Parallel algebraic modeling for stochastic optimization

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Abstract—We present scalable algebraic modeling software, StochJuMP, for stochastic optimization as applied to power grid economic dispatch. It enables the user to express the problem in a high-level algebraic format with minimal boilerplate. StochJuMP allows efficient parallel model instantiation across nodes and efficient data localization. Computational results are presented showing that the model construction is efficient, requiring less than one percent of solve time. StochJuMP is configured with the parallel interior-point solver PIPS-IPM but is sufficiently generic to allow straightforward adaptation to other solvers.

Keywords—optimization, parallel programming, high performance computing, mathematical model, Power system modeling, scalability

I. INTRODUCTION

Algebraic modeling languages (AMLs) for optimization are widely used by both academics and practitioners for specifying optimization problems in a human-readable, mathematical form, which is then automatically translated by the AML to the low-level formats required by efficient implementations of optimization algorithms, known as solvers. For example, a classical family of optimization problems is linear programming (LP), that is, minimization of a linear function of decision variables subject to linear equality and inequality constraints. As input, LP solvers expect vectors \( c, l, u, L, U \), and \( U \) and a sparse constraint matrix \( A \) defining the LP problem,
\[
\begin{align*}
\text{minimize} & \quad c^T x \\
\text{subject to} & \quad L \leq Ax \leq U \\
& \quad l \leq x \leq u,
\end{align*}
\]
where inequalities are componentwise.

In practice, it is tedious and error-prone to explicitly form a sparse constraint matrix and corresponding input vectors; this process often involves concatenating different classes of decision variables, each indexed over multidimensional symbolic sets or numeric ranges, into a single \( x \) decision vector. Instead, AMLs, which may be considered domain-specific languages in the vocabulary of computer science, handle these transformations automatically.

Notable AMLs include AMPL [1], GAMS [2], YALMIP [3], Pyomo [4], and CVX [5]. While commercial AMLs (AMPL and GAMS) offer standalone environments, open-source AMLs are typically embedded in high-level dynamic languages; CVX and YALMIP are both toolboxes for MATLAB, and Pyomo is a package for Python. Indeed, dynamic languages provide convenient platforms for AMLs, and domain-specific languages in general, because they make development easier (no need for a customized parser), and promote the ease of use by being accessible from a language that is already familiar to users and has its own tools for processing and formatting data.

Historically, speed has been a trade-off for using AMLs embedded in high-level dynamic languages, primarily because of their extensive use of operator overloading within an interpreted language. These AMLs may be orders of magnitude slower at generating the optimization model before passing it to a solver; in some reasonable use cases, this model generation time may perversely exceed the time spent in the solver.

The Julia programming language [6] presented an opportunity to address this performance gap. Lubin and Dunning developed JuMP [7], an AML embedded in Julia. By exploiting Julia’s metaprogramming and just-in-time compilation functionality, they achieved performance competitive with that of commercial AMLs (AMPL and Gurobi/C++) and an improvement of one to two orders of magnitude over open-source AMLs (Pyomo and PuLP [8]) in model generation time for a benchmark of LP problems.

In this paper, we present an extension to JuMP, called StochJuMP, for modeling a class of stochastic optimization problems, namely, two-stage stochastic optimization with recourse, a popular paradigm for optimization under uncertainty [9]. These problems are computationally challenging and at the largest scale require the use of specialized solvers employing distributed-memory parallel computing. These solvers require structural information not provided by standard AMLs, and because of memory limits, it may not be feasible to build up the entire instance in serial and then distribute the pieces. Instead, StochJuMP generates the model in parallel according to the data distribution across processes required by the specialized solvers. We present numerical results demonstrating that StochJuMP can effectively generate these structured models in a small fraction (≈ 1.5%) of the solution time. To our knowledge, these results are among the first reports of employing Julia on a moderately sized MPI-based cluster.
II. BLOCK-ANGULAR STRUCTURE

The structured optimization problems considered in this paper are known in the optimization community as dual block angular problems, which means that they have separable objective function and block angular, or half-arrow shaped constraints. In particular, we consider dual block angular problems with quadratic objective function and affine constraints, which can be mathematically expressed as

\[
\begin{align*}
\text{min} & \quad \frac{1}{2} x^T Q_0 x_0 + c_0^T x_0 + \sum_{i=1}^{N} \left( \frac{1}{2} x_i^T Q_i x_i + c_i^T x_i \right) \\
\text{s.t.} & \quad A x_0 = b_0, \\
& \quad T_i x_0 + W_1 x_1 = b_1, \\
& \quad T_2 x_0 + W_2 x_2 = b_2, \\
& \quad \vdots \\
& \quad T_N x_0 + W_N x_N = b_N,
\end{align*}
\]

(1)

The decision variables are the vectors \( x_i, i = \{0, 1, \ldots, N\} \). The data of the problem consist of the coefficients vectors \( c_i \), right-hand sides \( b_i \), symmetric matrices \( Q_i \), and rectangular matrices \( A, T_i, W_i (i = \{1, \ldots, N\}) \).

Problems of the form (1) arise in stochastic optimization with recourse. Stochastic programming problems with recourse (in (2) we show two-stage quadratic problems [10]) provide optimal decisions to be made now that minimize the expected cost in the presence of future uncertain conditions:

\[
\begin{align*}
\text{min} & \quad \frac{1}{2} x_0^T Q_0 x_0 + c_0^T x_0 + \mathbb{E}_\xi[G(x_0, \xi)] \\
\text{s.t.} & \quad A x_0 = b_0, x_0 \geq 0.
\end{align*}
\]

(2)

The recourse function \( G(x_0, \xi) \) is defined by

\[
G(x_0, \xi) = \min_y \frac{1}{2} y^T Q_\xi y + c_\xi^T y \\
\text{s.t.} \quad T_\xi x_0 + W_\xi y = b_\xi, y \geq 0.
\]

(3)

and the expected value \( \mathbb{E}[\cdot] \), which is assumed to be well defined, is taken with respect to the random variable \( \xi \), which contains the data \( Q_\xi, c_\xi, T_\xi, W_\xi, b_\xi \). Some of the elements in \( \xi \) can be deterministic. The matrix \( Q_\xi \) is assumed to be symmetric for all possible \( \xi \), \( W_\xi \), the recourse matrix, and \( T_\xi \), the technology matrix, are random matrices. The symmetric matrix \( Q_0 \), the matrix \( A_0 \), and the linear coefficients \( c_0 \) are deterministic. The variable \( x_0 \) is called the first-stage decision, which is a decision to be made now. The second-stage decision \( y \) is a recourse or corrective decision that one makes in the future after some random event occurs.

The recourse problems (2) take the form of the dual-block angular problems (1) when the probability distribution of \( \xi \) has finite support. When \( \xi \) does not have finite support, problems (1) arise from the sample average approximation (SAA) method, in which a sample \( (\xi_1, \xi_2, \ldots, \xi_N) \) is used to estimate \( \mathbb{E}_\xi[G(x_0, \xi)] \approx \frac{1}{N} \sum_{i=1}^{N} G(x_0, \xi_i) \) and the minimization and expectation operators are commuted [11]. In this case, the data \( (Q_i, c_i, T_i, W_i, b_i) \) of (1), \( i = \{1, \ldots, N\} \), correspond to the sample \( (\xi_1, \xi_2, \ldots, \xi_N) \) used to approximate (2), and \( x_i \) correspond to the decision \( y \) of the recourse problem (3) given by the sample \( \xi_i, i = \{1, \ldots, N\} \).

While the modeling framework presented in this paper can be used to specify any quadratic dual block angular problem (1), the optimization solver handles only convex problems; thus, the matrices \( Q_i, i = \{0, 1, \ldots, N\} \), are required to be positive semidefinite. In addition, the interior-point algorithm used by the solver requires the matrices \( A, W_1, \ldots, W_N \) to have full row rank.

III. PIPS-IPM

Solving practically sized instances of (1) requires the use of memory-distributed computing environments and specialized optimization solvers capable of efficiently exploiting the dual block angular structure. One such solver, PIPS-IPM, developed in the past few years by some of the authors of this paper, achieves data parallelism inside the numerical Mehrotra predictor-corrector optimization algorithm [12] by distributing the data and computations required by the optimization across computational nodes. The data distribution is done by partitioning the scenarios and assigning a partition to a computational node. The first-stage data are replicated on all nodes to avoid communication overhead. The computations follow a similar distribution scheme: the intrascenario computations (factorizations of sparse matrices, solving with the factors, matrix-matrix and matrix-vector products, and vector-vector operations) are performed in parallel, independently for each scenario, while the computations corresponding to the first-stage (factorizations of dense matrices, solve with the factors and matrix-vector and vector-vector operations) are replicated across all computational nodes [13].

Several algorithmic and implementation developments were targeted at reducing the time to solution and increasing the parallel efficiency of PIPS-IPM. In particular, the augmented incomplete factorization approach [14] considerably reduced the time to solution by making better use the cores inside each node and by fully exploiting the sparsity of the scenario data. Additional implementation developments, such as hardware-tuned communication and adoption of GPUs in the dense first-stage calculations [15], enabled real-time solutions of stochastic economic dispatch problems with very good parallel scaling on a variety of HPC platforms: Cray XK7, Cray XC30, IBM BG/P, and IBM BG/Q [15]. However, the algebraic specification and instantiation of the economic dispatch problems were done serially and required considerably longer time than the time spent in the optimization process. Removing this capability limitation and being able to process and instantiate the optimization models in times that are negligible, that is, a few percent of total solution time, are the main motivations behind the existing work.

IV. STOCHJUIMP

StochJuMP is an extension built on top of the JuMP algebraic modeling language [7]. Extending AMPLs to deal with structured problems such as multistage stochastic optimization presents additional challenges for expressing problem structure, both technically and conceptually. Conceptually, one must choose a syntax that is clean and offers an intuitive, high-level encapsulation of the problem. From a technical perspective, efficiently handling the highly structured problem in a parallelized setting requires care.

The SML project [16] is an extension built on top of AMPL for conveying multistage structure to solvers by using a new block keyword. SML is implemented as a pre- and
post-processor to AMPL itself; as a result, the subproblems must be constructed as intermediary AMPL problem files on disk. The more recent work on PSMG [17] parses SML files and provides parallel model instantiation, avoiding the memory bottleneck that can arise when attempting to construct massive problem instances on a single node. While PSMG aims to address the same technical challenge as StochJuMP, we emphasize that PSMG is a large C++ project that required considerable development effort, based on our discussions with the authors. The development of StochJuMP, on the other hand, was greatly accelerated by being written in a high-level dynamic language at no significant penalty in total solution time.

We will describe details of the implementation and the use of StochJuMP. Throughout, we will refer to the code in Listing 1, which is the entirety of the Illinois model used in the computational results section. For brevity, we omit the somewhat tedious code that reads the problem data from file into memory.

JuMP represents all data required to describe an optimization problem in a Model data type in Julia; StochJuMP extends this by appending two fields: parent, a reference to the parent block, and children, a vector of references to children blocks. Since each block is a bona fide Model object, all methods to describe variables and constraints apply to blocks in a JuMPStoch model. Additionally, using Julia’s scoping rules, it is possible to structure the model specification such that a child block can include variables from parent blocks in constraints. This allows the description of arbitrary nested block structure in the model, all within a very lightweight extension to the existing JuMP infrastructure.

As a brief primer on JuMP and StochJuMP syntax, we discuss the functions and macros that appear in the example code in Listing 1. The StochasticModel(NS) method defines a stochastic model container object with NS scenarios. Similarly, the StochasticBlock(m) method is a thin wrapper around a Model constructor, specifying that m is the parent model.

The macro @defVar defines a variable or dictionary of variables, attached to a particular model, for use in describing the problem constraints and objective. The first argument is the model object; the second argument is a description of the variable name, the index set used for indexing into the dictionary of variables, and appropriate variable bounds. For instance,

```julia
@defVar(m, 0 <= Pgen_f[i=GENTHE] <= np_capThe[i])
```

constructs a dictionary named Pgen_f, attached to the model m, for which indexing is defined by the iterable object GENTHE. The lower bound is zero for every component, and the ith entry has an upper bound from the ith entry in the array np_capThe.

The @addConstraint macro adds a set of constraints to a specified model. For instance,

```julia
@addConstraint(bl, t_w_con2[g=GENWIN],
    tw[g] == gen_cost_win[g] * PgenWin_f[f[g]])
```

adds a set of constraints to the model bl. The second argument specifies a constraint identifier that is indexed into in the same way as in @defVar. The third argument takes an algebraic description of constraints to be added. Similarly, the @setObjective macro takes an algebraic description of the objective function, as well as an associated Model object and a value specifying whether the model is a minimization or maximization problem.

Again, we stress that minimal additions to JuMP have been made for this feature set: no new types have been introduced. A series of Model objects has been constructed with corresponding constraints (containing variables owned by other Models), and the hierarchical structure of the model is disentangled immediately prior to passing the problem data to PIPS-IPM. Furthermore, the nested structure can be arbitrarily more complex than that of the two-stage example illustrated here.

The @second_stage macro abstracts data localization in the model specification, allowing concise, rank-agnostic code to be distributed across the cores. The first argument m specifies the Model object to which to add the second stage structure. The second argument node is a global index for the particular scenario assigned to a particular node. More specifically, the macro expands to construct the global indices assigned to a particular compute node, and wraps the body of @second_stage (denoted with the begin and end delimiters) in a loop over those particular values. In this example, the loop index is node.

Since the blocks in a StochJuMP model are specified with JuMP models themselves, one can easily adapt the existing JuMP functions to construct the problem data needed by a particular solver. The repository for StochJuMP is available at https://github.com/joehuchette/StochJuMP.jl. All told, the code to specify nested model structure takes less than 100 lines of Julia code, whereas the interface to PIPS-IPM requires roughly 300 lines.

V. COMPUTATIONAL RESULTS

We present here computational results and analyze the efficiency of the construction of a model arising in the optimization of power grid.

A. Model

To test StochJuMP, we consider the economic wind dispatch model presented in [18], which we briefly describe here. The model attempts to capture the impact that wind supply correlation information has on economic dispatch, a problem solved in real-time by power grid operators in order to set market prices. The model uses data describing the power transmission grid in the state of Illinois. The model problem is as follows.
Listing 1. An implementation in StochJuMP of the stochastic economic dispatch model described in Section V-A

\[ \min_{x, X(\omega)} \sum_{i \in G} (p_i x_i \mathbb{E}_\omega [p_i^+(X_i(\omega) - x_i) + p_i^-(X_i(\omega) - x_i)]) \]

subject to

\[ \tau_n(f) + \sum_{i \in G(n)} x_i = d_n, \quad \forall n \in \mathcal{N} \]
\[ \tau_n(F(\omega)) - \tau_n(f) + \sum_{i \in G(n)} (X_i(\omega) - x_i) = 0, \quad \forall n \in \mathcal{N}, \omega \in \Omega \]
\[ \mathcal{F}(f, \omega) = \mathcal{U}, \omega \in \Omega \]
\[ (x_i, X_i(\omega)) \in \mathcal{C}(\omega), i \in G, \omega \in \Omega \]

For this model, \( N \) is the number of nodes or buses, \( L \) is the set of transmission lines, \( C \) is the set of buses, and \( \mathcal{G} \) is the set of all energy suppliers. We use the subset \( \mathcal{G}(n) \) to denote all those providers connected to a given node \( n \). The forward dispatch values are \( x_i \) for each provider, and their spot quantity for a given scenario \( \omega \) is \( X_i(\omega) \). Forward power flow through a line \( \ell \) is \( f_\ell \). The demand for each node is \( d_n \), which is assumed to be deterministic and inelastic. The function \( \tau_n \) maps the flow vector to a node \( n \). We use \( v_1(n) \) and \( v_0(n) \) to denote the inflow and outflow lines, respectively, to node \( n \). Bid prices are denoted by \( p_i \), and \( p_i^+ \) and \( p_i^- \) denote bid prices for real-time corrections. More explicitly, supplier \( i \) is capable of selling additional power at price \( p_i^+ > p_i \) or of buying power at \( p_i < p_i^- \). The random scenarios \( \omega \) represent the randomness in the model and live in a probability space \((\Omega, \mathcal{F}, \mathbb{P})\). Using standard notation, \((z)_+ := \max z, 0 \) and \((z)_- := \min -z, 0 \).

Here, \( \mathcal{U} \) is a polyhedral set that constrains the maximum flow constraints on individual lines.

B. Weak-scaling experiment

We present a weak-scaling study on Blues, a cluster at Argonne National Laboratory with 310 compute nodes, each with two Sandy Bridge 2.6 GHz Pentium Xeon, for a total of 4,960 cores. The PIPS-IPM interior-point solver is used, with MA57 [19] as the underlying sparse linear algebra library.
The weak scaling study runs from 4 to 2,048 scenarios. The data for the first scenario are from empirical measurement, and the remaining scenarios are artificially generated from normally distributed perturbations of the first. We compare the model load time, modeling time, and solver computation time in Table I: explanations of the interpretations of these timing blocks are detailed below. Note that we use MPI barrier commands at the end of each timing block to ensure representative results.

Because of current limitations in Julia, external packages loaded in a script (e.g., StochJuMP) must be compiled before execution of a program, and this is recorded in the module load time field. As development of Julia continues, precompilation of packages will likely be introduced into the core language, removing this additional cost.

In order to force compilation of the appropriate methods, a common technique in Julia is the “JIT warm-up,” where a much smaller instance of a trial is performed first in order to ensure that all appropriate methods have been compiled by the just-in-time (JIT) compiler for a fair timing. Specifically, we perform a JIT warm-up run with a single scenario and abort just before calling PIPS-IPM to solve the problem. Note that this warm-up run includes reading and broadcasting data, building the model, and constructing the appropriate callbacks to pass to the solver. For reference, the JIT warm-up runs took roughly twice as long as the model generation time.

Model generation time records how long it takes for the internal representation to be built up in memory, as well as the transformations necessary to construct the data required by PIPS-IPM for the solve process. This is performed in parallel across the cluster, independently for each scenario.

Table II shows that model generation time takes at worst 1.5% of the solve time for all experiments in the weak scaling trial. Additionally, the model generation displays favorable scalable properties compared to the PIPS-IPM underlying solver. This timing includes the time it takes to read the problem data from file on a single core and broadcast it to all worker cores.

There are two regimes in the data presented: the small trials with 4 and 8 scenarios take roughly twice as long as larger instances across the timing metrics. We believe this result is attributable to the saturation of the 16 cores on each node. All experiments assigned one MPI process per core. Figure 1 shows this change in performance clearly, as well as the nice scaling behavior for StochJuMP.

VI. CONCLUSION

In this paper we presented StochJuMP, algebraic modeling software for multi-stage stochastic optimization. StochJuMP is built as an extension to JuMP, a fully-featured and performant algebraic modeling language embedded in the Julia programming language. It is implemented as a front-end for the PIPS-IPM solver and supports model instantiation in parallel. In a weak-scaling experiment we observe extremely good scaling and modeling times which are a very small fraction of solve times ($\approx 1.5\%$). In addition to performance, we emphasize the fast prototyping and development that made StochJuMP possible, due to Julia and the JuMP framework.

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\begin{table}[h]
\centering
\begin{tabular}{|c|ccc|}
\hline
N & Load Module & Generate Model & Solve \\
\hline
4  & 12.161 & 4.857 & 344.716 \\
8  & 13.661 & 5.234 & 369.796 \\
16 & 7.780  & 2.732 & 297.940 \\
32 & 7.570  & 2.732 & 297.940 \\
64 & 7.866  & 2.737 & 274.795 \\
128 & 12.486 & 2.770 & 360.695 \\
256 & 10.348 & 2.856 & 394.954 \\
512 & 13.163 & 3.157 & 458.392 \\
1024 & 13.056 & 3.414 & 888.038 \\
2048 & 13.568 & 3.521 & 464.705 \\
\hline
\end{tabular}
\caption{Weak scaling results (in seconds) for the Illinois power generation model.}
\end{table}

\begin{table}[h]
\centering
\begin{tabular}{|c|cc|}
\hline
N & Load Module & Generate Model \\
\hline
4  & 3.528 & 1.409 \\
8  & 3.694 & 1.415 \\
16 & 3.334 & 1.169 \\
32 & 2.541 & 0.917 \\
64 & 2.863 & 0.996 \\
128 & 3.462 & 0.768 \\
256 & 2.620 & 0.723 \\
512 & 2.871 & 0.689 \\
1024 & 1.470 & 0.384 \\
2048 & -      & 0.500 \\
\hline
\end{tabular}
\caption{Ratio of StochJuMP timings over solve time ($\times 100$).}
\end{table}

Fig. 1. Timing results for the weak-scaling trials.
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