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# Dippy - a simplified interface for advanced mixed-integer programming 

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# Dippy - a simplified interface for advanced mixed-integer programming 

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#### Abstract

Mathematical modelling languages such as AMPL, GAMS, and Xpress-MP enable mathematical models such as mixed-integer linear programmes (MILPs) to be expressed clearly for solution in solvers such as CPLEX, MINOS and Gurobi. However some models are sufficiently difficult that they cannot be solved using "out-of-the-box" solvers, and customisation of the solver framework to exploit model-specific structure is required. Many solvers, including CPLEX, Symphony and DIP, enable this customisation by providing "callback functions" that are called at key steps in the solution of a model. This approach traditionally involves either expressing the mathematical formulation in a low-level language such as C++ or Java, or implementing a complicated indexing scheme to be able to track model components such as variables and constraints between the mathematical modelling language and the solver's callback framework.

In this paper we present Dippy, a combination of the Python-based mathematical modelling language PuLP and the open source solver DIP. Dippy provides the power of callback functions, but without sacrificing the usability and flexibility of modelling languages. We discuss the link between PuLP and DIP and give examples of how advanced solving techniques can be expressed concisely and intuitively in Dippy.


## 1 Introduction

Using a high-level modelling language such as AMPL, GAMS, Xpress-MP or OPL Studio enables Operations Research practitioners to express complicated mixed-

[^0]integer linear programming (MILP) problems quickly and naturally. Once defined in one of these high-level languages, the MILP can be solved using one of a number of solvers. However these solvers are not effective for all problem instances due to the computational difficulties associated with solving MILPs (an NP-Hard class of problems). Despite steadily increasing computing power and algorithmic improvements for the solution of MILPs in general, in many cases problem-specific techniques need to be included in the solution process to solve problems of a useful size in any reasonable time.

Both commercial solvers such as CPLEX and Gurobi and open source solvers such as CBC, Symphony and DIP (all from the COIN-OR repository [1]) provide callback functions that allow user-defined routines to be included in the solution framework. To make use of these callback functions the user must first create their MILP problem in a low-level computer programming language (C, C++ or Java for CPLEX, C, C++, C\#, Java or Python for Gurobi, C or C++ for CBC, Symphony or DIP). As part of the problem defintion, it is necessary to create structures to keep track of the constraints and/or variables. Problem definition in C/C++/Java for a MILP problem of any reasonable size and complexity is a major undertaking and thus a major barrier to the development of customised MILP frameworks by both practitioners and researchers.

Given the difficulty in defining a MILP problem in a low-level language, another alternative is use a high-level mathematical modelling language. By carefully constructing an indexing scheme, constraints and/or variables in the highlevel language can be identified in the low-level callback functions. However implementing the indexing scheme can be as difficult as using the low-level language to define the problem in the first place and does little to remove the barrier to solution development.

The purpose of the research presented here is to provide a tool, Dippy, that supports easy experimentation with and customisation of advanced MILP solution frameworks. To achieve this aim we needed to:

1. provide a modern high-level modelling system that enables users to quickly and easily describe their MILP problems;
2. enable simple identification of constraints and variables in user-defined routines in the solution framework.

The first requirement is satisfied by the modelling language PuLP [3]. Dippy extends PuLP to use the Decomposition for Integer Programming (DIP) solver, and enables user-defined routines implemented in the same Python file as the MILP model to be accessed by the DIP callback functions. Variable scope in the Python computer programming language [5] can be used so that any constraints or variables defined in the MILP model are easily accessible in the user-defined routines. In addition to this, DIP is implemented so that the MILP problem is defined the same way whether branch-and-cut or branch-price-and-cut is being used - it hides the implementation of the master problem and subproblems. This makes it very easy to switch between the two approaches when experimenting with solution methods. All this functionality combines to overcome the barrier
described previously and provides researchers, practitioners and students with a simple and integrated way of describing problems and customising the solution framework.

The rest of this article is structured as follows. In section 2 we provide an overview of the interface between PuLP and DIP, including a description of the callback functions available in Python from DIP. In section 3 we describe how Dippy enables experimentation with improvements to DIP's MILP solution framework by showing example code for a common problem. We conclude in section 4 where we discuss how this project enhances the ability of researchers to experiment with approaches for solving difficult MILP problems. We also demonstrate that DIP (via PuLP and Dippy) is competitive with leading commercial (Gurobi) and open source (CBC) solvers.

## 2 Combining DIP and PuLP

Dippy is the primarily the "glue" between two different technologies: PuLP and DIP.

PuLP [3] is a mathematical modelling language and toolkit that uses Python. Users can define MILP problems and solve them using a variety of solvers including CPLEX, Gurobi and CBC. PuLP's solver interface is modular and thus can be easily extended to use other solvers such as DIP. For more details on PuLP see the PuLP project in the COIN-OR repository [1].

Decomposition for Integer Programming (DIP) [4] provides a framework for solving MILP problems using 3 different methods ${ }^{1}$ :

1. "branch-and-cut",
2. "branch-price-and-cut",
3. "decompose-and-cut".

In this paper we will restrict our attention to branch-and-cut and branch-price-and-cut.

Branch-and-cut uses the classic branch-and-bound approach for solving MILPs combined with the cutting plane method for removing fractionality encountered at the branch-and-bound nodes. This framework is the basis of many state-of-the-art MILP solvers including Gurobi and CBC. DIP provides callback functions that allow users to customise the solution process by adding their own cuts and running heuristics at each node.

Branch-price-and-cut uses Dantzig-Wolfe decomposition to split a large MILP problem into a master problem and one or more subproblems. The subproblems solve a pricing problem, defined using the master problem dual values, to add new variables to the master problem. Branch-and-cut is then used on the master problem.

[^1]The cut generation and heuristic callback functions mentioned previously can also be used for branch-price-and-cut. Extra callback functions enable the user to define their own routines for finding initial variables to include in the master problem and for solving the subproblems to generate new master problem variables. For details on the methods and callback functions provided by DIP see [4].

In addition to the DIP callback functions (see §2.1), we modified DIP to add another callback function that enables user-defined branching in DIP and so can be used in any of the solution methods within DIP.

### 2.1 Callback Functions

Advanced Branching We replaced chooseBranchVar in the DIP source with a new function chooseBranchset. This is a significant change to branching in DIP that makes it possible for the user to define:

- a down set of variables with (lower and upper) bounds that will be enforced in the down node of the branch; and,
- an $u p$ set of variables with bounds that will be enforced in the up node of the branch.

A typical variable branch on an integer variable $x$ with integer bounds $l$ and $u$ and fractional value $\alpha$ can be implemented by:

1. choosing the down set to be $\{x\}$ with bounds $l$ and $\lfloor\alpha\rfloor$;
2. choosing the up set to be $\{x\}$ with bounds of $\lceil\alpha\rceil$ and $u$.

However, other branching methods may use advanced branching techniques such as the one demonstrated in §3.1. From DIP, chooseBranchset calls branch_method in Dippy.

Customised Cuts We modified generateCuts (in the DIP source) to call generate_cuts in Dippy. This enables the user to examine a solution and generate any customised cuts as necessary. We also modified APPisUserFeasible to call is_solution_feasible in Dippy, enabling users to check solutions for feasibility with respect to customised cuts.

Customised Columns (Solutions to Subproblems) We modified the DIP function solveRelaxed to call relaxed_solver in Dippy. This enables the user to utilise the master problem dual variables to produce solutions to subproblems (and so add columns to the master problem) using customised methods. We also modified generateInitVars to call init_vars in Dippy, enabling users to customise the generation of initial columns for the master problem.

Heuristics We modified ApPheuristics (DIP) to call heuristics (Dippy). This enables the user to define customised heuristics at each node in the branch-andbound tree (including the root node).

### 2.2 Interface

The interface between Dippy (in Python) and DIP (in C++) is summarised in figure 1.


Figure 1: Key components of interface between Dippy and DIP.

The MILP is defined as a DipProblem and then solved using the solve command in Dippy, that passes the Python DipProblem object, prob, to DIP in C++. DIP solve creates a DippyDecompAlgo object that contains a DippyDecompApp object, both of which are populated by data from prob. As DIP solve proceeds branches are created by the DippyDecompAlgo object using chooseBranchSet which passes the current node's fractional solution xhat back to the branch_method function in the Dipproblem object prob. This function generates lower and upper bounds for the "down" and "up" branches and returns to DippyDecompAlgo::chooseBranchset. When DIP generates cuts, it uses the DippyDecompApp object's generateCuts function which passes the current node's solution sol to the DipProblem object's generate_cuts function. This function generates any customised cuts and returns a list, cuts, back to DippyDecompApp: : generateCuts. These interfaces are replicated for the other callback functions provided by Dippy.

## 3 Dippy in Practice

We will use the Capacitated Facility Location problem from the PuLP documentation [3] to demonstrate the implementation of customised branching rules, custom cuts, heuristics, and a column generation algorithm.

The solution of the problem determines where, amongst $m$ locations, to place facilities and also assigns production of $n$ products to these facilities in a way that (in this version) minimises the wasted capacity of facilities. Each product $j=1, \ldots, n$ has a production requirement $r_{j}$ and each facility has capacity $C$. Extensions of this problem arise often in MILP in problems including network design and rostering.

The MILP formulation of the capacitated facility location problem is straightforward. The decision variables are

$$
\begin{aligned}
x_{i j} & = \begin{cases}1 & \text { if product } j \text { is produced at location } i \\
0 & \text { otherwise }\end{cases} \\
y_{i} & = \begin{cases}1 & \text { if a facility is located at location } i \\
0 & \text { otherwise }\end{cases} \\
w_{i} & =\text { "wasted" capacity at location } i
\end{aligned}
$$

and the formulation is

$$
\begin{array}{lcl}
\text { min } & \sum_{i=1}^{m} w_{i} \\
\text { s.t. } \quad \sum_{i=1}^{m} x_{i j}=1, j=1, \ldots, n & \text { (each product produced) } \\
& \sum_{j=1}^{n} r_{j} x_{i j}+w_{i}=C y_{i}, i=1, \ldots, m & \text { (aggregate capacity at location } i \text { ) } \\
x_{i j} \leq y_{i}, i=1, \ldots, m, j=1, \ldots, n & \text { (individual production at location } i \text { ) } \\
& x_{i j} \in\{0,1\}, w_{i} \geq 0, y_{i} \in\{0,1\}, i=1, \ldots, m, j=1, \ldots, n
\end{array}
$$

Note that the contraints to for individual production at a location are not necessary for defining the solution, but tighten the MILP formulation by removing fractional solutions from the solution space.

### 3.1 Adding Customised Branching

In §2.1 we explained the modifications made to DIP and how a simple variable branch would be implemented. The DIP function chooseBranchset calls Dippy's branch_method at fractional nodes. The function branch_method has two inputs supplied by DIP:

1. prob - the DipProblem being solved;
2. sol - an indexable object representing the solution at the current node.

We define branch_method using these inputs in the same Python file as the model definition, allowing Dippy to access the variables from the original formulation and eliminating any need for complicated indexing.

We can explore custom branching rules by trying to reduce the symmetry in the solution space of the facility location problem. Inefficiencies arise from solvers considering multiple equivalent solutions that have identical objective function values and differ only in the subset of the identical facilities selected. One way to address this is to add a constraint that determines the order in which the facilities can be considered:

$$
y_{i} \geq y_{i+1}, i=1, \ldots, m-1
$$

```
# Ordering constraints
for index, location in enumerate(LOCATIONS):
    if index > 0:
        prob += use_vars[LOCATIONS[index-1]] >= use_vars[location]
```

These ordering constraints also introduce the opportunity to implement an effective branch on the number of facilities:

If $\sum_{i=1}^{m} y_{i}=\alpha \notin \mathbb{Z}$, then:

$$
\begin{array}{l|l}
\text { the branch down restricts } & \begin{array}{l}
\text { the branch up restricts } \\
\sum_{i=1}^{m} y_{i} \leq\lfloor\alpha\rfloor
\end{array} \\
\begin{array}{l}
\sum_{i=1}^{m} y_{i} \geq\lceil\alpha\rceil \\
\text { and the ordering means that } \\
y_{i}=0, i=\lceil\alpha\rceil, \ldots, m
\end{array} & \begin{array}{l}
\text { and the ordering means that } \\
y_{i}=1, i=1, \ldots,\lceil\alpha\rceil
\end{array}
\end{array}
$$

We can implement this branch in Dippy by writing a definition for the branch_method.

```
def choose_antisymmetry_branch(prob, sol):
    num_locations = sum(sol[use_vars[i]] for i in LOCATIONS)
    up = ceil(num_locations) # Round up to next nearest integer
    down = floor(num_locations) # Round down
    if (up - num_locations > tol) \
    and (num_locations - down > tol): # Is fractional?
        # Down branch: provide upper bounds, lower bounds are default
        down_branch_ub = dict([(use_vars[LOCATIONS[n]], 0)
                for n in range(int (down),
                len(LOCATIONS) ) ])
        # Up branch: provide lower bounds, upper bounds are default
        up_branch_lb = dict([(use_vars[LOCATIONS[n]], 1)
        for }n\mathrm{ in range(0, int(up))])
        # Return the advanced branch to DIP
        return {}, down_branch_ub, up_branch_lb, {}
```

By adding the ordering constraints we decrease the branch-and-bound tree size from 419 nodes to 77 nodes. Adding the advanced branching solves the problem very quickly, decreasing the tree size further to 3 nodes.

### 3.2 Adding Customised Cut Generation

To add user-defined cuts in Dippy, we first define a new procedure for generating cuts and (if necessary) a procedure for determining a feasible solution. Within Dippy, this requires two new functions, generate_cuts and is_solution_feasible. As in §3.1, Python's scoping rules allow us to access the solution values of variables in our problem. Both these functions have the same inputs as branch_method:

1. prob - the DipProblem being solved;
2. sol - an indexable object representing the solution at the current node.

If a solution is determined to be infeasible either by DIP (for example some integer variables are fractional) or by is_solution_feasible (which is useful for solving problems like the travelling salesman problem with cutting plane methods), cuts will be generated by generate_cuts and the in-built Cut Generator Library (CGL) (if enabled).

Marchand and Wolsey [2] define many types of cuts for MILP problems. One of these is the weighted inequality. For each facility location $i$ and some subset $S_{i}(\subseteq\{1, \ldots, n\})$ of the products we can calculate

$$
\mu_{i}=C-\sum_{j \in S_{i}} w_{j} x_{i j}
$$

and use it to generate a weighted inequality

$$
\sum_{j \in S_{i}} w_{j} x_{i j}+\sum_{j \notin S_{i}}\left(w_{j}-\mu_{i}\right)^{+} x_{i j} \leq C-\mu_{i}
$$

which forms a valid inequality for the facility location problem.
The cut generating function creates the subsets $S_{i}$ for each location from the fractional solution in a greedy way depending on the $x_{i j}$ values, and from these we generate a set of weighted inequality cuts. The code listing below shows how to build the set of cuts, and omits the generation of $S_{i}$ for the sake of brevity.

```
def generate_weight_cuts(prob, sol):
    # Define mu and T for each knapsack
    mu = {}
    S = {}
    for i in LOCATIONS:
        mu[i] = CAPACITY
        S[i] = []
```

```
# Generate the weight cuts from the sets found above
new_cuts = []
for i in LOCATIONS:
    if len(S[i]) > 0: # If an item assigned to this location
        con = LpAffineExpression() # Start a new constraint
        con += sum(REQUIREMENT[j] * assign_vars[(i, j)]
                                for j in S[i])
        con += sum(max(0, REQUIREMENT[j] - mu[i]) *
                                    assign_vars[(i, j)] for j in PRODUCTS
                                    if j not in S[i])
        new_cuts.append(con <= CAPACITY - mu[i])
# Return the set of cuts we created to DIP
return new_cuts
```

Adding the weighted inequality cut generator reduces the branch-and-bound tree size from 419 nodes to 77 nodes.

### 3.3 Adding Customised Column Generation

Using Dippy it is easy to transform a problem into a form that can be solved by either branch-and-cut or branch-price-and-cut. Branch-price-and-cut decomposes a problem into a master problem and a number of distinct subproblems. We can identify subproblems using the relaxation member of the DipProblem class. Once the subproblems have been identified, then they can either be ignored (when using branch-and-cut - the default method for DIP) or utilised (when using branch-price-and-cut - specified by turning on the dopriceCut option).

In branch-price-and-cut, the original problem is decomposed into a master problem and multiple subproblems [6]:

$$
\begin{array}{rcrcc}
\min & c_{1}^{\top} x_{1}+c_{2}^{\top} x_{2} & +\cdots+c_{K}^{\top} x_{K} \\
\text { subject to } & A_{1} x_{1} & +A_{2} x_{2} & +\cdots+A_{K} x_{K}=b \\
& & F_{2} x_{2} & & =f_{2} \\
& & & \ddots & \vdots  \tag{1}\\
& & & F_{K} x_{K}=f_{K} \\
& & \\
x_{1} \in \mathbb{Z}_{n_{1}}^{+}, x_{2} \in \mathbb{Z}_{n_{2}}^{+}, \ldots, x_{K} \in \mathbb{Z}_{n_{K}}^{+}
\end{array}
$$

In (1), there are $K-1$ subproblems defined by the constraints $F_{k} x_{k}=f_{k}, k \in$ $2, \ldots, K$. The constraints $A_{1} x_{1}+A_{2} x_{2}+\cdots+A_{K} x_{K}=b$ are known as linking constraints. Instead of solving (1) directly, column generation uses a convex combination of solutions $y^{k}$ to each subproblem $j$ to define the subproblem variables:

$$
\begin{equation*}
x_{k}=\sum_{l_{k}=1}^{L_{k}} \lambda_{l_{k}}^{k} y_{l_{k}}^{k} \tag{2}
\end{equation*}
$$

where $0 \leq \lambda_{l_{k}}^{k} \leq 1$ and $\sum_{l_{k}=1}^{L_{k}} \lambda_{l_{k}}^{k}=1$. By substituting (2) into the linking constraints and recognising that each $y_{l_{k}}^{k}$ satisfies $F_{k} x_{k}=f_{k}, x_{k} \in \mathbb{Z}_{n_{k}}^{+}$(as it is a solution of this subproblem), we can form the restricted master problem (RMP) with
corresponding duals $\left(\pi, \gamma_{1}, \ldots, \gamma_{K}\right)$ :

$$
\begin{align*}
& \min c_{1}^{\top} x_{1}+\sum_{l_{2}=1}^{L_{2}}\left(c_{2}^{\top} y_{l_{2}}^{2}\right) \lambda_{l_{2}}^{2}+\cdots+\sum_{l_{K}=1}^{L_{K}}\left(c_{K}^{\top} y_{l_{K}}^{K}\right) \lambda_{l_{K}}^{K} \\
& \text { subject to } A_{1} x_{1} \quad+\sum_{l_{2}=1}^{L_{2}}\left(A_{2} y_{l_{2}}^{2}\right) \lambda_{l_{2}}^{2}+\cdots+\sum_{l_{K}=1}^{L_{K}}\left(A_{K} y_{l_{K}}^{K}\right) \lambda_{l_{K}}^{K}=b \quad: \pi \\
& \sum_{l_{2}=1}^{L_{2}} \lambda_{l_{2}}^{2} \quad=1 \quad: \gamma_{1} \\
& \sum_{l_{K}=1}^{L_{K}} \lambda_{l_{K}}^{K}=1 \quad: \gamma_{K}  \tag{3}\\
& \sum_{l_{2}=1}^{L_{2}} y_{l_{2}}^{2} \lambda_{l_{2}}^{2} \quad \in \mathbb{Z}_{n_{2}}^{+} \\
& \sum_{l_{K}=1}^{L_{K}} y_{l_{K}}^{K} \lambda_{l_{K}}^{K} \in \mathbb{Z}_{n_{K}}^{+} \\
& x_{1} \in \mathbb{Z}_{n_{1}}^{+}, \lambda^{2} \in[0,1]_{L_{2}}, \ldots, \lambda^{K} \quad \in[0,1]_{L_{K}}
\end{align*}
$$

The RMP provides the optimal solution $x_{1}^{*}, x_{2}^{*}, \ldots, x_{K}^{*}$ to the original problem (1) if the necessary subproblem solutions are present in the RMP. That is, if $y_{l_{k}}^{k, *}, l_{k}=$ $1, \ldots, L_{k}, k=2, \ldots K$ such that $x_{k}^{*}=\sum_{l_{k}=1}^{L_{k}} \lambda_{l_{k}}^{k} y_{l_{k}}^{k, *}, k=2, \ldots, K$ have been included.

Given that $x_{k}^{*}, k=1, \ldots, K$ are not known a priori, column generation starts with an initial solution consisting of $x_{1}$ and initial sets of subproblem solutions. "Useful" subproblem solutions, that form columns for the RMP, are found by looking for subproblem solutions that provide columns with negative reduced cost. The reduced cost of a solution $y_{l_{k}}^{k}$ 's column, i.e., the reduced cost for $\lambda_{l_{k}}^{k}$, is given by $c_{k}^{\top} y_{l_{k}}^{k}-\pi^{\top} A_{k} y_{l_{k}}^{k}-\gamma_{k}$. To find a solution with minimum reduced cost we can solve:
$\mathcal{S}_{k}: \min \quad\left(c_{k}-\pi^{\top} A_{k}\right)^{\top} x_{k}-\gamma_{k} \quad$ (reduced cost for corresponding $\lambda^{k}$ ) subject to $\quad F_{k} x_{k}=f_{k} \quad$ (ensures that $y^{k}$ solves subproblem $k$ )

$$
\begin{equation*}
x_{k} \in \mathbb{Z}_{n_{k}}^{+} \tag{4}
\end{equation*}
$$

If the objective value of $\mathcal{S}_{k}$ is less than 0 , then the solution $y^{k}$ will form a column in the RMP whose inclusion in the basis would improve the objective value of the RMP. The solution $y^{k}$ is added to the set of solution used in the RMP. There are other mechanisms for managing the sets of solutions present in DIP, but they are beyond the scope of this paper.

Within DIP, hence Dippy, the RMP and relaxed problems $S_{k}, k=2, \ldots, K$ are not specified explicitly. Rather, the constraints for each subproblem $F_{k} x_{k}=f_{k}$ are specified by using the .relaxation [ $j$ ] syntax. DIP then automatically constructs the RMP and the relaxed problems $S_{k}, k=2, \ldots, K$. The relaxed subproblems
$S_{k}, k=2, \ldots, K$ can either be solved using the default MILP solver (CBC) or a customised solver. A customised solver can be defined by the relaxed_solver function. This function has 4 inputs:

1. prob - the DipProblem being solved;
2. index - the index $k$ of the subproblem being solved;
3. redCosts - the reduced costs for the $x_{k}$ variables $c_{k}-\pi^{\top} A_{k}$;
4. convexDual - the dual value for the convexity constraint for this subproblem $\gamma_{k}$.
In addition to subproblem solutions generated using RMP dual values, initial columns for subproblems can also be generated either automatically using CBC or using a customised approach. A customised approach to initial variable generation can be defined by the init_vars function. This function has only 1 input, prob, the DipProblem being solved.

Starting from the original capacitated facility location problem from section 3:

$$
\begin{array}{lcl}
\min & \sum_{i=1}^{m} w_{i} \\
\text { s.t. } & \sum_{i=1}^{m} x_{i j}=1, j=1, \ldots, n \quad \text { (each product produced) } \\
& \sum_{j=1}^{n} r_{j} x_{i j}+w_{i}=C y_{i}, i=1, \ldots, m & \text { (aggregate capacity at location } i \text { ) } \\
x_{i j} \leq y_{i}, i=1, \ldots, m, j=1, \ldots, n & (\text { disaggregate capacity at location } i) \\
& x_{i j} \in\{0,1\}, w_{i} \geq 0, y_{i} \in\{0,1\}, i=1, \ldots, m, j=1, \ldots, n
\end{array}
$$

we can decompose this formulation:

$$
\begin{aligned}
& \min \quad 1 w_{2} \cdots \quad+1 w_{m} \\
& \text { s.t. } I \mathrm{x}_{2} \quad \cdots+I \mathbf{x}_{m} \quad=1 \text { (each product produced) } \\
& r^{\top} \mathbf{x}_{2}-C y_{2}+1 w_{2} \quad=0 \text { (aggregate cap. at loc. } 2 \text { ) } \\
& I \mathbf{x}_{2}-e y_{2} \quad \leq 0 \text { (disaggregate cap. at loc. 2) } \\
& \begin{array}{l}
r^{\top} \mathbf{x}_{m}-C y_{m}+1 w_{m}=0 \text { (aggregate cap. at loc. } \mathrm{K} \text { ) } \\
+I \mathbf{x}_{m}-e y_{m} \leq 0 \text { (disaggregate cap. at loc. } \mathrm{K} \text { ) }
\end{array}
\end{aligned}
$$

where

$$
\mathbf{x}_{i}=\left(\begin{array}{c}
x_{i 1} \\
\vdots \\
x_{i n}
\end{array}\right), r=\left(\begin{array}{c}
r_{1} \\
\vdots \\
r_{n}
\end{array}\right) \text { and } e=\left(\begin{array}{c}
1 \\
\vdots \\
1
\end{array}\right) .
$$

Now the subproblems $F_{k} x_{k}=f_{k}, k=2, \ldots, K$ are

$$
\left[\begin{array}{ccc}
r^{\top} & -C & 1 \\
I & e &
\end{array}\right]\left[\begin{array}{c}
\mathbf{x}_{i} \\
y_{i} \\
w_{i}
\end{array}\right]=\left[\begin{array}{l}
0 \\
0
\end{array}\right]
$$

$$
c_{k}^{\top}=[0|0| 1], A_{k}=[I|0| 0],
$$

so $\mathcal{S}_{k}$ becomes

$$
\begin{array}{rcc}
\mathcal{S}_{i}: \min & \sum_{j=1}^{n}-\pi_{j} x_{i j} & +1 w_{i}-\gamma_{i} \\
\text { subject to } & \sum_{j=1}^{n} r_{j} x_{i j}-C y_{i} & +1 w_{i}=0 \\
& x_{i j}-y_{i} & \leq 0, j=1, \ldots, n \\
& x_{i j} . & u_{i}, \in\{0,1\}, j=1
\end{array}
$$

where $\pi_{j}$ is the dual variable for the assignment constraint for product $j$ in the RMP.

In Dippy, we define subproblems for each facility location using the .relaxation syntax for the aggregate and disaggregate capacity constraints:

```
# Aggregate capacity constraints
for i in LOCATIONS:
    prob.relaxation[i] += lpSum(assign_vars[(i, j)] * REQUIREMENT[j]
        for j in PRODUCTS) + waste_vars[i] \
                                == CAPACITY * use_vars[i]
# Disaggregate capacity constraints
for i in LOCATIONS:
    for j in PRODUCTS:
        prob.relaxation[i] += assign_vars[(i, j)] <= use_vars[i]
```

All remaining constraints (the assignment constraints that ensure each product is assigned to a facility) form the master problem when using branch-price-and-cut. To use branch-price-and-cut we turn on the doPriceCut option:

```
dippy.Solve(prob, {
    'TolZero': '%s' % tol,
    'doPriceCut': '1',
    'generateInitVars': '1', })
```

Note that symmetry is also present in the decomposed problem, so we add ordering constraints (described in $\S 3.1$ ) to the RMP :

```
# Ordering constraints
for index, location in enumerate(LOCATIONS):
    if index > 0:
        prob += use_vars[LOCATIONS[index-1]] >= use_vars[location]
```

Using branch-price-and-cut, the RMP takes about ten times as long to solve as the original formulation, and has a search tree size of 37 nodes. The generateInitVars option uses CBC by default to find initial columns for the RMP and then uses CBC to solve the relaxed problems. Dippy lets us provide our own approaches to solving the relaxed problems and generating initial variables, which may be able to speed up the overall solution process.

In the relaxed problem for location $i$, the objective simplified to $\min \sum_{j=1}^{n}-\pi_{j} x_{i j}+$ $1 w_{i}-\gamma_{i}$. However, the addition of the ordering constraints and the possibility of a Phase I/Phase II approach in the MILP solution process to find initial variables mean that our method must work for any reduced costs, i.e., the objective
becomes $\min \sum_{j=1}^{n} d_{j} x_{i j}+f y_{i}+g w_{i}-\gamma_{i}$. Although the objective changes, the constraints remain the same. If we choose not to use a location, then $x_{i j}=y_{i}=w_{i}=0$ for $j=1, \ldots, n$ and the objective is $-\gamma_{i}$. Otherwise, we use the location and $y_{i}=1$ and add $f$ to the objective. The relaxed problem reduces to:

$$
\begin{aligned}
\min & \sum_{j=1}^{n} d_{j} x_{i j}+g w_{i}-\gamma_{i} \\
\text { subject to } & \sum_{j=1}^{n} r_{j} x_{i j}+1 w_{i}=C \\
& x_{i j}, \quad w_{i} \in\{0,1\}, j=1, \ldots, n
\end{aligned}
$$

However, the constraint ensures $w_{i}=C-\sum_{j=1}^{n} r_{j} x_{i j}$, so we can reformulate as:

$$
\begin{array}{rc}
\min & \sum_{j=1}^{n}\left(d_{j}-g r_{j}\right) x_{i j}+f C-\gamma_{i} \\
\text { subject to } & C-\sum_{j=1}^{n} r_{j} x_{i j} \geq 0 \Rightarrow \sum_{j=1}^{n} r_{j} x_{i j} \leq C \\
& x_{i j}, \in\{0,1\}, j=1, \ldots, n
\end{array}
$$

This is a 0-1 knapsack problem with "effective costs" costs for each product $j$ of $d_{j}-g r_{j}$. We can use dynamic programming to find the optimal solution.

In Dippy, we can access the problem data, variables and their reduced costs, so the 0-1 knapsack dynamic programming solution is straightforward to implement and use:

```
noEmpty = dict([(loc, True) for loc in LOCATIONS])
def solve_subproblem(prob, key, redCosts, convexDual):
    loc = key
    # Calculate effective objective coefficient of products
    effs= {}
    for j in PRODUCTS:
        effs[j] = redCosts[assign_vars[(loc, j)]] \
                - redCosts[waste_vars[loc]] * REQUIREMENT[j]
    avars = [assign_vars[(loc, j)] for j in PRODUCTS]
    obj = [-effs[j] for j in PRODUCTS]
    weights = [REQUIREMENT[j] for j in PRODUCTS]
    # Use 0-1 KP to max. total effective value of products at location
    z, solution = knapsack01(obj, weights, CAPACITY)
```

```
# Get the reduced cost of the knapsack solution and waste
rc = redCosts[use_vars[loc]] -z + \
        redCosts[waste_vars[loc]] * CAPACITY
waste = CAPACITY - sum(weights[i] for i in solution)
# Return the solution if the reduced cost is low enough ...
if rc - convexDual < -tol:
    var_values = dict([(avars[i], 1) for i in solution])
    var_values[use_vars[loc]] = 1
    var_values[waste_vars[loc]] = waste
    var_tuple = (waste, rc - convexDual, var_values)
    return [var_tuple]
# ... or an empty location is "useful"
elif noEmpty[loc] and (-convexDual < -tol):
    noEmpty[loc] = False
    var_values = {}
    var_tuple = (0.0, -convexDual, var_values)
    return [var_tuple]
return []
```

Adding this customised solver reduces the solution time because it has the benefit of knowing it is solving a knapsack problem rather than a general MILP.

To generate initial facilities (complete with assigned products) we implemented two approaches. The first approach used a first-fit method and considered the products in order of decreasing requirement:

```
def first_fit_heuristic():
    # Sort the items in descending weight order
    productReqs = [(REQUIREMENT[j],j) for j in PRODUCTS]
    productReqs.sort(reverse=True)
    # Add items to locations, fitting in as much
    # as possible at each location.
    allLocations = []
    while len(productReqs) > 0:
        waste = CAPACITY
        currentLocation = []
        j = 0
        while j < len(productReqs):
        # Can we fit this product?
        if productReqs[j][0] <= waste:
                currentLocation.append(productReqs[j][1]) # index
                waste -= productReqs[j][0] # requirement
                productReqs.pop(j)
            else:
                # Try to fit next item
                j += 1
        allLocations.append((currentLocation, waste))
# Return a list of tuples: ([products],waste)
return allLocations
```

```
def first_fit(prob):
    locations = first_fit_heuristic()
    bvs = []
    index = 0
    for loc in locations:
    i = LOCATIONS[index]
    var_values = dict([(assign_vars[(i, j)], 1) for j in loc[0]])
    var_values[use_vars[i]] = 1
    var_values[waste_vars[i]] = loc[1]
    dv = (loc[1], var_values)
    bvs.append((i, dv))
    index += 1
    return bvs
```

The second approach simply assigned one product to each facility:

```
def one_each(prob):
    bvs = []
    for index, loc in enumerate(LOCATIONS):
        lc = [PRODUCTS[index]]
        waste = CAPACITY - REQUIREMENT[PRODUCTS[index]]
        var_values = dict([(assign_vars[(loc, j)], 1) for j in lc])
        var_values[use_vars[loc]] = 1
        var_values[waste_vars[loc]] = waste
        dv = (waste, var_values)
        bvs.append((loc, dv))
    return bvs
```

Using Dippy we can define both approaches at once and then define which one to use by setting the init_vars method:

```
prob.init_vars = first_fit
##prob.init_vars = one_each
```

These approaches define the initial sets of subproblem solutions $y_{l_{k}}^{k}, l_{k}=1$, $\ldots, L_{k}, k=1, \ldots, K$ for the initial RMP before the relaxed problems are solved using the RMP duals.

The effect of the different combinations of column generation, customised subproblem solvers and initial variable generation methods, both by themselves and combined with branching, heuristics, etc are summarised in Table 1. For this size of problem, column generation does not reduce the solution time significantly (if at all). However, we show in section 4 that using column branching enables DIP (via Dippy and PuLP) to be competitive with state-of-the-art solvers.

### 3.4 Adding Customised Heuristics

To add user-defined heuristics in Dippy, we first define a new procedure for node heuristics, heuristics. This function has three inputs:

1. prob - the DipProblem being solved;
2. xhat - an indexable object representing the fraction solution at the current node;
3. cost - the objective coefficients of the variables.

Multiple heuristics can be executed and all heuristic solutions can be returned to DIP.

```
def heuristics(prob, xhat, cost):
    sols = []
    if prob.root_heuristic:
        prob.root_heuristic = False # Don't run twice
        sol = first_fit()
        sols.append(sol)
    if prob.node_heuristic:
        sol = frac_fit(xhat)
        sols.append(sol)
    return sols
prob.heuristics = heuristics
prob.root_heuristic = True
prob.node_heuristic = True
```

A heuristic that solves the original problem may not be as useful when a fractional solution is available, so we demonstrate two different heuristics here: a "first-fit" heuristic and a "fractional-fit" heuristic.

In the facility location problem, an initial allocation of production to locations can be found using the same first-fit heuristic that provided initial solutions for the column generation approach (see §3.3). The first-fit heuristic iterates through the items requiring production and the facility locations allocating production at the first facility that has sufficient capacity to produce the item. This can then be used to provide an initial, feasible solution at the root node within the customised heuristics function.

```
def first_fit():
    # Use generic first-fit heuristic that is shared
    # between heuristics and initial variable generation
    allLocations = first_fit_heuristic()
    # Convert to decision variable values
    sol= {}
    for i in LOCATIONS:
        for j in PRODUCTS:
            sol[assign_vars[(i, j)]] = 0
        sol[use_vars[i]] = 0
        sol[waste_vars[i]] = 0
    index = 0
    for loc in allLocations:
        i = LOCATIONS[index]
        sol[use_vars[i]] = 1
        sol[waste_vars[i]] = loc[1]
        for j in loc[0]:
            sol[assign_vars[(i, j)]] = 1
        index += 1
    return sol
```

At each node in the branch-and-bound tree, the fractional solution (provided by xhat) gives an indication of the best allocation of production. One heuristic approach to "fixing" the fractional solution is to consider each allocation (of an item's production to a facility) in order of decreasing fractionality and use a firstfit approach.

```
def frac_fit(xhat):
    # Initialise solution
    sol = {}
    for i in LOCATIONS:
    for j in PRODUCTS: sol[assign_vars[(i, j)]] = 0
    sol[use_vars[i]] = 0
    sol[waste_vars[i]] = 0
# Get the list of non-zero fractional assignments
fracAssigns = [ (xhat[assign_vars[(i, j)]], (i, j))
                                    for i in LOCATIONS for j in PRODUCTS
                                if xhat[assign_vars[(i, j)]] > tol ]
fracAssigns.sort()
# Track which products and locations have been used
notAllocated = dict((j,True) for j in PRODUCTS)
notUsed = dict((i,True) for i in LOCATIONS)
while len(fracAssigns) > 0:
    fracX = fracAssigns.pop() # Get best frac. assignment left
    (i,j) = fracX[1]
    if notAllocated[j]:
        if notUsed[i]: # Create a new location if needed
            notUsed[i] = False
            sol[use_vars[i]] = 1
            sol[waste_vars[i]] = CAPACITY
        if REQUIREMENT[j] <= sol[waste_vars[i]]: # Space left?
            sol[assign_vars[(i, j)]] = 1
            notAllocated[j] = False
            sol[waste_vars[i]] -= REQUIREMENT[j]
# Allocate the remaining products
unallocated = [(REQUIREMENT[j],j) for j in PRODUCTS
                                    if notAllocated[j]]
unallocated.sort(reverse=True)
unused = [i for i in LOCATIONS if notUsed[i]]
while len(unallocated) > 0:
    waste = CAPACITY
    index = 0
    loc = unused.pop()
    while index < len(unallocated):
        (j_req, j) = unallocated[index]
        if j_req <= waste:
            unallocated.pop(index)
            sol[assign_vars[(loc, j)]] = 1
            waste -= j_req
        else: index += 1
    sol[use_vars[loc]] = 1
    sol[waste_vars[loc]] = waste
return sol
```

Running the first-fit heuristic before starting the branching process has little effect on the solution time and does not reduce the number of nodes. Adding the first-fit heuristic guided by fractional values increases the solution time slightly
and the number of nodes remains at 419. The reason this heuristic was not that helpful for this problem instance is that:

- the optimal solution is found within the first 10 nodes without any heuristics, so the heuristic only provides an improved upper bound for $<10$ nodes;
- the extra overhead of the heuristic at each node increases the solution time more than any decrease from exploring fewer nodes.


### 3.5 Combining Techniques

The techniques and modifications of the solver framework can be combined to improve performance further. Table 1 shows that it is possible to quickly and easily test many approaches for a particular problem, including combinations of approaches ${ }^{2}$. Looking at the results shows that the heuristics only help when the size of the branch-and-bound tree has been reduced with other approaches, such as ordering constraints and advanced branching. Approaches for solving this problem that warrant further investigation use column generation, the customised solver and either ordering constraints or the first-fit heuristic to generate initial variables. Tests with different data showed that the solution time for branch-price-and-cut doesn't increase with problem size as quickly as for branch-and-cut, so the column generation approaches are worth considering for larger problems.

## 4 Performance and Conclusions

In section 3 we showed how Dippy works in practice by making customisations to the solver framework for an example problem. We will use the Wedding Planner problem from the PuLP documentation [3] to compare Dippy to two leading solvers that utilise branch-and-cut: the open-source CBC and the commercial Gurobi. This particular problem is useful for comparing performance because it has a natural column generation formulation and can be scaled-up in a simple way, unlike the Facility Location problem which is strongly dependent on the specific instance being tested.

The Wedding Planner problem is as follows: given a list of wedding attendees, a wedding planner must come up with a seating plan to minimise the unhappiness of all of the guests. The unhappiness of guest is defined as their maximum unhappiness at being seated with each of the other guests at their table, making it a pairwise function. The unhappiness of a table is the maximum unhappiness of all the guests at the table. All guests must be seated and there is a limited number of seats at each table.

This is a set partitioning problem, as the set of guests $G$ must be partitioned into multiple subsets, with the members of each subset seated at the same table.

[^2]The cardinality of the subsets is determined by the number of seats at a table and the unhappiness of a table can be determined by the subset. The MILP formulation is:
$x_{g t}= \begin{cases}1 & \text { if guest } g \text { sits at table } t \\ 0 & \text { otherwise }\end{cases}$
$u_{t}=$ unhappiness of table $t$
$S=$ number of seats at a table
$U(g, h)=$ unhappiness of guests $g$ and $h$ if they are seated at the same table

$$
\min \begin{aligned}
\sum_{t \in T} u_{t} & \text { (total unhappiness of the tables) } \\
\sum_{g \in G} x_{g t} & \leq S, t \in T \\
\sum_{t \in T} x_{g t} & =1, g \in G \\
u_{t} & \geq U(g, h)\left(x_{g t}+x_{h t}-1\right), t \in T, g<h \in G
\end{aligned}
$$

Since DIP, and thus Dippy, doesn't require a problem to be explicitly formulated as a Dantzig-Wolfe decomposition, a change from DIP to CBC is trivial. The only differences are that:

1. A LpProblem is created instead of a DipProblem;
2. No .relaxation statements are used;
3. The LpProblem. solve method uses CBC to solve the problem.

To see if CBC and Gurobi would perform well with a column-based approach, we also formulated a problem equivalent to the restricted master problem from the branch-price-and-cut approach and generated and added all possible columns before the solving the MILP. Finally we used to Dippy to develop a customised solver and initial variable generation function for the branch-price-and-cut formulation in DIP. In total, six approaches were tested on problem instances of increasing size:

1. CBC called from PuLP;
2. CBC called from PuLP using a columnwise formulation and generating all columns a priori;
3. Gurobi called from PuLP;
4. Gurobi called from PuLP using a columnwise formulation and generating all columns a priori;
5. DIP called from Dippy using branch-price-and-cut without customisation;
6. DIP called from Dippy using customised branching, cuts and column generation callback functions.

In Table 2 and Figure 2 we see that ${ }^{3}$ :

- Gurobi is fastest for small problems;
- The symmetry present in the problem means the solution time of CBC and Gurobi for the original problem deteriorate quickly;
- The time taken to solve the columnwise formulation also deteriorates, but at a lesser rate than when using CBC or Gurobi on the original problem;
- Both DIP and customised DIP solution times grow at a lesser rate than any of the CBC/Gurobi approaches;
- For large problems, DIP becomes the preferred approach.

The main motivation for the development of Dippy was to alleviate obstacles to experimentation with and customisation of advanced MILP frameworks. These obstacles arose from an inability to use the description of a problem in a high-level modelling languag integrated with the callback functions in leading solvers. This is mitigated with Dippy by using the Python-based modelling language PuLP to describe the problem and then exploiting Python's variable scoping rules to implement the callback functions.

Using the Capacitated Facility Location problem we have shown that Dippy is relatively simple to experiment with and customise, enabling the user to quickly and easily test many approaches for a particular problem, including combinations of approaches. In practice Dippy has been used successfully to enable final year undergraduate students to experiment with advanced branching, cut generation, column generation and root/node heuristics. The Wedding Planner problem shows that Dippy can be a highly competitive solver for problems in which column generation is the preferred approach. Given the demonstrated ease of the implementation of advanced MILP techniques and the flexibility of a high-level mathematical modelling language, this suggests that Dippy is effective as more than just an experimental "toy" or educational tool. It enables users to concentrate on furthering Operations Research knowledge and solving hard problems instead of spending time worrying about implementation details. Dippy breaks down the barriers to experimentation with advanced MILP approaches for both practitioners and researchers.

[^3]

Figure 2: Comparing solver performance on the Wedding Planner problem. In this figure the times for generating the columns for "CBC with columns" and "Gurobi with columns" have been included in the total solve time. The time required for solving the original formulation sharply increases for both Gurobi and CBC (marked with crosses) but at different problem sizes. However the time for the column-wise formulation is similar for Gurobi and CBC. The time for DIP does not smoothly increase with problem size, but is consistently lower than Gurobi for instances with 16 or more guests.

## 5 Acknowledgments

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| Strategies | Time (s) | Nodes |
| :--- | :--- | :--- |
| Default (branch and cut) | 0.26 | 419 |
| + ordering constraints (OC) | 0.05 | 77 |
| + OC \& advanced branching (AB) | 0.01 | 3 |
| + weighted inequalities (WI) | 0.34 | 77 |
| + WI \& OC | 0.17 | 20 |
| + WI \& OC \& AB | 0.06 | 4 |
| + first-fit heuristic (FF) at root node | 0.28 | 419 |
| + FF \& OC | 0.05 | 77 |
| + FF \& OC \& AB | 0.01 | 3 |
| + FF \& WI | 0.36 | 77 |
| + FF \& WI \& OC | 0.14 | 17 |
| + FF \& WI \& OC \& AB | 0.05 | 3 |
| + fractional-fit heuristic (RF) at nodes | 0.28 | 419 |
| + RF \& OC | 0.05 | 77 |
| + RF \& OC \& AB | 0.01 | 3 |
| + WI \& RF | 0.38 | 77 |
| + WI \& RF \& OC | 0.14 | 17 |
| + WI \& RF \& OC \& AB | 0.05 | 3 |
| + FF \& RF | 0.28 | 419 |
| + FF \& RF \& OC | 0.05 | 77 |
| + FF \& RF \& OC \& AB | 0.01 | 3 |
| + WI \& FF \& RF | 0.38 | 77 |
| + WI \& FF \& RF \& OC | 0.14 | 17 |
| + WI \& FF \& RF \& OC \& AB | 0.05 | 3 |
| + column generation (CG) | 2.98 | 37 |
| + CG \& OC | 2.07 | 23 |
| + CG \& OC \& AB | 0.56 | 10 |
| + CG \& Customised subproblem solver (CS) | 2.87 | 37 |
| + CG \& CS \& OC | 1.95 | 23 |
| + CG \& CS \& OC \& AB | 0.44 | 10 |
| + CG \& first-fit initial variable generation (FV) | 3.96 | 45 |
| + CG \& CS \& FV | 3.72 | 45 |
| + CG \& CS \& FV \& OC | 1.70 | 18 |
| + CG \& CS \& FV \& OC \& AB | 0.22 | 3 |
| + CG \& one-each initial variable generation (OV) | 3.40 | 41 |
| + CG \& CS \& OV | 3.33 | 41 |
| + CG \& CS \& OV \& OC | 2.23 | 24 |
| + CG \& CS \& OV \& OC \& AB | 0.27 | 3 |
|  |  |  |

Table 1: Experiments for the Capacitated Facility Location Problem

| \# guests | Time (s) |  |  |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  | CBC | CBC \& gen vars | olumns solve | Gurobi | Gurobi \& gen vars | columns solve | DIP | Customised DIP |
| 6 | 0.07 | 0.01 | 0.06 | 0.04 | 0.01 | 0.05 | 0.90 | 0.33 |
| 7 | 0.07 | 0.01 | 0.12 | 0.04 | 0.01 | 0.11 | 1.77 | 0.57 |
| 8 | 0.90 | 0.01 | 0.27 | 0.07 | 0.01 | 0.25 | 4.78 | 0.57 |
| 9 | 2.54 | 0.01 | 0.57 | 0.09 | 0.01 | 0.55 | 2.11 | 0.78 |
| 10 | 3.83 | 0.01 | 1.23 | 0.13 | 0.01 | 1.15 | 5.60 | 0.94 |
| 11 | 6.48 | 0.01 | 2.46 | 0.14 | 0.01 | 2.36 | 8.62 | 0.91 |
| 12 | 26.73 | 0.01 | 4.64 | 0.34 | 0.01 | 4.55 | 25.17 | 2.80 |
| 13 | 53.18 | 0.01 | 8.57 | 0.39 | 0.01 | 8.28 | 14.86 | 1.40 |
| 14 | 70.51 | 0.01 | 15.27 | 0.38 | 0.01 | 14.65 | 20.09 | 2.66 |
| 15 | 97.79 | 0.01 | 26.26 | 0.47 | 0.01 | 25.07 | 33.52 | 1.59 |
| 16 | $>1000$ | 0.01 | 43.86 | 68.08 | 0.01 | 42.11 | 26.73 | 2.09 |
| 17 | - | 0.01 | 72.07 | 79.71 | 0.01 | 68.87 | 50.48 | 3.92 |
| 18 | - | 0.01 | 115.64 | 96.03 | 0.01 | 110.52 | 40.80 | 4.67 |
| 19 | - | 0.01 | 181.39 | 113.01 | 0.01 | 173.13 | 78.20 | 2.64 |
| 20 | - | 0.02 | 283.16 | >6000 | 0.01 | 270.08 | 61.86 | 5.31 |
| 21 | - | 0.02 | 434.60 | - | 0.02 | 418.04 | 77.66 | 4.23 |
| 22 | - | 0.02 | 664.87 | - | 0.02 | 639.04 | 134.76 | 4.63 |
| 23 | - | - | >1000 | - | - | >1000 | 149.82 | 9.16 |
| 24 | - | - | - | - | - | - | 110.24 | 6.51 |
| 25 | - | - | - | - | - | - | 202.59 | 8.80 |
| 26 | - | - | - | - | - | - | 185.21 | 18.47 |

Table 2: Experiments for the Wedding Planner Problem


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[^1]:    ${ }^{1}$ The skeleton for a fourth method (branch, relax and cut) exists in DIP, but this method is not yet implemented.

[^2]:    ${ }^{2}$ All tests were run using Python 2.7 .1 on a Windows 7 machine with an Intel Core 2 Duo T9500@2.60GHz CPU

[^3]:    ${ }^{3}$ All tests were run using Python 2.7.1 on a Dell XPS1530 laptop with an Intel Core 2 Duo CPU T9500@2.60GHz and 4 GB of RAM. We used CBC version 2.30.00, Gurobi version 4.5.1, and Dippy version 1.0.10.

