Dual Scattering Channel Schemes Extending the Johns Algorithm

Abstract. Dual scattering channel schemes extend the transmission line matrix numerical method (JOHNS’ TLM algorithm) in two directions. For one point, transmission line links are replaced by abstract scattering channels in terms of paired distributions (characteristic impedances are thus neither needed, nor in general defined, e.g.). In the second place, non-trivial cell interface scattering is admitted during the connection cycle. Both extensions open a wide field of applications beyond the range of classical time domain schemes, such as Yee’s FDTD method and TLM. A DSC heat propagation [diffusion] scheme in non-orthogonal mesh, wherein heat sources are directly coupled to a lossy Maxwell field, illustrates the approach.

Keywords: Time domain methods, finite differences, dual scattering channel schemes, TLM, DSC.

MSC-classes: 65M06, 78M20, 80M20

1. Introduction

It seems a paradox - and is just a typical process in mathematical analysis that a structure turns simple in a more general setting which at the same time widens its range of application. Accordingly - not very surprising - some ill-famed ‘intricacies of the propagator approach to TLM’ (sic. Rebel [1], p. 5) virtually vanish if some of its elements are taken as the building blocks of a more general scheme. In fact, constructing the latter on essentially these elements in a quasi axiomatic manner will prove such intricacies to be mere artefacts of an inadequate framework.

The choice of elements proposed in this paper ‘generalizes’ the Johns algorithm in two directions. In the first place, abstract scattering channels replace transmission lines, which have some unpleasant properties (section [2]). Secondly, non-trivial cell boundary (interface) scattering is permitted during the connection cycle. The schemes thus obtained are characterized by a non-trivial two-step (connection-reflection) cycle of iteration which exhibits certain duality relations - whence their name.

When P.B. Johns and co-workers introduced the transmission line matrix (TLM) numerical method in the 1970s [2] it was almost instantaneously
assimilated by the microwave engineering community. In the same audience
the method remained until today subject of assiduous study and extensive
publication. Three conferences explicitly focussed on TLM [3,4,5], and the
monographs of Christopoulos [6] and de Cogan [7] deal in detail with the
original ideas as well as with classical applications.

Familiarity with the transmission line picture, and the well known scat-
tering concept, certainly fostered the acceptance of the TLM method among
microwave engineers. On the other hand, just so the primary interest turned
of course on applications in their own discipline, rather than onto the inner
algorithmic structure as an object of mathematical analysis. Over the years
still a node more was routinely invented, with new dispersion characteristics
and/or equipped with still another ingenious stub, designed to model spe-
cial propagation or transport phenomena, in varied geometries or boundary
conditions. [8] stands somewhat exemplary for this line of research.

Mathematical questions addressing the inner structure of the TLM al-
gorithm and its potential generalizations have thus apparently been for a
long time of secondary interest. They have yet not been left completely
out of view. Chen, Ney and Hoefer [9] proved equivalence of the original
(expanded) TLM node without stubs to the Yee finite-difference grid
[13,14]. Recently, the non-trivial question of consistence of Johns’ symmet-
rical condensed node (SCN), cf. Johns [11], with Maxwell’s equations and
the intimately related problem of convergence to a smooth solution for
decreasing time step and grid spacing have been tackled, and in parts solved,
by Rebel [1]. His thesis presents, by the way, a thorough survey over the
ramifications of TLM until that time (year 2000), without perhaps spending
sufficient attention to its non-orthogonal mesh extensions.

From a quite general viewpoint, viz. widely independent of any par-
ticular physical interpretation, the structure of the stub loaded (deflected)
non-orthogonal TLM algorithm has been analysed in [16]. The present paper
goes even further and challenges the transmission line picture at all. The
latter, in universally imposing free wave propagation between cells (with
great benefit, at times), induces modeling limitations under circumstances
that are outlined in section 2. Many restrictions can be by-passed by replac-
ing transmission lines with abstract scattering channels in terms of ‘paired’
distributions.

Dual scattering channel schemes are characterized by a two-step updat-
ing cycle with certain duality relations between the two steps. The TLM
method with its familiar connection-reflection cycle is trivial as a dual
scheme in that the connection map reduces essentially to identity (viz. pure
transmission or total reflection) - again with modeling limitations. These
can be raised, anew, in permitting non trivial cell interface scattering dur-
ing the connection step of iteration.

One major merit of the transmission line method is unconditional sta-
bility [12]. Since this property is usually drawn upon the passivity of linear
transmission line networks, the question of stability needs a proper inves-
tigation for DSC schemes that do not use lines. That problem is studied
in \cite{22}, where it is shown that a wide class of DSC schemes are in fact unconditional stable. Due to the convolution type updating scheme (Johns’ cycle; cf. section 3) it is sufficient for stability that the reflection and connection maps share simple contraction properties (paraphrased as $\alpha$-passivity in \cite{22}). In summary, DSC schemes are unconditionally stable under quite general circumstances, and they are conceptually simple, though a set of technical definitions is of course ineluctable in a neat theory. Last but not least, nothing obscure nor 'intricate' should be associated anymore with the propagator approach.

2. Scattering channels

Any extension of the TLM method that includes heat transfer, fluid flow, or particle current, for instance, involves scattering channels other than transmission lines. The latter, for a non vanishing real part of the characteristic line impedance, inherently impose wave propagation between cells. Degenerate lines, with a purely imaginary impedance, still work in diffusion models, cf. \cite{6}, chap.7. Other types of transport or modes of propagation, such as for example the relativistic charged particle current treated in \cite{16}, are very unnaturally and more or less imperfectly modeled using transmission lines. There is good reason to get rid of lines in such and other cases within an extended framework.

A first step towards the definition of more general scattering channels in TLM has been undertaken in replacing transmission lines with abstract projections into in- and outgoing field components, cf. \cite{16}. It was postulated in this paper that the propagating fields ('link quantities') allow of a decomposition into a direct sum

\begin{equation}
\mathbf{z} = \mathbf{z}_{\text{in}} \oplus \mathbf{z}_{\text{out}},
\end{equation}

$\mathbf{z}_{\text{in}}$ and $\mathbf{z}_{\text{out}}$ representing the incident and outgoing fields, respectively. Moreover, it is essential in our understanding of TLM that the latter have a merely operational meaning in that only the total field $\mathbf{z}$ enters the dynamical model equations (cf. sections 3,4). In singular cases, a physical interpretation can yet still be given to $\mathbf{z}_{\text{in}}$, $\mathbf{z}_{\text{out}}$ on the basis of a special analysis, e.g. \cite{18}, Corollary 2.

For the Maxwell field model the technical passage from the transmission line formulation to the projection operator setting is outlined in \cite{16}, Appendix A.

In the classical TLM setup the connection map simply transfers without further modifications the quantities outgoing from a cell into quantities incident at neighbouring cells or rejected at some totally reflecting electric or magnetic wall. This is in perfect harmony with the behaviour of a propagating electromagnetic field, the components of which are tangential to the cell boundary (as the link quantities always are in a classical TLM cell, cf. \cite{21}) and that is thus not subject to refractive scattering at the cell face, even if the medium changes there.
The situation is clearly not thus simple for arbitrary propagating quantities. To circumvent any modeling restrictions, the connection cycle of a non-trivial DSC scheme comprises cell interface scattering from the outset. Nodal and cell face scattering thus enter a kind of duality relation that becomes visible, for instance, in an apparent symmetry of the model equations in their most general form \cite{21, 25, 26}. Nodal and cell boundary scattering may in fact be of equal importance and sometimes boundary scattering plays even the leading rôle in a DSC algorithm.

In the generalized setup, just as in the traditional TLM framework, scattering channels interconnect a node, viz. a suitably defined centre of a mesh cell, with ports at the cell boundary. The channels are yet no longer represented by transmission lines. With respect to a computed physical field in D-dimensional configuration space, they simply form a pair of scalar or vector valued distributions, transposed over a distance in space, which test the field within the cell and on its boundary. A DSC scattering channel will thus be defined, precisely, as a pair of continuous linear functions \((p, p^\sim)\) which act on a class of (suitably smooth real or complex) vector fields \(Z\) in configuration space, such that \(p\) has its support on a cell face and \(p^\sim\) is connected to \(p\) via pull back into the node, i.e.: Given any notions of centre of cell and face, as well as the spatial translation \(s : \mathbb{R}^D \to \mathbb{R}^D\) that shifts the centre of a cell (node) into the centre of the face where \(p\) has its support, then the nodal image \(p^\sim\) of \(p\) is defined as the distribution

\[
(p^\sim, Z) := (p \circ s^\top, Z) = (p, Z \circ s^{-1}),
\]

and the pair \((p, p^\sim)\) is called a scattering channel. Equivalently, a scattering channel can of course be identified with \((p, s)\) or even simply with the port \(p\), the pertinent shift and nodal image then being tacitly understood.

The concept should in fact not be handled in too rigid a fashion - and there is no need to do so. In certain applications the support of the port distribution may be extended over a neighbourhood of a face, or the node distribution may rather be thought of as a mean over the entire cell (in the way familiar from finite volume methods). Needless to say that the

\begin{figure}[h]
\centering
\includegraphics[width=0.3\textwidth]{fig1.png}
\caption{Ports on a cell face with their nodal images.}
\end{figure}

stressed duality between nodal and cell boundary scattering is not to be
misunderstood in the narrow sense of category theory. Here, it refers simply

to the observation that a set of propositions are valid, modulo symmetry in
certain terms, in the two scattering situations - which of course reflects the
paired distribution concept of scattering channel and the already mentioned
symmetry of the pertinent model equations in their most general form. A
parallel symmetry then clearly characterizes the structure of the reflection
and connection maps that solve these equations.

Cell boundary scattering is, for the rest, not thus new an option: Already

in the TLM model for superconducting boundary \[20\] cell face s-parameters
and boundary stubs have been introduced for solving the discretized London
equations, cf. also \textsuperscript{18}.\[18]

Despite the abolition of transmission lines, by their replacement with
abstract scattering channels the computed (‘physical’) fields can still be
represented – in the way familiar from the classical TLM method – as sums
of \textit{in- and out-going} scalar or vector fields

\[
\begin{align*}
z & = z_{\text{in}} + z_{\text{out}}.
\end{align*}
\]

No physical interpretation or propagation property is, however, in general

ascribed to \(z_{\text{in}}, z_{\text{out}}\). In fact, these quantities are merely operationally defined
by means of the well known Johns cycle of iteration

\[
\begin{align*}
z_{\text{in}} & = (\mathcal{C}[z_{\text{out}}] + e) 
\downarrow \hspace{1cm} t + \tau/2 
\uparrow \hspace{1cm} t + \tau 
\mathcal{R}[z_{\text{in}}] 
\uparrow \hspace{1cm} z_{\text{out}} 
\rightarrow \hspace{1cm} z_{\text{in}} + z_{\text{out}}.
\end{align*}
\]

\(\mathcal{R}\) and \(\mathcal{C}\) denote the node and cell face propagators (or so-called reflection
and connection maps - the latter including now cell boundary scattering,
and \(e = e(t)\) induces any excitation. Note again that \(z_{\text{in}}, z_{\text{out}}\) are so far
\textit{purely operational} quantities, i.e. only the total fields \(z\) enter the model
equations, while \(z_{\text{in}}, z_{\text{out}}\) are \textit{in general} bare of any physical meaning (a
physical interpretation in terms of an energy flow still exists within the
classical Maxwell field TLM model, cf. \textsuperscript{18}, Corollary 2).

As will be seen in the next section, the structures of the propagators \(\mathcal{R}\)
and \(\mathcal{C}\) are very similar in the general DSC scheme, thus reflecting the dual
role that nodal and boundary scattering play therein. In a sense, precised
in section \[4\] \(\mathcal{R}\) and \(\mathcal{C}\) are the discrete \textit{convolution integrals} that in every
Johns cycle strictly solve the model equations. In fact, the Johns cycle can
be looked at as basically a two-step convolution method for solving certain
types of explicit finite difference equations in time.

Note that the model equations can in principle be directly solved inasmuch
as they provide complete recurrence relations, cf. sections \[3\] \[4\]. The
scattering approach (using Johns’ cycle of convolutions) offers, however,
important advantages. Thus, it provides clear cut reliable stability criteria
that make the DSC algorithm unconditionally stable under very general circumstances [22].

3. The elements characterizing DSC schemes

So far, we dealt on a largely informal level with some typical traits of the TLM algorithm that either characterize DSC schemes in like manner, or which have to be modified in a specified way in order to attain a greater generality. We are still bound to keep our introductory promise and give a coherent description of DSC schemes in terms of some quasi axioms that condense their distinctive properties. Of course, we shall not really pursue axiomatics, here, in the sense of building a new theory on a complete set of basic assumptions. (Nor are we adopting a dogmatic attitude and going to fix a rigid framework that with certainty must be modified sooner or later in order to face a particular problem.) The emphasis is rather on compiling on a largely preliminary basis those formal elements that, in essence, lead to the peculiar structure of DSC schemes (in general), and of the TLM method (in particular), without being distracted by unnecessary information, such as mesh topology and geometry, s-parameters, etc., which characterize only a singular physical interpretation. A basic set of technical notions need simply to be clarified.

In the following, ‘simple’ definitions are visualized in writing the defined object(s) in italics, more crucial or technical ones are explicitly designated as Definition.

Until further notice, a mesh denotes a non-void finite family (i.e., an indexed set) of elements named cells, which are sets in their turn, and share the following properties. Each cell \( \zeta \) contains an element \( n_\zeta \), called its node, and a finite family \( \partial \zeta = \{ \partial_\zeta^{\iota} \} \), called the (cell) boundary. The latter is built up of elements \( \partial_\zeta^{\iota} \), named faces, which are sometimes simply written \( \iota \) in the place of \( \partial_\zeta^{\iota} \).

**Definition 1** A mesh \( M \) is called regular, if and only if it satisfies the following requirements of simplicity (S) and connectedness (C):

- (S) Every node belongs to exactly one cell and every face to at most two cells in \( M \).
- (C) For every two cells \( \zeta^i, \zeta^j \in M \), there exists a connecting sequence \( s = (\zeta^k)_{k=0}^{k} \in M^N \), such that \( \zeta^0 = \zeta^i, \zeta^k = \zeta^j \) and every two subsequent cells \( \zeta^k, \zeta^{k+1} \) in \( s \) have a common face, for \( 0 \leq k < k \).

Also - certainly not too misleading: any two cells with a common face are called adjacent or neighbouring cells, and the common face a connecting face or interface.

By a first postulate, DSC meshes are always regular meshes.

The state space is any product of real or complex normed linear spaces labelled by mesh cells

\[
S = \prod_{\zeta \in M} S_\zeta
\]
In addition, we require that a DSC state space always contains a non-void subspace $\mathcal{P} \subset \mathcal{S}$, named the space of propagating fields, which on every $\mathcal{S}_\zeta$ reduces to a product of 'squared' spaces in the following precise sense

$$\mathcal{P}_\zeta = \mathcal{P} \cap \mathcal{S}_\zeta = \prod_{\iota \in \partial \zeta} P^2_{\iota, \zeta}.$$  

In less formal language: every propagating field $z$ splits over the cells into a sequence of pairs

$$z_\zeta = (z_\iota, z_\iota^\sim)_{\iota \in \partial \zeta}$$

labelled by the faces of the cell boundary. In other words, there is a canonical automorphism of normed linear spaces on $\mathcal{P}$, which on every $\mathcal{P}_\zeta$ reduces to

$$nb : \mathcal{P}_\zeta \rightarrow \mathcal{P}_\zeta$$

$$ (z_\iota, z_\iota^\sim) \mapsto (z_\iota^\sim, z_\iota) .$$

$nb$ is obviously involutary ($nb^2 = Id$), and is called the node-boundary map.

The components $z_\iota$ and $z_\iota^\sim$ in (10), (11) are named the port (or face) component, and the node component, respectively, of $z = (z_\iota, z_\iota^\sim)$. They are usually written $z^p = z_\iota = \pi^p(z)$ and $z^n = z_\iota^\sim = \pi^n(z)$ with projections $\pi^p, \pi^n$ that are canonically extended over the entire space $\mathcal{P}$.

Let $J := \bigcup_{\zeta \in M} \partial \zeta$ be the set of all faces in $M$ (remember that $\partial \zeta$ is defined as a union of faces). Then, in virtue of (10), $\mathcal{P}$ splits completely into subspaces

$$\mathcal{P} = \prod_{\iota \in J} \mathcal{P}_\iota$$

with $\mathcal{P}_\iota := \prod_{\zeta; \iota \in \partial \zeta} P^2_{\iota, \zeta}$.

A DSC process is a step function of time

$$pr : [0, T) \rightarrow \mathcal{S} ,$$

such that $\pi^p \circ pr(t)$ and $\pi^n \circ pr(t - \tau/2)$ are constant on every time interval $[\mu \tau, (\mu + 1)\tau), \mu \in \mathbb{N}$, where they are defined. In other words, port components of a DSC process switch at integer multiples of the time step $\tau$ while node quantities switch at odd integer multiples of $\tau/2$.

Given a process, a state $z$ with its entire history up to time $t$ is usually written as a 'back in time running' sequence

$$[z](t) := (z(t - \mu \tau))_{\mu \in \mathbb{N}} ,$$

expanding so the domain of definition eventually to the negative time axis in the trivial way, i.e. $z(s) := 0$ for $s < 0$.

By this convention, we assign thus to index $\mu$ the (varying) state back in the past from present time $t$

$$[z]_\mu(t) := z(t - \mu \tau) ,$$
rather than the (fixed) state \(z(\mu \tau)\) - which has the technical advantage that \(\mu\) so is directly related to a time difference (and eventually to an order of a finite difference equation in time), rather than to an absolute time (which is quite uninteresting, in general).

Functions defined on back in time running sequences, such as (11), are called causal functions (or propagators). Any such map is a discrete analogue to a causal Green’s function integral, as has already been outlined in [18].

Let \(X_n^0\) denote the set of all sequences with an arbitrary, but finite, number of non-vanishing elements in a linear space \(X\). For every mesh cell \(\zeta\) and face \(\iota\) in \(M\) consider then the subspaces of propagating fields \(P_n^\zeta := \pi^n(\mathcal{P}_\zeta), P_p^\iota := \pi^p(\mathcal{P}_\iota)\).

**Definition 2** A reflection map (in \(\zeta \in M\)) is a (possibly time dependent) causal operator

\[
R_\zeta : (P_n^\zeta)_0^N \rightarrow P_n^\zeta \\
[z^n] \mapsto R_\zeta[z^n],
\]

and a connection map (in \(\iota \in J\)) is a (likewise possibly time dependent) causal operator

\[
C_\iota : (P_p^\iota)_0^N \rightarrow P_p^\iota \\
[z^p] \mapsto C_\iota[z^p].
\]

Also, a DSC system over \(M\) is a pair \((C, R)\) consisting of any two families of connection and reflection maps. Note that sometimes also the families are called the connection and reflection map. For the sake of algorithm stability, these maps (in whatever meaning) should share certain contraction properties studied in [22].

An excitation is only a distinguished process with values in the mesh boundary states. More precisely, let \(B := \{\iota \mid \iota \in J \text{ and } \iota \text{ is not an interface}\}\) be defined as the mesh boundary, then an excitation is a process

\[
e : [0, T) \rightarrow \prod_{\iota \in B} \mathcal{P}_\iota \\
t \mapsto e(t) = \pi^p \circ e(t),
\]

i.e. \(e\) is a port process, and hence switches at entire multiples of the time step \(\tau\), and \(e\) generates non-interface (mesh boundary) states.

**Definition 3** The DSC process generated by \((C, R)\) and excited by \(e\) is the unique process \(z(t) = (z^p, z^n)(t)\) which at every time \(t \in [0, T)\) satisfies

\[
\begin{align*}
z^p(t) &= nb \circ z^n_{in}(t + \tau/2) + z^p_{out}(t) \\
z^n(t) &= z^n_{in}(t) + nb \circ z^p_{out}(t + \tau/2)
\end{align*}
\]
the right-hand side being recursively defined through cyclic iteration of

\begin{align}
 z_n^\text{in}(t + \tau/2) & := nb \circ [ C[z^\text{out}_n](t) + e(t) ] \\
 z^\text{out}(t + \tau) & := nb \circ R[z_n^\text{in}](t + \tau/2) \\
 t & := t + \tau
\end{align}

(in that order) with, initially, \( z_n^\text{in}(0) = z^\text{out}_n(0) = 0 \).

Remember that \( nb \) denotes the node-boundary map (8).

In (18) \( C \) and \( R \) stand of course for application of all propagators \( C_\iota \) and \( R_\iota \) in the pertinent families (over \( J \) and \( M \), respectively). Note that the order of application within the families is unimportant in virtue of the pairwise disjointness of all \( P_\iota \), and of all \( P_\iota \) - which obviously implies that either \( C \) and \( R \) are completely parallelizable at every time step.

It follows immediately that \( z^n \) and \( z_n^\text{in} \) thus defined are node processes, hence switch at odd integer multiples of \( \tau/2 \), while \( z^p, z^\text{out} \) are port processes (so they carry their superscripts aright).

Equations (17), (18) can still be simplified to

\begin{align}
 z_{n+}^p &= z_{n+}^\text{in} + z_{n+}^\text{out} \\
 z^p(t) &= C[z^\text{out}_p](t) + e(t) \\
 z^\text{out}_n(t + \tau/2) &= R[z_{n}^\text{in}](t + \tau/2)
\end{align}

writing as usual \( z_{n+}^p \) and \( z_{n+}^\text{out} \) for

\begin{align}
 z^p_{n+}(t) & := nb \circ z_{n+}^\text{in}(t + \tau/2) \\
 nb \circ z^p_{n+}(t) & =: z_{n+}^\text{out}(t - \tau/2)
\end{align}

Comparing this to the TLM usage we note that in (16) port quantities \( z^p_{n+}, z_{n+}^\text{out} \) are first introduced. With these are then node quantities \( z_{n+}^\text{in}, z_{n+}^\text{out} \) identified (modulo the time shifts \( \pm \tau/2 \), just as in (15)) without yet explicitly mentioning the node-boundary isomorphism \( nb \). We shall sometimes follow this usage and omit the symbol \( nb \) where this cannot lead to confusion.

Nothing has been said, so far, about physical interpretations or any implemented dynamical equations. In fact, the characteristic structure of the DSC algorithm is entirely laid down with the given definitions. As will be seen in the next section, typical features and facts, some quite familiar from TLM, are derived straight away with only the above elements.

With respect to the dynamical model equations - which the algorithm has ultimately to solve and that determine the propagators \( R_\iota, C_\iota \), cf. section 4 - we reiterate the important general agreement that only total fields \( z^p, z^n \), not, however, their incident and outgoing components separately, shall enter these equations. Accordingly, we consider only model equations between total fields. Quite generally, and modulo further restrictions (inferred in the next section), the DSC model equations should be of the types

\begin{align}
 F^n[z^n][z^p] & \equiv 0 \\
 F^p[z^p][z^n] & \equiv 0
\end{align}
with causal functions $F_n$, $F_p$ and shortly $[z_{\pm}] := [z](t \pm \tau/2)$. The $\tau/2$ time shifts synchronize node and cell boundary switching in (21), such that the equations can be strictly solved, for every time $t \in [0, T)$ and are well-posed, in this sense. Of course, time shifts by $-\tau/2$ would also lead to synchronization. The resulting equations would, however, conflict with the causality property of $R$ and $C$.

Inspection of equations (21) shows that $F_n$ affects only the reflection cycle, while $F_p$ has impact only on the connection cycle. We are now dealing with the model equations in some more detail.

4. The model equations

The physical interpretation of a DSC system fixes, intuitively speaking, the terms in that states in $P$ are read as physical fields. More deliberately, certain states in $P$ are interpreted as distributional values (finite integrals, e.g.) of physical fields, which are localized in a mesh cell system.

Any interpretation requires, hence, in the first instance a geometric realization of the underlying regular mesh, wherein the relations between abstract cells, nodes, and boundary faces which characterize $M$ are translated into relations between geometric objects, bounded subsets of $\mathbb{R}^D$, such as (in general) polyhedral mesh cells with their faces, e.g.

Given any geometric realization of $M$, a physical interpretation of a DSC system over $M$ is, precisely, a family $I = \{I^\iota_\zeta\}_{\zeta \in M, \iota \in \partial \zeta}$ of continuous linear functions

\[(22) \quad I^\iota_\zeta : E^\iota_\zeta \rightarrow P_\zeta ,\]

each defined on a space $E^\iota_\zeta$ of smooth $m$-component vector fields in $D$-dimensional configuration space (i.e. $E^\iota_\zeta \subset C^\infty(\mathbb{R}^D)^m$; $m \in \mathbb{N}$ depending on $(\iota, \zeta)$), such that $I^\iota_\zeta$ has its distributional support on a cell face and its range in $P^p_\zeta = \pi^p(P_\zeta)$.

Note that index $\iota$ in (22) may optionally be read as a cell face label or as a multiindex referring to a set of ports on the same face. Since we are dealing with vector-valued distributions (with range in $P^p$) and the support of every $I^\iota_\zeta$ is required to be localized on a cell face (which can be weakened to at least associated to a face), there is essentially no difference in reading $z^p_\zeta = I^\iota_\zeta(Z)$ as a cell face state vector, or as an array of components (labelled by port indices) of such a vector. Thus, index $\iota$ in (22) may be thought of as implicitly labelling a subset of ports on face $\iota \in \partial \zeta$.

Attention is also drawn to the fact that the functions $I^\iota_\zeta$ are not required to be surjective onto $P^p_\zeta$ (i.e. not every state in $P^p_\zeta$ must be directly related do a distribution in $I$). There is, for instance, no need to exclude from $P^p_\zeta$ any function or linear combination of fields in different spaces $E^\iota$ (which may represent a spatial finite difference of fields, as in the approximate gradient of our sample model in section 5).

The evaluation of nodal fields goes quasi pick-a-pack with $I$ by applying the scattering channel concept of section 3.

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Let $s_\zeta^\iota$ denote the translational shift in $\mathbb{R}^D$ from any node $n_\zeta$ into the (centre of the) face where $I_\zeta^\iota$ has its support, cf. fig 4 and assume without loss of generality that $S_\zeta^\iota : Z(x) \mapsto Z(x+s_\zeta^\iota)$ is an inner map in $E_\zeta^\iota$ (otherwise take the closure of $E_\zeta^\iota$ under such transformations). Then clearly holds

**Proposition 1** For every $I_\zeta^\iota \in I$ there exists exactly one function $I_\zeta^\sim^\iota : E_\zeta^\iota \to P_\zeta^\iota$ (note $I_\zeta^\sim^\iota \notin I$), such that the following diagram is commutative

\[
\begin{array}{ccc}
P_\zeta^p & \xrightarrow{nb} & P_\zeta^n \\
I_\zeta^\iota & \uparrow & I_\zeta^\sim^\iota \\
E_\zeta^\iota & \xrightarrow{S_\zeta^\iota} & E_\zeta^\iota
\end{array}
\]

i.e. $I_\zeta^\sim^\iota \circ S_\zeta^\iota = nb \circ I_\zeta^\iota$.

**Proof** Mere retrospection of definitions.

In the terminology of section 2 is $I_\zeta^\sim^\iota$ the nodal image of the port(s) $I_\zeta^\iota$, and the pair of distributions $(I_\zeta^\sim^\iota, nb \circ I_\zeta^\sim^\iota)$ forms a scattering channel.

The dynamical DSC model equations are, in the line of the preceding, to be read as finite difference equations in time between states $z = (z_\zeta^p, z_\zeta^n) \in P$ that have an interpretation as distributional values of physical fields $Z \in E_\zeta^\iota$,

\[
z_\zeta^p = I_\zeta^\iota(Z) , \quad z_\zeta^n = I_\zeta^\sim^\iota(Z) .
\]

Unlike classical FD equations between pointwise evaluated physical fields, the DSC model equations interrelate in many cases finite integrals over a line segment or face, e.g. pointwise evaluation (with a Dirac measure as distribution) not excluded. The DSC approach is, in this respect, by far more versatile than the classical finite difference time domain method. So, the former permits, for instance, of generalizing to a non-orthogonal mesh Johns’ TLM method [11] in much a simpler way, cf. [17], than the FDTD approach allows for Yee’s method of approximation to Maxwell’s equations [13].

A central principle underlying DSC schemes is near-field interaction. Like causality, this is already implicit in the (domains of) definition of the reflection and connection maps. Near-field interaction simply spells that only the fields in the immediate neighbourhood of a state $z$ - precisely only those in $P_\zeta$ if $z \in P_\zeta^n$, and those in $P$, if $z \in P_\zeta^p$, cf. [6, 9], along with their history, of course - determine the evolution of that state on the next updating step (note, this refers to fields evaluated in $P$ and not, for instance, to an exterior potential, which may still induce a time dependence of $R$ or $\mathcal{C}$).

In other words, an updated nodal state depends only on states of the pertinent cell, including its boundary, while the evolution of a port state
is determined by states of the respective face and by nodal states of the adjacent cells.

It follows that the model equations (21) split into the two families

\[(25) \quad \mathcal{F}_\zeta[z^n_\zeta,z^p_\zeta] \equiv 0, \quad z_\zeta \in \mathcal{P}_\zeta, \quad \zeta \in M\]

\[(26) \quad \mathcal{F}_\iota[z^p_\iota,z^n_\iota] \equiv 0, \quad z_\iota \in \mathcal{P}_\iota, \quad \iota \in J.\]

(Remaining aware of the dependence upon \(\zeta\) and \(\iota\) of these equations, we can in general omit the subscripts, if there is no danger of confusion.)

Since the following analysis runs perfectly parallel for the dual equations, from now on it is confined to the implications of (25). - The reader may write down parallel statements for dual equations, at times, by exchanging port for nodal and incident for outgoing quantities, starting, for instance, with the cell boundary version (with \(\mathcal{C}\) in the place of \(\mathcal{R}\)) of the following:

**Definition 4** Let \(I = [0, T]\) be a finite interval (i.e. \(T \in \mathbb{R}_+\)). Then we shall say that \(\mathcal{R}\) generates solutions of the model equations (25) on \(I\), if and only if for every sequence of incident nodal fields \([z^n_{in}]\) the (obviously unique) process \(z = z_{in} + z_{out}\) which is recursively given by

\[(27) \quad z^n_{in}(t - \tau/2) = nb z^n_{in}(t) \]

\[z^n_{out}(t) = \mathcal{R}[z^n_{in}](t) \]

\[z^p_{out}(t + \tau/2) = nb z^p_{out}(t)\]

algebraically solves equations (25) (identically on \(I\)). Sometimes, we then simply say that \(\mathcal{R}\) solves the model equations on that (finite \(I\)) interval.

**Remark 1**

(i) Note that Definition 4 refers to a purely algebraic property of \(\mathcal{R}\) that is not yet related to any stability questions, for instance (cf. also the Remark to Theorem 1).

(ii) If \(\mathcal{R}\) solves equations (25) on \(I\), then in particular every process generated by \((\mathcal{C}, \mathcal{R})\) in the sense of Definition 4 solves (25) on \(I\), since every such process obviously satisfies (27) (cf. 24).

The least awkward (yet fortunately frequently encountered) situation is brought about with homogeneous linear equations, i.e. for the evolution of nodal states

\[(28) \quad \mathcal{F}^n[z^n_{in},z^p] = \sum_{\mu=0}^\infty \phi_\mu z^n(t + \tau/2 - \mu\tau) + \psi_\mu z^p(t - \mu\tau) \equiv 0,\]

with linear and possibly time dependent operators

\[\phi_\mu : \mathcal{P}^n \rightarrow J, \quad \psi_\mu : \mathcal{P}^p \rightarrow J\]
that map into any common linear space $\mathcal{I}$ wherein $\mathcal{F}$ has its range. Similar dual equations $\mathcal{F}_p[z^n_\mu][z^n] \equiv 0$ determine the linear evolution of the port states during the connection cycle.

Many, if not almost all (viz. all but a finite number) of the $\phi_\mu$, $\psi_\mu$ may be zero. Any maximum $\mu \in \mathbb{N} \cup \{\infty\}$, such that $\phi_\mu \neq 0$ or $\psi_\mu \neq 0$, is called the (dynamical) order of the model equations, and in general equals the order in time of the integro-differential equations which physically describe the underlying dynamical problem in terms of smooth fields.

The following statement provides a theoretical means for computing the reflection map of equation (28) recursively.

**Theorem 1** Let $\phi_0 : \mathcal{P}^n \rightarrow \mathcal{I}$ be bijective (i.e. one-to-one and onto). Then $\mathcal{R}$ solves equation (28) on a finite interval $[0, T)$, if and only if for every $t \in [0, T)$

\[
\mathcal{R}[z^n_{\text{in}}](t) = z^n_{\text{out}}(t) = (-\phi_0)^{-1} \sum_{\mu=0}^{\infty} \left\{ (\phi_\mu + \psi_\mu nb) z^n_{\text{in}}(t - \mu \tau) + (\phi_{\mu+1} + \psi_\mu nb) z^n_{\text{out}}(t - \tau - \mu \tau) \right\}.
\]

**Proof** Substituting in (28) for $z^n_{\text{in}}$ the right-hand sides of (17) and using equations (20) yields for $t < T$ on a time domain trivially extended over the negative axis

\[
\sum_{\mu=0}^{\infty} \left\{ \phi_\mu \left[ z^n_{\text{in}}(t + \tau/2 - \mu \tau) + z^n_{\text{out}}(t + \tau/2 - \mu \tau) \right] + \psi_\mu nb \left[ z^n_{\text{in}}(t + \tau/2 - \mu \tau) + z^n_{\text{out}}(t - \tau/2 - \mu \tau) \right] \right\} \equiv 0,
\]

which for invertible $\phi_0$ and $t' := t + \tau/2$ is equivalent to the recursion formula of the theorem.

**Remark 2** For every $t < T < \infty$ the sum in Theorem 1 is actually finite, hence convergence is not a question as long as one abstains from considering limits. For finite $T$, the theorem conveys purely algebraic relations inherited from the structure of the DSC process as laid down by [17, 18] in section 4.

The question of stability of a process for $T \rightarrow \infty$ is being addressed below.

If $\phi_0$ is not bijective, then the model equations (28) are incomplete in that they do not determine a well-defined and unique reflection propagator in every cell. We therefore require that the uniqueness condition

(U) \quad $\phi_0 : \mathcal{P}^n \rightarrow \mathcal{I}$ is bijective \quad (i.e. a linear isomorphism )

be always satisfied. Clearly, (28) then defines an explicit scheme.

The model equations of the classical TLM method discretize Maxwell’s equations. So, they are linear and first order in time, viz. of the type

(29) \quad $\phi_0 z^n(t + \tau/2) + \phi_1 z^n(t - \tau/2) + \psi_0 z^p(t) ,$

with time independent operators $\phi_0,1$ and $\psi_0$, which are derived in [17, ?], for instance. Of the same type - viz. linear and of first dynamical order - are the discretized diffusion equations. Theorem 1 applied to this special situation yields
Corollary 41 The (for a given time step unique) reflection map that solves on a finite interval the first order linear equations \( (29) \) with time independent \( \phi_{0,1} \) and \( \psi_0 \) is

\[
R[z^n_{in}](t) = z^n_{out}(t) = \]

\[
= K z^n_{in}(t) + L \sum_{\nu=1}^{\infty} N^{\nu-1} M z^n_{in}(t-\nu \tau),
\]

wherein

\[
K = -Id - \phi_0^{-1} \psi_0 nb
\]

\[
L = -\phi_0^{-1}
\]

\[
M = + \phi_1 + (\phi_1 + \psi_0 nb) \left( -\phi_0^{-1} \right) \left( \phi_0 + \psi_0 nb \right)
\]

\[
N = -(\phi_1 + \psi_0 nb) \phi_0^{-1}.
\]

Proof By induction, applying Theorem 1 to an incident Dirac pulse

\[
z^n_{in}(t) := z \chi_{[0, \tau]}(t - \tau/2)
\]

with any fixed state vector \( z \in \mathbb{P}^n \); \( \chi_I \) denotes the characteristic function of interval \( I \) (which equals 1 for every argument in \( I \) and 0 elsewhere). By linearity the statement then holds for arbitrary incident processes \( z^n_{in}(t) \), each of them being writable as a superposition of Dirac processes that start at subsequent time steps.

Remark 3 A sufficient condition for convergence of the propagator series \( (30) \) (applied to any finite incident pulse \( z^n_{in} \)) in the limit \( t \to \infty \) is obviously

\[
\| N \| < 1,
\]

where \( \| ... \| \) is any submultiplicative norm, e.g. the Hilbert (spectral) norm

\[
\| N \|_H := max \{ |\lambda| : \lambda \text{ eigenvalue of } N \}\]

In fact, the latter condition is sufficient for algorithm stability of the Maxwell field TLM model, as shown in [21,17]. Any first order linear process is clearly stable, if the Hilbert norms of \( K, L, M, N \) are bounded by 1 (strictly for \( K \) and \( N \)), since the propagator \( R \) then is contractive. This is in general ensured with bounds for the time step (if necessary, in combination with a transformation \( (34) \)).

For linear model equations of any finite order, recursion formulae that generalize \( (30) \) are easily derived from Theorem 1 e.g. [16], equation \( (32) \).

Corollary 42 With \( K, L, M, N \) as above, the DSC process solving \( (29) \) permits a representation as a 'deflected' scattering process

\[
\left( \frac{z^n_{out}(t)}{d(t)} \right) = \left( \begin{array}{c} K L \\ M N \end{array} \right) \left( \begin{array}{c} z^n_{in}(t) \\ d(t - \mu \tau) \end{array} \right),
\]

the deflection \( d(t) \) being completely recursively defined with the (universally maintained) initial conditions \( d_{t<0} = 0 \).
Proof Immediate consequence of Corollary 41

Remark 4 Note that there remains an arbitrariness in the definition of the operators $L$, $M$, and $N$ in Corollaries 41, 42. In fact, any invertible transformation $I : J \rightarrow J$ together with the simultaneous replacements of $L$, $M$, and $N$ by, respectively,

\begin{align*}
L^- &:= L \circ I^{-1}, \\
M^- &:= I \circ M, \quad \text{and} \\
N^- &:= I \circ N \circ I^{-1}
\end{align*}

clearly do not alter the propagator $R$, i.e. generates the same DSC process.

The corollaries offer solutions of first order linear equations with time independent operator coefficients which are complete in the sense of (U). They thus cover the entire field of classical TLM (with connection maps that are trivial in reducing essentially to identity).

In certain situations it may be useful, or necessary, to integrate some new (possibly non-linear) interactions into a given DSC model. Sometimes this can be carried out by adding suitable coupling terms to the equations of the yet existing model. We therefore consider perturbed model equations of the type

\begin{align*}
F_\sim [z_n^+][z_p] &\equiv F_n[z_n^+][z_p] + J[z_n^-][z_p] = 0
\end{align*}

(35)

(here exemplary for nodal perturbations), wherein $J$ denotes any causal map into $J$. The $(-\tau/2)$ time shift in the first argument of $J$ again synchronizes port and node switching. Note that the time shift is negative, here. This is to ensure that the perturbation $J$ cannot destroy the uniqueness conditions (U) in the case of a linear function $F_n$, and to preserve explicitness of the updating relations, in general. The importance of this condition becomes clear in the proof of the following formula.

Proposition 2 (Deflection Formula) Let the reflection map $R$ generate solutions of the linear equations (28) on a finite interval $I = [0, T)$. Then $R_\sim$ solves the perturbed equations (35) on $I$, if and only if the so-called deflection

\begin{align*}
D &:= R_\sim - R
\end{align*}

satisfies recursively

\begin{align*}
\phi_0 D_{|t+\tau/2} &= - J[z_n^+][z_p]_t \\
&\quad - \sum_{\mu \in \mathbb{N}} (\phi_{\mu+1} + \psi_{\mu} nb) D_{|t-\tau/2-\mu\tau}.
\end{align*}

Proof By definition, $R_\sim$ solves equations (35) on $I$, if and only if for every incident sequence $[z_{in}^n]$ and

\begin{align*}
\z_{out} := R_\sim [z_{in}^n], \\
z := z_{in} + \z_{out}
\end{align*}

holds

\begin{align*}
F_\sim [z_n^+][z_p] &= F_n[z_n^+][z_p] + J[z_n^-][z_p] = 0.
\end{align*}
However, with $w^n_{\text{in}} := R[z^n_{\text{in}}]$, $w := z_{\text{in}} + w_{\text{out}}$

(hence $D = z^n_{\text{out}} - w^n_{\text{out}}$) and using (28), this is the case iff

$$-J[z^n_{\text{in}}][z^p]_t = \sum_{\mu \in \mathbb{N}} \left( \phi_{\mu|t}(z^n_{\text{in}}|t + \tau/2 - \mu \tau + D|t + \tau/2 - \mu \tau) + \psi_{\mu|t}(z^n_{\text{in}}|t + \tau/2 - \mu \tau + D|t + \tau/2 - \mu \tau) \right)_{|t + \tau/2 - \mu \tau}.$$

In virtue of the linearity of the $\phi_{\mu}$, $\psi_{\mu}$, the latter identity holds iff

$$-J[z^n_{\text{in}}][z^p]_t = \mathcal{T}[w^n_{\text{in}}][w^p]_t + \phi_{0|t}(D|t + \tau/2 + \sum_{\mu \in \mathbb{N}} (\phi_{\mu + 1|t} + \psi_{\mu|t} n) D|t - \tau/2 - \mu \tau),$$

which is the recurrence relations of the proposition.

**Corollary 1 (Deflected processes)** Let $R$ solve (28) on a finite interval $I$ and $\phi_0 : \mathcal{P} \to \mathcal{I}$ be any bijective operator (that thus satisfies the completeness conditions (U)). Then

$$D_{|t + \tau/2} := -\phi^{-1}_{0|t} \left( J[z^n_{\text{in}}][z^p]_t + \sum_{\mu \in \mathbb{N}} (\phi_{\mu + 1|t} + \psi_{\mu|t}) D_{|t - \tau/2 - \mu \tau} \right)$$

with initial conditions $D_{|t < 0} \equiv 0$ defines recursively a causal operator $D$, such that $R \sim := R + D$ solves equations (35) on $I$.

Concluding this section, we stress once again that Theorem 1 and the ensuing propositions and corollaries apply just as well to the connection cycle, i.e. to cell interface scattering, provided the replacements (26) by (29), $R$ by $\mathcal{C}$, $\mathcal{P}_c$ by $\mathcal{P}_0$, port by node superscripts, and incoming by outgoing fields (and vice-versa) are simultaneously made. - Note, however, that any excitations may temporarily violate the model equations at a mesh boundary face. The model developer is encouraged to care for physically consistent excitations.

**5. A heat propagation scheme in non-orthogonal mesh**

The physical interpretation underlying the following application relates a smoothly varying (viz. in time and space continuously differentiable, $C^1$-) temperature field $T$, evaluated as $T^p$ at the face centre points and as $T^n$ in the nodes of a of non-orthogonal hexahedral mesh, to total states $z^p_{\mu,n}$ of a DSC model. In fact, we derive the model equations for the connection and reflection cycles of a DSC heat propagation (diffusion) scheme.
the equations are linear and of dynamical order 0 and 1, respectively - as will be seen - they can be processed, following the guidelines of the last section. In the end, we display some computational results of a dispersion test carried out with this model.

In order to simplify the notation we follow Einstein’s convention to sum up over identical right-hand (!) sub and superscripts within all terms where such are present (summation is not carried out over any index that also appears somewhere at the left-hand side of a pertinent symbol - thus, in $(-1)^{\kappa}a_\kappa^{\lambda}b_\lambda \kappa c$ the sum is made over $\lambda$ but not over $\kappa$). The shape of a

![Hexahedral cell](image)

**Fig. 2.** Non-orthogonal hexahedral mesh cell. (a) Edge vectors. (b) Node vectors.

![Temperature points](image)

**Fig. 3.** Face vectors and temperature points (nodal section).

hexahedral cell is completely determined by its 12 edge vectors $(\mu,e)_{\mu=0,...,11}$. Also, with the labelling scheme of fig. 2, node vectors $(\mu,b)_{\mu=0,1,2}$ and face vectors $(i,f)_{i=0,...,5}$ are defined as

$$
\mu b := \frac{1}{4} \sum_{\nu=0}^{3} (4\nu+\nu) e^\nu, \quad \mu = 0, 1, 2
$$

and

$$
i f := \frac{(-1)^i}{4} \left( (8+2\nu) e^\nu + (9+2(\nu+(-1)^i)) e \right) \land
\land (4+2\nu) e^{(\nu+2)} e \land (5+2\nu) e
$$

$$
i = 0, ..., 5
$$
(edge vector indices cyclic modulo 12 and $\wedge$ denoting the wedge (‘cross’) product in $\mathbb{R}^3$).

At every face $\iota \in \{0, \ldots, 5\}$ of a mesh cell and for any given $\tau \in \mathbb{R}_+$ the following time shifted finite temperature differences in directions $\mu b$ ($\mu = 0, 1, 2$) form a vector valued function

\begin{equation}
\nabla^B T_\mu(t, \tau) := \begin{cases} 
2 (-1)^t (T^m|_{t-\tau/2} - T^p|_{t}) & \text{if } \mu = \lfloor t/2 \rfloor \\
(2\mu+1T^p - 2\mu T^p)|_{t-\tau} & \text{if } \mu \neq \lfloor t/2 \rfloor
\end{cases}
\end{equation}

($[x]$ denotes the integer part of $x \in \mathbb{R}$). The time increments are chosen to attain technical consistence with the updating conventions of DSC schemes. They do not destroy convergence, as easily seen: In fact, in the centre point of face $\iota$ the vector $\nabla^B T$ approximates in the first order of the time increment $\tau$, and of the linear cell extension, the scalar products of the node vectors with the temperature gradient $\nabla T$. Let, precisely, for a fixed centre point on face $\iota$ and $\epsilon \in \mathbb{R}_+$ the $\epsilon$-scaled cell have edge vectors $\iota e := \epsilon \iota e$. Let also $\nabla^B T_\mu$ denote function (37) for the $\epsilon$-scaled cell (with node vectors $\mu b := \epsilon \mu b$). Then at the fixed point holds

\begin{equation}
< \mu b, \text{grad}(T) > = \mu b \cdot \nabla T = \lim_{\epsilon \to 0} \lim_{\tau \to 0} \frac{1}{\epsilon} \nabla^B_{\epsilon} T_\mu,
\end{equation}

as immediately follows from the required $C^1$-smoothness of the temperature field $T$.

To recover, in the same sense and order of approximation, the gradient $\nabla T$ from (37), observe that for every orthonormal basis $(\nu u)_{\nu=0, \ldots, m-1}$ of $V = \mathbb{R}^m$ or $\mathbb{C}^m$, and for an arbitrary basis $(\mu b)_{\mu=0, \ldots, m-1}$ with coordinate matrix $\beta^\mu = < \nu u, \mu b >$, the scalar products of any vector $a \in V$ with $\mu b$ are

\begin{equation}
< \mu b, a > = \sum_{\nu=0}^{m-1} < \mu b, \nu u > < \nu u, a > = \beta^\nu \alpha^\nu .
\end{equation}

(At the right-hand side, and henceforth, we follow Einstein’s convention to sum up over identical sub and superscripts within terms where such are present). It follows that

\begin{equation}
\alpha^\nu = \gamma^\mu_\nu \alpha^B_\mu, \quad \text{with } (\gamma^\mu_\nu) : = ((\beta^\nu)*)^{-1} .
\end{equation}

Loosely speaking, the scalar products of any vector with the basis vectors $\mu b$ transform into the coordinates of that vector with respect to an orthonormal basis $\nu u$ by multiplication with matrix $\gamma = (\beta^*)^{-1}$, where $\beta^\nu = < \nu u, \mu b >$, i.e. $\beta$ is the matrix of the coordinate (column) vectors $\mu b$ with respect to the given ON-basis $\nu u$, and $\gamma$ is the adjoint inverse of $\beta$. 
This applied to the node vector basis $\mu b$ and $(38)$ yields the approximate temperature gradient at face $\iota$ as

$$\nabla T_\nu = \gamma^\mu_\nu \nabla B T_\mu.$$  

Thus, the heat current into the cell through face $\iota$ with face vector components $s_{\iota f} \nu = \langle s_{\iota f}, \nu u \rangle$, $\nu \in \{0, 1, 2\}$, is

$$\iota J = \lambda H s_{\iota f} \cdot \nabla T = \lambda H s_{\iota f} \gamma^\mu_\nu s_{\iota f} \nabla B T_\mu = \iota s^\mu \nabla B T_\mu,$$

$\lambda H$ denoting the heat conductivity in the cell.

The heat current through every interface is conserved, i.e. between any two adjacent cells $\zeta$, $\chi$ with the common face labelled $\iota$ in cell $\zeta$ and $\kappa$ in $\chi$ applies

$$\zeta J = - \chi J.$$  

Also, the nodal temperature change in cell $\zeta$ is

$$\frac{dT_\mu}{dt} = \frac{1}{c_v V} ( S + \sum_{\iota=0}^5 \iota J ).$$

where $c_v$ denotes the heat capacity (per volume), $V$ the cell volume, and $S$ any heat source(s) in the cell.

We finally introduce quantities $\iota z_{\mu n} (t)$ $(\iota = 0, ..., 5; \mu = 0, 1, 2)$, which still smoothly vary in time with the temperature $T$ (and that are hence not yet DSC states, but will later be updated as such)

$$\iota z_{\mu n} (t) := \begin{cases} 
2(-1)^\iota T_\mu |_t & \text{if } \mu = [\iota/2] \\
(2\mu+1)T_\mu - 2\mu T_\mu |_{t-\tau/2} & \text{else}
\end{cases}$$

and

$$\iota z_{\mu p} (t) := \begin{cases} 
2(-1)^\iota T_\mu |_t & \text{if } \mu = [\iota/2] \\
\iota z_{\mu n} (t - \tau/2) & \text{else}.
\end{cases}$$

From $(37, 42)$ follows

$$\iota J |_{t+\tau/2} = \iota s^\mu (\iota z_{\mu n} |_{t} - 2(-1)^{\iota n} \delta_{[\iota/2]} \iota z_{\mu n} |_{t+\tau/2}) = \iota s^\mu (\iota z_{\mu n} |_{t} - \delta_{[\iota/2]} \iota z_{\mu p} |_{t+\tau/2}).$$

This, with $(43)$ and continuity of the temperature at the interface, $\iota z_{\mu p} |_{t+\tau/2} = \iota T_\mu |_{t+\tau/2}$, together with $(46)$ imply

$$\iota z_{\mu p} |_{t+\tau/2} = \begin{cases} 
\iota T_\mu |_{t+\tau/2} & \mu = [\iota/2] \\
\iota z_{\mu n} |_{t} & \text{else}.
\end{cases}$$
which form a complete set of recurrence relations for \( z^p \) (given \( z^n \)) and so can be taken as model equations for the connection cycle of a DSC algorithm.

Equations (44) discretely integrated in the time balanced form with increment \( \tau \) yield

\[
T^n_{|t+\tau/2} = T^n_{|t-\tau/2} + \frac{\tau}{c_p V} \left( S + \sum_{\iota=0}^{5} J_{|t} \right),
\]

i.e., with (45, 46, 47), the recurrence relations

\[
i_{|t+\tau/2} = \begin{cases} 
\iota \left( z^n_{|t-\tau/2} + \frac{(-1)^{\iota}}{2} \tau \right) + \sum_{\kappa=0}^{5} \kappa S_{|t-\tau/2} - \frac{\delta_{|t}^{[\kappa]}}{2} \iota \left( z^p_{|t} \right) & \text{if } \mu = |t/2| \\
-\frac{1}{2} \left( 2\mu+1 \iota \left( z^p_{|t} \right) + 2\mu \iota \left( z^p_{|t} \right) \right) & \text{else},
\end{cases}
\]

which provide a complete set of model equations for the reflection cycle of a DSC algorithm. Note that the first line, modulo the factor 2 \((-1)^{\iota}\), always updates the nodal temperature. Of course, this has to be carried out only once per cell and iteration cycle. In this - typical - example the dual state space concept of DSC (needed by Johns’ cycle) creates a redundancy, which is a price for process parallelizability within either parts of the connection-reflection cycle.

Equations (48, 50) can be directly taken as updating relations for total quantities of a DSC scheme. Alternatively, they may be further processed in deriving reflection and connection maps, along with stability bounds for the time step. The proceeding is canonical and amounts in essence to a straightforward transcription of the model equations along the lines of Theorem 4 and corollaries.

It is particularly easy to couple this heat conduction model - within one and the same mesh - to a Maxwell field TLM model in the non-orthogonal setting [17]. In fact, with the node vector definition in [18], the total node voltages are just the scalar products of \( \mu b \) with the electric field, hence the dielectric losses and heat sources per cell are

\[
S = \frac{1}{2} \sigma V E^\nu U^\nu = \frac{1}{2} \sigma V \sum_{\nu} \left| \gamma^\nu U^\nu \right|^2,
\]

for a frequency domain (complex) TLM algorithm, cf. [16]; \( \sigma = 2\pi f \epsilon \tan(\delta) \) denotes the effective loss current conductivity at frequency \( f \) in a mesh cell of absolute permittivity \( \epsilon \) and dielectric loss factor \( \tan(\delta) \); \( \gamma = (\beta^*)^{-1} \) as in [10]. In SPINNER’S Maxwell field solver the model couples, in addition, to magnetic and skin effect losses.

Finally, Fig 4 displays the result of a dispersion test, computed in a square mesh using non-orthogonal cells. It turns out that the heat conduction properties of the mesh are highly insensitive to cell shape and orientation (as of course should be the case).
Fig. 4. Transverse heat conduction over a square mesh using non-orthogonal cells. A Heaviside temperature step is imposed on one side and the transient temperature computed at the opposite side, assuming adiabatic boundary conditions on all but the heated sides. DSC results (+) are plotted over analytical solution (curve).
(a) The mesh. (b) Horizontal. (c) Vertical propagation.

6. Conclusions

Johns’ TLM algorithm can be extended with benefit in two major directions by replacing transmission line links between cells with abstract scattering channels in terms of paired distributions and in admitting non-trivial cell interface scattering. Executing this program lead us in this paper to a new class of Dual Scattering Channel schemes which offer enhanced modeling potentiality and canonical techniques for stable algorithm design. SPINNER’s implementation of a heat propagation scheme coupled to a lossy Maxwell field illustrates the approach. The connection and reflection cycles of a DSC process are (either) completely parallelizable, which can be turned into account in computational performance. DSC schemes open a challenging field to future research. Applications to fluid dynamics are presently under examination.

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SPINNER RF Lab, Erzgiessereistr. 30, DE-80335 München, Germany
E-mail address: s.hein@spinner.de