Unconditionally stable integration of Maxwell’s equations

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1. Introduction

Maxwell’s equations from electromagnetism model interrelations between electric and magnetic fields. The equations form a time-dependent system of six first-order partial differential equations (PDEs). The equations appear in different forms, such as in the compact curl notation.
\[
\begin{align*}
\partial_t B &= -\nabla \times E, \\
\varepsilon \partial_t E &= \nabla \times (\mu^{-1}) B - \sigma E - J_E.
\end{align*}
\] (1.1)

Here \( B \) and \( E \) represent the magnetic and electric field, respectively. \( J_E \) is a given source term representing electric current density and \( \varepsilon, \mu \) and \( \sigma \) are (tensor) coefficients representing, respectively, dielectric permittivity, magnetic permeability and conductivity. The equations are posed in a three-dimensional spatial domain and provided with appropriate boundary conditions. If the equations are posed in domains without conductors, the damping term \(-\sigma E\) is absent. If in addition the source \( J_E \) is taken zero, we have a prime example of a conservative wave equation system.

Numerical methods for time-dependent PDEs are often derived in two stages (the method of lines approach). First, the spatial operators are discretized on an appropriate grid covering the spatial domain, together with the accompanying boundary conditions. This leads to a time-continuous, semi-discrete problem in the form of an initial-value problem for a system of ordinary differential equations (ODEs). Second, a numerical integration method for this ODE system is chosen, which turns the semi-discrete solution into the desired fully discrete solution on the chosen space–time grid. In this paper we focus on the second numerical integration stage, as in [7]. While in [7] the focus was on methods treating the curl terms explicitly, here we address the question whether fully implicit and exponential time integration eliminating any temporal step size stability restriction can be feasible and efficient.

As in [7] we start from the general space-discretized Maxwell problem

\[
\begin{pmatrix}
M_u & 0 \\
0 & M_v
\end{pmatrix}
\begin{pmatrix}
u' \\
v
\end{pmatrix} =
\begin{pmatrix}
0 & -K \\
-K^T & -S
\end{pmatrix}
\begin{pmatrix}
u' \\
v
\end{pmatrix} + \begin{pmatrix} j_u \\
0
\end{pmatrix},
\] (1.2)

where \( u = u(t) \) and \( v = v(t) \) are the unknown vector (grid) functions approximating the values of the magnetic flux \( B \) and electric field \( E \), respectively. The matrices \( K \) and \( K^T \) approximate the curl operator and the matrix \( S \) is associated with the dissipative conduction term. Throughout \( S \) can be assumed symmetric positive semi-definite, \( M_u \) and \( M_v \) are symmetric positive definite mass matrices possibly arising from a spatial finite element or compact finite difference discretization. The functions \( j_u(t) \) and \( j_v(t) \) are source terms. Typically, \( j_u \) represents the given source current \( J_E \), but \( j_u \) and \( j_v \) may also contain boundary data. We do allow \( u \) and \( v \) to have different dimensions which can occur with certain finite-element methods, see e.g. [40], and assume \( u \in \mathbb{R}^m, v \in \mathbb{R}^n \) with \( m, n \geq 1 \) and \( M_u \in \mathbb{R}^{m \times m}, M_v \in \mathbb{R}^{n \times n}, K \in \mathbb{R}^{m \times m}, S \in \mathbb{R}^{n \times n} \). The ODE system (1.2) is generic in the sense that spatial discretization of \((H,E)\)-formulations of the Maxwell equations also lead to this form, see Section 4 of [7].

We allow the space grids underlying (1.2) to be non-Cartesian. This has an important consequence in that it excludes the well-known unconditionally stable alternating direction implicit–finite difference time domain method attributed to [50,51], see also [13,17,20,35] and references therein. We will instead focus on conventional fully implicit integration (Section 3) and on exponential integration (Section 4). This means that we need efficient solvers from the field of numerical linear algebra. For solving the systems of linear algebraic equations arising with implicit integrators we will use the conjugate gradient (CG) iterative method with preconditioning (Section 3). For exponential integration we will consider Krylov subspace iteration (Section 4). Both for the theory behind CG and Krylov iteration we refer to the text books [43,48]. Seminal papers on Krylov subspace iteration for matrix functions are [14,15,22,23,29,41,47].

2. Stability and conservation properties

To begin with, we recall from [7] some stability and conservation properties of the semi-discrete system (1.2). Let \( w \in \mathbb{R}^{m+n} \) denote the solution vector composed by \( u \in \mathbb{R}^m \) and \( v \in \mathbb{R}^n \). A natural norm for establishing stability and conservation is the inner-product norm

\[
\|w\|^2 = \|u\|_{M_u}^2 + \|v\|_{M_v}^2, \quad \langle u, u \rangle_{M_u} = \langle M_u u, u \rangle, \quad \langle v, v \rangle_{M_v} = \langle M_v v, v \rangle.
\] (2.1)

where \( \langle \cdot, \cdot \rangle \) denotes the \( L_2 \) inner product. As \( S \) is symmetric positive semi-definite, for the homogeneous part of (1.2) follows:
\[
\frac{d}{dt} \|w\|^2 = -2 \langle Sv, v \rangle \leq 0
\]  
(2.2)

showing stability in the \( L_2 \) sense and (energy) conservation for a zero matrix \( S \). It is desirable that integration methods respect these properties, either exactly or to sufficiently high accuracy.

For the purpose of analysis a formulation without mass matrices equivalent to (1.2) is obtained as follows. Introduce the Cholesky factorizations \( L_{Mc}L_{Mc}^T = M_u \) and \( L_{Mc}L_{Mc}^T = M_v \). Then
\[
\begin{pmatrix}
\tilde{u}' \\
\tilde{v}'
\end{pmatrix} = \begin{pmatrix} 0 & -\tilde{K} \\
\tilde{K}^T & -\tilde{S} \end{pmatrix} \begin{pmatrix} \tilde{u} \\
\tilde{v}
\end{pmatrix} + \begin{pmatrix} j_u \\
j_v
\end{pmatrix},
\]  
(2.3)

where \( \tilde{u} = L_{Mc}^T u, \tilde{v} = L_{Mc}^T v \) and
\[
\tilde{K} = L_{Mc}^{-1} KL_{Mc}^{-T}, \quad \tilde{S} = L_{Mc}^{-1} SL_{Mc}^{-T}, \quad j_u = L_{Mc}^{-1} j_u, \quad j_v = L_{Mc}^{-1} j_v.
\]  
(2.4)

Next introduce the transformed inner-product norm
\[
\|\tilde{w}\|^2 = \|\tilde{u}\|^2 + \|\tilde{v}\|^2, \quad \|\tilde{u}\|^2 = \langle \tilde{u}, \tilde{u} \rangle, \quad \|\tilde{v}\|^2 = \langle \tilde{v}, \tilde{v} \rangle
\]  
(2.5)

for the vector \( \tilde{w} \) composed of \( \tilde{u} \) and \( \tilde{v} \). For the homogeneous part of (2.3) then follows immediately:
\[
\frac{d}{dt} \|\tilde{w}\|^2 = -2 \langle \tilde{S} \tilde{v}, \tilde{v} \rangle \leq 0,
\]  
(2.6)

while the norm is preserved under the transformation, that is, \( \|\tilde{w}\|^2 = \|w\| \) and \( \langle \tilde{S} \tilde{v}, \tilde{v} \rangle = \langle Sv, v \rangle \). We note that the transformed system is introduced for analysis purposes only and that our numerical methods will be applied to system (1.2).

If in (1.1) the conductivity coefficient \( \sigma \) and the permittivity coefficient \( \varepsilon \) are constant scalars instead of space-dependent tensors \( (3 \times 3 \) matrices), then the matrices \( M_v \) and \( S \) from (1.2) can be assumed identical up to a constant, implying that the matrix \( \tilde{S} \) introduced in (2.3) becomes the constant diagonal matrix
\[
\tilde{S} = \alpha I, \quad \alpha = \frac{\sigma}{\varepsilon}.
\]  
(2.7)

This enables the derivation of a two-by-two system for the sake of further analysis. Introduce the singular-value decomposition \( \tilde{K} = U \Sigma V^T \) where \( U \in \mathbb{R}^{m \times m} \) and \( V \in \mathbb{R}^{n \times n} \) are orthogonal and \( \Sigma \) is a diagonal \( m \times n \) matrix with nonnegative diagonal entries \( s_1, \ldots, s_m \) satisfying
\[
s_1 \geq s_2 \geq \cdots \geq s_r > s_{r+1} = \cdots = s_m = 0.
\]  
(2.8)

Here \( r \leq m \) is the (row) rank of \( \tilde{K} \) and the \( s_i \) are the singular values of the matrix \( \tilde{K} \) (the square roots of the eigenvalues of \( \tilde{K} \tilde{K}^T \)). The transformed variables and source terms
\[
\tilde{u}(t) = U^T u(t), \quad \tilde{v}(t) = V^T v(t), \quad j_u(t) = U^T j_u(t), \quad j_v(t) = V^T j_v(t)
\]  
(2.9)

satisfy the equivalent ODE system
\[
\begin{pmatrix}
\tilde{u}' \\
\tilde{v}'
\end{pmatrix} = \begin{pmatrix} 0 & -\Sigma \\
\Sigma^T & -\alpha I \end{pmatrix} \begin{pmatrix} \tilde{u} \\
\tilde{v}
\end{pmatrix} + \begin{pmatrix} j_u \\
j_v
\end{pmatrix},
\]  
(2.10)

where \( I \) is the \( n \times n \) identity matrix. Note that the matrix transformation induced by (2.9) is a similarity transformation, so that the matrices of systems (2.3) and (2.10) have the same eigenvalues. Further, \( \|\tilde{w}\|^2 = \|\tilde{u}\|^2 + \|\tilde{v}\|^2 \) due to the orthogonality of \( U \) and \( V \). Thus, if (2.7) applies, the stability of a time integration method may be studied for the homogeneous part of (2.10), provided also the method is invariant under the transformations leading to (2.10).

Since the matrix \( \Sigma \) is diagonal, the homogeneous part of (2.10) decouples into \( r \) two-by-two systems
\[
\begin{pmatrix}
\tilde{u}' \\
\tilde{v}'
\end{pmatrix} = \begin{pmatrix} 0 & -s \\
s & -\alpha \end{pmatrix} \begin{pmatrix} \tilde{u} \\
\tilde{v}
\end{pmatrix}, \quad s = s_k > 0, \quad k = 1, \ldots, r,
\]  
(2.11)

\( m - r \) scalar equations \( \tilde{u}' = 0 \), and \( n - r \) scalar equations \( \tilde{v}' = -\alpha \tilde{v} \). From the viewpoint of time integration stability, these equations are canonical for Maxwell equation systems of which the conductivity
coefficient \( \sigma \) and the permittivity coefficient \( \varepsilon \) are constant scalars. For stability analysis we thus arrive at the two-by-two test model
\[
\begin{pmatrix}
\hat{u}' \\
\hat{v}'
\end{pmatrix} = \begin{pmatrix}
0 & -s \\
s & -\alpha
\end{pmatrix} \begin{pmatrix}
\hat{u} \\
\hat{v}
\end{pmatrix}, \quad s \geq 0, \; \alpha \geq 0.
\] (2.12)

Stability for this test model is equivalent to stability for (2.10), which in turn is equivalent to stability for the original semi-discrete Maxwell system (1.2), provided the conductivity coefficient \( \sigma \) and the permittivity coefficient \( \varepsilon \) are constant scalars. The eigenvalues of (2.12) are
\[
(-\alpha \pm \sqrt{\alpha^2 - 4s^2})/2.
\]
Assuming sufficiently small and large singular values \( s_k \) in (2.8), the spectra of (2.3) and (2.10) thus are cross-shaped with real eigenvalues between \(-\alpha\) and 0 and complex eigenvalues with real part \(-\alpha/2\) and imaginary parts \(\pm \sqrt{4s^2/2^2 - \alpha^2/2}\).

3. Implicit integration

We will examine fully implicit time stepping for (1.2) for the second-order implicit trapezoidal rule (ITR). This method has the right stability and conservation properties for Maxwell’s equations and shares the numerical algebra challenge with many other implicit methods, such as diagonally-implicit Runge–Kutta methods. So numerical algebra conclusions drawn for ITR are also valid for related higher-order methods. In this paper, we focus on second-order methods because the order of the spatial discretization scheme for the 3D example problem used for testing is also limited to two. Before discussing ITR we first recall an economical second-order implicit–explicit method called CO2 (Composition 2nd-order) in [7] which serves as a reference method.

3.1. The implicit–explicit method CO2

Method CO2 is given by
\[
\begin{align*}
M_u u_{n+1/2} - u_n \\
M_v v_{n+1} - v_n \\
M_u u_{n+1} - u_{n+1/2}
\end{align*}
\]
\[
\begin{array}{c}
= -\frac{1}{2}Kv_n + \frac{1}{2}j_u(t_n), \\
= K^T u_{n+1/2} - \frac{1}{2}S(v_n + v_{n+1}) + \frac{1}{2}(j_v(t_n) + j_v(t_{n+1})), \\
= -\frac{1}{2}Kv_{n+1} + \frac{1}{2}j_u(t_{n+1}).
\end{array}
\] (3.1)

Like ITR this method is a one-step method stepping from \((u_n, v_n)\) to \((u_{n+1}, v_{n+1})\) with step size \(\tau\). Here \(u_n\) denotes the approximation to the exact solution \(u(t_n)\), etc., and \(\tau = t_{n+1} - t_n\). The subindex \(n\) should not be confused with the length of the vector \(v\) in (1.2). CO2 is symmetric and treats the curl terms explicitly and the conduction term implicitly. Of practical importance is that the third-stage derivative computation can be copied to the first stage at the next time step to save computational work. Per time step this method thus is very economical. Apart from the mass matrices (see Remark 3.1), the method requires a single explicit evaluation of the full derivative per integration step which is the least possible.

In contrast to ITR, method (3.1) is not unconditionally stable and a sharp step size bound for stability for the general system (1.2) is not known up to now. However, for Maxwell problems for which (2.7) holds stability can be concluded from the \(2 \times 2\)-model (2.11). Let \(z_\delta = \tau s_{\text{max}}\). The resulting step size bound is then valid for (1.2) and is given by
\[
z_\delta < 2 \quad \text{if } \alpha = 0 \text{ and otherwise } z_\delta \leq 2.
\] (3.2)

Hence the conduction puts no limit on \(\tau\). Recall that \(\alpha = 0\) in the absence of conduction and that \(s_{\text{max}}\) here is to be taken as the maximal square root of the eigenvalues of \(\hat{K}\hat{K}^T\). Because \(K\) approximates the first-order curl operator these eigenvalues are proportional to \(h^{-2}\) where \(h\) represents a minimal spatial grid size. So for time stepping stability, a relation \(\tau \sim h\) for \(h \to 0\) is required. On fine space grids and long time intervals this may lead to large amounts of time steps.
It is this observation which underlies the question whether implicit or exponential integration is feasible and competitive so as to enhance time stepping efficiency. For the derivation and further discussion of this method we refer to [7] where it was called CO2 as it is of second order and based on COMposition of a certain partitioned Euler rule. One of the results in [7] states that the second-order behavior is maintained in the presence of time-dependent boundary conditions (stiff source terms). A similar result will be proven in the appendix (Section 5) for the exponential integrator EK2 derived in Section 4. Finally, with regard to time stepping CO2 bears a close resemblance to the popular time-staggered Yee-scheme [49] and as such is a natural candidate for a reference method.

**Remark 3.1.** The mass matrices naturally give rise to implicitness such that we encounter at each integration step one linear system solution with $M_u$ and $M_v + \frac{1}{2} \tau S$. Systems with mass matrices can be (approximately) solved in an efficient way. This can be achieved either by fast solvers (sparse direct or preconditioned iterative) or by special mass lumping techniques. For mass lumping of the finite-element discretization used in Section 3.5, see e.g. [2, 19]. For keeping our assessments as general as possible we will use the original non-lumped form. Throughout this paper (so also for the other integration methods) we will use sparse Cholesky factorization to realize the mass matrix inversions. For constant $\tau$ the factorization should only be carried out once at the start of the integration leaving only sparse forward–backward substitutions during the time stepping.

### 3.2. The implicit trapezoidal rule ITR

Denote (1.2) by

$$Mw' = Aw + g(t),$$

(3.3)

where

$$w = \left( \begin{array}{c} u \\ v \end{array} \right), \quad M = \left( \begin{array}{cc} M_u & 0 \\ 0 & M_v \end{array} \right), \quad A = \left( \begin{array}{cc} 0 & -K \\ K^T & -S \end{array} \right), \quad g(t) = \left( \begin{array}{c} j_u \\ j_v \end{array} \right).$$

(3.4)

ITR then reads

$$M w_{n+1} - \frac{\tau}{2} w_n + Aw_n + g(t_n) + g(t_{n+1}).$$

(3.5)

This classical implicit method mimics the stability and conservation property (2.2). That is, for zero sources,

$$\frac{\|w_{n+1}\|^2 - \|w_n\|^2}{\tau} = -2 \left( S \frac{v_{n+1} + v_n}{2}, \frac{v_{n+1} + v_n}{2} \right) \quad \forall \tau > 0.$$

(3.6)

Hence (3.5) is unconditionally stable (and conservative for zero $S$). Like for CO2 the method is second-order consistent, also for stiff source terms emanating from time-dependent boundary functions. From that perspective the method is ideal, however, at the costs of solving each time step the linear system

$$\left( M - \frac{1}{2} \tau A \right) w_{n+1} = \left( M + \frac{1}{2} \tau A \right) w_n + \frac{1}{2} \tau g(t_n) + \frac{1}{2} \tau g(t_{n+1}).$$

(3.7)

for the matrix

$$M - \frac{1}{2} \tau A = \left( \begin{array}{cc} M_u & \frac{1}{2} \tau K \\ -\frac{1}{2} \tau K^T & M_v + \frac{1}{2} \tau S \end{array} \right).$$

(3.8)

Sparse LU-decomposition will become too costly in memory for large-scale 3D simulations. We therefore focus on iteration whereby we rewrite (Schur complement) the linear system (3.7) to an equivalent form which is significantly more amenable for iterative solution. Note that (3.7) and (3.8) can be brought to the saddle point form by changing the sign of the second block row in (3.8). The Schur complement approach for solving saddle point systems, possibly in combination with CG, is well known (see e.g. [3]).
Let \( r_u, r_v \) denote the right-hand sides of (3.7) belonging to \( u, v \). Suppressing the time index \( n + 1 \) in \( u_{n+1}, v_{n+1} \) this system then reads
\[
M_u u + \frac{1}{2} \tau K v = r_u,
\]
\[
-\frac{1}{2} \tau K^T u + M_v v + \frac{1}{2} \tau S v = r_v.
\]
(3.9)

Since the mass matrix \( M_u \) is symmetric positive definite, we can multiply the first equation from left by \( \frac{1}{2} \tau K M_u^{-1} u \). Then adding the two equations yields the equivalent system
\[
\mathcal{M} u = r_u,
\]
\[
\mathcal{M} v = r_v + \frac{1}{2} \tau K^T M_u^{-1} r_u,
\]
(3.10)

wherein \( u \) has been eliminated from the second equation. The \( n \times n \)-matrix \( \mathcal{M} \) is given by
\[
\mathcal{M} = M_v + \frac{1}{2} \tau S + \frac{1}{4} \tau^2 K^T M_u^{-1} K.
\]
(3.11)

So we can first solve \( v \) from the second equation and subsequently \( u \) from the first. Hereby we assume that the three inversions for \( M_u \) are carried out through sparse Cholesky decomposition, entirely similar as for method CO2. Of main interest is that \( \mathcal{M} \) is symmetric positive definite which calls for the iterative conjugate gradient (CG) method.

3.3. CG convergence

Let us first assess the convergence of the CG method. For this purpose we employ the transformation underlying system (2.3) which can be shown to be equivalent to Cholesky factorization preconditioning with the mass matrix \( M_v \), see also Section 3.4. The counterpart of (3.10) then reads
\[
\tilde{u} + \frac{1}{2} \tau \tilde{K} \tilde{v} = \tilde{r}_u,
\]
\[
\tilde{\mathcal{M}} v = \tilde{r}_v + \frac{1}{2} \tau \tilde{K}^T \tilde{r}_u
\]
(3.12)

with the straightforward definition of \( \tilde{r}_u, \tilde{r}_v \) and
\[
\tilde{\mathcal{M}} = I + \frac{1}{2} \tau \tilde{S} + \frac{1}{4} \tau^2 \tilde{K}^T \tilde{K}.
\]
(3.13)

CG is a natural choice as it optimal in the following sense [48]: for any initial guess \( \tilde{v}_0 \) it computes iterants \( \tilde{v}_i \) which satisfy the polynomial relation
\[
\tilde{v}_i - \tilde{v} = P_i(\tilde{\mathcal{M}})(\tilde{v}_0 - \tilde{v})
\]
(3.14)

such that in the \( \tilde{\mathcal{M}} \)-norm the iteration error \( \| \tilde{v}_i - \tilde{v} \|_{\tilde{\mathcal{M}}} \) is minimal over the set of all polynomials \( P_i \) of degree \( i \) satisfying \( P_i(0) = 1 \). This iteration error satisfies the well-known bound
\[
\| \tilde{v}_i - \tilde{v} \|_{\tilde{\mathcal{M}}} \leq 2 \left( \frac{\sqrt{\kappa} - 1}{\sqrt{\kappa} + 1} \right)^i \| \tilde{v}_0 - \tilde{v} \|_{\tilde{\mathcal{M}}},
\]
(3.15)

where \( \kappa \) is the spectral condition number of \( \tilde{\mathcal{M}} \), that is, \( \kappa = \lambda_{\text{max}}/\lambda_{\text{min}} \) being the quotient of the maximal and minimal eigenvalue of \( \mathcal{M} \). This upper bound does not reflect the celebrated superlinear convergence [45] of CG which makes it a truly successful solver. However, the bound does provide a rough assessment of the potential of the combination ITR–CG in relation to CO2. Hereby it is noted that in good approximation a single CG iteration with matrix \( \mathcal{M} \) is cost wise equal to a single CO2 step.

\[\text{footnote}{1}\text{The subindex } i \text{ should not be confused with the subindex } n \text{ used to denote a time level } t_n.\]
Would $\tilde{S}$ and $\tilde{R}^T \tilde{K}$ commute, the condition number can be derived directly from the spectra of $\tilde{S}$ and $\tilde{R}^T \tilde{K}$. In the general case commutation will be rare. Therefore, we next assume that we do have a Maxwell problem for which condition (2.7) holds. Then we have commutation and the eigenvalues $\lambda$ of $\tilde{M}$ are given by

$$\lambda = 1 + \frac{1}{2} \tau \alpha + \frac{1}{4} \tau^2 s^2,$$

(3.16)

where $s^2$ is an eigenvalue of $\tilde{R}^T \tilde{K}$ the square root of which also features in (2.11). Hence

$$\lambda_{\text{min}} = 1 + \frac{1}{2} \tau \alpha + \frac{1}{4} \tau^2 s_{\text{min}}^2, \quad \lambda_{\text{max}} = 1 + \frac{1}{2} \tau \alpha + \frac{1}{4} \tau^2 s_{\text{max}}^2.$$

(3.17)

Regarding ITR we are only interested in step sizes $\tau$ such that $z_s = \tau s_{\text{max}} \gg 2$ because otherwise method CO2 will be more efficient, see the step size bound (3.2). Since $s_{\text{max}}$ is proportional to $h$ which represents the minimal spatial grid size, we then may neglect the contribution $\tau \alpha$ and approximate $\kappa$ by

$$\kappa \approx 1 + \frac{1}{4} z_s^2$$

(3.18)

showing that one CG iteration reduces the initial iteration error in the $\tilde{M}$-norm by

$$\nu(z_s) = \frac{\sqrt{1 + \frac{1}{4} z_s^2} - 1}{\sqrt{1 + \frac{1}{4} z_s^2} + 1} \sim 1 - \frac{4}{z_s} \sim e^{-4 / z_s}, \quad z_s \to \infty.$$  

(3.19)

Unfortunately, this reduction factor is by far too low for ITR implemented with CG to become a competitive method. To see this the following argument suffices. For $z_s \gg 2$ the number of CG iterations for an overall reduction factor $\epsilon$ is approximately $j = -\frac{1}{4} z_s \ln(\epsilon)$. Because each iteration is in good approximation as expensive as a single integration step with method CO2, we can afford $j$ steps with CO2 with step size $\tau / j$ (provided we have stability of CO2), that is, if $z_s / j \lesssim 2$. Inserting $j$ this appears to hold for all $\epsilon \lesssim e^{-2} \approx 10^{-1}$. When iterating with CG an error reduction of the initial error by a factor 10 is of course quite poor and we can conclude that the computational effort is better spent in applying CO2 with a step size $\tau / j$. This will lead to a notable smaller time stepping error for comparable effort since ITR and CO2 are both of second order. Clearly, ITR will not be competitive to CO2 unless superlinear CG convergence, not reflected by (3.15), takes place and/or CG is applied with a more efficient preconditioner.

3.4. CG implementation

CG was implemented for the following reformulation of the ITR scheme (3.7):

$$\begin{pmatrix} M_u & \frac{1}{2} K \\ -\frac{1}{2} K^T & M_v + \frac{1}{2} S \end{pmatrix} \begin{pmatrix} \Delta u \\ \Delta v \end{pmatrix} = \begin{pmatrix} b_u \\ b_v \end{pmatrix},$$

(3.20)

where

$$\begin{pmatrix} b_u \\ b_v \end{pmatrix} = \begin{pmatrix} 0 \\ \tau K^T \end{pmatrix} \begin{pmatrix} u_n \\ v_n \end{pmatrix} + \frac{\tau}{2} \begin{pmatrix} j_u(t_n) + j_u(t_{n+1}) \\ j_v(t_n) + j_v(t_{n+1}) \end{pmatrix}$$

(3.21)

and $u_{n+1} = u_n + \Delta u$, $v_{n+1} = v_n + \Delta v$. Hereby the linear system was treated with the Schur complement as described above. Writing (3.7) in this form is beneficial since this makes the zero vector a natural initial guess for the iteration process and saves one matrix–vector multiplication which is otherwise needed for the initial residual.\(^2\)

For an efficient usage it is important to choose a proper stopping criterion for CG. Too many iterations would mean a waste of effort, whereas too few might cause loss of stability.\(^3\) Using, for convenience,
the same notation for $\Delta u$ and $\Delta v$, solving system (3.7) approximately with residual $r_{cg}$ effectively means that the found $\Delta v$ is a solution of the perturbed linear system

$$M_{\Delta v} = \frac{\tau}{2} K^T M^{-1}_u b_u + b_v + r_{cg},$$  \hspace{1cm} (3.22)

where $M$ is defined in (3.11) and the approximate solution $\Delta u, \Delta v$ of (3.20) satisfies

$$\begin{pmatrix} M_u & \frac{\tau}{2} K \\ -\frac{\tau}{2} K^T & M_v + \frac{\tau}{2} S \end{pmatrix} \begin{pmatrix} \Delta u \\ \Delta v \end{pmatrix} = \begin{pmatrix} b_u \\ b_v \end{pmatrix} + \begin{pmatrix} 0 \\ r_{cg} \end{pmatrix}.$$  \hspace{1cm} (3.23)

We stop CG as soon as for a certain constant $\delta$

$$\| r_{cg} \|_2 \leq \frac{\tau}{2} K^T M^{-1}_u b_u \|_2 \delta,$$  \hspace{1cm} (3.24)

which means that the inexact ITR–CG scheme (3.23) can be seen as a perturbed form of the exact ITR scheme (3.20) where the perturbations are kept bounded. The purpose of this inequality is to enforce $r_{cg}$ to be a fraction of the local truncation error of ITR for component $v$ which we aim by means of an educated guess for $\delta$. Note that $r_{cg}$ just becomes the local truncation error upon substitution of the exact ODE solution. Choosing $\delta$ too large implies of course loss of ITR accuracy, whereas a too small $\delta$ wastes matvecs. We will give actual values of $\delta$ when we report our test results.

For the CG solution of the Schur complement system with the matrix $M$ we have used two preconditioners. The first one is the sparse Cholesky factorization of the mass matrix $M_v$, the second is the incomplete-Cholesky (IC) factorization with the drop tolerance $\varepsilon = 10^{-6}$ [32,42] applied to the matrix

$$M_v + \frac{\tau}{2} S + \frac{\tau^2}{4} K^T K$$  \hspace{1cm} (3.25)

generated from $M$ by deleting $M^{-1}_u$. The mass matrix preconditioner is readily available and as argued earlier, for ITR the costs of one mass matrix preconditioned CG iteration are roughly the same as the costs of one time step with CO2. This is because one CG iteration requires just one matvec with the preconditioned matrix (and several vector updates).

The IC($\varepsilon$) preconditioner requires significant set up time. For example, for the fourth grid of Table 3.1 given in Section 3.5 the preparation cost required a CPU time sufficient for performing 90–100 matvecs with the preconditioned matrix $M$. An attractive property of the IC($\varepsilon$) preconditioner compared to the mass matrix preconditioner is a higher level of sparsity. For example, for $\varepsilon = 10^{-6}$ the sparsity is at least twice as large as for the Cholesky factors of the mass matrix. During integration the IC($\varepsilon$) preconditioner therefore is slightly cheaper due to the higher level of sparsity. In our experiments, we found little differences between numbers of iterations for the mass matrix and IC($\varepsilon$) preconditioner. We therefore will report only iteration numbers for the first one. Note that the eigenvalues of the mass matrix preconditioned $M$ are given by (3.16) if we do have a Maxwell problem for which condition (2.7) holds.

3.5. Comparing ITR and CO2

In this section, we compare the fully implicit integrator ITR, equipped with the above described preconditioned CG implementation, to method CO2.

3.5.1. A 3D Maxwell test problem

The comparison is based on tests with a three-dimensional (3D) Maxwell problem we earlier used in [7]. This problem is given in the $(H,E)$ formulation

$$\begin{align*}
\mu \partial_t H &= -\nabla \times E, \\
\varepsilon \partial_t E &= \nabla \times H - \sigma E - J
\end{align*}$$  \hspace{1cm} (3.26)

with independent variables $(x,y,z) \in \Omega \subset \mathbb{R}^3$, $t \in [0,T]$, and initial and boundary conditions

---

4 Here and in the remainder $\| \cdot \|_2$ denotes the discrete inner-product ($L_2$) norm.
The coefficients $\mu, \varepsilon$ and $\sigma$ are taken constant in time and space and $\vec{n}$ denotes the outward unit normal vector to the boundary $\partial \Omega$. The boundary functions $E_{bc}$ and $H_{bc}$ vary in space and time. Specifically, $\Omega = [0, 1]^3$ and $T = 10$ and we choose the source current $J = J(x, y, z, t)$ such that the Maxwell system (3.26) allows a specific exact solution

$$E(x, y, z, t) = \alpha(t)E_{stat}(x, y, z), \quad H(x, y, z, t) = \beta(t)H_{stat}(x, y, z),$$

where the scalar functions $\alpha, \beta$ and the vector functions $E_{stat}, H_{stat}$ satisfy $\mu \beta'(t) = -\alpha(t)$ and $H_{stat} = \nabla \times E_{stat}$. The source function $J$ is then defined as

$$J(x, y, z, t) = -\varepsilon \alpha'(t) + \sigma \alpha(t))E_{stat}(x, y, z) + \beta(t)\nabla \times H_{stat}(x, y, z)$$

with

$$E_{stat}(x, y, z) = \begin{pmatrix} \sin \pi y \\ \sin \pi x \\ \sin \pi y \end{pmatrix}, \quad H_{stat}(x, y, z) = \begin{pmatrix} \pi \sin \pi x (\cos \pi y - \cos \pi z) \\ \pi \sin \pi y (\cos \pi x - \cos \pi z) \\ \pi \sin \pi z (\cos \pi x - \cos \pi y) \end{pmatrix},$$

$$\alpha(t) = \frac{3}{\mu} \sum_{k=1}^{3} \cos \omega_k t, \quad \beta(t) = -\frac{1}{\mu} \sum_{k=1}^{3} \sin \omega_k t$$

and $\omega_1 = 1, \omega_2 = 1/2, \omega_3 = 1/3$. Further, we take $\mu = 1, \varepsilon = 1$ and either $\sigma = 0$ or $\sigma = 60\pi$ (this corresponds with values encountered in real applications).

This 3D Maxwell problem is spatially discretized with first-order, first-type Nédélec edge finite elements on tetrahedral unstructured grids [34,36,37]. Although it is formulated with $H$ and $E$ as primary variables, the resulting semi-discrete system belongs to class (1.2). In [7], we observed first-order spatial convergence for $E$ and second order for $H$. We have used the grids numbered four and five listed in Table 3.1 which displays grid parameters and step size information for CO2. To save space we refer to [7] and references therein for a more complete description of this test problem and its spatial discretization.

### 3.5.2. Test results

Table 3.2 reports computational costs in terms of matvecs for CO2 and ITR–CG for the fourth and fifth grid mentioned in Table 3.1. Two cases are distinguished, the zero conduction coefficient $\sigma = 0$ and the nonzero conduction coefficient $\sigma = 60\pi$, see Section 3.5.1. For both cases we have chosen $\delta = 0.05$ in the stopping criterion (3.24) and step sizes $\tau$ for ITR–CG much larger than the limit step size of CO2. For the chosen values the temporal errors remain smaller than the spatial ones, justifying the use of ITR–CG with respect to the full discretization error.

Our first observation is that the number of CG iterations per ITR time step grows only sublinearly with the time step size $\tau$, in particular for $\sigma = 60\pi$. For this reason ITR can become faster than CO2 for

<table>
<thead>
<tr>
<th>Grid</th>
<th>Number of edges</th>
<th>Longest edge $h_{\text{max}}$</th>
<th>Shortest edge $h_{\text{min}}$</th>
<th>CO2 time step restriction</th>
<th>CO2 time step used</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>105</td>
<td>0.828</td>
<td>0.375</td>
<td>0.47</td>
<td>0.2</td>
</tr>
<tr>
<td>2</td>
<td>660</td>
<td>0.661</td>
<td>0.142</td>
<td>0.18</td>
<td>0.1</td>
</tr>
<tr>
<td>3</td>
<td>4632</td>
<td>0.359</td>
<td>0.0709</td>
<td>0.079</td>
<td>0.05</td>
</tr>
<tr>
<td>4</td>
<td>34608</td>
<td>0.250</td>
<td>0.0063</td>
<td>0.028</td>
<td>0.025</td>
</tr>
<tr>
<td>5</td>
<td>85308</td>
<td>0.118</td>
<td>0.0139</td>
<td>0.014</td>
<td>0.0125</td>
</tr>
</tbody>
</table>
Table 3.2

Computational costs of CO2 (applied with maximal \( \tau \)) versus the costs of ITR–CG (applied with different \( \tau \)); stopping criterion parameter \( \delta = 0.05 \).

<table>
<thead>
<tr>
<th>Grid</th>
<th>CO2</th>
<th>ITR/mass</th>
<th>ITR/mass</th>
<th>ITR/mass</th>
<th>ITR/mass</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>( \tau = 0.025 )</td>
<td>( \tau = 0.0625 )</td>
<td>( \tau = 0.125 )</td>
<td>( \tau = 0.25 )</td>
<td>( \tau = 1.0 )</td>
</tr>
<tr>
<td></td>
<td>( \sigma = 0 ) # matvecs per t.step</td>
<td>( \sigma = 0 ) total # matvecs per t.step</td>
<td>( \sigma = 60 \tau ) # matvecs per t.step</td>
<td>( \sigma = 60 \tau ) total # matvecs</td>
<td></td>
</tr>
<tr>
<td>Grid 4</td>
<td>1</td>
<td>4.94</td>
<td>8.99</td>
<td>15.95</td>
<td>29.6</td>
</tr>
<tr>
<td></td>
<td>400</td>
<td>790</td>
<td>719</td>
<td>638</td>
<td>296</td>
</tr>
<tr>
<td></td>
<td>1</td>
<td>2.00</td>
<td>2.01</td>
<td>2.98</td>
<td>4.60</td>
</tr>
<tr>
<td></td>
<td>400</td>
<td>320</td>
<td>161</td>
<td>119</td>
<td>77</td>
</tr>
<tr>
<td>Grid 5</td>
<td>1</td>
<td>31.52</td>
<td>47.5</td>
<td>57.8</td>
<td></td>
</tr>
<tr>
<td></td>
<td>800</td>
<td>1261</td>
<td>950</td>
<td>578</td>
<td></td>
</tr>
<tr>
<td></td>
<td>1</td>
<td>5.3</td>
<td>6.65</td>
<td>7.6</td>
<td></td>
</tr>
<tr>
<td></td>
<td>800</td>
<td>212</td>
<td>133</td>
<td>76</td>
<td></td>
</tr>
</tbody>
</table>

Fig. 3.1. The full \( L_2 \) error (left \( E \)-field, right \( H \)-field) versus time for CO2 and ITR on the fourth grid, \( \sigma = 0, \delta = 0.05 \). ITR uses the mass matrix preconditioner. The costs for these runs are 400 matvecs for CO2, 638 for ITR with \( \tau = 0.25 \) and 296 for ITR with \( \tau = 1.0 \) (see Table 3.2).

sufficiently large \( \tau \) if \( \delta \) is chosen properly (which appears to hold for \( \delta = 0.05 \)). Taking \( \delta \) 10 times smaller results for the fourth grid and \( \sigma = 0 \) in the matvec sequence \((1088, 1020, 945, 827, 668)\), showing a greater expense than CO2 for the larger step sizes. Likewise, for \( \sigma = 60\tau \) we find the sequence \((345, 250, 158, 117, 69)\), showing only a small expense growth for \( \delta \) 10 times smaller. As anticipated, the numbers increase as the grid gets finer. However, as the grid gets finer, the maximum allowable time step for CO2 does decrease too. This is also the case on the finest fifth grid even though it is more uniform than the fourth one, see Table 3.1.

Our second observation concerning Table 3.2 is that the number of CG iterations per time step for \( \sigma = 60\tau \) is significantly smaller than for \( \sigma = 0 \). The reason is that for the current test problem \( M_V \) and \( S \) are identical up to a constant, see Section 2. Hence, for growing \( \sigma \), the eigenvalues of the mass-preconditioned matrix given by (3.16) get more clustered around \( 1 + \alpha \tau / 2 \) and the condition number \( \lambda_{\text{max}} / \lambda_{\text{min}} \) decreases.

Note that in the ITR scheme one needs to repeatedly solve the linear system (3.20) where the matrix remains the same and only the right-hand side changes per time step. This suggests that computational effort can be saved by reusing the information generated by CG. One possible way of doing this is Method 2 of [18] which essentially consists of storing an orthonormal basis spanning the successive CG solutions and starting every new CG process with a projection on the stored subspace. As evidenced in [18], Method 2 can lead to a significant saving in the total number of iterations. We have tested the method for this problem but have not observed any improvement. This is because the right-hand
side of (3.20) changes quite significantly from one time step to another, thus making the projection procedure futile.

For \( \delta = 0.05, \sigma = 0 \) and the fourth grid, Fig. 3.1 shows the time evolution of full (space and time) errors in \( \| \cdot \|_2 \) for CO2 and ITR. We see that the errors are comparable and more or less independent of \( \tau \) which illustrates that the spatial error dominates. This is the sort of situation where implicit time stepping can be competitive. Our test with \( \sigma = 0 \) (undamped case) shows no distinct advantage when counting numbers of matvecs. On the other hand, the test with \( \sigma = 60\pi \) is no doubt promising and warrants further investigation with a more sophisticated CG implementation, finer space grids and more test examples including variable conduction.

4. Exponential integration

The implicit trapezoidal rule ITR is a conventional method in the sense that it is a representative of the Runge–Kutta and linear multistep methods. The so-called exponential integration methods form another class being built on linearization and direct use of accurate, unconditionally stable approximations to the exponential operator. For this reason they are of potential interest to the Maxwell equations. Exponential integrators do have a long history \([10,21,22,27,30,31,38,46]\) and have undergone a revival during the last decade, see e.g. \([4,9,11,24,26,33]\). An important reason for this revival is renewed attention for the Krylov subspace iteration technique for approximating the exponential and the so-called derived \( \psi \)-functions. In this section, we will also use Krylov subspace iteration.

4.1. The exponential integrator EK2

For formulating our exponential integrator we rewrite the semi-discrete system (1.2) as

\[
\dot{w} = F(t, w), \quad F(t, w) = Jw + f(t), \tag{4.1}
\]

where \( J = M^{-1}A \) and \( f(t) = M^{-1}g(t) \) and \( w, M, A \) and \( g(t) \) are defined as in (3.4). For this ODE system we consider the second-order exponential integrator

\[
w_{n+1} = w_n + \tau \varphi_1(\tau)f(t_n, w_n) + \tau \varphi_2(\tau)(f(t_{n+1}) - f(t_n)), \tag{4.2}
\]

where \( \varphi_1(z) = (e^z - 1)/z \) and \( \varphi_2(z) = (\varphi_1(z) - 1)/z \). This second-order method follows from linearly interpolating \( f \) over \([t_n, t_{n+1}]\) in the variation of constants formula

\[
w(t_{n+1}) = e^{\tau J}w(t_n) + \int_0^\tau e^{(\tau - s)J}f(t_n + s)ds \tag{4.3}
\]

and subsequently computing the resulting integrals analytically. The first paper we know of where this interpolating approach with exact, analytic computation of integrals has been used is \([10]\). Formula (4.2) can be found there. A closely related, somewhat later contribution, is \([31]\). In the recent literature this approach is sometimes called exponential time differencing, see e.g. \([11,39]\). In \([39]\), exponential integration has been applied to the Maxwell equations. Note that (4.2) becomes ITR for zero \( J \) and \( f \) is allowed to depend on \( w \). A second-order method closely related to (4.2) where \( f^* \) is used reads

\[
w_{n+1} = w_n + \tau \varphi_1(\tau)f(t_n, w_n) + \tau^2 \varphi_2(\tau)f'(t_n). \tag{4.4}
\]

This method belongs to a class of exponential Runge–Kutta–Rosenbrock methods \([8,26]\).

In the literature many formulas of higher order are proposed. Here we restrict ourselves to using the second-order method (4.2) because we wish to compare to the second-order method CO2 and the spatial discretization of our test example does not exceed order two either. Per integration step this method requires the approximation of two vectors \( \varphi(z)h \) representing \( \varphi_1(\tau)f(t_n, w_n) \) and \( \varphi_2(\tau)f(t_{n+1}) - f(t_n) \) for which we use Krylov subspace iteration, similar as in \([24,26]\) and in related work on exponential integration. In the remainder of the paper we will refer to (4.2) as method EK2 (Exponential Krylov 2nd-order). More background information on EK2 supporting its choice in the current investigation is given in the Appendix of this paper.
4.2. Krylov subspace iteration

Let \( e_1 \) be the first unit vector in \( \mathbb{R}^{n+m} \) (\( n + m \) is the dimension of the matrix \( J \)). Krylov subspace iteration for \( \psi(\tau f) b \) computes an approximation \( d \approx \psi(\tau f) b \) through

\[
d = V_k \psi(\tau H_k) e_1 \cdot \|b\|,
\]

where \( V_k = [v_1, \ldots, v_k] \) is the \((n + m) \times k\) matrix containing the Arnoldi (or Lanczos) basis of the \( k\)th Krylov subspace with respect to \( \tau f \) and \( H_k \) is an \( k \times k \) upper Hessenberg matrix. So \( \psi(\tau H_k) \) replaces \( \psi(\tau f) \) which explains the success of this method as long as \( k \ll n + m \), because then \( \psi(\tau H_k) \) is relatively cheap to compute, e.g. through the Schur decomposition. The costs of building \( d \) mainly consists of \( k \) matrix–vector multiplications with \( \tau f \) within the Arnoldi process. Hereby it is noted that one such multiplication costs about the same as one single integration step with method CO2. So when comparing EK2 to CO2 with regard to CPU time, the latter can be applied with a \( k \) times smaller step size.

A practical drawback is that matrix \( V_k \) must be kept in storage before \( d \) can be formed. Hence if \( n + m \) is large as is the case in large-scale 3D simulations, the storage requirement for \( k \) vectors of dimension \( n + m \) can be substantial. For example, a worst-case estimate for skew-symmetric matrices with uniformly distributed eigenvalues says that \( k \) can get as large as \( \|\tau f\| \) before the iteration error starts to decay [23]. It is obvious that this may require too much storage if we allow \( \|\tau f\| \gg 1 \) which after all is the main purpose of using an exponential integrator like EK2. Fortunately, in applications one often obtains convergence for \( k \) substantially smaller than \( \|\tau f\| \). If not one can split the time interval in subintervals and use restarts, although at the expense of additional work (the software package from [44] does this automatically). For the theory behind Krylov subspace iteration for matrix functions we refer to the research monograph [48] and to the seminal papers [14,15,23,29,41,47] and references therein.

4.3. Krylov implementation

Like for CG we need a proper stopping criterion for the Arnoldi process. Consider the step with (4.2) from \( t_n \) to \( t_{n+1} \) starting in \( w_n \) and write in short

\[
w_{n+1} = w_n + \tau \Phi_1 + \tau \Phi_2.
\]

We stop after \( k_1, k_2 \) iterations for \( \Phi_1, \Phi_2 \) approximating \( w_{n+1} \) by

\[
\hat{w}_{n+1} = w_n + \tau \Phi_1^{(k_1)} + \tau \Phi_2^{(k_2)}.
\]

Ideally, \( \|w_{n+1} - \hat{w}_{n+1}\| \) is smaller than the unknown local truncation error for \( w_{n+1} \) which we represent by the quantity \( \tau \|w_n\|\delta \) for a certain constant \( \delta \). So we require

\[
\|w_{n+1} - \hat{w}_{n+1}\| \leq \tau \|w_n\|\delta,
\]

which holds if

\[
\|\Phi_i - \Phi_i^{(k_i)}\| \leq \frac{1}{2} \|w_n\|\delta, \quad i = 1, 2.
\]

The number of iterations \( k_i, i = 1, 2 \), is now chosen as follows. We assume for \( i = 1, 2 \) separately that (4.9) is satisfied if, in the \( L_2 \) norm, \( p_3 \) times in succession

\[
\|\Phi_i^{(k_i)} - \Phi_i^{(k_i-1)}\|_2 \leq \frac{1}{2} \|w_n\|_2\delta,
\]

where \( p_3 \) is an integer we can choose. Like for ITR we use constant \( \tau \) and have not implemented a local error estimator. So also here we make an educated guess for \( \delta \) and assume that (4.10) works properly. In our experiments this turned out to be the case, even with \( p_3 = 1 \) which we have chosen henceforth. In our tests all occurring matrix functions \( \psi(\tau H_k) \) have been computed exactly using the exponential operator. Finally, we note that \( \Phi_2 = \mathcal{O}(\tau) \) because of the difference \( f(t_{n+1}) - f(t_n) \). This means that normally this term will require less Krylov subspace iterations than the first one which is confirmed in the experiments.
Table 4.1
Computational costs of CO2 (applied with maximal $\tau$) versus the costs of EK2 (applied with different $\tau$); stopping criterion parameters $\delta = 10^{-3}, p_1 = 1$.

<table>
<thead>
<tr>
<th>$\tau$</th>
<th>$\sigma = 0$ # matvecs per t.step</th>
<th>$\sigma = 0$ total # matvecs</th>
<th>$\sigma = 60\tau$ # matvecs per t.step</th>
<th>$\sigma = 60\tau$ total # matvecs</th>
</tr>
</thead>
<tbody>
<tr>
<td>Grid 4</td>
<td>CO2 0.025 1</td>
<td>400</td>
<td>1</td>
<td>400</td>
</tr>
<tr>
<td></td>
<td>EK2 0.0625 14.93</td>
<td>2388</td>
<td>11.48</td>
<td>1836</td>
</tr>
<tr>
<td></td>
<td>EK2 0.125 21.96</td>
<td>1757</td>
<td>13.7</td>
<td>1096</td>
</tr>
<tr>
<td></td>
<td>EK2 0.25 35.45</td>
<td>1418</td>
<td>16.35</td>
<td>654</td>
</tr>
<tr>
<td></td>
<td>EK2 0.5 62.2</td>
<td>1252</td>
<td>21.55</td>
<td>431</td>
</tr>
<tr>
<td></td>
<td>EK2 1.0 116</td>
<td>1160</td>
<td>29.6</td>
<td>296</td>
</tr>
<tr>
<td>Grid 5</td>
<td>CO2 0.0125 1</td>
<td>800</td>
<td>1</td>
<td>800</td>
</tr>
<tr>
<td></td>
<td>EK2 0.25 61.88</td>
<td>2475</td>
<td>25.88</td>
<td>1035</td>
</tr>
<tr>
<td></td>
<td>EK2 0.5 116.5</td>
<td>2330</td>
<td>37.10</td>
<td>742</td>
</tr>
<tr>
<td></td>
<td>EK2 1.0 196.8</td>
<td>1968</td>
<td>53</td>
<td>530</td>
</tr>
</tbody>
</table>

4.4. Comparing EK2 and CO2

We have repeated the experiments of Section 3.5.2 with ITR replaced by EK2, again focusing on the comparison to method CO2 in terms of workload expressed in matvecs. For the chosen step size range the spatial error again dominates (so Fig. 3.1 also applies to EK2) justifying our focus on workload without referring to the temporal errors. Workload is found in Table 4.1 for $\delta = 10^{-3}$ and $p_1 = 1$, see (4.10). The $\sigma = 0$ test indicates that for problems without damping EK2 will be more costly in matvecs when compared to CO2, let alone the much larger memory demand. Lowering or increasing $\delta$ will not change much for the larger step sizes. For example, for $\sigma = 0$ and grid 4 we find for $\delta = 10^{-2}$ and $\delta = 10^{-4}$ the total matvec sequences (1900, 1457, 1222, 1132, 1075) and (2942, 2043, 1592, 1363, 1230).

The $\sigma = 60\tau$ test is much more favorable for EK2. We see that for step sizes far away from the critical CO2 limit method EK2 becomes competitive in terms of matvecs, similar to what we have observed for ITR. For EK2, however, the gain is less substantial and given the large memory demand this method will likely not to be of great practical interest when it comes to truly large-scale computations. A positive point of EK2 is that for the range of step sizes used its temporal error behavior turned out to be very good. Of course, would the temporal error dominate, method CO2 will be hard to beat as it is optimally efficient (just one matvec per time step).

5. Concluding remarks

Maxwell’s equations (1.1) provide a prime example of a damped wave equation system. After spatial discretization such systems are commonly integrated in time by implicit–explicit methods, such as method CO2 defined by (3.1) which is prototypical for Maxwell’s equations. CO2 is symmetric and thus of second order and requires just one derivative evaluation per time step which makes it very economical. However, the step size is limited by stability which may turn out restrictive, for example when the spatial error dominates for step sizes larger than the incurred step size limit. In such cases implicit time stepping, for which no such limit exists, may come into sight.

In the setting of the generic semi-discrete system (1.2) we have examined the feasibility of implicit time stepping for two different techniques:

(i) The conventional integrator ITR (Implicit Trapezoidal Rule, see Section 3) combined with preconditioned CG (Conjugate Gradient) iteration. Experiments with the 3D problem posed in Section 3.5.1 indicate that in the absence of conduction (no damping) our ITR–CG implementation based
on either Schur–complement mass matrix or incomplete-Cholesky preconditioning falls short. To truly become competitive with CO2 for problems without conduction, more effective preconditioners are needed. Whether these exist for the linear systems we are facing, is an open question. On the other hand, for our test problem with conduction the experiments were no doubt promising for the ITR–CG implementation. This warrants further investigation to the effectiveness of implicit time stepping for problems with conduction.

(ii) The exponential integrator EK2 (Exponential Krylov 2nd order, see Section 4) combined with Arnoldi-based Krylov subspace iteration to deal with the $\phi$ functions. For this combination we have reached similar conclusions as for ITR–CG. For conduction free problems CO2 remains the method of choice, whereas with conduction EK2 can become competitive, but most likely not more efficient than a well-designed ITR–CG implementation. Given, in addition, the substantial memory demand, we consider this method less promising for truly large-scale Maxwell computations. Subsequent tests with an implementation based on the equivalent EK2 formulation

\[ w_{n+1} = w_n + \tau F(t_n, w_n) + \tau \phi_2(\tau f(t_n, w_n)) + f(t_{n+1}) - f(t_n) \]

are in line with this conclusion. Due to the fact that (5.1) requires Krylov subspace iteration for $\phi_2$ only, we were able to reduce the number of iterations on average by about 45%. This substantial reduction, however, is still insufficient for EK2 to become truly competitive to ITR–CG, as can be concluded by comparing Table 3.2 with Table 4.1 after accounting for the 45% reduction.

Acknowledgement

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A. Appendix on the exponential integrator EK2

A.1. Connection with the Adams–Moulton method

EK2, that is method (4.2), can also be seen to belong to the class of $(k+1)$st order multistep methods

\[ w_{n+1} = R(\tau_n J_n)w_n + \sum_{l=0}^{k} \tau_n \beta_l(\tau_n J_n) [w_{n+1-l} - J_n w_{n+1-l}] \]

where $F$ may be nonlinear in $w, J_n$ is an arbitrary matrix, $R(z) = e^{z + \mathcal{O}(z^{k+2})}$, $z \to 0$ and

\[ \sum_{l=0}^{k} q_{l-1} \beta_l(z) = \phi_j(z), \quad j = 1, \ldots, k + 1, \]

\[ \phi_1(z) = (R(z) - 1)/z, \quad \phi_{j+1}(z) = (j \phi_j(z) - 1)/z, \quad j = 1, \ldots, k, \]

\[ q_l = (t_{n-l} - t_n)/\tau_n, \quad \tau_n = t_{n+1} - t_n, \quad l = -1, 0, \ldots, k - 1. \]

Putting $k = 1, R(z) = e^{z}, \tau_n = \tau$ and $J_n = J$, a simple calculation leads us to EK2. Method (A.1) is a generalization of the classical, variable step size, Adams–Moulton method. The precise formulation (A.1) and (A.2) stems from [27, 46]. An earlier closely related Adams–Bashforth paper is [38]. As a further example we give the fixed-step fourth-order method from class (A.1) which for system (4.1) can be written as

\[ w_{n+1} = w_n + \tau \varphi_1(\tau f(t_n, w_n)) + \tau \varphi_2(\tau f) D_{n,2} + \tau \varphi_3(\tau f) D_{n,3} + \tau \varphi_4(\tau f) D_{n,4}. \]
Evaluating derivatives of $f$ at $t = t_n$, the $D_{nj}$ satisfy

$$D_{n2} = \frac{1}{3} fn+1 + \frac{1}{2} fn - fn-1 + \frac{1}{6} fn-2 = r f^{(1)} + \frac{1}{12} r^4 f^{(4)} + O(\epsilon),$$

$$D_{n3} = \frac{1}{2} fn+1 - fn + \frac{1}{2} fn-1 = \frac{1}{2} r^2 f^{(2)} + \frac{1}{24} r^4 f^{(4)} + O(\epsilon),$$

$$D_{n4} = \frac{1}{6} fn+1 - \frac{1}{2} fn + \frac{1}{2} fn-1 - \frac{1}{6} fn-2 = \frac{1}{6} r^3 f^{(3)} - \frac{1}{12} r^4 f^{(4)} + O(\epsilon).$$

So the $D_{nj}$ act as correction terms of decreasing size $O(\epsilon^{j-1})$ which can be exploited in computing the vectors $\psi(tf) D_{nj}$ by means of the Krylov method.

### A.2. Stiff source terms

The source function $f(t)$ of (4.1) may grow without bound if the spatial grid is refined due to contributions from time-dependent boundary functions (stiff source term). For Maxwell’s equations these contributions are proportional to $h^{-1}$ for $h \to 0$ where $h$ is the spatial grid size. Stiff source terms may cause order reduction, that is, the actual order observed under simultaneous space–time grid refinement can be smaller than the ODE order observed on a fixed space grid. Assuming sufficient differentiability of the exact solution $w(t)$ we will prove that method EK2 is free from order reduction for any $f(t)$ and any stable $J$ with its spectrum in $\mathbb{C}^-$ (so not necessarily emanating from Maxwell’s equations).

First we expand the right-hand side of EK2 at $t = t_n$ for $w_n = w(t_n)$. By eliminating $f(t_n)$ and $f(t_{n+1})$ through the relation $f(t) = w(t) - Jw(t)$ this yields

$$\hat{w}_{n+1} = w + \tau \psi_1 w' + \tau \psi_2 \sum_{j=1}^{6} \frac{1}{j!} \tau^j \left( w^{(j+1)} - J w^{(j)} \right),$$

where $w = w(t_n)$, etc., and $\psi_k = \psi_k(tf)$. Using the definition of $\psi_2$ we next eliminate the Jacobian $J$ from this expansion and reorder some terms. This yields

$$\hat{w}_{n+1} = w + \tau w' + \left( \frac{1}{2} + \psi \right) \tau^2 w'' + S,$$

where $\psi = \psi_2 - \frac{1}{2} \psi_1$ and

$$S = \sum_{j=3}^{6} \left( \frac{1}{j!} (I - \psi_1) + \frac{1}{(j-1)!} \psi_2 \right) \tau^j w^{(j)}.$$

In what follows remainder terms $O(\tau^p)$ are assumed independent of $f$ and $f$ implying proportionality to only $\tau^p$ for $\tau \to 0$ and $\|J\|, \|f\| \to \infty$. The local truncation error $\delta_n = w(t_{n+1}) - \hat{w}_{n+1}$ thus can be expressed as

$$\delta_n = -\psi \tau^2 w'' - S + O(\epsilon^3),$$

where the term $O(\epsilon^3)$ is fully determined by solution derivatives. Further, because $J$ is stable, the matrix functions $\psi_k$ featuring in $S$ are bounded. This means that $S = O(\epsilon^3)$ so that

$$\delta_n = -\psi \tau^2 w'' + O(\epsilon^3).$$

The matrix function $\psi$ is also bounded implying $\delta_n = O(\epsilon^2)$. Consequently, when adding up all preceding local errors towards the global error $\epsilon_{n+1} = w(t_{n+1}) - \hat{w}_{n+1}$ in the standard way through the recursion

$$\epsilon_{n+1} = e^{t_{n+1}} \epsilon_n + \delta_n,$$

we will loose one power of $\tau$ and predict first-order instead of second-order convergence. We can come around this non-optimal result by adopting the approach of Lemma II.2.3 from [28]. Write
\[ \delta_n = (I - e^{tJ})\xi_n + \mathcal{O}(\tau^3), \quad \xi_n = -(I - e^{tJ})^{-1}\psi(t\tau^2)w''(t_n). \]  

\text{(A.11)}

Introducing \( \tilde{\epsilon}_n = \epsilon_n - \xi_n \) we can write

\[ \tilde{\epsilon}_{n+1} = e^{tJ}\tilde{\epsilon}_n + \delta_n, \quad \delta_n = -(\tilde{\epsilon}_{n+1} - \xi_n) + \mathcal{O}(\tau^3). \]  

\text{(A.12)}

Since \( J \) is a stable Jacobian, the matrix function featuring in \( \xi_n \) is bounded which implies that \( \tilde{\epsilon}_n = \mathcal{O}(\tau^2) \) and \( \tilde{\epsilon}_{n+1} - \xi_n = \mathcal{O}(\tau^3) \) giving \( \delta_n = \mathcal{O}(\tau^3) \). Now we can add up all preceding local errors in the standard way to conclude second-order convergence for method EK2. We here tacitly assumed that \( \tilde{\epsilon}_0 = 0 \) so that \( \tilde{\epsilon}_0 = -\tilde{\epsilon}_0 = \mathcal{O}(\tau^2) \). This convergence result holds for any stable Jacobian \( J \) and any source function \( f(t) \) eliminating the possibility of order reduction due to contributions from time-dependent boundaries. With a slight change the proof is also valid for the alternative method \( \text{(4.4)} \).

**Example.** We will illustrate the above convergence result for EK2 with a simple yet instructive numerical example. By way of contrast so as to emphasize that when it occurs order reduction may work out badly, we will also apply the method

\[ \bar{w}_{n+1} = e^{tJ}(\bar{w}_n + \frac{1}{2}J\bar{f}(t_n)) + \frac{1}{2}J\bar{f}(t_{n+1}). \]  

\text{(A.13)}

This exponential integration method is obtained from the variation of constants formula \( \text{(4.3)} \) by directly approximating the integral term with the quadrature trapezoidal rule, rather than first interpolating and integrating the obtained terms analytically. The method can also be obtained through time splitting. As an ODE method it is second-order consistent and even symmetric. However, it suffers from order reduction. In fact, for \( \tau \to 0 \) and \( \|J\|, \|\bar{f}\| \to \infty \) it is not even convergent which we will illustrate numerically. Also, unlike EK2, the method is not exact for constant \( f \).

We have integrated the \( 2 \times 2 \)-system (Prothero–Robinson type model from stiff ODEs [12])

\[ W' = \begin{pmatrix} 0 & -s \\ s & 0 \end{pmatrix} W + f(t), \quad f(t) = g'(t) - \begin{pmatrix} 0 & -s \\ s & 0 \end{pmatrix} g(t), \quad g(t) = e^{tJ} \begin{pmatrix} 1 \\ 0 \end{pmatrix}. \]  

\text{(A.14)}

Putting \( w(0) = [1,1]^T \) yields for any \( J \) the solution \( w(t) = [e^t, e^{tJ}]^T, t \geq 0 \). So we can take \( s \) as large as we wish to illustrate the order reduction phenomenon. Fig. A.1 shows convergence results for \( s = 10, r = \frac{1}{2}2^{-j} \) and \( s = 5 \cdot 2^j, r = \frac{1}{2}2^{-j} \) where \( j = 1, \ldots, 10 \). So in the first case \( \|J\| \to 0 \) and \( \|\bar{f}\| \to 0 \) whereas in the second case \( \|J\| \) and \( \|\bar{f}\| \) are fixed and thus \( \|\bar{f}\| \) and \( \|\bar{f}\| \) are increasing. With the first case we test normal ODE convergence and with the second case order reduction. We plot maximum absolute errors at \( t = 1 \) versus \( \tau \) for EK2 (left plot) and \( \text{(A.13)} \) (right plot). The dashed line is the exact order two line, +-marks refer to \( s \) fixed and o-marks to increasing \( s \). EK2 is shown to converge in the right manner for both cases whereas in both cases \( \text{(A.13)} \) is much less accurate and in particular suffers from severe order reduction in the second case even resulting in lack of convergence.

Fig. A.1. Maximum absolute errors at \( t = 1 \) for EK2 (left plot) and \( \text{(A.13)} \) (right plot). The dashed line is the exact order two line, +-marks refer to \( s \) fixed and o-marks to increasing \( s \).
References


