Level Crossing Minimization [\[16\]](#page-9-9) uses direct transitivity with  $\bigcirc$  symmetry [breaking,](#page-3-2)  $\Box$  [cycle constraints,](#page-4-1) and  $\Box$  [leaf node collapse.](#page-4-2)

Our ensemble method ran six combinations in parallel, taking the layout from whichever fnished frst. The table below lists these combinations by their transitivity type [\(direct transitivity](#page-3-7) or [vertical tran](#page-3-6)[sitivity\)](#page-3-6), and the switches combined. These combinations were selected because they performed strongly in the all-combinations experiment.



The case study was run on a desktop computer with an Intel Core i7-8700K CPU with 6 cores, 32 GB RAM, Windows 10, & Gurobi 10. The number of processing threads available to the solver was limited to 1 for all ensemble members, and 6 for all other methods. Our ensemble method found an optimal control fow diagram for the implementation of the Linux chmod command in an average of 2.34 seconds over fve repeated trials, see [Tab. 4.](#page-8-0) Part of this optimal layout is pictured in [Fig. 1.](#page-0-0) This control flow graph has 511 nodes and 581 edges after layering and inserting [dummy nodes](#page-1-1) and edges—all control fow graphs included had a large number of layers but reasonable layer widths, allowing for quick solving even for the relatively huge graph size.

Our recommended method never took longer than 1.5 seconds to fnd the optimal drawing, which has 16 crossings. The advantage of the ensemble method is that it leverages our fnding in [Sec. 4.3](#page-6-1) that different formulations perform better on different input graphs. For chmod, the fastest-performing ensemble member was model 2, which used only  $\Box$  [symmetry breaking.](#page-3-2) More complicated models worked much better than this for echo and cp, on the other hand, which were solved fastest by models 5 and 4 respectively. Each member solved at least one of the graphs very quickly, but speed often varied by up to  $5\times$  slower than the fastest member, particularly for cp. As for the other models evaluated, the access to  $6\times$  more processing power did not seem to improve performance much if at all, likely due to the relatively quick execution times and the solver not needing to visit many branch-and-bound nodes [\[21\]](#page-9-10). chmod and echo were solved quickly by OkLCM [\[16\]](#page-9-9), but cp was solved the quickest by the vertical transitivity-based formulations such as SL [\[9\]](#page-9-0).

## **6 LIMITATIONS**

Like all methods that use ILP, the key limitation of our approach is the upper bound on the size of graphs which can be solved in a reasonable amount of time. In our experiments, graphs with up to 100 nodes in the original graph could typically be solved quickly with our techniques. Graphs with more than 300 nodes could not often be solved within a 5-minute cutoff time for the edge densities studied in our experiments.

It should be noted, however, that the number of edges and nodes is not fully indicative of the complexity of a problem. For clarity of presentation, we have categorized graphs throughout this paper by their [total nodes.](#page-2-1) Although it is more consistent than reporting based on the number of nodes in the original graph, it is still far from perfect as a predictor of runtime. We have studied some additional factors (see appendices at <https://osf.io/5vq79>), but more work is needed. There is also little indication in our results of what causes certain techniques to perform much better on some graphs than others, an example being the differences in performance between direct transitivity and vertical position transitivity for the control fow graphs studied in [Sec. 5.](#page-7-2)

Furthermore, it is impossible to perfectly predict the runtime of the ILP solver given only knowledge of the input graph because ILP solvers contain internal randomization such as for selecting branching variables. This randomization causes some variability in optimization time with repeated trials [\[15,](#page-9-33) [34\]](#page-9-34). Sometimes, the solver gets lucky and fnds a solution very quickly, when the solver would take much longer on average. Conversely, the solver can get unlucky and take much longer than average. We overcome this variance in the experiments by using a large number of graphs, and in practice by recommending an ensemble method.

# **7 CONCLUSION AND FUTURE WORK**

Algorithms that arrange node-link graph visualizations with consideration of human factors can assist users in comprehending complex graphs more quickly. With the aim of scaling optimal layouts to larger graphs, we present in this paper an adaptable framework that yields empirically faster layout optimization, and analyzed nine established and innovative techniques using two different solvers. The result is a comprehensive benchmark characterizing the degree to which the nine techniques, two types of transitivity formulation, and two ILP solvers impact the running time of ILP solutions for layered crossing minimization. Our recommended approach can quickly generate optimal layouts for most graphs with up to 150 nodes and works in a reasonable amount of time for many graphs with even more nodes. Our implementation is available as open-source code, as are our benchmark results and datasets used for our empirical study.

Future work. More research can be done to evaluate techniques for the scalability of aesthetic criteria besides crossing minimization, such as planarity, edge length, edge bundling, and node groupings. The experiments described in this paper can be replicated using additional ILP solvers, ensemble methods such as the one in [Sec. 5,](#page-7-2) and larger graphs over longer runtimes, to better recommend software to use for the problem and further investigate the trends observed. Additionally, more experimentation is in order to better understand the relationship between the input graph and the solver runtime. Work could be done to integrate our fast crossing minimization algorithm with Devkota et al.'s *CFGConf*  for even more understandable, useful control flow graph generation.

# **SUPPLEMENTAL MATERIALS**

All supplemental materials can be found on OSF at <https://osf.io/5vq79>. We provide:

- 1. A copy of our paper, including all appendices and supplemental figures.
- 2. Open-source GitHub repository of our implementation in Python, including a LayeredOptimizer class which takes as input a layered graph, and allows the user to choose what switches to use, then generates the optimal drawing.
- 3. All result data and datasets used in our experiments, as well as code used to sample said datasets.
- 4. All results fgures:
	- (a) 21 individual switch evaluation fgures, one for each switch on each transitivity formulation.
	- (b) 24 total fgures which extend [Fig. 6](#page-7-0) to all switches.
	- (c) Full optimal layouts for the three control fow graphs in the case study [\(Sec. 5\)](#page-7-2).

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