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# PARALLEL-IN-TIME MAGNUS INTEGRATORS

# B. T. KRULL AND M. L. MINION \*

**Abstract.** Magnus integrators are a subset of geometric integration methods for the numerical solution of ordinary differential equations that conserve certain invariants in the numerical solution. This paper explores temporal parallelism of Magnus integrators, particularly in the context of nonlinear problems. The approach combines the concurrent computation of matrix commutators and exponentials within a time step with a pipelined iteration applied to multiple time steps in parallel. The accuracy and efficiency of time parallel Magnus methods up to order six are highlighted through numerical examples and demonstrate that significant parallel speedup is possible compared to serial methods.

11 **Key words.** Ordinary differential equations, nonlinear ordinary differential equations, Magnus 12 expansions, isospectral flows, Lax pairs, parallel-in-time

## 13 AMS subject classifications. 34L30, 65L05, 65Y05

1 2

14 **1.** Introduction. The solution of ordinary differential equations (ODEs) is a 15 well established field with applications across the spectrum of scientific disciplines. Numerical methods date back at least to Euler's work in 1768 [5], and the accuracy, 16stability, and efficiency of various methods is well studied (see for example Refs. [8, 9]). In more recent years, the study of specialized numerical methods for ODEs that 18 preserve certain mathematical properties of the numerical solution has seen increased 20 interest. Examples include methods for Hamiltonian systems that numerically conserve invariants of the true dynamical system such as the energy or angular momentum. 21 More generally, the true solution of an ODE posed in N-dimensional space may reside 22 for all time on a manifold  $\mathcal{M}$  of dimension d < N, and the goal is to devise a method for which the numerical solution will also remain on  $\mathcal{M}$ . Such methods are referred 24 to in general as *geometric integrators*. The interested reader is encouraged to consult 2526 Ref. [7] for a comprehensive introduction to the subject.

As a concrete example, consider the ODE given by

28 (1.1) 
$$\frac{d}{dt}Y(t) = F(Y(t)), \qquad Y(0) = Y_0,$$

where Y and F(Y) are both  $N \times N$  matrices. Depending on the form of F, certain properties of the initial value  $Y_0$  may be preserved for all time, such as the determinant, orthogonality, idempotency, or the spectrum of eigenvalues. In general, standard numerical methods such as linear multistep or Runge-Kutta methods will not produce solutions that conserve such properties [16].

Magnus integrators are a subset of geometric integrators based on the expansion 34 proposed by Wilhelm Magnus in 1954 [14]. The Magnus expansion is closely tied to the concept of Lie groups and algebras due to the presence of matrix commutators. 36 also known as Lie brackets. The discussion concerning solutions to Eq. (1.1) can in fact be generalized to differential equations on Lie groups (see e.g. Ref. [12]), 38 however in this work we strictly use matrix valued solutions. Magnus integrators can 39 also be viewed as a type of exponential integrator (see e.g. Ref. [11] for a review) 40 since they require that the matrix exponential be evaluated. The comprehensive 41 review of the Magnus expansion and applications by Blanes, et. al. [1] provides 42

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a description of several physical applications to which the Magnus expansion has
been applied including nuclear, atomic, and molecular dynamics; nuclear magnetic
resonance; quantum field theory and high energy physics; electromagnetism; optics;
geometric control of mechanical systems; and the search for periodic orbits.

In general, numerical Magnus integrators are constructed by applying quadrature 47 rules of suitable order to a truncation of the Magnus expansion. The review [12] presents 48 various types of Magnus integrators up to sixth-order, and methods of order up to eight 49are considered in Ref. [2]. As the order increases, the number of commutator terms 50required in the Magnus expansion grows quickly, hence in these papers and others, a detailed discussion of how to minimize the number of commutators required for a given order is presented. So called *commutator-free* Magnus integrators (e.g. Refs. [3, 18]) 53 54have also been proposed that replace the need to compute matrix commutators with additional matrix exponentials. This can reduce the total computational cost of the method depending on the relative cost of computing commutators versus matrix 56 exponentials. In this paper, an additional avenue for reducing the time to solution for Magnus integrators is investigated, namely parallelization in time. 58

The study of parallel numerical methods for ODEs dates back at least to Nievergelt 59 in 1964 [17], and the field has seen an increase in activity in the last 15 years. (See Ref. [6] for a recent review.) The standard classification of parallel methods for ODEs 61 includes parallelism across the method, across the problem, and across the time steps 62 [4]. In this work we demonstrate the utility of parallelization across both the method 63 and the time steps for Magnus integrators for both linear and nonlinear equations. 64 65 Special attention is paid to schemes for solving nonlinear differential equations for isospectral flow problems, although the methodology that is described can be applied 66 more generally. 67

The remainder of this paper is organized as follows. Section 2 presents the 68 mathematical background behind the Magnus expansion and Magnus integrators 69 followed by some specific Magnus integrators based on Gaussian collocation in section 3. 70 71The parallelization strategies for these integrators is presented in section 4. Numerical results comparing the efficiency of different parallel methods is presented in section 725. The results demonstrate that significant parallel speedup over serial methods is 73 possible and show that parallel higher-order methods are superior in terms of accuracy 74and time to solution compared to lower-order methods. 75

76 2. Mathematical Preliminaries. In this section, a review of the mathematics
 77 behind the construction of Magnus integrators for both linear and nonlinear problems
 78 is reviewed.

79 **2.1. Matrix Calculus and Differential Equations.** Given a constant matrix 80  $A \in \mathbb{F}^{N \times N}$ , where  $\mathbb{F}$  can be either  $\mathbb{R}$  or  $\mathbb{C}$ , the exponential of A is defined by the power 81 series

82 (2.1) 
$$e^A = \sum_{n=0}^{\infty} \frac{A^n}{n!}.$$

83 Given the time-dependent vector  $y(t) \in \mathbb{F}^N$ , the solution to the differential equation

84 (2.2) 
$$y'(t) = Ay(t), \qquad y(0) = y_0$$

85 is given by

86 (2.3) 
$$y(t) = e^{At}y_0.$$

This is easily shown by differentiating the power series definition (2.1) of the exponential on the right hand side term-by-term to give

89 (2.4) 
$$\frac{d}{dt}e^{At} = Ae^{At}.$$

Now consider the more general case of a time-dependent matrix A(t). Again

91 differentiating definition (2.1) term-by-term and using the product rule yields (2.5)  $d_{-A(t)} = A(t)A'(t) + A'(t)A(t) - A(t)'A(t)^2 + A(t)A'(t)A(t) + A(t)^2A'(t)$ 

92 
$$\frac{d}{dt}e^{A(t)} = A'(t) + \frac{A(t)A'(t) + A'(t)A(t)}{2!} + \frac{A(t)A(t)^2 + A(t)A'(t)A(t) + A(t)^2A'(t)}{3!} \dots$$

93 The right hand side of (2.5) can be rearranged to give

94 (2.6) 
$$\frac{d}{dt}e^{A(t)} = \exp_{A(t)}(A'(t))e^{A(t)},$$

95 where the operator on the right is defined by

96 (2.7) 
$$\operatorname{dexp}_{A(t)}(X(t)) = X(t) + \frac{[A(t), X(t)]}{2!} + \frac{[A(t), [A(t), X(t)]]}{3!} \dots,$$

97 and brackets  $[\cdot, \cdot]$  denote the matrix commutator [A, X] = AX - XA.

98 Now consider the linear system of ODEs

99 (2.8) 
$$y'(t) = A(t)y(t), \qquad y(0) = y_0.$$

100 To find a solution, suppose that it can be written in the form

101 (2.9) 
$$y(t) = e^{\Omega(t)} y_0$$

102 for some matrix  $\Omega(t)$ . Using Eq. (2.6),  $\Omega(t)$  satisfies

103 (2.10) 
$$\operatorname{dexp}_{\Omega(t)}(\Omega'(t)) = A(t), \qquad \Omega(0) = 0.$$

104 The same formal derivation can be applied to a nonlinear system of equations

105 (2.11) 
$$y'(t) = A(y(t), t)y(t), \qquad y(0) = y_0.$$

106 Again, if the solution to this equation is to take the form of Eq. (2.9), then  $\Omega(t)$ 107 satisfies

108 (2.12) 
$$\operatorname{dexp}_{\Omega(t)}(\Omega'(t)) = A(y(t), t), \qquad \Omega(0) = 0.$$

109 In the next section, methods for finding  $\Omega(t)$  are considered.

110 **2.2. Magnus Expansion.** In 1954, Magnus introduced an explicit expression 111 for the solution of Eq. (2.8), which is reviewed here [14]. The first step is the inversion 112 of the operator  $dexp_{\Omega(t)}$ , which gives a differential equation for  $\Omega(t)$ 

113 (2.13) 
$$\Omega'(t) = \sum_{n=0}^{\infty} \frac{B_n}{n!} \mathrm{ad}_{\Omega(t)}^n(A(t)), \qquad \Omega(0) = 0,$$

114 where the  $B_n$  are the Bernoulli numbers and

115 (2.14) 
$$\operatorname{ad}_{\Omega}^{k}(X) = [\Omega, \operatorname{ad}_{\Omega}^{k-1}X], \quad \operatorname{ad}_{\Omega}^{0}(X) = X.$$

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116 Next, a Picard-type iteration is applied to Eq. (2.13)

117 (2.15) 
$$\Omega^{k+1}(t) = \int_0^t \sum_{n=0}^\infty \frac{B_n}{n!} \mathrm{ad}_{\Omega^k(t)}^n(A(t)).$$

118 Collecting terms in Eq. (2.15) yields an infinite series for  $\Omega(t)$ 

119 (2.16) 
$$\Omega(t) = \Omega^{(1)}(t) + \Omega^{(2)}(t) + \Omega^{(3)}(t) + \dots$$

120 where

121 (2.17) 
$$\Omega^{(1)}(t) = \int_0^t d\tau A(\tau)$$

122

123 (2.18) 
$$\Omega^{(2)}(t) = \frac{1}{2} \int_0^t d\tau_2 \int_0^{\tau_2} d\tau_1[A(\tau_2), A(\tau_1)]$$

<sup>124</sup> (2.19)

125 
$$\Omega^{(3)}(t) = \frac{1}{6} \int_0^t d\tau_3 \int_0^{\tau_3} d\tau_2 \int_0^{\tau_2} d\tau_1 [A(\tau_3), [A(\tau_2), A(\tau_1)]] + [[A(\tau_3), A(\tau_2)], A(\tau_1)]$$
126

127 
$$\Omega^{(4)}(t) = \frac{1}{12} \int_0^t d\tau_4 \int_0^{\tau_4} d\tau_3 \int_0^{\tau_3} d\tau_2 \int_0^{\tau_2} d\tau_1 \left[ \left[ [A(\tau_4), A(\tau_3)], A(\tau_2) \right], A(\tau_1) \right] + \left[ A(\tau_4), \left[ [A(\tau_3), A(\tau_2)], A(\tau_1) \right] + \left[ A(\tau_4), \left[ A(\tau_3), [A(\tau_2), A(\tau_1)] \right] + \left[ A(\tau_4), [A(\tau_3), [A(\tau_2), A(\tau_1)] \right] \right] \right]$$

130 (2.20) + 
$$[A(\tau_3), [A(\tau_2), [A(\tau_1), A(\tau_4)]]]$$
.

Each subsequent term in the series contains an additional integration operator, commutators of one higher order, as well as an increasing number of commutator terms. The reader is referred to the original work of Magnus [14] or the extensive review [1] for further details. To summarize, the Magnus expansion gives an explicit formula for the solution of the linear equation given by Eq. (2.8) in the form of the exponential of a matrix defined by an infinite series given by Eq. (2.16).

2.3. The Magnus Expansion for Nonlinear Problems. The same formal 137138 procedure used in the last section to construct the solution to the linear problem Eq. (2.8) can also be applied to the nonlinear system Eq. (2.11). One can still represent 139the solution in terms of the exponential of the function  $\Omega(t)$ , and the only difference 140 is that in the Magnus expansion terms given above in Eqs. (2.17)-(2.20), the terms 141 $A(\tau)$  must be replaced with  $A(y(\tau), \tau)$ , which by the definition of the solution is 142 $A(e^{\Omega(\tau)}y_0,\tau)$ . Although this may appear at first a small change in notation, the 143 implication is quite important. For the nonlinear problem, the Magnus expansion 144145 does not give an explicit formula for the function  $\Omega(t)$  as in the linear case. Instead, the result is an equation for  $\Omega(t)$  involving an infinite expansion of terms containing 146integrals of commutators dependent on  $\Omega(t)$ . The central insight of this paper is that 147this equation for  $\Omega(t)$  can be solved efficiently by a fixed point iteration that is readily 148

149 amenable to parallelization in the time direction.

**2.4.** Isospectral Flows. A special type of matrix differential equation for which the eigenvalues of the solution are independent of time is called *isospectral flow*. Problems of this form exist in application domains including electronic structure, wave dynamics, and linear algebra. Isospectral flow is often associated with the concept of a Lax pair: two matrices or operators A(t), Y(t) dependent on time that satisfy the

155 Lax equation

156 (2.21) 
$$Y'(t) = [A(Y,t), Y(t)], \quad Y(0) = Y_0,$$

157 where  $Y(t), A(Y,t) \in \mathbb{F}^{N \times N}$ .

158 It is straightforward to show that the solution to Eq. (2.21) can be written in the 159 form of the transformation

160 (2.22) 
$$Y(t) = (e^{\Omega(t)})Y_0(e^{\Omega(t)})^{-1}$$

161 where  $\Omega(t)$  is defined by the Magnus expansion with respect to A(t). Since the form of 162 Y(t) takes on a similarity transformation, the eigenvalues of Y(t) do not change in time, 163 hence the term *isospectral*. In the special case where A is Hermitian (or self-adjoint) 164 and Y is skew-Hermitian (or skew-adjoint), the exponential  $e^{\Omega(t)}$  is unitary, which 165 reduces Eq. (2.22) to

166 (2.23) 
$$Y(t) = (e^{\Omega(t)})Y_0(e^{\Omega(t)})^{\dagger}.$$

167 3. Numerical Methods Based on the Magnus Expansion. In this section, the process for constructing numerical methods for differential equations based on the 168 Magnus expansion is discussed. In general, numerical methods are constructed by 169designing appropriate quadrature rules for the Magnus expansion truncated to a given 170 order. The presentation here is focused on collocation type schemes based on Gaussian 171quadrature rules. As proved in Ref. [13], quadrature rules for the terms in the Magnus 172173 expansion based on s Gauss-Legendre quadrature nodes are sufficient for constructing a method of order 2s. Here, methods of order two, four, and six are considered 174175using both Gauss-Legendre and Gauss-Lobatto quadrature nodes. These methods correspond to quadrature rules applied to one, two, and four terms, respectively, in 176Eq. (2.16). 177

Considerable attention in the literature on Magnus integrators is devoted to designing methods requiring the minimum number of function evaluations and matrix commutators for a given order of accuracy [13, 1, 12]. In the context of time parallelization, the manner in which the cost of commutators and function evaluations are counted must reflect the fact that much of the work can be done in parallel, and the minimum parallel cost is not necessarily achieved by a direct parallelization of the serial method with the fewest number of commutators.

185 Methods for linear equations are discussed first, followed by a discussion of 186 additional considerations for nonlinear problems in section 3.3.

3.1. Quadrature Rules for the Magnus Expansion. In this section, the spe-187 188 cific types of quadrature rules used in the numerical methods are described. Quadrature rules based on Gauss-Lobatto or Gauss-Legendre quadrature rules using either two or 189190 three quadrature nodes are considered here. Table 1 lists the specific nodes used for each choice as well as the accompanying classical weights. For a method of a given 191 order, each term in the truncated Magnus expansion must be approximated using the 192function values  $A_m = A(t_m)$  (or  $A_m = A(y(t_m), t_m)$  for nonlinear problems) at the 193194quadrature nodes  $t_m$  corresponding to the quadrature nodes scaled to the time step

Name	Order	Nodes	$q^{(1)}$		
Lob-2	2	0, 1	$\frac{1}{2}, \frac{1}{2}$		
Lob-3	4	$0, \frac{1}{2}, 1$	$\frac{1}{6}, \frac{4}{6}, \frac{1}{6}$		
Leg-3	6	$\frac{1}{2} - \frac{1}{2}\sqrt{\frac{3}{5}}, \ \frac{1}{2}, \ \frac{1}{2} + \frac{1}{2}\sqrt{\frac{3}{5}}$	$\frac{5}{18}, \frac{8}{18}, \frac{5}{18}$		
TABLE 1					

Quadrature nodes and weights for Gauss-Legendre and Gauss-Lobatto rules.

interval  $[t_n, t_{n+1}]$ . For the schemes described below, the same quadrature nodes are used at each term in the expansion in Eq. (2.16).

197 First consider the approximation to the first term of the expansion  $\Omega^{(1)}(t)$  on 198 the interval  $[t_n, t_{n+1}]$  with  $\Delta t = t_{n+1} - t_n$ . Approximating the integral by Gaussian 199 quadrature gives

200 (3.1) 
$$\Omega^{(1)}(t_{n+1}) = \int_{t_n}^{t_{n+1}} A(t) \ dt \approx \Delta t \sum_{j=1}^M q_j^{(1)} A_j = \Omega_{n+1}^{(1)}$$

where M is the number of quadrature nodes. This is classical quadrature, and the well-known coefficients  $q_j^{(1)}$  are given for completeness in Table 1. In order to obtain a fourth-order method, the second term in the Magnus expansion

In order to obtain a fourth-order method, the second term in the Magnus expansion must be included. The simplest approximation to  $\Omega_{n+1}^{(2)}$  sufficient for fourth-order accuracy requires the calculation of only a single commutator term

206 (3.2) 
$$\Omega_{n+1}^{(2)-1} = \Delta t^2 q^{(2)-1} [A_1, A_3],$$

with  $q^{(2)-1} = 1/12$ . The method denoted Lob-4-1 (where the 1 denotes one commutator term) uses this approximation. To compute  $\Omega^{(2)}$  to the accuracy required for a sixth-order method, three nodes can be used and it is necessary to compute three commutators

211 (3.3) 
$$\Omega_{n+1}^{(2)-3} = q_1^{(2)-3}[A_1, A_2] + q_2^{(2)-3}[A_1, A_3] + q_3^{(2)-3}[A_2, A_3]$$

212 with the values

213  $q_j^{(2)-3} = [-7.1721913818656e-2, -3.5860956909328e-2, -7.1721913818656e-2].$ 

Despite the increased computational cost of two additional commutators, in a parallel implementation all three commutators can be computed simultaneously.

To achieve sixth-order accuracy, the first four terms of the Magnus expansion must be included. The sixth-order method denoted Leg-6 approximates the  $\Omega^{(3)}$  term using three Gauss-Legendre nodes following the discussion in Ref. [12]. Specifically,

219 
$$\Omega_{n+1}^{(3)} = \Delta t^3 ([q_{1,1}^{(3)}A_1 + q_{1,2}^{(3)}A_2 + q_{1,3}^{(3)}A_3, [A_1, A_2]] +$$

220 
$$[q_{2,1}^{(3)}A_1 + q_{2,2}^{(3)}A_2 + q_{2,3}^{(3)}A_3, [A_1, A_3]] +$$

$$[q_{3,1}^{(3)}A_1 + q_{3,2}^{(3)}A_2 + q_{3,3}^{(3)}A_3, [A_2, A_3]])$$

223 The values of the coefficients  $q_{i,j}^{(3)}$  are the same as those in Ref. [12], and in matrix

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224 form are (3.6)

$$q^{(3)} = \begin{bmatrix} 3.4538506760729e - 3 & -5.5849500293944e - 3 & -7.1281599059377e - 3 \\ 1.6534391534391e - 3 & 0.0 & -1.6534391534391e - 3 \\ 7.1281599059377e - 3 & 5.5849500293945e - 3 & -3.4538506760729e - 3 \end{bmatrix}$$

Exploiting the linear property of commutators ([A, X] + [A, Y] = [A, X + Y]) allows one to combine terms that share the same inner single commutator and reduce the number of commutators from nine to three. Note that the single commutator terms, i.e.  $[A_1, A_2]$ , are computed during the formation of the  $\Omega_{n+1}^{(2)}$  term and need not be computed again.

The fourth term in the Magnus expansion can be approximated using a low-order quadrature for a sixth-order method. Following the discussion in Ref. [2], the fourth term is approximated by

234 (3.7) 
$$\Omega_{n+1}^{(4)} = \Delta t^4 q^{(4)} [B_0, [B_0, [B_0, B_1]]]$$

235 where  $q^{(4)} = 1/60$  and

236 (3.8) 
$$B_i = \Delta t \sum_{j=1}^{3} q_j^{(1)} (t_j - 0.5)^i A_j$$

237 with  $q_j^{(1)}$  given in the last row of Table 1.

In section 5, numerical examples are presented for five different Magnus integrators.

Table 2 lists the specific discretization of each term included for a given method. The overall order of each method is determined either by the number of terms used in the expansion, or the order of the quadrature rules. For example, the methods Leg-2, Leg-

4-3, and Leg-6 use the same quadrature nodes, but differ in the number of terms used in the expansion, while Lob-2 and Leg-2 use different nodes, but are both second-order

because only one term in the expansion is used.

Name	Order	Nodes	Ω		
Lob-2	2	Lob-2	$\Omega^{(1)}$		
Leg-2	2	Leg-3	$\Omega^{(1)}$		
Lob-4-1	4	Lob-3	$\Omega^{(1)} + \Omega^{(2)-1}$		
Leg-4-3	4	Leg-3	$\Omega^{(1)} + \Omega^{(2)-3}$		
Leg-6	6	Leg-3	$\Omega^{(1)} + \Omega^{(2)-3} + \Omega^{(3)} + \Omega^{(4)}$		
TABLE 2					

Description of the numerical schemes.

244

3.2. The Matrix Exponential and Solution Update. Once all of the quadrature approximations are applied and the value of  $\Omega_{n+1}$  is computed, the solution can be updated by

248 (3.9) 
$$y_{n+1} = e^{M_{n+1}} y_n.$$

There are many approaches to computing the product of a matrix exponential and a vector of the form  $e^A y$  [15], some of which explicitly compute the term  $e^A$  and some which only approximate the product  $e^A y$ . The choice of method is problem dependent and does not affect the discussion of time parallelism of the methods. In the numerical examples presented here, the scaling-and-squaring method from [10] is used to form the matrix exponential explicitly. **3.3. Considerations for Nonlinear ODEs.** Consider now problems of the form

257 (3.10) 
$$y' = A(y,t)y$$

where non-linearity is introduced through the y-dependence of A. The terms in the Magnus expansion approximations introduced above now depend on the solution through  $A_m$ , and cannot simply be evaluated. The numerical solution at each quadrature node  $t_m$  will be denoted  $y_m$ , and hence  $A_m = A(y_m, t_m)$ . The values of  $y_m$  at each quadrature node are computed by

263 (3.11) 
$$y_m = e^{\Omega_m} y_n,$$

where  $\Omega_m$  is an approximation to the Magnus expansion on the interval  $[t_n, t_m]$ . The construction of  $\Omega_m$  is discussed below.

A simple fixed-point Picard-type iteration is used to simultaneously solve for the values  $\Omega_m$  and  $y_m$ . The iterative scheme is initialized by setting  $y_m^{k=1} = y_n$  at each node m, where k denotes the iteration. The solution at each quadrature node is updated by

270 (3.12) 
$$y_m^{k+1} = e^{\Omega_m^k} y_n.$$

271 Then  $\Omega_m^{k+1}$  is computed using values  $A(y_m^{k+1}, t_m)$  as described below.

To compute  $\Omega_m^k$ , the process for constructing the quadrature rules in section 3.1 272 needs to be applied to each quadrature node. Evaluating the values  $A(y_m^k, t_m)$  at 273each node  $t_m$  is straight-forward but needs to be performed each iteration. In all 274cases considered here, the same matrix commutators are used for each quadrature 275rule, so computing the commutators is also identical to the linear case. The significant 276277 difference is that a quadrature rule for each term in the Magnus expansion must be computed for each interval  $[t_n, t_m]$  rather then just  $[t_n, t_{n+1}]$  as in the linear case. The 278coefficients for each of the terms are included in Appendix A. 279

280 Once each  $\Omega_m^k$  is computed, the matrix exponential can be computed for each 281 node, and a new solution is obtained at each node by Eq. (3.12). The solution is 282 considered converged when the maximum absolute value of the residual

283 (3.13) 
$$R_m^k = e^{\Omega_m^k} y_n - y_m^k = y_m^{k+1} - y_m^k$$

is less than a predefined tolerance. In a traditional implementation of this iteration using Gauss-Legendre nodes, it is not necessary to compute the values  $\Omega_{n+1}^k$  and  $y_{n+1}$ at the end of the time step during the iterations; however, when pipelining of iterations is employed as discussed in the next section,  $y_{n+1}$  is computed each iteration to update the initial condition for the next time step.

**3.4.** Considerations for Isospectral Flows. Consider now problems of the form of Eq. (2.21). The procedure for constructing Magnus integrators follows exactly that laid out in the previous section except that the solution is defined at quadrature nodes by

293 (3.14) 
$$Y_m^{k+1} = e^{\Omega_m^k} Y_n e^{-\Omega_m^k}$$

and likewise for the computation of  $Y_{n+1}^{k+1}$  from  $\Omega_{n+1}^k$ .

2954. Parallelization in Time for Magnus Integrators. In this section, we 296investigate the theoretical computational cost of the Magnus integrators introduced in the previous section in both serial and parallel settings. We consider both parallelization 297across the method and parallelization across the time steps. In the following discussion, 298 299 it is assumed that arbitrarily many processors are available for a given problem and the cost of communication between processors is ignored. It is also assumed that the 300 matrix exponential is formed explicitly as is done for the the numerical results given 301 in Section 5. 302

4.1. The Linear Case. First consider linear problems where A(t) does not depend on the solution y. For each of the N time steps, the following tasks must be performed:

- L1. Evaluate  $A_m = A(t_m)$  for each quadrature node  $t_m$
- L2. Compute commutators necessary for each term in the truncated Magnus expansion
- 309 L3. Apply quadrature rules to compute  $\Omega(t_{n+1})$
- 310 L4. Form the exponential of  $\Omega(t_{n+1})$

306

307

308

311 L5. Compute the solution  $y_{n+1}$  from  $y_n$  by matrix multiplication

Denote by  $C_A$  the computational cost of computing A(t) for a given time. Then if M 312 quadrature nodes are used, L1 has a computational cost of  $MC_A$ . Next let  $n^p$  denote 313 the number of commutators required to compute the *p*th term in the Magnus expansion 314 and  $C_C$  the cost of computing one commutator. Assuming that each commutator 315 in the p + 1 term can be formed with one additional commutator applied to a term 316from term p, the total number of commutators to compute is simply  $n^1 + \ldots n^P$ . 317 Denoting this sum by  $N_C$ , the cost of L2 is  $N_C C_C$ . Task L3 requires only that a 318 linear combination of the terms computed in L2 be computed. We can denote this cost 319 by  $N_C C_L$ , where  $C_L$  is the cost of adding a term in the linear combination. Denote 320 by  $C_E$  the cost of the matrix exponential, and hence the cost of L4 is  $C_E$ . Likewise 321 denote by  $C_M$  the cost of multiplying the solution by a matrix which corresponds to 322 the cost of task L5. Putting these together, the serial cost for N time steps is 323

324 (4.1) 
$$\mathbf{C}_{\mathbf{S}} = N(MC_A + N_C(C_C + C_L) + C_E + C_M).$$

Now consider the parallelization of the method for the linear problem across the time steps. In task L1, each function evaluation can be done concurrently, so that the cost is reduced from  $MC_A$  to  $C_A$ . For task L2, all the commutators of a given order can be computed concurrently, so that cost is reduced from  $N_CC_C$  to  $PC_C$ . Task L3 can be done with cost  $\log_2(N_C)C_L$ , and the cost of task L4 and L5 remains the same. Next consider the cost when both parallelization across the method and across the time steps is employed. Given sufficient processors, tasks L1-L4 can all be computed

on all time steps concurrently. Only task L5 must be done serially so that the total cost using both forms of parallelism becomes

334 (4.2) 
$$\mathbf{C}_{\mathbf{P}} = C_A + PC_C + \log_2(N_C)C_L + C_E + NC_M.$$

Clearly this is a significant reduction in computational cost. If the cost of computing the commutators and matrix exponential (L2 and L4) dominate the other terms, the theoretical parallel speedup approaches N.

An important point to make about this counting is that the cost per step when using parallelization across the method depends very little on the number of quadrature nodes or commutators used in each term since only the cost of task L3 depends on

these factors. Hence higher-order methods are only modestly more expensive per 341 342step than lower-order methods and there is less benefit from reducing the number of

commutators required in each term of the Magnus expansion since multiple terms can 343

be computed in parallel. Furthermore, for a given accuracy, higher-order methods will 344

typically require fewer time steps (i.e. smaller N). 345

**4.2.** The Nonlinear Case. For nonlinear problems, the theoretical accounting 346 of cost must be modified somewhat. Since we are using an iterative procedure to 347 compute  $\Omega(t_m)$ , the following steps must be done for each iteration in each time step: 348 N1. Evaluate  $A(y_m^k, t_m)$  for each quadrature node  $t_m$ 349

- N2. Compute commutators necessary for each term in the truncated Magnus 350 expansion 351
  - N3. Apply quadrature rules to compute  $\Omega_m^k$  at each quadrature node  $t_m$

N4. Form exponential of  $\Omega_m^k$  at each quadrature node  $t_m$ . N5. Compute  $y_m^{k+1}$  at each quadrature node by matrix multiplication by  $\Omega_m^k$ . 354

The main difference between these tasks and the linear case is that N2-N5 are done 355356 for each quadrature node instead of only once. For simplicity, the serial cost of these steps will be assumed to be M times that of the linear case. Hence, denoting by  $K_S$ 357 358 the number of iterations required for each step in a serial implementation, the serial cost for the nonlinear Magnus method iteration becomes 359

360 (4.3) 
$$\mathbf{C}_{\mathbf{S}} = NK_{S}M(C_{A} + N_{C}(C_{C} + C_{L}) + C_{E} + C_{M}).$$

As will be shown below, the number of iterations required for convergence  $K_S$  depends 361 on  $\Delta t$  in a nontrivial way. 362

If we allow parallelization across the method, the tasks above can all be computed 363 concurrently at each quadrature node, and hence the cost of each iteration for the 364 nonlinear method is essentially that of one step in the linear case using parallelization 365 across the method. 366

367 (4.4) 
$$\mathbf{C}_{\mathbf{I}} = C_A + PC_C + \log_2(N_C)C_L + C_E + C_M.$$

The speedup across the method is bounded by M. 368

Now consider parallelization across the time steps. The simplest way that this can 369 be accomplished is to pipeline the iterations. We first divide the N time steps into 370blocks of size  $N_P$  with each step in a block assigned to a group of processors indexed by 372  $n_p$ . At each time step  $n_p$  for each iteration k, the initial condition is assigned the final value from iteration k-1 of the time step  $n_p-1$ . For each block,  $N_P-1$  pipelined 373 iterations are required before the last processor has a consistent initial condition. After 374 this initialization step, assume  $K_P$  additional iterations are needed for convergence on every time step in the block. As in the serial case  $K_P$  depends in general on the 376 time step  $\Delta t$  and now also on  $N_P$ , increasing as  $N_P$  increases and decreasing at  $\Delta t$ 377 decreases. Furthermore  $K_P \geq K_S$ . 378

The parallel cost on each block will be  $(N_P - 1 + K_P)\mathbf{C}_{\mathbf{I}}$  compared to  $(N_P K_S)\mathbf{C}_{\mathbf{I}}$ 379 when no parallelization across the time steps is applied (i.e.  $N_P = 1$ ). The potential 380 speedup from parallelization across the time steps is then 381

382 (4.5) 
$$\mathbf{S} = \frac{N_P K_S}{N_P - 1 + K_P} = \frac{K_S}{1 + (K_P - 1)/N_P}.$$

Clearly the speedup is bounded by the number of serial iterations required and the 383 best speedup will occur when the quantity  $K_P/N_P$  remains small as  $N_P$  increases.

384385 This ratio will be investigated in the numerical examples.

352

386 5. Numerical Examples. In this section, the performance of the different 387 Magnus integrators introduced in section 3 is examined in both serial and parallel settings. First, the performance of the methods is considered as applied to a Toda 388 lattice problem from the literature. Serial results demonstrating the relative accuracy 389 and efficiency of the various methods are presented. In section 5.1.3, some preliminary 390 results on parallelization of the methods are presented. In section 5.2, the parallel 391 performance of the methods is considered on a problem motivated by real-time time-392 dependent density functional theory. 393

5.1. Test Case 1: The Periodic Toda Lattice. The numerical methods will first be evaluated on the test problem of a *d*-particle periodic Toda lattice [19], a one-dimensional chain whose dynamics are governed by nonlinear nearest-neighbor interactions. The equations of motion are a Hamiltonian system for positions  $q_j$  and momenta  $p_j$  (assuming unit masses)

399 (5.1) 
$$q'_j = p_j$$

400 (5.2) 
$$p'_{i} = e^{-(q_{j}-q_{j-1})} - e^{-(q_{j+1}-q_{j})}$$

In order to cast the dynamics in terms of a Lax pair as in Eq. (2.21), one uses theFlaschka change of variables

404 (5.3) 
$$\alpha_j = \frac{1}{2} e^{-(q_{j+1} - q_j)/2}$$

405 (5.4) 
$$\beta_j = \frac{1}{2} p_j,$$

407 which leads to definitions of Y and A

(5.5)

408 
$$Y = \begin{bmatrix} \beta_1 & \alpha_1 & 0 & \dots & \alpha_d \\ \alpha_1 & \beta_2 & \alpha_2 & \ddots & \vdots \\ 0 & \alpha_2 & \ddots & \ddots & 0 \\ \vdots & \ddots & \ddots & \ddots & \alpha_{d-1} \\ \alpha_d & \dots & 0 & \alpha_{d-1} & \beta_d \end{bmatrix}, A(Y) = \begin{bmatrix} 0 & -\alpha_1 & 0 & \dots & \alpha_d \\ \alpha_1 & 0 & -\alpha_2 & \ddots & \vdots \\ 0 & \alpha_2 & \ddots & \ddots & 0 \\ \vdots & \ddots & \ddots & \ddots & 0 \\ \vdots & \ddots & \ddots & \ddots & -\alpha_{d-1} \\ -\alpha_d & \dots & 0 & \alpha_{d-1} & 0 \end{bmatrix}.$$

409 The numerical example considered here is an 11-particle periodic Toda lattice 410 taken from Ref. [21] with the initial conditions

411 
$$q(0) = [0, \dots, 0]^T$$

412 (5.6) 
$$p_j(0) = \begin{cases} 4, & 1 \le j \le 4, \\ 0, & j \ge 5. \end{cases}$$

The periodic Toda lattice is not asymptotically free and has considerably complicated motion. Figure 1 shows the position and momenta of the 11 particles up to  $t_{final} = 10.0$ .

417 **5.1.1. Simulation parameters.** The following numerical experiments are all 418 performed on this 11-particle periodic Toda lattice with initial conditions as in Eq. (5.6) 419 and a fixed  $t_{final} = 10.0$ . A Picard iteration tolerance of  $10^{-12}$  is used on the maximum 420 absolute value of the residual at the end of the timestep. The reference solution is 421 taken as the Leg-6 method with  $\Delta t = 2^{-15} t_{final}$  or 32768 steps. The reported error

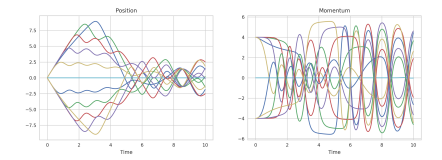


FIG. 1. Symmetric periodic 11-particle Toda lattice solutions with initial conditions as in Eq. (5.6).

422 is defined as the matrix 2-norm of the absolute value of the difference between the423 solution in question and the reference solution.

The methods have been implemented using the LibPFASST<sup>1</sup> library. Communication between pipelined iterations is done in LibPFASST using MPI, and the variable  $N_P$  corresponding to the number of parallel steps is also the number of MPI ranks. For parallelization across the method, OpenMP is used to parallelize the steps in the nonlinear iteration. All timing results were performed on a single compute node of NERSC's Cray XC30 supercomputer, Edison, which contains 24 hardware cores and 64 GB of memory.

431 **5.1.2.** Serial Results. We first perform convergence tests to examine the error with respect to  $\Delta t$  for different Magnus methods. Figure 2 shows the behavior for 432 each method summarized in Table 2. Each method displays the proper convergence 433 rate for a range of  $\Delta t$ . For the second-order methods, note that the error for Leg-2 434 is significantly smaller than that of Lob-2, which implies the dominant error term 435 for Lob-2 is due to the quadrature rule as opposed to the truncation of the Magnus 436437 expansion. The Leg-2 method requires more serial work in this case due to the use of three quadrature nodes instead of two. The difference between the fourth-order 438 methods is less significant. Leg-4-3 is more accurate than Lob-4-1 (with a higher serial 439 cost), but since the main difference between the two methods is how the second term 440 in the Magnus expansion is treated, the difference between the two is smaller than 441 442 for the second-order methods. Leg-6 is clearly more accurate than the other methods. Note that only about nine significant digits of accuracy is attainable for the reference 443 solution for this problem using double precision due to the sensitivity of the solution 444 to perturbations. 445

To better demonstrate the relative computational cost of each method, Figure 3 446 shows the total serial wall-time versus number of steps for the experiment above. As 447 expected, Lob-2 is the method with the shortest time to solution. Lob-4-1, the simplest 448 fourth-order method that can be constructed, is actually less expensive than the second-449 450order Leg-2 scheme despite the fact that Lob-4-1 requires a matrix commutator. For the small problem size used here, the matrix commutators are relatively inexpensive, 451 and the fact that matrix exponentials must be computed at three internal quadrature 452nodes for Leg-2 more than makes up for the lack of commutator terms. Leg-4-3 and 453

<sup>&</sup>lt;sup>1</sup>LibPFASST is available at https://github.com/libpfasst/LibPFASST.

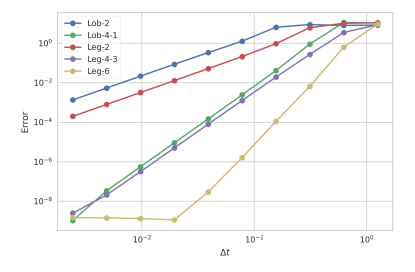


FIG. 2. Error at  $t_{final} = 10.0$  versus  $\Delta t$  for the Toda lattice test case.

Leg-6 are unsurprisingly the most expensive in serial.

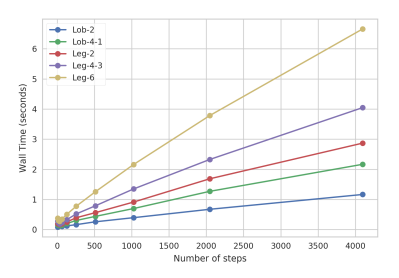
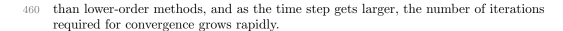


FIG. 3. Total wall-time for the solution for the Toda lattice test case for fixed  $t_{final} = 10.0$ .

454

Note that the cost of the methods displayed in Figure 3 does not grow exactly linearly with the number of time steps. This is due to the fact that the number of Picard iterations needed to converge to the tolerance depends on the time step  $\Delta t$ . Figure 4 shows the average number of iterations over all time steps for each method as a function of  $\Delta t$ . Note the higher-order methods require moderately fewer iterations



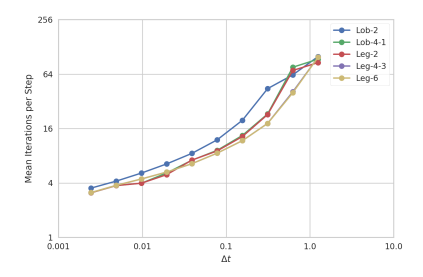


FIG. 4. Average number of iterations as a function of  $\Delta t$  for serial methods on the Toda lattice test case.

The three figures provided above demonstrate that it is not necessarily trivial to 462 choose a method and time step that will provide a solution to a given accuracy with 463 the least computational effort. To illustrate this, Figure 5 shows the accuracy versus 464 wall-clock time for the Toda lattice test. Reading from left to right for a given accuracy 465 shows in increasing order, the methods with the fastest time-to-solution. While Lob-2 466 is by far the cheapest method, it only the fastest method for simulations where the 467 error is O(1). Lob-4-1 is the most efficient for error tolerances to about  $10^{-6}$ , after 468 which Leg-6 becomes the most efficient. For an error of about  $10^{-6}$ , Leg-6 and Lob-4-1 469are more than an order of magnitude more efficient than the second-order methods. 470

471 5.1.3. Parallel in Time Results. In this section, the relative performance of parallel Magnus integrators is explored. We first consider the speedup due to 472parallelization over time steps by pipelining the Picard iterations. As discussed in 473 Section 4, the theoretical speedup from pipelining is bounded by the number of serial 474 iterations required for the method and depends on how the total number of parallel 475iterations required to reach convergence grows as the number of time parallel steps 476 is increased. As in the serial case, the number of iterations required depends on the 477 time step but now depends also on the number of parallel time steps in the pipeline. 478 Figure 6 demonstrates this dependence for the Toda lattice test using method Leg-6 by 479480plotting the average number of iterations required for convergence for different  $\Delta t$  and parallel steps, denoted by  $N_P$ . As in the serial case, the number of iterations required 481 482 decreases with decreasing  $\Delta t$  and here it also increases for fixed  $\Delta t$  as  $N_P$  increases.

483 A second way to display the convergence behavior of the pipelined iteration is to 484 plot the residual after each iteration for each time rank. Figure 7 shows this data for 485 the Leg-6 method using a time step of  $\Delta t = 10/128$  in the top panel and  $\Delta t = 10/1024$ 486 in the bottom panel for 16 parallel time steps. As discussed in section 4.2, the speedup

## 14

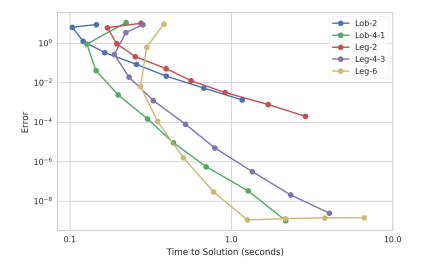


FIG. 5. Error versus time to solution for the serial Toda lattice test case. Each point on a line, read from top to bottom, represents a two-fold increase in the number of steps.

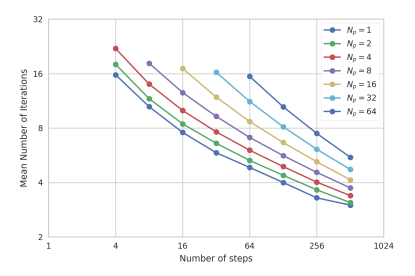


FIG. 6. Average number of pipelined iterations to reach residual of  $10^{-12}$  for the parallel Toda lattice test case using the Leg-6 method.

achievable from pipelining depends on the ratio  $K_P/N_P$ . In this example it is clear that  $K_P/N_P$  is decreasing for large  $N_P$  as more pipelined time steps are used.

The nontrivial dependence of the parallel iterates on both  $\Delta t$  and  $N_P$  makes it difficult to predict which method with which parameters will minimize the time to solution for a given accuracy. As in the serial case, it is instructive to consider the accuracy versus wall-clock for different methods with different number of time-parallel

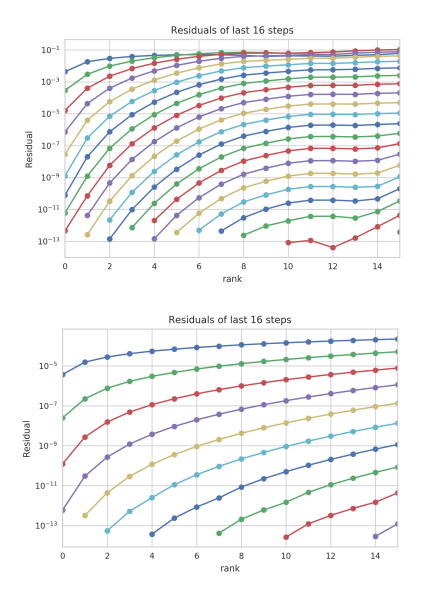


FIG. 7. Residual convergence for each processor in a 16 MPI task simulation using the secondorder Legendre method Leg-6 with 128 steps (above) and 1024 steps (below). Every reoccurrence of a given color is another 6 iterations.

493 steps. Figure 8 displays this information for each of the methods with increasing 494 number of processors. Across a single method, e.g. Leg-6, there's an optimal value 495 of parallelization due to the fact that the increased number of iterations required in 496 the pipeline algorithm starts to cost more than it gains. It is also likely for this test 497 case that the small problem size implies that communication latency is not negligible. 498 Nevertheless, it is again clear that higher-order parallel method gives a shorter time 499 to solution than lower-order alternatives. The second-order methods are only less

500 expensive than the higher-order methods when no digits of accuracy are computed.

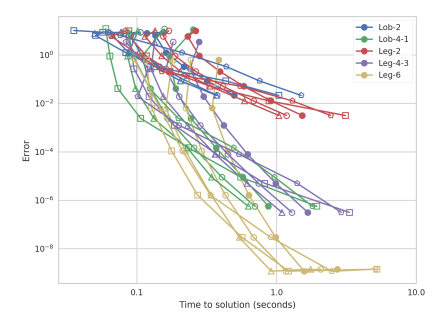


FIG. 8. Error versus time to solution for an 11-particle periodic Toda lattice example. Each color represents a different method and each marker represents a different number of time-parallel steps,  $N_p = 1, 2, 4, 8, 16$ . Closed circles represent the serial computation, open circles  $N_p = 2$ , triangles  $N_p = 4$ , squares  $N_p = 8$ , and pentagons  $N_p = 16$ .

501Finally, we present preliminary results using parallelization across the method and 502across the time steps. Figure 9 shows the error versus time to solution for the serial implementation and time- and method-parallel methods. The particular configuration 503 of  $N_p = 8$  and 3 OpenMP threads uses the entirety of 1 hardware node on NERSC's 504Edison Cray XC30 supercomputer with no hyperthreading. The additional parallel 505across method provides an additional factor of at best two using simple OpenMP 506 507parallel do loops over nodes. Using both types of parallelism in this context gives up to a factor of 4.7 in the overall compute time compared to serial methods using 50824 total processors. Note that compared to serial second-order methods, the parallel 509sixth-order method can compute a solution in less time with more than four orders of 510magnitude lower error. 511

512 **5.2. Test Problem 2: An RT-TDDFT Proxy.** The movitating application 513 for the parallel Magnus methods introduced here is the simulation of electron dynamics 514 using real-time time-dependent density functional theory (RT-TDDFT). In this section 515 we study the parallel performance of the time-parallel Magnus methods on a test 516 problem with a similar but simpler structure than RT-TDDFT.

517 **5.2.1. Physical Background.** RT-TDDFT describes the time evolution of the 518 density matrix *P* through the von Neumann equation of motion

519 (5.7) 
$$P' = -i[F(P), P],$$

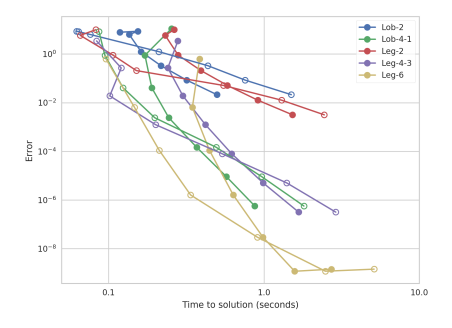


FIG. 9. Error versus time to solution for an 11-particle periodic Toda lattice example. Solid markers indicate serial calculations and open markers indicate time- and method-parallel runs with 8 MPI tasks and 3 OpenMP threads.

520 where F(P) is the Fock matrix. The size of the matrices N scales with the number of electrons times the number of basis functions used to represent the electron density. 521522 The evaluation of F(P), the construction of the Fock matrix, requires for each entry in P the approximation of one- and two-electron integrals which formally requires  $O(N^4)$ 523 work. Furthermore, the time scales on which electron dynamics occur are typically 524525 on the order of tens or hundreds of atto-seconds, hence even simulations of moderate 526 sized molecules at the femto-second range are extremely computationally expensive. In the RT-TDDFT simulation suite in the NWChem code [20], Eq. (5.7) is integrated 527 using a serial second-order Magnus integrator, and part of the motivation of this paper 528 is to improve the efficiency of this choice. 529

530 **5.2.2. The Proxy Problem.** To give an example of the performance of the 531 parallel Magnus methods without requiring the considerable infrastructure necessary 532 to compute the true Fock matrix, we consider a test problem inspired by a simplified 533 one-dimensional RT-TDDFT problem. Here P is again an  $N \times N$  complex matrix, 534 and

535 (5.8) 
$$F(P)_{i,j} = \sum_{i=1}^{N} \sum_{j=1}^{i} \left( Zf_1(i,j) + \sum_{m=1}^{N} \sum_{n=1}^{N} E(f_2(i,j,m,n) - \frac{1}{2}f_2(i,n,m,j)) \right)$$

The functions  $f_1$  and  $f_2$  correspond to the one- and two-electron integrals in RT-DTDFT, with the  $f_1$  term corresponding to the single electron Hamiltonian and the two  $f_2$  terms corresponding to the Coulomb and exchange operators respectively. The simplified forms here are derived by considering single Gaussian basis functions in one 540 dimension with single point approximations of the integrals. Specifically

541 (5.9) 
$$f_1(i,j) = -\frac{P_{i,j}\bar{P}_{j,i}}{r^2}\exp(-\frac{(x(i)-x(j))^2}{2})$$

542 with r = (x(i) + x(j))/2. Similarly (5.10)

543 
$$f_2(i,j,m,n) = \frac{P_{m,n}\bar{P}_{n,m}P_{i,j}\bar{P}_{j,i}}{r^2}\exp(-\frac{(x(i)-x(j))^2}{2})\exp(-\frac{(x(m)-x(n))^2}{2})$$

with r = ((x(i) + x(j))/2 - (x(m) + x(n))/2). Note that the computation of F(P)in the problem is formally  $O(N^4)$  whereas the matrix commutator terms are  $O(N^3)$ . The matrix exponential when computed by power series is also  $O(PN^3)$  where P is the number of terms in expansion. Hence in this application (as in full RT-TDDFT simulations), the computation of F(P) is the dominant cost for even moderate N. In the numerical tests, the spatial locations are uniformly spaced on [-1,1] so that x(i+1) - x(i) = 2/(N-1). The values for Z and E are 0.5 and 0.05 respectively.

5515.2.3. Parallel in Time Results. For the test problem described above, we run the parallel Magnus algorithms Lob-2, Lob-4-1, and Leg-6 on a 20 particle RT-TDDFT proxy problem as described above, using a final time of the run T = 0.05. For each choice of method, runs using 48, 96, 144, and 192 time steps are compared for different 554numbers of parallel processors  $N_p = 1, 2, 4, 8, 16$  and 24. A uniform time step is used 555in all cases, which is justified since the character of the dynamics does not change in 556time. As above, computations are run on 1 hardware node on NERSC's Edison Cray 557 XC30 consisting of 24 cores. For these comparisons, only the parallelism across time 558 steps is used. Unlike the results in the previous section, the convergence tolerances 559 are adjusted by case to avoid unnecessary iterations. In practice this means that 560the tolerances are larger for less accurate simulations since additional SDC iterations 561 will reduce the residual but not the numerical error determined by the underlying 562quadrature rule. The tolerances here were chosen using knowledge of the error in the 563 simulations and are shown in table 3. Dynamically choosing the optimal tolerance for 564serial or parallel SDC methods is an open problem.

	48	96	144	196		
Lob-2	1e - 4	1e - 4	2e - 5	1e - 5		
Lob-4-1	1e - 5	1e - 6	5e - 7	1e - 7		
Leg-6	1e - 6	1e - 8	1e - 9	1e - 10		
TABLE 3						

Residual tolerances for the RT-TDDFT proxy problem for different methods and number of steps.

565

Fig. 10 compares the accuracy versus total computational time for each variation 566 and demonstrates the attractiveness of both higher-order methods and time parallelism. 567 The reported error in each case is the maximum absolute error along the diagonal of 568 569 P (which in RT-TDDFT are the relevant quantities). It is clear from the timings that significant parallel speedup is attainable for methods of all order. As expected, the 570 571 smaller the error tolerance, the larger the possible parallel speedup since there are in general more iterations to amortize over processors. Note that serial second-order methods are more expensive than the fastest fourth-order methods. In particular, 573 the fastest serial fourth-order method runs in less time than the fastest serial second-574575 order method because fewer iterations are needed for convergence (despite the smaller

residual tolerance). Parallel sixth-order methods run faster than serial second-order
methods while producing smaller errors by several orders of magnitude. In general,
the optimal choice of method and parallelization depends on the error tolerance.

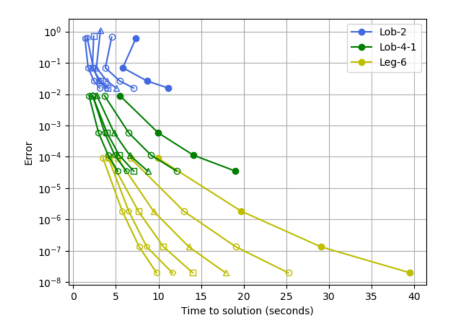


FIG. 10. Error versus time to solution for a 20-particle RT-TDDFT proxy example. Each color represents a different method and each marker represents a different number of time-parallel steps,  $N_p = 1, 2, 4, 8, 16, 24$ . Closed circles represent the serial computation, open circles  $N_p = 2$ , triangles  $N_p = 4$ , squares  $N_p = 8$ , pentagons  $N_p = 16$ , hexagons  $N_p = 24$ .

578

**6.** Discussion. Much of the initial work on Magnus integrators is focused on linear problems where only a single evaluation of the Magnus expansion is required for each time step. In contrast, this paper explores the accuracy and cost of Magnus integrators applied to nonlinear problems. Nonlinear Magnus methods require solving an implicit equation involving the Magnus expansion to obtain the solution in each timestep, and the methods proposed here use a simple fixed point iteration for this solution that can be readily parallelized in time.

586 One conclusion presented here is that in both the linear and nonlinear case, 587 straight-forward parallelization across the method is possible, leading to higher-order 588 methods with only marginally higher computational cost than lower-order methods. 589 This is complimentary to previous results in the literature where considerable effort is 590 placed on reducing the complexity of each of the terms in regards to the number of 591 commutators required.

The second level of parallelism described in this work, namely parallelization across the time steps through pipelined iterations, can further decrease the overall time-to-solution for nonlinear problems. There is a non-trivial relationship between the number of pipelined time steps, the time step size, and the number of iterations required for convergence of the iteration, hence it is not easy to predict *a priori* which choice of parameters will lead to the shortest wall clock time given a desired level of accuracy. Nevertheless, for the test cases considered here, in most situations, the parallel sixth-order methods require the least computation time and are far superior to serial second-order methods.

It is important to note that the results here are preliminary and are performed on 601 two test cases of moderate size. In general, the possible parallel speedup attainable 602 for a given problem will depend on the sensitivity of the problem to perturbations, 603 the relative cost of the operations such as computing commutators or the matrix 604 exponential, and the ratio of computation to communication costs. In future work, 605 the authors will use this parallel methodology to investigate real-time electronic 606 dynamics, where the calculation of the right-hand side values is more expensive than 607 both commutators and matrix exponentials as in the second numerical example. This 608 609 paper addresses only the use of time-parallelism to reduce the wall clock time of serial Magnus methods, and not important issues like the best way to compute the matrix 610 exponential, the optimal choice of time step, nor the dynamic stopping criteria for 611 iterations. Such issues are present in both serial and parallel implementations of 612 Magnus methods and are most likely problem dependent. 613

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Appendix A. Quadrature Rules for Intermediate Nodes. The necessary
 weights for implementing each of the methods Lob-4-1, Leg-2, Leg-4-3, and Leg-6 for
 nonlinear problems are presented here. For Lob-2, no additional rules are necessary.
 For the computation of the single integrals in the first Magnus term, the necessary
 rules take the form

626 (A.1) 
$$\Omega_m^{(1)} = \Delta t \sum_{j=1}^M q_{m,j}^{(1)} A_j,$$

and the coefficients  $q_{m,j}^{(1)}$  correspond to the classical collocation schemes (see e.g. [9]) For Lob-3, only one additional rule is needed at the midpoint  $t_2$ ,

629 (A.2) 
$$q_{2,j}^{(1)} = \left[\frac{5}{24}, \frac{1}{3}, -\frac{1}{24}\right].$$

630 For Leg-3, rules for each node are required, and the  $q_{m,i}^{(1)}$  are given by

631 (A.3) 
$$q_{m,j}^{(1)} = \begin{bmatrix} \frac{5}{36} & \frac{2}{9} - \frac{\sqrt{15}}{15} & \frac{5}{36} - \frac{\sqrt{15}}{30} \\ \frac{5}{36} + \frac{\sqrt{15}}{24} & \frac{2}{9} & \frac{5}{36} - \frac{\sqrt{15}}{24} \\ \frac{5}{36} + \frac{\sqrt{15}}{30} & \frac{2}{9} + \frac{\sqrt{15}}{15} & \frac{5}{36} \end{bmatrix}.$$

For the second term  $\Omega^{(2)}$  there are two versions described in the linear case by Eqs. (3.2) and (3.3). In the first case, only one additional coefficient is needed, namely  $q_2^{(2)-1} = 1/48$ . In the second case, we have

635 (A.4) 
$$\Omega_m^{(2)-3} = q_{m,1}^{(2)-3}[A_1, A_2] + q_{m,2}^{(2)-3}[A_1, A_3] + q_{m,3}^{(2)-3}[A_2, A_3],$$

$$\begin{array}{ll} & \text{with the values of } q_{m,j}^{(2)-3} \text{ given by} \\ & (\text{A.5}) \\ & (\text{A.6}) \\ & (\text{a}_{m,2,1}^{(3)}A_1 + q_{m,1,2}^{(3)}A_2 + q_{m,1,3}^{(3)}A_3, [A_1, A_2]] + \\ & (\text{A.6}) \\ & (\text{a}_{m,2,1}^{(3)}A_1 + q_{m,2,2}^{(3)}A_2 + q_{m,2,3}^{(3)}A_3, [A_1, A_3]] + \\ & (\text{A.6}) \\ & (\text{a}_{m,3,1}^{(3)}A_1 + q_{m,3,2}^{(3)}A_2 + q_{m,3,3}^{(3)}A_3, [A_2, A_3]]. \\ & (\text{A.6}) \\ & (\text{A.6}) \\ & (\text{A.7}) \\ & (\text{A.6}) \\ & (\text{A.7}) \\ & (\text{A.7}) \\ & (\text{A.8}) \\ & (\text{A.9}) \\ &$$

Finally, for the fourth term,  $\Omega_m^{(4)}$  is computed as in Eqs. (3.7) and (3.8), with  $q^{(4)} = 1/60$  and  $q_j^{(1)}$  in (3.8) replaced with  $q_{m,j}^{(1)}$  from Eq. (A.3).

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## REFERENCES

- [1] S. BLANES, F. CASAS, J. A. OTEO, AND J. ROS, The Magnus expansion and some of its
   applications, Physics Reports, 470 (2009), pp. 151–238.
- [2] S. BLANES, F. CASAS, AND J. ROS, Improved High Order Integrators Based On The Magnus
   *Expansion*, Bit, 40 (2000), pp. 434–450.
- [3] S. BLANES AND P. C. MOAN, Fourth-and sixth-order commutator-free Magnus integrators for linear and non-linear dynamical systems, Applied Numerical Mathematics, 56 (2006), pp. 1519–1537.
- [4] K. BURRAGE, Parallel methods for ODEs, Advances in Computational Mathematics, 7 (1997),
   pp. 1–3.
- [5] L. EULER, Institutiones calculi integralis, no. v. 2, Acad. Imper. scientiarum, 1768.
- [6] M. J. GANDER, 50 years of time parallel time integration, in Multiple Shooting and Time
   Domain Decomposition Methods, T. Carraro, M. Geiger, S. Körkel, and R. Rannacher, eds.,
   Cham, 2015, Springer International Publishing, pp. 69–113.
- [63] [7] E. HAIRER, C. LUBICH, AND G. WANNER, Geometric numerical integration : structure-preserving
   algorithms for ordinary differential equations, Springer, 2006.
- [8] E. HAIRER, S. P. NØRSETT, AND G. WANNER, Solving ordinary differential equations I. nonstiff
   problems, Mathematics and Computers in Simulation, 29 (1987), p. 447.
- [9] E. HAIRER AND G. WANNER, Solving Ordinary Differential Equations II : Stiff and Differential Algebraic Problems, Springer Berlin Heidelberg, 1991.
- [10] N. J. HIGHAM, The Scaling and Squaring Method for the Matrix Exponential Revisited, SIAM
   Journal on Matrix Analysis and Applications, 26 (2005), pp. 1179–1193.
- [11] M. HOCHBRUCK AND A. OSTERMANN, Exponential integrators, Acta Numerica, 19 (2010),
   pp. 209–286.
- [12] A. ISERLES, H. Z. MUNTHE-KAAS, S. P. NØRSETT, AND A. ZANNA, *Lie-group methods*, Acta
   Numerica, (2000), pp. 215–365.
- [13] A. ISERLES AND S. P. NØRSETT, On the solution of linear differential equations in Lie groups,
   Philosophical Transactions: Mathematical, Physical and Engineering Sciences, 357 (1999),
   pp. 983–1019.
- [14] W. MAGNUS, On the exponential solution of differential equations for a linear operator, Com munications on pure and applied ..., VII (1954), pp. 649–673.
- [15] C. MOLER AND C. VAN LOAN, Nineteen Dubious Ways to Compute the Exponential of a Matrix,
   Twenty-Five Years Later, SIAM Review, 45 (2003), pp. 3–49.
- [16] H. MUNTHE-KAAS AND B. OWREN, Computations in a free Lie algebra, Philosophical Transac tions of the Royal Society A: Mathematical, Physical and Engineering Sciences, 357 (1999),
   pp. 957–981.
- [17] J. NIEVERGELT, Parallel methods for integrating ordinary differential equations, Communications
   of the ACM, (1964), pp. 731–733.
- [18] M. THALHAMMER, A fourth-order commutator-free exponential integrator for nonautonomous
   differential equations, SIAM J. Numer. Anal., 44 (2006), pp. 851—864 (elect.
- [19] M. TODA, Vibration of a chain with nonlinear interaction, Journal of the Physical Society of Japan, 22 (1967), pp. 431–436.
- [20] M. VALIEV, E. BYLASKA, N. GOVIND, K. KOWALSKI, T. STRAATSMA, D. W. H.J.J. VAN DAM,
   J. NIEPLOCHA, E. APRA, T. WINDUS, AND W. DE JONG, NWChem: a comprehensive and scalable open-source solution for large scale molecular simulations, Comput. Phys.
   Commun., (2010), pp. 1477–1489.
- [21] A. ZANNA, Numerical solution of isospectral flows, PhD thesis, University of Cambridge, 1998.