Implementation of a reaction-diffusion process in the Abaqus Finite Element software

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Résumé :

Pour augmenter les possibilités du logiciel Abaqus, nous proposons une stratégie permettant d’activer les degrés de liberté cachés du logiciel, et d’inclure des phénomènes couplés supplémentaires. À titre d’illustration, nous appliquons cette approche à la simulation d’un processus de diffusion-réaction, le modèle de Gray-Scott, qui génère des structures spatio-temporelles complexes. Plusieurs configurations ont été calculées et comparées aux résultats de la littérature afin d’analyser le potentiel de notre stratégie et d’Abaqus à permettre la prise en compte de phénomènes complexes dans Abaqus.

Abstract :

To increase the Abaqus software capabilities, we propose a strategy to force the software to activate hidden degrees of freedom and to include extra coupled phenomena. As an illustration, we apply this approach to the simulation of a reaction diffusion process, the Gray-Scott model, which exhibits very complex patterns. Several setups have been considered and compared with available results to analyze the abilities of our strategy and to allow the inclusion of complex phenomena in Abaqus.

Mots clefs : reaction-diffusion, finite elements, user subroutines, Gray-Scott model

1 Introduction

Simulating the effect of impurities on the integrity of structures leads to account for several interactions between, e.g., the mechanical fields, the impurities transport and trapping, the thermal fields, etc. The simulation of all these phenomena simultaneously is a complex task, especially when strong couplings are involved or investigated: in the hydrogen embrittlement of metals [1], or in the hydrolysis of polymers [2,3], for instance, mobile species are adsorbed, transported through the material, and trapped on specific sites whose density is time and space dependent [4,5] (e.g., through
the development of plasticity for hydrogen in metals [6]). Furthermore, mechanical fields can be affected by these species because of induced deformations or through modifications of mechanical properties [7,8].

Numerous studies account for such interactions in finite element codes, in various application fields (metal/hydrogen, water/polymer, metal/lithium ions, see [9-17] among others), but very few developments include several phenomena in the computations [18,19], especially in the commercial finite element codes, due to their inherent limitations in terms of available degrees of freedoms at each node. Such an inclusion may, however, be of importance, e.g., to model the behavior of structures in the presence of both impurities and evolving thermal boundary conditions [20]. The aim of this work is thus to introduce some developments performed in Abaqus to solve coupled mechanical-multidiffusion finite element problems. This paper is limited to a reaction-diffusion process between two species, which is solved by using a coupled mechanical-diffusion scheme (‘coupled temp-displacement’ in Abaqus) that allows further developments to account for the mechanical fields as well. First, the multidiffusion implementation strategy is presented, and then an application to the Gray-Scott reaction-diffusion model is presented to illustrate the new capabilities [21,22].

2 Introduction of a multidiffusion process in Abaqus

In order to solve a complex problem with mechanics and multidiffusive fields in a finite element (FE) software, it is mandatory (i) to have a finite element formulation that includes as many degrees of freedom (DOFs) per node as the number of unknown fields, and (ii) to introduce the correct weak formulation for all of these DOFs for solving the problem. Introducing extra DOFs is complex; one may exploit the unused mechanical DOFs (rotations, numbered from 3 to 6, or the third displacement component in 2D problems), or add extra features to the elements (see [23,24] for phase field implementation in Abaqus) through a user element (UEL) routine [25]. One approach of particular interest has been proposed by Chester [26] to solve coupled thermo-chemo-mechanical problems in polymers (this work has been applied in [27] for a simple adsorption process). In this work, a UEL has been developed that activated an extra DOF (not numbered between 1 and 6, for displacements and rotations, nor NT11), in addition to the introduction of a relevant weak formulation as specified in [25]. Such DOFs are included by default in the Abaqus element library for ‘coupled temp-displacement’ procedures, but they are hidden and cannot be accessed through the CAE interface or input files1. These DOFs, numbered from 12 to 30, correspond to NT (for ‘Nodal Temperature’) variables. Once activated by the UEL routine, their boundary conditions can be imposed in the input file and their values (NT, HFL, etc.) can be required in the output database file.

It is worth noting that all the studies mentioned above, where an UEL was used to redefine the problem, have also superimposed additional layers of elements taken from the Abaqus library in order to visualize the results. As demonstrated in [29], it is possible to go further and extend the Abaqus finite element formulation by superimposing a user element to an Abaqus element: the terms that are not included by default in the formulation are introduced through the UEL routine and the Abaqus material library. The approach chosen in the present study combines the advantages of keeping the features of the Abaqus libraries (materials, elements, etc.) and of adding extra terms and DOFs in the finite element formulation by using a superimposed UEL. Thus, the implementation work is optimized because the mechanical behavior needs not being redefined. Even if a multidiffusion process only is considered here, the ultimate goal of a fully coupled mechanical-multidiffusion problem has been kept in mind during the developments.

3 Implementation process

Our strategy is presented in Figure 1: several element layers sharing the same nodes are defined, and a ‘coupled temp-displacement’ procedure is used. In this example, the three UEL layers have the same

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1 Their presence can be inferred from [28], sections 28.3.6 and 28.6.5, in the ‘Output’ subsection.
numbers of DOFs and, assuming that DOFs 11, 12, and 13 represent diffusion DOFs between which a reaction may occur, all user elements layers share the same UEL routine with different parameters. Each layer, in this example, has a specific role:

(i) Layer 1: the Abaqus element (with mechanical DOFs 1 to 3, and 11) involves the mechanical behavior, one diffusion phenomenon (related to DOF 11), and its effects on the mechanical behavior. The problem is strongly coupled (i.e., the diffusion and the mechanical problems are solved simultaneously), but no effect of mechanics on diffusion is possible here (except with developments beyond the scope of this work).

(ii) Layer 2: this UEL layer activates DOF 12 through its related weak formulation (here, diffusion, but it could be any other physical or chemical process), and the coupling between mechanics and DOF 11 (for no complete strong thermo-mechanical coupling is included by default in Abaqus).

(iii) Layer 3 has the same role as layer 2, but for DOF 13.

(iv) Layer 4 defines only the relation between DOFs 12 and 13.

Figure 1. Principle of the implementation of a multidiffusion process.

It is worth noting that other approaches can be considered in the superimposition process (for instance, a single UEL can be used to activate DOFs 12 and 13, and to introduce all the ingredients needed in Abaqus). Each element layer leads to the computation of a specific stiffness matrix, performed either by Abaqus or by the UEL, as shown below:
In the case presented in Figure 1, the stiffness matrices are 16×16 and 20×20 for the Abaqus element and for the UEL, respectively. At the end of the superimposition process, the stiffness matrix of the global problem is 24×24: due to the activation of the extra nodes, the initial Abaqus element stiffness matrix has increased significantly, without any other user manipulation than the activation of hidden DOFs.

This strategy is applied below, where only DOFs 12 and 13 are considered, for illustration. The transient ‘coupled temp-displacement’ procedure is used, even if there is no coupling between DOFs (12,13) and (1,2,3,11) in the present work.

4 Application

The Gray-Scott model is considered here as a test reaction-diffusion process to be implemented.

4.1 The Gray-Scott model

The Gray-Scott (GS) reaction-diffusion model represents a particular case of Turing systems [30], where the reactions of three chemical species are focused on. These species, \( U \), \( V \), and \( P \), define an autocatalytic system so that [21,22]

\[
\begin{align*}
U + 2V & \rightarrow 3V \\
V & \rightarrow P
\end{align*}
\]

(2)

The space-time evolution of species \( U \) and \( V \) can be obtained by solving the following system of differential equations:

\[
\begin{align*}
\frac{\partial u}{\partial t} &= D_u \Delta u - uv^2 + F(1 - u) \\
\frac{\partial v}{\partial t} &= D_v \Delta v + uv^2 - (F + k)v
\end{align*}
\]

(3)

where \( u \) and \( v \) denote the concentrations of species \( U \) and \( V \), respectively, \( D_u \) and \( D_v \) represent their diffusion coefficients, \( F \) is the feed rate for \( U \) and \( k \) the kill rate for \( V \). This reaction has been widely studied as a simple model to reproduce the patterns observed in several chemical reactions (not to speak of natural patterns [31]), as illustrated in Figure 2.

CIMA (chlorite-iodide-malonic acid) reaction in various experimental conditions [32].

FIS (ferrocyanide-iodate-sulfite) reaction in various experimental conditions [33].
CDIMA (chlorine dioxide, iodine, malonic acid) reaction after 12, 20, 35 and 52 minutes [34].

Belousov–Zhabotinsky (BZ) reaction [35]

Figure 2. Examples of chemical patterns.

4.2 Numerical implementation

The patterns induced by the GS model have been the subject of numerous studies from the seminal work by Pearson [36] (see, e.g., [37-42]), including many for entertainment purposes, and a classification of the GS patterns has been proposed (see Figure 3), depending on the \((F, k)\) values. Consequently, many implementations of the GS reaction can be found, based on finite differences and forward Euler integration scheme for efficiency reasons ([43-45], among others, and the very complete webpage of R. Munafò [46]), mainly in 2D. Very few [47-49] apply the finite element method, especially Abaqus. One study [50] includes mechanical coupling, but no indication on the implementation process is given, nor if extra DOFs have been introduced, unfortunately.

We have implemented the GS reaction in Abaqus by introducing DOFs 12 and 13; the details of the RHS vector and of the AMATRIX matrix have been adapted from [48] by considering constant diffusion coefficients, in particular. Computations have been performed with the ‘coupled temp-displacement’ procedure, even if no mechanical nor temperature field is computed. In order to evaluate the ability of our implementation to simulate a GS process accurately, all the results are compared with those given by the Python script written by D. Bennewies [44].

Figure 3. (a) Types of patterns obtained with the GS reaction, and (b) their position in the \((F, k)\) plane (using \(D_u = 2D_v = 2 \times 10^{-3}\)) as defined in [36]. For \((F, k)\) points where no pattern is specified, a constant homogeneous field for \(u\) as well as for \(v\) is expected.

\(^2\) For instance, ‘Gray-Scott reaction diffusion’ keywords in YouTube gives 779 results.
4.3 Configuration studied

The configuration studied is a square domain 2.5×2.5 mm², which is meshed with 250×250 fully integrated linear square elements (i.e., with an element size of 0.01×0.01 mm²), over which as many user elements are superimposed for the activation of DOFs 12 and 13 (representing the concentrations of $U$ and $V$, respectively) and for the integration of the reaction-diffusion process. A transient ‘coupled temp-displacement’ procedure is applied. Periodic boundary conditions are prescribed to DOFs 12 and 13 along the border of the domain, as set in [44]. The following initial conditions for $u$ and $v$ are defined using a DISP user subroutine:

$$
\begin{align*}
  x \in \Omega & \Rightarrow u = 0.5 - 0.01\delta(x); \\
  x \notin \Omega & \Rightarrow u = 1 \\
  x \in \Omega & \Rightarrow v = 0.25 + 0.01\delta(x); \\
  x \notin \Omega & \Rightarrow v = 0
\end{align*}
$$

where $\Omega$ is a rectangular domain $0.125(1+\delta) \times 0.125(1+\delta)$ with $\delta \in [0,1]$ a random perturbation. Finally, $D_u$ and $D_v$ have been set to $10^{-5}$ mm²/s and $2.10^{-5}$ mm²/s, respectively. Several $(F,k)$ parameters have been considered, as listed in Table 1.

<table>
<thead>
<tr>
<th>$k$</th>
<th>0.006</th>
<th>0.022</th>
<th>0.026</th>
<th>0.046</th>
<th>0.062</th>
</tr>
</thead>
<tbody>
<tr>
<td>Expected pattern [44,46]</td>
<td>Propagating wavefronts (Type $\xi$)</td>
<td>Chaotic oscillations (Type $\beta$)</td>
<td>Solitons (Type $\lambda$)</td>
<td>Worms (Type $\mu$)</td>
<td>Negatons (Type $\pi$)</td>
</tr>
</tbody>
</table>

4.4 Results

The Abaqus results for $u$ (NT12) and $v$ (NT13) are presented in Figure 4(a) to 8(a), with the corresponding Python reference results for $u$ shown on (b). All Abaqus computations have been performed with a constant time increment of 10 s, while the python’s one is equal to 1 s. It can be observed that our implementation in the Abaqus code is able to reproduce quite well the results obtained with another software, for various configurations.

It may be noted that the U-skate geometries exhibited by Munaf $[41,46]$ could not be generated, as in [44].

![Figure 4. (a) $u$ and $v$ fields obtained with Abaqus and (b) $u$ field computed with python following [44], using $(F,k) = (0.006,0.037)$ at $t = 800$ s.](image)
An important feature observed in our simulations is a non constant velocity of the pattern front, with a strong influence of the \((F,k)\) parameters. This behavior is consistent with results obtained by other methods, especially in [44]. From the Figure 4 to 8, it might be observed that the front velocity computed by Abaqus has the same order of magnitude than the one obtained using Python.

We have also investigated the effects of the element size and of the time increment (see [46] for a more complete investigation of the time increment influence). The influence of the element size is illustrated in Figure 9 for \((F,k) = (0.006,0.037)\). When the element size increases, the generated
pattern is strongly influenced by the mesh structure and tends to a square rather than a circle. Moreover, the velocity of the pattern front is increased because of a rapidly vanishing $V$ field that annihilates the reaction process.

![Pattern images](image)

Figure 9. Influence of the element size on the Abaqus results at $t = 800$ s for $u$ (left) and $v$ (right), with $(F, k) = (0.006, 0.037)$.

In contrast, decreasing the time increment has no influence on the Abaqus results and on their consistency with [44], except for $(F, k) = (0.022, 0.049)$ where the intensities of the pattern oscillations decrease and a steady state is finally reached for $t$ about 3400 s. For this configuration, the influence of the time increment is shown in Figure 10: when it is decreased from 10 s to 1 s, no steady state is reached with Abaqus up to 5000 s and chaotic oscillations are observed, as in [44].

![Reference pattern](image)

(a) Reference pattern for $u$ [44]

(a) Element size: 0.012 mm$^2$, time increment: 10 s
6 Conclusion

An appropriate application of user elements allows the extension of Abaqus capabilities, including the modification of library elements, the activation of hidden DOFs, and the addition of various physical processes with or without couplings. This study has been focused on the activation of DOFs and on the addition of chemical reactions in Abaqus. An application to the Gray-Scott model has been made successfully. However, this model, though spectacular, has very complex features in term of spatio-temporal evolution, intimately linked with the used parameters. This complexity leads to some difficulties in the definition of the finite element setup in terms of time increment and mesh. Further work will extend the proposed approach to 3D simulations, reactions involving 3 species or more, and mechanical coupling.

To include mechanical interactions, especially, it will be only necessary to introduce in the UELs the related contribution to the weak formulation. Furthermore, a 4th layer might be added to include the coupling between DOF 11 and mechanical fields. Equation (1) thus becomes

\[
\begin{array}{c}
\begin{bmatrix}
K_{uu} & K_{u,v} \\
K_{v,u} & K_{vv}
\end{bmatrix} + \\
\begin{bmatrix}
0 & 0 \\
0 & 0
\end{bmatrix} + \\
\begin{bmatrix}
0 & K_{u,11} \\
K_{v,11} & 0
\end{bmatrix} + \\
\begin{bmatrix}
0 & 0 \\
K_{11,2} & 0
\end{bmatrix}
\end{array}
\]

(5)

Acknowledgments

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