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# Direct Verification of Linear Systems with over 10000 Dimensions (Experience Report)

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#### Abstract

We evaluate a recently-proposed reachability method on a set of high-dimensional linear system benchmarks taken from model order reduction and presented in ARCH 2016. The approach uses a state-set representation called a generalized star set and the principle of superposition of linear systems to achieve scalability. The method was previously shown to have promise in terms of scalability for direct analysis of large linear systems. For each benchmark, we also compare computing the basis matrix, a core part of the reachability method, using numerical simulations versus a matrix exponential formulation. The approach successfully analyzes systems with hundreds of dimensions in minutes, and can scale to systems that have over 10000 dimensions with a computation time ranging from tens of minutes to tens of hours, depending on the desired time step.

# 1 Introduction

Reachability analysis attempts to compute, given a set of initial states, the set of states a system can enter. Safety verification can then be done by performing intersections between the unsafe states and reachable set. In this paper, we examine reachability for affine, time-invariant systems, with system dynamics expressed as  $\dot{x} = Ax + c$ . We only consider the continuous post operation, that is, we do not evaluate guards or other features of a hybrid automaton [3]. Nonetheless, continuous post is a core component of many hybrid systems reachability methods.

When faced with a high-dimensional safety verification problem, a powerful approach is abstraction [5, 4, 10]. Abstraction methods try to simplify a complex system to a simpler one, such that properties of interest, such as safety, are preserved. For example, abstraction methods can reduce certain high-dimensional systems to ones with less dimensions [20]. The reason to do this is because there are scalability limits to verification approaches, for example, the number of dimensions that can be handled in a reasonable amount of time. The methods evaluated here are orthogonal to these abstraction approaches, and deal with the direct verification problem. Nonetheless, note that abstraction is still typically followed up by direct verification on

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a simplified model. Finally, it is common to combine abstraction and direct verification in a loop, using counter-examples from direct verification to guide the abstraction process [12]. To enable such counter-example abstraction refinement (CEGAR), the direct verification method must produce counter-example traces.

Reachable state set computation tools typically track states using data structures such as polyhedra [15], zonotopes [17], support functions [18], Taylor models [11], or combinations of these [2]. Recently, a data structure and associated reachability method was proposed which stores states in a data structure called a generalized star set [14]. The reachability method we evaluate is based on this technique, which essentially uses individual executions and superposition to compute the reachable states.

The implementation we used only reasons about the states reachable at discrete time steps. It does not look at states between time steps. However, it is capable of generating counterexample traces whenever the unsafe states are deemed reachable. We call this the *simulationequivalent reachable set*, since it consist of all the states visited by every fixed-step simulation.

Since the dynamics are time-invariant, however, if some external method is first used to bound the states reachable at any time within first time step, then this larger set can be used as the initial states with the unmodified technique to prove properties about the system at all times [13]. For example, for small time steps the system's Lipschitz constant may be used to produce such a bound on all the states reached between the first two time steps. This aspect of the reachability problem is not the focus of this paper.

We perform our evaluation using a benchmark suite consisting of nine large-scale linear systems [19]. These benchmarks were taken from "diverse fields such as civil engineering and robotics." The difficulty of these benchmarks was rated by the benchmark authors as "low" to "challenge", and the models range from tens of dimensions to over ten thousand dimensions. Using these benchmarks, we evaluate two approaches which can be used to compute a generalized star set's basis matrix, which is a key component when computing its reachability. This is discussed in the next section.

## 2 Computing a Star's Basis Matrix

Here, we discuss the computation of the basis matrix of a generalized star set with linear predicates (or simply star). We do not review the full reachability computation with this approach, which is available in earlier work [14].

The generalized star set approach for reachability computation takes advantage of the superposition principle of linear systems in order to compute the reachable set. At each instant in time, if one knows how each orthonormal unit vector in the standard basis has evolved since the initial time, one can reconstruct the state reachable from an arbitrary point. This is done by taking linear combinations of the states reached by the trajectories which originated at each unit vector.

For example, say in a 2-d space the initial point  $\begin{pmatrix} 1 \\ 0 \end{pmatrix}$  after some time t evolves under the linear differential equation  $\dot{x} = Ax$  and reaches point  $\begin{pmatrix} a \\ b \end{pmatrix}$ . The reachable state of  $\begin{pmatrix} 2 \\ 0 \end{pmatrix}$  then could be easily computed by taking twice the state that  $\begin{pmatrix} 1 \\ 0 \end{pmatrix}$  reached, which is  $\begin{pmatrix} 2a \\ 2b \end{pmatrix}$ . Knowing how  $\begin{pmatrix} 1 \\ 0 \end{pmatrix}$  and  $\begin{pmatrix} 0 \\ 1 \end{pmatrix}$  evolve is sufficient to be able to reconstruct where any initial point has evolved to at time t. If the dynamics are affine,  $\dot{x} = Ax + c$ , then one more simulation is required from the origin, in addition to the n simulations that are performed on the linear part of the system  $\dot{x} = Ax$ .



Figure 1: Plot of states reached by any fixed-step simulation (simulation-equivalent reachable set) in the harmonic oscillator system, using a step size of  $\frac{\pi}{4}$  (left), and the star's LP formulation at time  $\frac{\pi}{4}$  (right). The values circled in red are the star's basis matrix at time  $\frac{\pi}{4}$ , which gets updated at each step.

If we are interested in where a set of states, rather than a single point, can evolve to at time t, we can construct a linear program (LP) that encodes the constraints on the initial conditions in addition to how each unit vector has evolved at time t. Linear conditions can also be added to encode unsafe state conditions, and the LP will then be feasible if and only if an unsafe state is reachable from some initial state.

At each time step, the value of t changes, and so "how each unit vector has evolved at time t" will change. There are n initial unit vectors, and each one reaches an n-dimensional point, so these can be combined to form what is called the star's  $n \times n$  basis matrix.

The computation of this basis matrix at each time step dominates the runtime of the safety verification method. On the proposed benchmarks, we will evaluate two methods to compute the basis matrix. First, however, we present an example of the star-based reachability approach.

**Harmonic Oscillator Example.** A 2-d harmonic oscillator has the dynamics  $\dot{x} = y$ ,  $\dot{y} = -x$ . Take the initial states to be  $x \in [-6, -5]$  and  $y \in [0, 1]$ . Simulations of this system rotate clockwise around the origin. The simulation-equivalent reachable sets, and the associated LP formulation at time step  $\frac{\pi}{4}$  is shown in Figure 1. If there was a linear unsafe error condition, it could be added to this LP. The LP would then be feasible if, and only if, unsafe states were reachable. The values in the basis matrix at time  $\frac{\pi}{4}$  are the red encircled values in the LP formulation in the figure. The basis matrix gets updated at each time step, while the rest of the constraints remain the same. The initial conditions are encoded in rows 3-6. The basis matrix at time 0 is the identity matrix,  $\begin{pmatrix} 1 & 0 \\ 0 & 1 \end{pmatrix}$ . After  $\frac{\pi}{4}$  time, the point which started at state  $\begin{pmatrix} 1 \\ 0 \end{pmatrix}$  evolves to  $\begin{pmatrix} 0.707 \\ -0.707 \\ 0.707 \end{pmatrix}$  and the point which started at  $\begin{pmatrix} 0 \\ 1 \end{pmatrix}$  goes to  $\begin{pmatrix} 0.707 \\ 0.707 \\ 0.707 \end{pmatrix}$ . The basis matrix at time  $\frac{\pi}{4}$  is therefore  $\begin{pmatrix} 0.707 \\ -0.707 \\ 0.707 \end{pmatrix}$ , as circled in red in the figure. At the next time step, time  $\frac{\pi}{2}$ , the basis matrix would be updated to  $\begin{pmatrix} 0 \\ -1 \\ 0 \end{pmatrix}$ . Error states could be encoded by adding additional rows to the LP constraints imposing a linear condition on x and y. If an error state is reachable, the LP solution would provide concrete assignments to  $x, y, \alpha_1$ , and  $\alpha_2$ . The corresponding error trace, then, starts at the point  $(\alpha_1, \alpha_2)$ , and, after evolving for t time, reaches the unsafe point (x, y).

As mentioned before, computing the basis matrix for each time step dominates the runtime of the safety verification method, and so it is important that its computation is optimized. In the original work [14], the computation of this basis matrix was done using simulations. However, notice that, since the solution to a linear system  $\dot{x} = Ax$  is  $x(t) = e^{At}x(0)$ , the basis matrix can also be computed using the matrix exponential. In particular the basis matrix at time t is equal to  $e^{At}$ . For example, in the harmonic oscillator system above, the dynamics matrix is  $A = \begin{pmatrix} 0 & 1 & 0 \\ -0.707 & 0.707 \end{pmatrix}$ , and the basis matrix at time  $\frac{\pi}{4}$  evaluates to  $e^{A(\pi/4)} = \begin{pmatrix} 0.707 & 0.707 \\ -0.707 & 0.707 \end{pmatrix}$ , which is the same as the numerical simulation approach.

Further, rather than recomputing the basis matrix at each step, notice that  $e^{A2t} = e^{At} \cdot e^{At}$ . Thus, a single computation of the matrix exponential can be performed at the desired time step, and then the resultant matrix can be multiplied using standard matrix multiplication to get the basis matrix at the next time step. This can be repeated until the time horizon is reached. In the next section, we compare these methods for computing the basis matrix on a set of benchmark systems.

### 3 Benchmark Results

The simulation-equivalent reachability method has been shown to have promise in terms of scalability [8, 9]. In this section, we evaluate the approach on a recently-proposed benchmark suite for linear systems [19], which includes a system with over 10000 dimensions. For each benchmark, we also considered a variant with a weakened or strengthened unsafe condition, so that each system would have both a safe and an unsafe case. In order to evaluate the accuracy of the method, when an unsafe state is reached, we output the initial state and final states extracted from the LP which are deemed to be unsafe. Then, we perform a numerical simulation with higher accuracy parameters from the exact initial state and check how close the final point is to the expected final point. We report the relative error of these two points (CE Error).

Both the speed and the accuracy of the overall verification approach depends on how accurately the basis matrix is computed. For numerical simulation, we tried to choose absolute and relative tolerances in the simulation engine so that the method is close to the accuracy of the matrix exponential approach. In our case, we found that simulation tolerances of  $10^{-12}$  yielded errors close to the double-precision matrix exponential method.

In communications with the benchmark authors, we found out that the error conditions were chosen based on a finite number of simulations of the models, while holding the inputs constant. To model this, for each input  $u_i$  we add a new variable to the model with the appropriate initial range and differential equation with  $\dot{u}_i = 0$ . This model transformation was automated using the Hyst model transformation tool [6]. Each model also includes a time variable with  $\dot{t} = 1$ . Thus, for example, while the original benchmark for the Motor system had 8 variables, 2 inputs, and a time variable, we report this as an 11-dimensional system.

For the benchmarks, we use the original time bound of 20. We consider different step sizes, from 0.1 (200 steps) down to the step size suggested by the benchmark authors of 0.001 (20000 steps). We used the hypy library [7] in order to script the process of running all the benchmarks with all the parameters.

Our evaluation uses a new tool, Hylaa<sup>1</sup>, which is written mostly in Python. Matrix exponential is computed using scipy and the sparse.linalg.expm function, which uses a Padé approximation optimized for sparse matrices, which we found was faster on the systems in this benchmark set than the dense-matrix equivalent. Numerical simulations are done with scipy's odeint function, which can simulate both stiff and non-stiff systems with using lsoda from the Fortran library odepack. Matrix multiplication was performed using the numpy library's dot function. The implementation parallelizes both the matrix multiplication (for matrices larger than  $150 \times 150$ , which was empirically derived), as well as the simulations, across all the cores in the system. The measurements were performed on an Intel i7-3930K processor (6 cores, 12 threads) running at 3.5 GHz with 24 GB RAM.

For large matrices with a large number of time steps, it is infeasible to store the complete simulation results in memory. For example, a single basis matrix for a 10000 dimensional model takes about 800 megabytes of RAM ( $10000 \times 10000$  entries times 8 bytes per entry). With a time step of 0.001 and the time bound of 20, storing all 20000 of these would require over 15 terabytes of memory. To reduce the memory requirement, the simulations are not always performed for the full time-bound, and can be split into several parts. This does have the effect of slightly slowing down the computation, since the internally variable-step simulations must be reinitialized each time. The length of the simulations performed at one time is computed based on the amount of memory made available for the computation. A positive side-effect of this is that it allows for early termination when an unsafe state can be reached versus if the entire simulation needed to be computed beforehand. The matrix exponential approach, in contrast, computes one step at a time, and so will terminate on exactly the step where an unsafe state is reached.

The results for each model, safety condition, basis matrix computation method, and for the various step sizes, are shown in Table 1. The star-based reachability method is confirmed as scalable, and is able to perform analysis on all of the benchmarks, often taking only minutes for systems with hundreds of dimensions.

Notice that with the original error conditions (indicated by an asterisk next to the model name in the table), the Beam model (350 dimensions) can actually reach the unsafe states. This error condition was not known prior to analysis with Hylaa. Since the original error conditions were derived based on simulations, such a situation is possible where the simulation which demonstrates the error states was not tested. This shows the incompleteness of a sampled simulation-based analysis, and motivates more rigorous approaches. If one wanted to exhaustively simulate all the corners of the initial states, since 49 of the variables are initially intervals, one would need to run  $2^{49} = 5.6 \cdot 10^{14}$  simulations. For this system, our parallel implementation averaged 1-2 simulations per second, depending on the time step. With this simulation rate, exhaustive analysis would possible after 9 to 18 million years.

The effects of the time step are also apparent in the counter example time (CE Time column). In the MNA1 model (588 dimensions), using a time-step of 0.1 results in a counter-example at time 16.6, which is reduced to 16.56 for a time step of 0.01, and then finally a time 16.554 for a 0.001 time step. The counter-example time, however, is unaffected by the basis matrix computation method on these models, which confirms that they are basically computing the same thing.

In terms of correctness, both approaches, as well as all values of the time step are able to correctly verify or find counter-examples traces to the unsafe error states for all the models, with the exception of the Building (50 dimensions) model when using a time step of 0.1. Examining the plot of the reachable states of this system, the reason becomes apparent. As shown in

<sup>&</sup>lt;sup>1</sup>http://stanleybak.com/hylaa

Model	Dims	Unsafe Error Condition	Method	Step Size	Runtime	Safe?	CE Error	CE Time
Motor*	11	$x_1 \in [0.35, 0.4] \land x_5 \in [0.45, 0.6]$	Simulation	$0.1 \\ 0.01$	0.5s 0.8s	$\checkmark$		
				0.001	2.6s			
			Matrix Exp		0.4s	V		
				$0.01 \\ 0.001$	0.7s 3.9s	$\checkmark$		
Motor	11	$x_1 \in [0.3, 0.4] \land x_5 \in [0.4, 0.6]$	Simulation		0.5s		$1.6 \cdot 10^{-11}$	0.1
		1 - [ , ] 0 - [ , ]		0.01	0.6s		$6.2 \cdot 10^{-13}$	0.04
				0.001	0.6s		$2.2 \cdot 10^{-12}$	0.037
			Matrix Exp	0.1	0.5s		$9.3 \cdot 10^{-13}$	0.1
				0.01	0.5s		$9.9 \cdot 10^{-13}$	0.04
				0.001	0.6s		$1.4 \cdot 10^{-12}$	0.037
Building*	50	$x_{25} \ge 0.006$	Simulation		2.1s	√		
				0.01	2.5s	~		
			Matrix Exp	0.001	7.6s 0.6s			
			Matrix Exp	0.01	0.0s 1.4s	v √		
				0.001	8.9s	√		
Building	50	$x_{25} \ge 0.004$	Simulation	0.1	1.9s	~		
				0.01	2.0s		$1.2 \cdot 10^{-9}$	0.07
				0.001	2.8s		$2.2 \cdot 10^{-9}$	0.07
			Matrix Exp	0.1	0.5s	· √		
				0.01	0.5s		$1.1 \cdot 10^{-9}$	0.07
				0.001	0.4s		$1.1 \cdot 10^{-9}$	0.07
PDE*	86	$y_1 \ge 12$	Simulation	0.1	0.7s	$\checkmark$		
				0.01	1.9s	V		
			Matrix Exp	0.001	13.4s			
			Matrix Exp	0.1	0.7s 2.2s	√ √		
				0.001	17.9s	~		
PDE	86	$y_1 \ge 10.75$	Simulation	0.1	0.6s		$1.2 \cdot 10^{-12}$	0.1
		31		0.01	0.8s		$3.2 \cdot 10^{-12}$	0.03
				0.001	4.0s		$1.3 \cdot 10^{-11}$	0.021
			Matrix Exp	0.1	0.4s		$1.3 \cdot 10^{-12}$	0.1
				0.01	0.6s		$1.3 \cdot 10^{-12}$	0.03
				0.001	0.5s		$1.2 \cdot 10^{-12}$	0.021
Heat*	202	$x_{133} \ge 0.1$	Simulation	0.1	4.1s	$\checkmark$		
				0.01	7.7s	√		
			· · · · · · · · · · · · ·	0.001	56.6s			
			Matrix Exp	0.1	2.1s 12.0s	$\checkmark$		
				0.001	1m57s	<b>√</b>		
Heat	202	$x_{133} \ge 0.02$	Simulation	0.1	4.1s		$3.9 \cdot 10^{-11}$	15.7
				0.01	6.9s		$6.7 \cdot 10^{-12}$	15.67
				0.001	47.7s		$8.9 \cdot 10^{-12}$	15.67
			Matrix Exp	0.1	1.6s		$2.4 \cdot 10^{-11}$	15.7
				0.01	9.8s		$2.5 \cdot 10^{-11}$	15.67
				0.001	1 m 30 s		$2.5 \cdot 10^{-11}$	15.67
				0.001	111005			
ISS*	274	$y_3 \notin [-0.0005, 0.0005]$	Simulation	0.1	7m23s	√		
ISS*	274	$y_3 \notin [-0.0005, 0.0005]$	Simulation	0.1 0.01	7m23s 7m14s	√ √		
ISS*	274	$y_3 \notin [-0.0005, 0.0005]$		0.1 0.01 0.001	7m23s 7m14s 7m44s			
ISS*	274	$y_3 \notin [-0.0005, 0.0005]$	Simulation Matrix Exp	$\begin{array}{c} 0.1 \\ 0.01 \\ 0.001 \\ 0.1 \end{array}$	7m23s 7m14s 7m44s 2.9s	√ . √		
ISS*	274	$y_3 \notin [-0.0005, 0.0005]$		0.1 0.01 0.001	7m23s 7m14s 7m44s			
			Matrix Exp	$\begin{array}{c} 0.1 \\ 0.01 \\ 0.001 \\ 0.1 \\ 0.01 \\ 0.001 \\ 0.001 \end{array}$	7m23s 7m14s 7m44s 2.9s 11.5s 1m39s	✓ . ✓ ✓		0.5
ISS*	274	$y_3 \notin [-0.0005, 0.0005]$ $y_3 \notin [-0.00017, 0.00017]$		$\begin{array}{c} 0.1 \\ 0.01 \\ 0.001 \\ 0.1 \\ 0.01 \\ 0.001 \\ 0.001 \\ \end{array}$	7m23s 7m14s 7m44s 2.9s 11.5s 1m39s 7m11s	✓ . ✓ ✓	 1.1·10 <sup>-6</sup>	0.5
			Matrix Exp	$\begin{array}{c} 0.1 \\ 0.01 \\ 0.001 \\ 0.1 \\ 0.01 \\ 0.001 \\ \end{array}$	7m23s 7m14s 7m44s 2.9s 11.5s 1m39s 7m11s 7m8s	✓ . ✓ ✓	$1.1 \cdot 10^{-6}$ $7.9 \cdot 10^{-7}$	0.5 0.5 0.498
			Matrix Exp	$\begin{array}{c} 0.1 \\ 0.01 \\ 0.001 \\ 0.1 \\ 0.001 \\ 0.001 \\ 0.001 \\ 0.001 \\ 0.001 \\ 0.001 \\ 0.001 \\ \end{array}$	7m23s 7m14s 7m44s 2.9s 11.5s 1m39s 7m11s	✓ . ✓ ✓	 1.1·10 <sup>-6</sup>	0.5

 Table 1: Benchmark results. Stars (\*) indicate each benchmark's original specification.

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				0.001	3.2s		$6.3 \cdot 10^{-7}$	0.498
Beam	350	$x_{89} \ge 2100$	Simulation	$\begin{array}{c} 0.1 \\ 0.01 \\ 0.001 \end{array}$	4m38s 4m47s 7m0s	$\checkmark$		
			Matrix Exp	0.1 0.01 0.001	5.2s 28.7s 4m24s	√ √ √		
Beam*	350	$x_{89} \ge 1000$	Simulation	$0.1 \\ 0.01 \\ 0.001$	4m28s 4m39s 6m30s		$\begin{array}{c} 1.4 \cdot 10^{-12} \\ 1.5 \cdot 10^{-12} \\ 8.8 \cdot 10^{-13} \end{array}$	$16.1 \\ 16.05 \\ 16.041$
			Matrix Exp		5.1s 23.7s 3m28s		$\begin{array}{c} 1.3 \cdot 10^{-11} \\ 2.2 \cdot 10^{-11} \\ 2.1 \cdot 10^{-12} \end{array}$	$   \begin{array}{c}     16.011 \\     16.1 \\     16.05 \\     16.041   \end{array} $
MNA1*	588	$x_1 \ge 0.5$	Simulation	0.1 0.01 0.001	9m49s 10m22s 20m41s	\ \ \		
			Matrix Exp		22.6s 3m0s 30m8s	· · · · · · · · · · · · · · · · · · ·		
MNA1	588	$x_1 \ge 0.2$	Simulation	0.1 0.01 0.001	9m45s 10m19s 18m20s		$\begin{array}{c} 1.2 \cdot 10^{-10} \\ 1.2 \cdot 10^{-10} \\ 1.2 \cdot 10^{-10} \end{array}$	$16.6 \\ 16.56 \\ 16.554$
			Matrix Exp		20.2s 2m31s 24m58s		$7.4 \cdot 10^{-11}$ 7.4 \cdot 10^{-11} 7.4 \cdot 10^{-11} 7.0 \cdot 10^{-11}	16.6 16.56 16.554
FOM*	1008	$y_1 \ge 45$	Simulation	0.1 0.01 0.001	7m3s 9m39s 45m15s	\ \ \		
			Matrix Exp		30.3s 4m51s 48m49s	· · · · · · · · · · · · · · · · · · ·		
FOM	1008	$y_1 \ge 7$	Simulation	0.1 0.01 0.001	7m4s 4m27s 2m39s		$3.3 \cdot 10^{-10} \\ 1.7 \cdot 10^{-10} \\ 1.7 \cdot 10^{-10}$	0.2 0.07 0.069
			Matrix Exp		2.5s 3.3s 12.5s		$\begin{array}{c} 7.5 \cdot 10^{-12} \\ 4.4 \cdot 10^{-12} \\ 4.6 \cdot 10^{-12} \end{array}$	0.2 0.07 0.069
MNA5*	10923	$x_1 \ge 0.2 \lor x_2 \ge 0.15$	Simulation	0.1 0.01 0.001	$\begin{array}{c} 41\mathrm{m}28\mathrm{s}\\ 3\mathrm{h}51\mathrm{m}\\ 24\mathrm{h}2\mathrm{m} \end{array}$	√ √		
		· · · · · · · · · · · · · · · · · · ·	Matrix Exp		$ \begin{array}{c} 2412 \text{m} \\ 14 \text{h} 3 \text{m} \\ (>5 \text{d}) \\ (>53 \text{d}) \end{array} $	· ·		
MNA5	10923	$x_1 \ge 0.1 \lor x_2 \ge 0.15$	Simulation	0.1 0.01 0.001	5m57s 21m40s 2h11m		$9.2 \cdot 10^{-8} 2.0 \cdot 10^{-7} 2.0 \cdot 10^{-7}$	2.0 1.92 1.919
			Matrix Exp		1h25m 13h29m (>5d)		$9.2 \cdot 10^{-8} \\ 2.0 \cdot 10^{-7}$	2.0 1.92

Figure 2, a fixed-step simulation with a large time step, in fact, does not reach error states, which only happens around time 0.07. Traditional reachability analysis methods, such as those implemented in SpaceEx [16], Flow\* [11], or CORA [1], do reason in between time steps and would not mark such systems as safe. Nonetheless, due to the scalability and generation of counter-examples, we believe there is a place for simulation-equivalent reachability.

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Figure 2: Plot of  $x_{25}$  over time of the first 0.25 seconds for the Building (50 dimensions) model produced in SpaceEx. Notice that fixed-time simulations with a time step of 0.1 do not enter the unsafe states ( $x_{25} \ge 0.004$ ).

In terms of the computation time of the basis matrix, both approaches are viable for most of the systems. The runtime of the simulation method is sensitive to the internal variable timesteps the simulation engine can use, which in turn depends on the system's dynamics. This effect can be seen with the ISS model (274 dimensions), which is completed much faster using the matrix exponential approach. For the largest, MNA5 system (10923 dimensions), however, the matrix exponential method becomes extremely slow when there are a large number of steps. This is because matrix multiplication, which is done at each step to compute the basis matrix, takes about 230 seconds with our parallel implementation. The estimated times in parenthesis in the table are based on this value, multiplied by the number of steps required. With this estimation method, the 0.1 time step variant (200 steps) should take about 200  $\cdot$  230 seconds, which equals 12.8 hours, which is close to the measured value of about 14 hours.

One rough intuition as to why the simulation approach may outperform the matrix exponential method on the largest systems with many steps is that the matrix exponential approach requires a matrix multiplication at each step, which takes  $O(n^3)$ , or slightly better. Performing a single step of an Euler method simulation<sup>2</sup> involves computing the update equation x' := x + Ax. If A is sparse (which it is for the larger benchmarks), this operation may be computed in less than  $O(n^2)$ , and so n of these updates (one for each unit vector) may be possible faster than  $O(n^3)$ . Essentially, the simulation approach can take advantage of the sparsity of the A matrix, whereas the matrix exponential method requires multiplying by  $e^{At}$ , which is a dense matrix.

While a simulation method is expected to take slightly longer with more intermediate steps, we expect the number of time steps to have more impact on the matrix exponential approaches, where the number of matrix multiplications is exactly equal to the number of time steps. This trend is apparent for the larger systems, such as the safe variant of the MNA1 model (588 dimensions). Here, the matrix exponential approach increases from 20 seconds with a time step of 0.1, to 151 seconds (2m31s) with a time step of 0.01, to 1498 seconds (18m20s) with a time

 $<sup>^{2}</sup>$ The actual numerical simulation method performed by odeint uses a more complicated update, and may require multiple internal steps per desired time step. These, however, are constant factors.

step of 0.001. This is close to a factor of ten increase in runtime each time the number of steps increases by a factor of ten. The simulation method does take longer with more intermediate steps on this system, but the relationship is less than linear, increasing from 589 seconds (9m49s) to 622 seconds (10m22s) to 1241 seconds (20m41s) as the time step is reduced.

The counter-example accuracy is not too affected by the time step. This is because the simulation engine will adjust its internal step size based on the (fixed) accuracy parameter. It is possible in the matrix exponential method that numerical errors could accumulate during repeated matrix multiplications when there are a large number of time steps, but for these systems, we did not observe this making a difference on the counter-example accuracy. Note that the counter-example reported in the CE Error column is relative error, and it is small in all cases. Additionally, some of the input models are provided with less than 5 digits of precision, which may lead to errors in the analysis result even if the reachability was exact.

In terms of models which do reach unsafe states, the matrix exponential approach usually takes less time to find a counter-example. This is because, as an optimization, simulations are run in batches over many steps rather than a single step at a time. The matrix exponential methods, however, perform one step at a time. For larger systems, where the simulation length is limited for memory reasons as described earlier, we do observe that the analysis time is reduced for the simulation approach when compared with the case where no unsafe states are reachable. This is why the unsafe variant of the FOM model (1008 dimensions) takes less time to compute as the time step is reduced (the simulation lengths are reduced since there are more intermediate time steps).

Finally, since the runtime is dominated by the computation of the basis matrix, and this computation is independent of the initial states or error states, its result could be reused. This would allow the user to make arbitrary changes to the initial states and unsafe conditions, and then quickly compute simulation-equivalent reachability without having to do the expensive basis matrix computation. This was useful in order to find the strengthened error conditions where there are reachable unsafe states for each of the models, since it allowed us to perform many runs of the tool in a short amount of time. For the measurements, however, we ran all the computations from scratch.

#### 4 Conclusion

We analyzed nine high-dimensional benchmark systems using simulation-equivalent reachability analysis. Our main result confirms that the recently proposed technique for generalized starbased analysis of linear systems is capable of scaling to extremely large systems. Further, the counter-example error traces constructed are of high accuracy, as measured by post-analysis, high-accuracy simulations. We also evaluated two methods for the computation of the basis matrix, which dominates the runtime of the approach. We compared using a matrix exponential approach versus numerical simulations. Generally, both can be used, although for the largest systems with many steps, numerical simulations were faster.

There are other aspects that may be a factor when choosing the basis matrix computation method. From an interface perspective, using simulations allows more fine-grained control of the error than the matrix exponential. However, the matrix exponential method may be superior if a quick safety check is desired at a known specific time instant, rather than at all multiples of the time step. A hybrid approach is also possible, where simulations are used to compute the first step's basis matrix, and then matrix multiplication is used for the remaining steps. The runtime of this method is likely similar to the matrix exponential methods, since the runtime is dominated by matrix multiplication, especially when there are a large number of steps.

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