COMPARISON OF DATA TRANSFER METHODS BETWEEN TWO DIFFERENT MESHES.

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Abstract. Many problems solved with the Finite Element Method require more than one mesh (i.e. one specific mesh for each Physics or a remeshing is needed). The Data Transfer Method used, has a great importance in the capacity to solve the problem and in the reliability of the solution. In general, the data is composed of two kinds of fields (defined thanks to the nodal values or at the integration points). In this paper, the more used Data Transfer Method is compared with the Data Transfer Methods based on a Weak Form (using Mortar Element or Finite Volume).

1 Introduction

The numerical resolution with the Finite Elements Method of many multi-physical problems require more than one mesh. On the one hand, some problems require one specific mesh for each Physics (i.e. one mesh for the mechanical part and another one for the thermal part). On the other hand, in some cases, during the computation a remeshing is needed. In all these cases, the Data Transfer Method used to transfer information from one mesh to another has a great importance in the capacity to solve the problem and in the reliability of the solution. In general, two kinds of fields have to be transferred: the first is defined thanks to the nodal values (primary field) and the second one is defined at the integration points (secondary field). Currently, despite the research effort, no Transfer Method has been recognized as the best. Each method has important disadvantages.

This paper compares the more used Data Transfer Method (Element Transfer Method [1, 2]) with the Data Transfer Methods based on the Weak Form (using Mortar Element [3, 4] or Finite Volume [5, 6, 7]).
2 Definition of the problem

In this paper a problem is computed with the Finite Element Method. During the computation, a Data Transfer is needed from the mesh call old mesh to the one call new mesh. Some primary and secondary fields known on the old mesh are needed on the new mesh. The field $P$ is known on the old mesh thanks to the primary field $P^{\text{old}}$. The old mesh is composed of $n_{\text{ele}}^{\text{old}}$ elements, $n_{\text{node}}^{\text{old}}$ nodes and the nodal values of $P^{\text{old}}$ is noted $P^{\text{old}}_n$. The value of $P^{\text{old}}$ on each element $e^{\text{old}}$ is written as:

$$P^{\text{old}} = \sum_{j=1}^{n_{\text{elem}}^{\text{old}}} N_j P_j^{\text{old}}$$  \hspace{1cm} (1)

where $N_j$ is the shape function of the node $j$ in the element $e^{\text{old}}$ and $n_{\text{elem}}^{\text{old}}$ the number of node of the element. The field $S$ is estimated on the old mesh by the secondary field $S^{\text{old}}$. The value of $S^{\text{old}}$ on each element $e^{\text{old}}$ is defined by means of the value at the integration points (noted $S^{\text{old}}$). The fields $P$ and $S$ are evaluated on the new mesh thanks to the primary field $P^{\text{new}}$ and the secondary field $S^{\text{new}}$, respectively. The aim of the Data Transfer Method is to define on the new mesh, the nodal values of $P^{\text{new}}$ ($P_n^{\text{new}}$) and the value at the integration points of $S^{\text{new}}$ ($S^{\text{new}}$). The properties of the Data Transfer Method should be:

- weak numerical diffusion,
- conservation of the extrema,
- easily treatment of the boundaries.

3 Transfer Methods

The Data Transfer Method makes the link between the two discretisations. The reliability of the field on the new mesh is directly linked with the Data Transfer Method used.

3.1 Element Transfer Method

The Element Transfer Method (ETM) is the most commonly used. The computation of the field on the new mesh is done in two steps:

- Firstly, for each characteristic point (node or integration point) of the new mesh a search is done to find the element of the old mesh $e^{\text{old}}$ in which the characteristic point lies inside.

- For the secondary field $S^{\text{old}}$, the nodal values of the element $e^{\text{old}}$ ($S_n^{\text{old}}$) are computed by extrapolation of the values at the integration points of this element ($S^{\text{old}}$). So, in the general case, these nodal values are different for each element. Then, the
value of the field on this characteristic point is computed by interpolation of the nodal values of the element $e^{old}$ ([1, 2]), as:

$$P_{new} = \sum_{j=1}^{n_{elem}} N_j P_j^{old} = P^{old}(x)$$

or:

$$S_{new} = \sum_{j=1}^{n_{elem}} N_j e S_j^{old}$$

This method does not deal with the elements of the new mesh. These elements have no influence on the value of the field. The transfer is done from the old mesh to a characteristic point of the new mesh. On the one hand, for the secondary field, this method does not conserve the extrema because of the extrapolation on the nodal values. On the other hand, the extrema are conserved for the primary field, but due to geometrical approximations, some nodes on the boundary can be outside of the old mesh. So, a special treatment is required for the boundaries.

### 3.2 Mortar Element Transfer Method

The Mortar Element Transfer Method (METM) is based on a weak conservation form of the field (using Mortar Element [3]). The field on the new mesh are not directly computed at the characteristic points. But, it is evaluated considering that the integral of the difference between the value of the fields on the new mesh and the value on the old mesh is null ([3, 4, 8]). This integral is done over the new mesh. To compute this integral, for each element, the nodal values of the secondary field ($S^{old}$ or $S^{new}$) are defined (in function of the values at the integration points of the element). The nodal values are noted $e S^{old}$ for an element of the old mesh and $e S^{new}$ for an element of the new mesh. The value of these fields on each element is defined like the primary field (see equation 1, but in the general case, these nodal values are different for each element). The value of the primary field $P^{new}$ on the new mesh is done thanks to the relation:

$$\sum_{e^{new}=1}^{n_{new}} \int_{e^{new}} (P^{new} - P^{old}) f de = 0$$

and the relation between the secondary field $S^{new}$ and $S^{old}$ is defined for each element $e^{new}$ of the new mesh by:

$$\int_{e^{new}} (S^{new} - S^{old}) f de = 0$$

where $f$ is a weighting function defined on each element $e^{new}$ (like a primary field). The nodal value of the function $f$, $f_{\bullet}$, can take any value.
The transfer relation of the secondary field (equation (5)) can be written like:

\[
\sum_{A=1}^{n_{new}} \left( \sum_{B=1}^{n_{new}} N_{AB}^{1}(e^{new}) S_{B}^{new} - N_{A}^{2}(e^{new}, S^{old}) \right) = 0. \tag{6}
\]

Where \( N_{AB}^{1}(e^{new}) \) and \( N_{AC}^{2}(e^{new}, S^{old}) \) are the mortar elements linked with the element \( e^{new} \) defined as:

\[
N_{AB}^{1}(e^{new}) = \int_{e^{new}} N_{A} N_{B} \, de \quad N_{AC}^{2}(e^{new}, S^{old}) = \int_{e^{new}} N_{A} S^{old} \, de \tag{7}
\]

where \( N_{i} \) is the shape function of node \( i \) (A or B) in the element \( e^{new} \). \( N_{AC}^{2} \) is the coupling term.

The transfer relation of the primary field (equation (4)) can be written like:

\[
\sum_{A=1}^{n_{new}} \left( \sum_{B=1}^{n_{new}} N_{AB}^{1}(e^{new}) P_{B}^{new} - \sum_{C=1}^{n_{old}} N_{AC}^{2}(e^{new}, S^{old}) \right) f_{A} = 0. \tag{8}
\]

Where \( N_{AB}^{1} \) and \( N_{AC}^{2} \) are the mortar elements defined as:

\[
N_{AB}^{1} = \sum_{e^{new}=1}^{n_{new}} N_{AB}^{1}(e^{new}) \quad N_{AC}^{2} = \sum_{e^{new}=1}^{n_{new}} N_{AC}^{2}(e^{new}, N_{C}) \tag{9}
\]

where \( N_{C} \) is the shape functions of the node \( C \) in the corresponding element \( e^{old} \).

### 3.2.1 Computation of mortar element

The first Mortar Element \( N_{AB}^{1} \) is computed by numerical integration over the element of the new mesh \( e^{new} \) (because is a product of two shape functions of this element). The evaluation of the coupling term (the second Mortar Element, \( N_{AC}^{2} \)) is more complex. Because, in the general case, the field \( S^{old} \) is not continue on each element of the new mesh. In addition, the sum of the shape functions \( N_{old}^{C} \) on each element of the old mesh is not a polynomial function on each element of the new mesh \( (e^{new}) \). A numerical and an exact integration are used to compute this Mortar Element.

**Numerical integration** The mortar element is computed by numerical integration over each element of the new mesh. For the element of the new mesh \( e^{new} \), the computation is done with \( n^{ip} \) integration points. The numerical integration supposes that the value of the field on the old mesh can be evaluated by a polynomial function on each element of the new mesh.
Exact integration  Each element $e^{\text{new}}$ of the new mesh is divided in $n^{\text{sub}}_e$ elements, as each sub-element ($e^{\text{sub}}$) is only over one element of the old mesh. So, on each element $e^{\text{sub}}$, $N^{\text{odd}}_C$ is a polynomial function. Finally, the mortar element is computed exactly by numerical integration over each sub-element $e^{\text{sub}}$ (because is an integration of polynomial function). The exact integration of mortar element considers all intersections between the element of the new mesh and the elements of the old mesh. However, with the numerical integration, the mortar elements are evaluated in function of the intersection in which the integration point lies inside. So, the intersections than are smaller than the influence area of the integration point can be ignored.

3.2.2 Evaluation of the field on the new mesh

Global solving (GS)  The relation between the nodal value of the field $S^{\text{new}}$ on each element of the new mesh and the value of the field $S^{\text{old}}$ (equation (6)) can be written as:

$$
\begin{bmatrix}
N^1_{11}(e^{\text{new}}) & \cdots & N^1_{1n^{\text{elem}}}(e^{\text{new}}) \\
\vdots & \ddots & \vdots \\
N^1_{n^{\text{elem}}1}(e^{\text{new}}) & \cdots & N^1_{n^{\text{elem}}n^{\text{elem}}}(e^{\text{new}})
\end{bmatrix}
\begin{bmatrix}
S^{\text{new}}_1 \\
\vdots \\
S^{\text{new}}_{n^{\text{elem}}}
\end{bmatrix}
= 
\begin{bmatrix}
N^2_{1}(e^{\text{new}}, S^{\text{old}}) \\
\vdots \\
N^2_{n^{\text{elem}}}(e^{\text{new}}, S^{\text{old}})
\end{bmatrix}
$$

The size of this equation is equal to the number of node of the element of the new mesh. The value at each integration point is equal to the interpolation of the nodal values. This equation can be solved for each element of the new mesh.

For the primary field, the equation (8) can be written as:

$$
\begin{bmatrix}
N^1_{11} & \cdots & N^1_{1n^{\text{new}}} \\
\vdots & \ddots & \vdots \\
N^1_{n^{\text{new}}1} & \cdots & N^1_{n^{\text{new}}n^{\text{new}}}
\end{bmatrix}
\begin{bmatrix}
T^{\text{new}}_1 \\
\vdots \\
T^{\text{new}}_{n^{\text{new}}}
\end{bmatrix}
= 
\begin{bmatrix}
\sum_{C=1}^{n^{\text{old}}} N^2_C T^{\text{old}}_C \\
\vdots \\
\sum_{C=1}^{n^{\text{old}}} N^2_{n^{\text{new}}C} T^{\text{old}}_C
\end{bmatrix}
$$

The size of this equation is equal to the number of node of the new mesh. In addition, the solution of this equation cannot certify the conservation of the extrema.

Local solving (LS)  To obtain a local system, a diagonal matrix is used. The value of the diagonal term is equal to the sum of the line (or the column, because the matrix is symmetric). This is totally equivalent to the row-sum technique used to lump mass matrix in explicit time integration method.

So the value of the field $S$ on each node of the element $e^{\text{new}}$ is done by:

$$
S^{\text{new}}_A = \frac{N^2_A(e^{\text{new}}, S^{\text{old}})}{\sum_{B=1}^{n^{\text{elem}}} N^1_{AB}(e^{\text{new}})}
$$

The value at each integration point of the element $e^{\text{new}}$ is computed by interpolation of the nodal values.
And, the value of the field $P$ on each node is done by:

$$P_{A}^{\text{new}} = \frac{\sum_{C=1}^{n_C^{\text{old}}} N_{AC}^2 P_{C}^{\text{old}}}{\sum_{B=1}^{n_B^{\text{new}}} N_{AB}^1}$$

(13)

With this method, the weak conservation of the field is done in a cell composed of the elements of the new mesh including the node $A$. This technique increases the area of computation of nodal value and in the same time the numerical diffusion. But, in opposition of the global solving, the local solving conserves the extrema.

To sum up, the evaluation of the field is done at the node of the new mesh on a function of the shape function of the elements of the new mesh and the value of the field on the elements of the old mesh.

### 3.3 Finite Volume Transfer Method

With the Finite Volume Transfer Method (FVTM), each field is rebuild thanks to a finite volume mesh (called old finite volume mesh on the old finite element mesh and new finite volume mesh on the new finite element mesh). The finite volumes are called cells. For the primary field, the cells are based on the node of the finite elements (see figure 1). On the finite volume mesh build for a secondary field, the cells are based on the integration points of the elements of the finite element mesh (see figure 1). So, each field is transferred from one old finite volume mesh to a new finite volume mesh. The same procedure is used to transfer the primary and the secondary field (to more information see [5]).

![Figure 1: Finite element mesh and finite volume mesh based on integration points and on the nodes](image)

The value of the field $\varphi^{\text{new}}$ (primary or secondary) on one cell is equal to the value of the field at the corresponding characteristic point (node or integration point) of the finite
element mesh. The value of the of $\varphi^{\text{new}}$ on one cell $c^{\text{new}}$ of the new finite volume mesh ($\varphi^{\text{new}}$) is done by:

$$\varphi_{c^{\text{new}}}^{\text{new}} = \int_{c^{\text{new}}} \varphi_{c^{\text{old}}}^{\text{old}} \, dc.$$  

(14)

Where $\varphi_{c^{\text{old}}}^{\text{old}}$ is the value of the field on the old finite volume mesh. Like with the mortar element, a numerical or an exact integration can be used to evaluate the coupling between the cells of the old and the new mesh.

**Numerical integration** The value of the field $\varphi^{\text{new}}$ on the cell $c^{\text{new}}$ is defined by numerical integration over this cell. This computation supposes that the field on the old finite volume mesh can be evaluated by a polynomial function on this cell.

**Exact integration** A super-mesh is built, each cell $c^{\text{new}}$ of the new mesh is divided in $n_{\text{sub}}^{c^{\text{new}}}$ sub-cells, like each of them corresponds only to one cell of the old mesh. For each cell $c^{\text{new}}$ the value of $\varphi^{\text{new}}_{c^{\text{new}}}$ is done by:

$$\varphi_{c^{\text{new}}}^{\text{new}} = \sum_{c^{\text{sub}}}^{n_{c^{\text{sub}}}} V_{c^{\text{sub}}} \times \varphi_{c^{\text{old}}}^{\text{old}}.$$  

(15)

Where $\varphi_{c^{\text{old}}}^{\text{old}}$ is the value of the cell of the old mesh corresponding to the sub-cell $c^{\text{sub}}$. $V_{c^{\text{sub}}}$ is the value of the volume (the surface in two dimensions) of the sub-cell $c^{\text{sub}}$. The exact integration of coupling considers all intersections between the cell of the new mesh and the cells of the old mesh.

4 Examples

The difference between these Data Transfer Methods is shown on two dimensional academic examples. These examples expose the numerical diffusion and the evaluation of the data on the boundaries. The meshes are composed of quadrilateral elements. The evaluation of the Mortar Element $N_{AB}^1$ is done by numerical integration using two Gauss points in each direction. For the numerical integration, the coupling elements (Mortar Element $N_{AC}^2$ or coupling between cells) are evaluated with five Gauss points in each direction. For the exact integration, the evaluation of the coupling elements is done using six Gauss points on each triangle of the subdivision (to exact integration of quadratic function). These two examples are be used to compare the transfer of a primary field with the Transfer Element Method and the Transfer with Mortar Element in [9].

4.1 Numerical diffusion

The numerical diffusion lies to the Data Transfer Operator is studied with this example. A primary and a secondary field are known on the square. The square’s sides are meshed by 30 elements. A rotation of $\pi/8$ is apply on this square. The Data is transferred from
the moved mesh to the initial mesh. The initial value of the field is of a hundred inside a circle and null outside. The centre of the circle is identical to the centre of the square and the radius is the half of the side’s square.

![Figure 2: Numerical diffusion after twenty transfers of the secondary field (4 Gauss points)](image)
The comparison of the numerical diffusion is done after twenty transfers (see figures 2(a), 2(b), 2(c) and 2(d)). These figures show that the numerical diffusion is minimized with the METM with global solving and the FVTM (the both with numerical computation of coupling terms). But the METM with global solving does not conserve the extrema, this Transfer Method introduces oscillations around a steep variation (see figure 2(b)). In this case, the difference between the numerical and the exact integration of coupling terms is not significant.

4.2 Influence of boundaries

To study the influence of the boundaries, a mesh of a disc with a hole is used. Like the first problem, the transfer is done after rotation of \( \frac{\pi}{8} \) from the moved mesh to the initial mesh. The half circle is divided in fifteen elements and the radius in ten. The exact value of the field is a linear function of the abscissa (from zero to hundred). The difficulty is that in the general case, the nodes on the boundaries of the new mesh are not inside an element of the old mesh.

Figures 3 show the value of the field after sixteen transfers (one revolution). This problem proves that the ETM applied to a primary field requires a special technique to deal with the boundaries. The nodes located on the boundaries of the new mesh do not lie inside any element of the old mesh. This problem does not appear with the transfer of a secondary field (see figures 3), because the computation of the field is done on the integration points of the new mesh and these points generally lie inside of an element of the old mesh. In addition, the computation of the coupling terms by numerical integration does not consider the part of the elements that is outside of the other mesh. This explains that the METM with global solving does not introduce any error after the transfer (see figure 3(b)). The error after the transfer with the other Transfer Methods is a numerical diffusion and not a wrong evaluation of the field on the boundaries (see figures 3(c), 3(a) and 3(d)). The exact integration of the coupling terms introduces an error of space discretisation of the boundaries. The parts of the element of the old mesh that do not lie inside any element of the new mesh are not considered on the Mortar Elements. In the same time, the parts of the element of the new mesh that do not lie inside any element of the old mesh are considered and the value of the field inside is null. So, the integral of the field over the new mesh is not equal to the integral over the old mesh. This error impairs the quality of the solution.
Figure 3: Numerical diffusion after sixteen transfers of the secondary field (4 Gauss points)

5 Conclusion and future works

In conclusion, this paper presents a comparison of Data Transfer Methods between two different meshes. The Data Transfer Methods based on the Weak Form (the Mortar
Element Transfer Method and the Finite Volume Transfer Method) are compared to the more used, the Element Transfer Method. With these methods (METM and FVTM) the value of the field at one characteristic point (a node or an integration point) of the new mesh is a function of the value of field on the old mesh and the elements of the new mesh. This paper shows that with the numerical integration of the coupling terms, these methods deal with complex boundaries without any specific procedure. In addition, the METM with global solving minimizes the numerical diffusion, but the global computation can introduce oscillations around steep variations of the field. So, this method cannot conserve the extrema. On the other hand, the local computation increases the smoothing of the field.

These Data Transfer Methods are applied to couple the simulation of the friction stir welding and the numerical simulation after the end of the welding. A remeshing is done after the end of the welding to continue the computation on a more appropriate mesh.

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