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On some aspects of the CNEM implementation in 3D in order to simulate high speed machining or shearing

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Abstract

This paper deals with the implementation in 3D of the Constrained Natural Element Method (CNEM) in order to simulate material forming involving large strains. The CNEM is a member of the large family of mesh-free methods, but is at the same time very close to the finite element method. The CNEM’s shape function is built using the constrained Voronoï diagram (the dual of the constrained Delaunay tessellation) associated with a domain defined by a set of nodes and a description of its border.

The use of the CNEM involves three main steps. First, the constrained Voronoï diagram is built. Thus, for each node, a Voronoï cell is geometrically defined, with respect of the boundary of the domain. Then, the Sibson-type CNEM shape functions are computed. Finally, the discretization of a generic variational formulation is defined by invoking an ”stabilized conforming nodal integration”. In this work, we focus especially on the two last points. In order to compute the Sibson shape function, five algorithms are presented, analyzed and compared, two of them are developed. For the integration task, a discretization strategy is proposed to handle domains with strong non-convexities. These approaches are validated on some 3D benchmarks in elasticity under the hypothesis of small transformations. The obtained results are compared with analytical solutions and with finite elements results. Finally, the 3D CNEM is applied for addressing two forming processes: high speed shearing and machining.

Keywords: Meshless methods, Natural neighbour Galerkin methods, 3D constrained natural element method, Forming processes simulation
1. Introduction

The "meshless" methods are alternative approaches to the finite elements method to solve partial differential equations (PDE). As finite elements, the meshless methods allow to approximate fields with values stored on points, called nodes, distributed on the studied domain. With finite elements the approximation is defined element by element, every element being defined by all the nodes connected to it. Generally, for a same set of nodes several meshes are possible. That leads the approximation to a mesh dependence. With meshless methods the connectivity between nodes is only defined by the notions of neighborhood or domains of influence. For these approaches, the construction of the approximation is thus not dependent any more on an any meshing but only on the relative position of the nodes. An important motivation of the meshless approaches is to avoid numerical problems resulting from the mesh, for example mesh distortion when studied domain is subjected to large strains. The meshless methods can be used advantageously for the numerical simulation of manufacturing processes like extrusion, injection, machining or blanking for example.

The finite difference method generalized for arbitrary cloud of nodes [1] was the first step toward the meshless methods. Other methods were then proposed: the "Smooth Particle Hydrodynamics" (SPH) [2] (an overview of this approach is given in [3]), the "Diffuse Element Method" (DEM) [4] [5] [6], the "Element Free Galerkin Method" (EFGM) [7], the "Reproducing Kernel Particle Method" (RKPM) [8], the "Natural Element Method" (NEM) [9], and the "Method of Finite Spheres" [10].

The NEM is based on the natural neighbor (NN) interpolant introduced by Sibson [11]. The NN interpolant uses a geometric construction: the Voronoi diagram associated with the cloud of nodes spread on the studied domain. Unlike the other meshless approaches: i) the support of NN shape functions are automatically defined, ii) the value of NN shape functions associated with internal nodes are null on the border of the domain. This last property, only valid for convex domains, is particularly interesting because it allows a direct imposition of the boundary conditions, exactly as in the finite elements framework. To extend these property to non-convex domains two strategies exist. The first one, based on the alpha forms [12], propose a very simple way to describe the border of the domain if this last one remains weakly non-convex. The second one, the Constrained Natural Element Method (CNEM) [13], handles the case of strongly non-convex do-
mains by building the interpolation on the Constrained Voronoï diagram, constrained by the border of the domain. The major inconvenience of NN shape functions is their significant calculation time.

This article has two main objectives. First of all it aims at clearing up certain properties inherent to the Sibson’s shape functions in the 3d for possibly non-convex domains. Secondly it gives a presentation and a comparison, in term of performance and applicability, also for non-convex 3d domains, of different needed algorithms. This paper gives also the headlines of the implementation of the CNEM in a finite transformations context and shows results for two kinds of cutting manufacturing processes. The modelling of metal cutting, well known as being particularly complex [14], gives interesting applications for the proposed approach.

2. Overview of the constrained natural neighbor (CNN) interpolation

The natural neighbor interpolant is based on a geometrical construction, the Voronoï diagram. This Voronoï diagram is associated with a cloud of nodes distributed on the studied domain. The nodes are points of the domain where the degrees of freedom (DOF) of the interpolation are defined. The NN interpolation is a linear combination of NN shape functions, one for each node. The NN shape functions are local in space and interpolating (the value of the interpolation on a node is equal to the DOF of the node).

The Voronoï diagram is the dual of the Delaunay tessellation associated with the same cloud of nodes. However, unlike to shape functions whose construction are based on a tessellation, the NN shape functions only depend on the space distribution of the nodes. Figure 1 illustrates this aspect for a fixed nodal distribution and at a given point $x$. In this figure, the value of the shape function associated with each node is proportional to the diameter of the red disk. For the NN shape functions, the neighbors of $x$ (nodes of the domain for whom the associated shape functions are not equal to zero) are the closest nodes to $x$, nodes $j$, $h$, $k$, and $g$. For this interpolation, the more a node is close to $x$, the more his influence (the associated value of the shape function) is important. We can note that the value of the shape functions associated with the neighbors of $x$ are almost identical, because, for this example, $x$ is roughly at the same distance from them. This is not the same for the finite elements linear shape functions. For a Delaunay tessellation represented in Figure 1 (on the right), the neighbors of $x$ are only $j$, $g$ and
$k$ (for $k$ the associated shape function value is practically zero). For this tessellation $h$ is not a neighbor of $x$ although $h$ being at a similar distance from $x$.

Figure 1: Values of neighbors Sibson NN shape functions (left) - Finite elements linear shape functions (right)

Compared with other meshless shape functions, the NN shape functions have a significant advantage. On the domain border, the value of the interpolated field by NN shape functions depends only on the value of this field on border node, and linearly (figure 2). In other words, the internal nodes have no influence on border of the domain, the associated shape functions vanished on the border. This property is only valid for convex domain.

Figure 2: Values of neighbors Sibson NN shape functions close to convex border

For a **non-convex domain**, on or close to a **non-convex border**, the interpolated field may not only depend on internal nodes but also on nodes being located on the other side of this border (figure 3 on the left).
constrained natural neighbor shape functions CNN [15] were introduced to cure this gap. The CNN shape functions are an extension of the NN shape functions for a non-convex domain: the calculation of NN shape functions is based on the Voronoï diagram, the calculation of CNN shape functions is based on the Constrained Voronoï diagram. The constrained Voronoï diagram is constrained by the border of the domain. Figure 3 illustrates the contribution of the CNN shape functions close to a non-convex border (on the left) comparatively to the NN shape functions (on the right).

3. Computation of CNN shape functions

In this section, for convenience reasons, we often refer to the two-dimensional case ($n = 2$), followed in italic by a generalization to the three-dimensional case ($n = 3$) or more ($n > 3$). We consider a domain $\Omega$ from $R^n$ the border of which is denoted $\Gamma$. We also consider a set, $S$, of nodes spread over $\Omega$ and its border $\Gamma$.

3.1. Voronoï diagram and Delaunay tessellation

Let $p$ be a node of $S$. The Voronoï domain [16] associated with $p$, noted $Vor_S(p)$, is the set of points of $R^n$ closer to $p$ than any others node $q$ of $S$ or equal distance:

$$Vor_S(p) = \{ x \in R^n : d(x,p) \leq d(x,q), \forall Q \in S, q \neq p \} \quad (1)$$

where $d(.,.)$ is the Euclidean distance between two points. The Voronoï diagram is the dual of the Delaunay tessellation [17]. The circumcircle (circumsphere, circum-hyper-sphere) of each triangle (tetrahedron, simplex) does not contain any node inside itself.
Let us define some vocabulary:

- **Voronoi vertex**: center of the circle (*sphere, hyper-sphere*) which circumscribes a *Delaunay triangle* (*tetrahedron, simplex*). This vertex is thus associated with three generator nodes (\((n+1)\) in \(nd\)), it is in connection with three other vertices (\((n+1)\) in \(nd\)), those associated with adjacent *Delaunay triangles* (*tetrahedron, simplex*).

- **Voronoi edge**: segment connecting two adjacent vertices. Each edge is associated with two geneartor nodes (\(n\) in \(nd\)).

- **Voronoi face**: median plan (*hyper-plan*) between two neighboring nodes (*generator nodes*).

- **Voronoi cell**: area of *Voronoï*.

### 3.2. Constrained Voronoï diagram and constrained Delaunay tessellation

The definition of constrained Voronoï diagram is based on the visibility criterion [18]: a node \(q\) is known as visible from another node \(p\), and reciprocally, if, and only if, the segment \([p, q]\) do not pass through the domain border \(\Gamma\) of \(\Omega\), and is not outside this last.

In Figure 4, the segments \([a, b]\), \([a, c]\), and \([a, d]\) pass through \(\Gamma\). The nodes \(b\), \(c\), and \(d\) are thus not visible from \(a\) and reciprocally. In addition, the segment \([e, c]\) is outside the domain, the nodes \(e\) and \(c\) are thus not visible from each other.

![Figure 4: The visibility criterion](image)

The definition of the constrained Voronoï area associated with node \(p\), \(\text{Vor}_S^\Gamma(p)\), found classically in the literature [19], is as follow:
\( \text{Vor}^S_\Gamma (p) = \{ x \in \mathbb{R}^n : d(x, p) \leq d(x, q), \forall q \in S, q \neq p, x \text{ visible from } p \text{ and } q \} \) (2)

This definition restricts \( \text{Vor}^S_\Gamma (p) \) to its internal domain part. In order to be able to compute shape functions thereafter, this area should not be \textit{truncated} by the domain border.

A more extended definition is:

\( \text{Vor}^S_\Gamma (p) = \{ x \in \mathbb{R}^n : d(x, p) \leq d(x, q), \forall q \in S, q \neq p \land p \text{ visible from } q \} \) (3)

In Figure 5 is represented, on the left, the (no constrained) Voronoï cell associated with node \( p \). On the right-hand side drawings, two constrained

Figure 5: Voronoï cell of a node \( p \), no-constrained (left), constrained (right)

cells associated with the same node \( p \) are present: on the top the cell resulting form the definition (2), on the bottom the cell resulting from the definition
The border of the domain is defined by the green segments. In the definition (3), the visibility criterion is applied only to nodes of the domain whereas in the definition (2) the visibility criterion is applied to all the points of the domain.

It must be noticed that the Voronoï diagram of a cloud of nodes, is the constrained Voronoï diagram, constrained by the convex hull of this cloud of nodes. For a non-convex domain, the convex hull of the cloud is not anymore its border. This explains the difference between the two diagrams, constrained and not constrained, for non-convex domains.

The constrained Voronoï diagram (CVD) is the dual of the constrained Delaunay tessellation (CDT) [20]. For this tessellation the circum circle (sphere, hyper-sphere) of each triangle (tetrahedron, simplex) does not contain any visible node from the nodes of this triangle. In 2d the constrained Delaunay tessellation always exists, for convex domain or not. This is unfortunately not true anymore in 3d for certain non-convex domains. An example is given with the Schönhhardt polyhedron [21] (figure 6). A solution in these cases consists in adding some nodes on the border of the domain.

![Figure 6: Schönhhardt polyhedron](image)

In our implementation, the construction of the CVD is done through its dual, the CDT. To build this CDT we use the algorithm proposed by Si and Gärtner [22] and implemented in TETGEN code [23] by the author. The input data of this algorithm are the nodes of the domain and the border of the domain described in the form of PLC (Piecewise Linear Complex) [24].
In our case, for a 3d domain, the tessellation of the border is a very simple PLC where all the faces are triangles.

3.3. Definition of the Sibson CNN shape functions

Let \( x \) be a point inside \( \Omega \), the point where we want to evaluate the interpolation, such as \( x \notin S \wedge x \notin \Gamma \). We denote \( V_S^\Gamma \) the constrained Voronoi diagram of \( \Omega \), \( V_{S+x}^\Gamma \) the constrained Voronoi diagram of \( \Omega \) with addition of \( x \) as node and \( D_{S+x}^\Gamma \) the corresponding constrained Delaunay tessellation, \( c'_x \) the Voronoi cell associated with \( x \) in \( V_{S+x}^\Gamma \).

The constrained natural neighbors of \( x \) are the nodes of \( \Omega \) whose Voronoi cell are directly close to the cell of \( x \) in \( V_{S+x}^\Gamma \), in other words, the nodes of \( \Omega \) connected to \( x \) by a Delaunay edge in \( D_{S+x}^\Gamma \).

Let say that \( v_i \) is such a neighbor, and \( c_{v_i} \) its associated Voronoi cell in \( V_S^\Gamma \). Similarly to the NN shape function [11][25][26], the value of Sibson CNN shape function associated with \( v_i \) at point \( x \), denoted \( \phi_{v_i}^s(x) \), is given by:

\[
\phi_{v_i}^s(x) = \frac{\|c'_x \cap c_{v_i}\|_L}{\|c'_x\|_L},
\]

where \( \|\cdot\|_L \) is the algebraic Lebesgue measure (length in 1d, area in 2d, volume in 3d...). For CNN shape functions, unlike NN shape functions, this measure can be negative or equal to zero for strongly non-convex 3d domains. Hence, for these domains, there are points where shape functions are non-definite. We return on this aspect in Appendix A.

A diagram illustrating the computation of Sibson shape functions in 3d is given Figure 7. In Figure 8 we can see the difference resulting from the use of constrained Voronoi diagram versus Voronoi diagram on a Sibson shape function (2d) (for an internal node near to non-convex border (top), border node (bottom)).

3.4. Properties of CNN shape functions

a) Delta Kronecker property: like NN shape functions, the CNN shape functions are interpolating:

\[
\phi_{n_i}(n_j) = \delta_{ij}, \; \forall (n_i, n_j) \in S^2
\]
Figure 7: Computation of Sibson shape functions in 3d
Figure 8: Contour of Sibson shape functions: NN (left) versus CNN (right)
\(b)\) **Partition of unity:** by construction CNN shape functions check the partition of unity:

\[
\sum_{i=1}^{n(x)} \phi_{v_i}(x) = 1, \ \forall \ x \in \Omega \tag{6}
\]

where \(n(x)\) is the number of neighbors of \(x\).

c) **Local coordinate property and linear consistency:** Sibson\cite{11,25,26} and Piper\cite{27} showed that NN shape function reproduce exactly coordinates of this point, and by extention CNN also verifies:

\[
x = \sum_{i=1}^{n(x)} \phi_{v_i}(x) x_i \tag{7}
\]

Properties b) and c) imply that CNN shape functions reproduce exactly the linear functions.

d) **Linearity on the border:** on \(\Gamma\), the CNN shape functions are strictly linear \cite{15}. For recall, this border is: the convex hull of the nodes of \(\Omega\) if we uses a Voronoï diagram, the real border of \(\Omega\), if we uses a **constrained** Voronoï diagram.

Properties a) and d) imply that on a segment (triangle in 3d) of the border, the CNN interpolation depends only on the DOF of the nodes of this segment (triangle).

e) **Support of CNN shape functions:** the support of a CNN shape function associated with a given node \(i\) of \(\Omega\), is the set of the circles (sphere, hyper-sphere) circumscribed with the Delaunay triangles (tetrahedron, simplexes) connected to \(i\). This support is truncated by \(\Gamma\) (see Figure 8, right).

f) **Continuity:** Sibson NN or CNN shape functions are \(C^0\) on the nodes, \(C^1\) on the Delaunay circles (sphere, hyper-sphere), and \(C^\infty\) everywhere else \cite{28}.

3.5. Computation of Sibson CNN shape functions

We denote by \(P_{\text{int}}\) the polyhedron defining the common domain of \(c'_x\) and \(c_v\) (\(P_{\text{int}} = c'_x \cap c_v\)), \(v\) being one of the neighbors of \(x\) in \(V_{S+T}^T\). The computation of \(\phi_{v_i}^s(x)\) is made in two steps: the first one is the local modification of \(V_S^T\) by the insertion of the point \(x\), the second one concerns the calculation of \(\|P_{\text{int}}\|_{L}\).
For the first step the use of an incremental algorithm is very well adapted. Two similar algorithms can be used [29]: the Bowyer algorithm [30], which works with Voronoï vertices, or the Watson algorithm [31] which works with Delaunay tetrahedrons. These two algorithms, initially developed for the construction of VD/DT, can be used in a incremental way on an existing CVD/CDT. Both of them maintain the constrained character of the initial diagram $V^T_S$ or Delaunay tesselation $D^T_S$. The firts step can be split in three main sub-steps:

1. Search for the tetrahedron which contains the point $x$. For that it is possible to use a gradient descent based on Voronoï graph (Schmitt and Borouchaki [32]), or that based on the Delaunay graph (Green and Sibson [33]). In the case of non-convex domains this two search can fail if search starting from a no visible tetrahedron by the point $x$. It is thus necessary to implement other types of search, such frontal search, or octree search.

2. Search for all the tetrahedrons whose circumscribed sphere contains the point $x$ and break them. This search is local, it is done by visiting the nearest tetrahedrons to the first one.

3. Remesh obtained cavity by connecting each cavity triangle to the $x$.

The second step can be done in different ways:

a) By Lasserre recursive algorithm

Alfaro et al. [34], like Sambridge et al. [35][36], propose to use the recursive Lasserre algorithm. The Lasserre algorithm can be used to compute the volume of convex polyhedrons described in a H-representation way (volumes delimited by a set of half-planes).

In the case of convex domain, the Voronoï cells are convex and, as being the intersection of two convex polyhedrons, $P^{int}$ is also convex. $P^{int}$ can then be described as a volume delimited by a set of half-planes. This set consists of all the faces of $c_v$ (cell of $v$ in $V^T_S$), and the common face to $c'_v$ (cell of $v$ in $V^T_{S+x}$) and $c'_x$ (see Figure 9). The Lasserre recursive algorithm can therefore be used to compute the volume of $P^{int}$. This algorithm can be succinctly described as follow:

Let $P$ a convex polyhedron in $\mathbb{R}^n$, delimited by $m$ half-planes. Each one of these half-planes can be represented by an inequation, which can be translated in a matrix form by the inequation:
\[ A \cdot x \leq b, \]  
\[ V(n, A, b) = \frac{1}{n} \sum_{i=1}^{m} \frac{b_i}{\|a_i\|} V_i(n-1, A, b), \]  

where \( A \) is an \( m \times n \) matrix, \( x \) the column of the \( n \) components of the point \( x \), and \( b \) a column of \( m \) terms.

The volume measurement of \( P \), denoted \( V(n, A, b) \), can be calculated by the following recursive formula:

The Volume measurement of \( P^{\text{int}} \) can also be defined as the measurement of volume variation of its associated Voronoï cell after the insertion of the
point \( x \) (see Figure 10).
This approach can be applied only if the cell of each neighbour node is non-infinite. If it is not the case, it is impossible to calculate the measurement of its volume, but considering that we need only for the measurement of the variation of volumes of the cells after insertion of the point \( x \), and not of their volume, this problem can be solve by truncating the infinite cells (addition of faces).

\[ \text{c) and d) By topological approaches} \]

It is possible to rebuild the topology of \( P_{\text{int}} \) polyhedron (faces, edges, vertices) with only c) CVD connectivity [37] or d) CDT connectivity [38]. No computations of geometrical intersections are needed. In both of these approaches the faces of the \( P_{\text{int}} \) polyhedron are classified into three categories (see Figure 11):
- \( f_i c'_x \), the face of \( c'_x \) cell having as second node \( v \),
- \( F_{\text{trq}} \), the set of \( c_v \) cell faces truncated by the edges of \( f_i c'_x \),
- and possibly \( F_{\text{trq}} \), the set of \( c_v \) cell faces being completely inside \( c'_x \).

\[ \text{e) By Watson algorithm} \]

Watson [39] proposes to use the Lawrence polyhedron decomposition [40] for the computation of \( \|P_{\text{int}}\|_L \). Taking a polyhedron \( P \), and an hyperplane
Figure 11: Topological decomposition of the $P^\text{int}$ polyhedron
\( \pi \) which is not parallel to any face of \( P \), its volume measurement can then be calculated by summing \textit{algebraically} the volume measures of the simplexes \( \text{sp}_{s}^{\text{Law}} \). \( \text{sp}_{s}^{\text{Law}} \) are build for each vertex \( s \) of \( P \) by taking in addition to the hyperplane \( \pi \), the hyperplanes associated with the faces of \( P \) having \( s \) as vertex. Each one of these hyperplanes divides space into two signed parts, the positive one includes \( P \). The sign of the volume measurement of a simplex \( \text{sp}_{s}^{\text{Law}} \) is the sign of the part of the space in which it is located, which is the product of the signs of this space part compared to each hyperplane associated with \( \text{sp}_{s}^{\text{Law}} \) \textit{(the hyperplane \( \pi \) is not considered)}. In Figure 12, we can see an example of the Lawrence polyhedron decomposition in \( 2d \).

Watson takes, for his algorithm, the \( \pi \) plane as the median plane between the point \( x \) and \( v \). An illustration of this is given in Figure 13.

The Watson algorithm is not applicable on evaluation point \( x \) being located on a face of any Delaunay tetrahedron. If the calculation of the interpolation is needed at such a point an other algorithm must be used.
3.6. Efficiency comparison of the five proposed algorithms

The five methods of Sibson shape function calculation were implemented (C, C++ languages). For the method based on the Lasserre recursive algorithm, we use the Vinci code [41], in which an improved version of this algorithm is implemented, the r-Las algorithm [42].

For the comparison, we takes a unit cube of dimension (1×1×1) with 10,000 random distributed nodes inside. The shape functions are evaluated at the barycentre of each Delaunay tetrahedron. Only the points for which the five methods are applicable are taken into account.

Given times (in table 1) are average times corresponding to 1000 evaluations. The total computing time includes:
- $T_{cav}$: time taken for the cavity construction (identical for the five algorithms).
- $T_{sf}$: needed time for shape functions computation once the cavity is built.

The Watson algorithm is clearly the more powerful. For points where this algorithm doesn’t work (points located on or close to a Delaunay face), the topological algorithm based on the CDT is recommended. In practice the rate of failure of the Watson algorithm for random evaluation points is about $\frac{1}{1000}$.

For time measurements the characteristics of the used computer are:
4. Numerical integration of CNN shape functions for the CNEM

Numerical integration over Ω are needed in order to solve mechanical problems. In the NEM, and thus by extension in the CNEM, two numerical integrations are generally used: the Gauss integration, performed on the Delaunay tetrahedrons, and the stabilized conforming nodal integration (SCNI). As CNN shape functions are rational, the first one requires a high number of Gauss points in order to minimize the integration error. To improve this direct nodal integration, Chen et al. [43][44] proposed the SCNI approach for the RK interpolation in the RKPM method [45]. The SCNI approach was taken again by González et al. [46] for the NEM method.

In the SCNI approach a strain smoothing stabilization is performed to stabilize the nodal integration. The SCNI is based on the assumed strain method in which an average gradient (called stabilized gradient) is introduced for each node \( n_i \):

\[
\tilde{\nabla} A_i = \frac{1}{\| \Omega_i \|_L} \int_{\Omega_i} \nabla A \, d\Omega_i,
\]

where \( A \) is field interpolated over \( \Omega \) and \( \Omega_i \) a sub-domain surrounding \( n_i \).

Using the divergence theorem, (10) can be rewrite as:

\[
\tilde{\nabla} A_i = \frac{1}{\| \Omega_i \|_L} \int_{\Gamma_i} A \, \tilde{n} \, d\Gamma_i,
\]

where \( \Gamma_i \) is the border of \( \Omega_i \) and \( \tilde{n} \) the outward normal on \( \Gamma_i \). This allows to pass from a volume integral to a surface integral, and to integrate directly the field rather than its gradient.

<table>
<thead>
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<th>Algorithm</th>
<th>Lasserre</th>
<th>Vol-comp</th>
<th>Topo-DVC</th>
<th>Topo-TDC</th>
<th>Watson</th>
</tr>
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<td>( T_{cav} (m.s) )</td>
<td></td>
<td></td>
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<tr>
<td>( T_{sf} (m.s) )</td>
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<td>544</td>
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<td>51</td>
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<tr>
<td>( T_{sf} , \text{Alg} )</td>
<td>82</td>
<td>11</td>
<td>8</td>
<td>4</td>
<td>1</td>
</tr>
</tbody>
</table>

Table 1: Comparison of the computation efficiency of \( \| P^{\text{int}} \|_L \)

- OS: Microsoft Windows XP,
- Compiler: Microsoft (R) Incremental Linker Version 7.10.3077,
- Processor: Mobile Intel Pentium M 745 - 1800 MHz,
- RAM: PC2700 DDR SDRAM.
In a large transformations context, SCNI avoid fields projection between two configurations during a reactualization, all variables \( (\text{displacement, strain, stress}) \) being associated with the nodes (colocation approach).

For the integration of the weakforms it is necessary that the set of \( \Omega_i \) forms a partition of \( \Omega \). In the case of convex, or slightly non-convex domain, the use of the Voronoï cells is a “natural” partition of the domain as they remain convex at interior domain (Figure 14). It is nevertheless necessary to truncate the cells which overflow domain border. This is the case of cells whose nodes are on the border and for some nodes close to the border. A fast and robust algorithm is presented in [37] to do this operation.

For strongly non-convex domains, the partition based on nodal Voronoï cells is no longer valid. For such domains, cells associated with border nodes, or close to it, can be non-convex inside the domain.

For the non-convex domain depicted in Figure 15 the phenomenon occurs at node \( n \). The cell associated with \( n \) (visible in Figure 16) is non-convex inside \( \Omega \) (the surrounded point with a circle belongs to the auto-intersection curve of the cell associated with node \( n \)). This problem can be avoided by using another nodal partition. The most simplest consists in subdividing each Delaunay tetrahedron in four parts. The set of all the parts of tetrahedron attached to a node \( n_i \) defines the sub-domain \( \tilde{\Omega}_i \) associated with \( n_i \) (Figure 17). The set of \( \tilde{\Omega}_i \) is a partition of \( \Omega \).
Figure 15: Example of non-convex domain
Figure 16: Example of a Voronoi cell having a non-convexity inside the domain

Figure 17: Domain partition for non-convex domains
5. 3d elastic Benchmarks

In order to validate the CNEM in 3d, we carried out simulations for two standard problems in isotropic elasticity (small displacement and small strain): the hollow sphere under pressure and the cylindrical excavation in an infinite medium subjected to constraints ad infinitum. An analytical solution exists for both of them.

We do not give here a description of the implementation of the CNEM in elasticity but the reader can refer to Yvonnet et al. [13].

When time calculation are examined, we use the sparse iterative solver MKL [47] for the resolution of the generated linear system. The characteristics of the used computer are the following:
- Operating System: Linux RedHat 64 bits,
- Compiler: Intel c++ compiler 9.1,
- Processor: Intel Itanium 2 Monticito - 1,6GHz - 1 core used,
- RAM: 32G - DDR2 - 667MHz.

5.1. Hollow sphere under internal pressure

The exact solution of the problem (stresses) in spherical coordinates \((r, \theta, \phi)\), is given by:

\[
\sigma_{rr} = -\frac{R_{int}^3}{R_{ext}^3 - R_{int}^3} \left( \frac{R_{ext}^3}{r^3} - 1 \right) P
\]

\[
\sigma_{\theta\theta} = \sigma_{\phi\phi} = -\frac{R_{int}^3}{R_{ext}^3 - R_{int}^3} \left( \frac{R_{ext}^3}{2r^3} + 1 \right) P
\]

\[
\sigma_{r\theta} = \sigma_{r\phi} = \sigma_{\theta\phi} = 0
\]

where \(P\) is the pressure inside the sphere, \(R_{int}\) is the internal sphere radius and \(R_{ext}\) is the external sphere radius.

For simulations, the pressure inside the sphere is taken equal to 1 MPa and due to the symmetry of the problem only the eighth of the sphere is considered (Figure 18). Figure 19 shows the distribution of Von Mises equivalent stress (CNEM simulation with Sibson SCNI integration for 300 000 nodes). In this Figure, the equivalent stress field is not smoothed. The stress is constant on each node cell.
Figure 18: Hollow sphere under internal pressure – Boundary conditions

Figure 19: Hollow sphere – Von Mises equivalent stress (No. nodes $\approx 300,000$, SCNI - Sibson shape function)
5.2. Cylindrical excavation in an infinite medium

This is a plane strain problem with a bi-axial load but we solve it in 3d. The exact solution of this problem is the following [48] (cylindrical coordinates \((r, \theta, z)\)):

\[
\sigma_{rr} = \frac{\sigma_{1\infty} + \sigma_{2\infty}}{2} \left(1 - \left(\frac{a}{r}\right)^2\right) + \frac{\sigma_{1\infty} - \sigma_{2\infty}}{2} \left(1 - 4\left(\frac{a}{r}\right)^2 + 3\left(\frac{a}{r}\right)^4\right) \quad (15)
\]

\[
\sigma_{\theta\theta} = \frac{\sigma_{1\infty} + \sigma_{2\infty}}{2} \left(1 + \left(\frac{a}{r}\right)^2\right) - \frac{\sigma_{1\infty} - \sigma_{2\infty}}{2} \left(1 + 3\left(\frac{a}{r}\right)^4\right) \cos(2\theta) \quad (16)
\]

\[
\sigma_{r\theta} = -\frac{\sigma_{1\infty} - \sigma_{2\infty}}{2} \left(1 + 2\left(\frac{a}{r}\right)^2 - 3\left(\frac{a}{r}\right)^4\right) \sin(2\theta) \quad (17)
\]

\[
\sigma_{zz} = \nu (\sigma_{rr} + \sigma_{\theta\theta}) \quad (18)
\]

\[
\sigma_{rz} = \sigma_{\theta z} = 0 \quad (19)
\]

For the stresses ad infinitum we take \(\sigma_{1\infty} = 0\) and \(\sigma_{2\infty} = 1\) MPa. Only a quarter of the domain is studied (the problem has two symmetry plans). The boundary conditions are illustrated in Figure 20 and the calculated Von Mises equivalent stress field is depicted in Figure 21.

![Figure 20: Cylindrical excavation – Boundary conditions](image-url)
Figure 21: Cylindrical excavation – Von Mises equivalent stress (No. nodes ≈ 370,000, SCNI, Sibson shape function)

5.3. Convergence curves

For each of the two problems, and for a given homogeneous nodal distribution, following simulations were carried out:
- degree 0 Gauss integration with linear shape function on Delaunay mesh (Gauss FEM-L M-D),
- degree 0 Gauss integration with linear shape function on an optimized mesh (Gauss FEM-L M-O),
- degree 4 Gauss integration with Sibson shape function (Gauss Sibson),
- SCNI with linear shape function on Delaunay mesh (SCNI FEM-L M-D),
- SCNI with linear shape function on an optimized mesh (SCNI FEM-L M-O),
- SCNI with Laplace shape function (SCNI Laplace),
- SCNI with Sibson shape function (SCNI Sibson).

For SCNI, calculations are made on nodal volumes resulting from the quarter tetrahedron discretization and by taking one Gauss point integration per face (quadrangular).

Figure 22 and Figure 23 show the convergence curves of the error between computed and analytic solution for hollow sphere and cylindrical excavation respectively. The error is based on strain energy:

\[
\|e\| = \sqrt{\frac{\int_\Omega (\sigma - \hat{\sigma}) : C^{-1} : (\sigma - \hat{\sigma}) d\Omega}{\int_\Omega \hat{\sigma} : C^{-1} : \hat{\sigma} d\Omega}},
\]

(20)
where $C$ is the elasticity tensor, $\sigma$ is the computed stress tensor, and $\hat{\sigma}$ is the exact stress tensor.

For the two studied examples, although having the same rate of convergence than the linear finite elements, the CNEM divides the total error by 3. This profit of precision is mainly due to the use of the stabilized gradient. In fact, the stabilized gradient gives comparable results on linear interpolation and on CNN interpolation. We can also see that the rate of convergence obtained with the CNEM, with a Gauss integration over the Delaunay tetrahedrons, drop off when the number of nodes increase. This is due to the under integration of the Sibson shape function gradients.

Figure 24 gives, for the hollow sphere, and for the largest number of nodes ($\approx 300,000$), the computation time $t_p$ for stiffness matrix preparation [integration + assembly + boundary conditions imposition] and the computation time $t_r$ for the resolution of the generated linear system. The relative error $\|e\|$ is also present in the same figure. It is interesting to note that $t_p$ is approximatively five time greater for Sibson approximation than for finite element approximation. This is due to the calculation of shape functions, and more precisely due to the calculation of stabilized gradients.

The bandwidth of the stiffness matrix is larger for sibson function than for linear interpolation on tetraedron. This is due to the larger connectivity
of nodes with the CNEM. It is then not surprising that the resolution for CNEM takes larger time of resolution than FEM.

5.4. Study of the integration efficiency

As we saw in the section 4, the use of Voronoï cells based discretization is not always possible. We thus proposed to use the quarters of the tetrahedrons. In order to see whether the latter does not induce additional errors, the hollow sphere under pressure problem was treated with the two discretizations. The domain being slightly non-convex, the Voronoï cells based discretization is possible. Knowing that the Sibson shapes functions are rational, the number of integration point taken for the stabilized gradient computation can influence the error. In order to make sure that the error of integration is converged for each discretization, we have taken various numbers of integration points.

Following simulations were carried out:

a) For quarters of tetrahedron discretization based:
   - 1 integration point for 3 faces,
   - 1 integration point per faces (barycenter),
   - 1 Gauss integration point per triangle,
   - 13 Gauss integration points per triangle (integrand of order 7).
b) For Voronoï cells discretization based:
-1 integration point per faces (barycenter),
-1 Gauss integration point per triangle,
-13 Gauss integration points per triangle (26 per quadrangle).

The previous triangles are obtained by meshing the faces of elementary volumes attached to each node. These faces are quadrangles if we takes the quarters of the tetrahedron discretization based, and polygons if we takes the Voronoï cells discretization based. The study was done for Sibson shape function.

The obtained results are summarized in Figure 25. We can see that the quarters of tetrahedron discretization based does not induce more error than the Voronoï cells discretization based. Moreover, the error of integration with only one integration point per face is sufficient (identic error with that obtained with 26 points per face). Hence, we have chosen this integration for all CNEM simulations.
6. CNEM in finite transformations

In this section, we interest us in the CNEM in the context of large strains nonlinear elastoplastic, including dynamics, thermomechanical coupling, as well as localization in the shearing process. For this purpose, the constitutive equations of the model as well as algorithmic procedures are briefly described. A larger description can be find in [49].

6.1. Elastoplasticity formulation

For the integration of the constitutive equation we followed the approach proposed by [50]. The basic hypothesis underlying this approach to finite strains elastoplasticity is the multiplicative split of the deformation gradient, $F$, into elastic and plastic parts:

$$F = F^e \cdot F^p$$

(21)

This assumption, firstly proposed by Lee [51], admits the existence of a local unstressed intermediate configuration. The elastic part of the behavior
is based on a hyperelastic formulation which uses the Eulerian (or spatial) logarithmic strain tensor $E^e = \ln(U^e)$. $U^e$ comes from the polar decomposition of the elastic part of the deformation gradient: $F^e = R^e \cdot U^e$.

Within the incremental numerical procedure for solving the history dependent problems, a numerical scheme to verify the material constitutive equation is needed to update stresses as well as the internal variables. We use J2 plasticity with a predicting correcting approach which has the same format that the standard return mapping used in the infinitesimal theory [52, 53, 54, 55].

6.2. Kinematic aspects

Large deformations generate large relative movements between nodes in particular when shearing occurs. To handle the modification of neighborhoods of such nodes we use an updated Lagrangian procedure.

After an updating at time $t^*$, the interpolation of the total displacement of a particle $m$ at $t$, denoted $\tilde{u}(m)_{0\rightarrow t}$, is defined by:

$$\tilde{u}(m)_{0\rightarrow t} = \tilde{u}(m)_{0\rightarrow t^*} + \tilde{u}(m)_{t^*\rightarrow t}$$ (22)

$$\tilde{u}(m)_{t^*\rightarrow t} = \sum_{i=1}^{n^*} \phi^*_i(x^*(m)) \cdot u(n_i)_{t^*\rightarrow t}$$ (23)

where $x^*(m)$ are the coordinates of $m$ at $t^*$, $n^*$ is the number of natural neighbors of $m$ at $t^*$, $\phi^*_i(x)$ are the shape functions defined on the node repartition at $t^*$ and $u(n_i)_{t^*\rightarrow t}$ are the displacements of the natural neighbor $n_i$ of $m$ at $t^*$.

The gradient of the transformation from 0 to $t$ can then be write:

$$F_{0\rightarrow t} = F_{t^*\rightarrow t} \cdot F_{0\rightarrow t^*}$$ (24)

6.3. Treatment of the additional points

As specified in section 3.2, the construction (or reconstruction when updating the configuration) of the CVD can require the addition of new points. These new points are always added on the border of the domain, and thus belong to one of the triangles describing this border.

We suggest handling these new points as “slaves” of the three nodes of the triangular facet on which they are situated. Their movement is thus defined as a linear combination of the displacements of these three “master” nodes.
\[
\mathbf{u}_{\text{slave}} = \sum_{i=1}^{3} \lambda_i \mathbf{u}_i
\]  

(25)

where \( \mathbf{u}_i \) is the displacement of each “master” node, and \( \lambda_i \) the associated barycentric coefficient location (\( \sum_{i=1}^{3} \lambda_i = 1 \)).

This approach presents several advantages: there is no additional unknown and the value of the time step (in the chosen conditionally stable explicit time integration scheme) is not affected. The volume of the “slave” cells is distributed, in proportion to \( \lambda_i \), on its “masters” nodes.

6.4. Explicit thermomechanical Lagrangian procedure

The principle of virtual work may be formulated as follows:

\[
\int_{\Omega_t} \rho(t) \ddot{\mathbf{u}} \ddot{\mathbf{d}}\Omega^t + \int_{\Omega_t} \mathbf{b} \cdot \mathbf{d} \ddot{\mathbf{d}}\Omega^t + \int_{\partial \Omega_t^e} \mathbf{t} \cdot \mathbf{d} \ddot{\Gamma}^t \forall \mathbf{\eta} \in \psi
\]  

(26)

where \( \rho \) is the density, \( \mathbf{b} \) and \( \mathbf{t} \) represent the body forces and applied tractions respectively, \( \mathbf{\sigma}^t \) is the Cauchy stress tensor at time \( t \), \( \mathbf{\eta} \) is the virtual displacement, and \( \psi \) is the space of virtual displacements. The introduction of the CNEM discretization in the variational form (26), transported in the last update configuration, results in the discrete set of algebraic time dependent equations which may be expressed, in matrix form, as:

\[
\mathbf{M} \ddot{\mathbf{u}}(t) = \mathbf{F}^{\text{ext}}(t) - \mathbf{F}^{\text{int}}(\mathbf{u}_n, t)
\]  

(27)

where \( t \) is the time, \( \mathbf{M} \) denotes the mass matrix, \( \mathbf{F}^{\text{int}}(\mathbf{u}, t) \) the internal forces vector, while \( \mathbf{F}^{\text{ext}}(t) \) is the external forces vector. The use of the SCNI quadrature results in a diagonal matrix for \( \mathbf{M} \). The diagonal terms are given by \( m_i = \rho_0 \Omega_i \), with \( \Omega_i \) the area (volume in 3D) of the Voronoï cell related to node \( n_i \).

To solve (27) the classical explicit central differences incremental scheme is used. In case of contact a predictor/corrector strategy is used according to the prescribed displacements or tractions.

The weak form of the heat balance can be expressed as:

\[
\int_{\Omega^t} \rho(t)c(t) \dot{T} \varphi d\Omega^t + \int_{\Omega^t} k(t) \nabla T \cdot \nabla \varphi d\Omega^t = \int_{\Omega^t} r^t \varphi d\Omega^t + \int_{\partial \Omega^t} q^t \varphi d\Gamma^t \forall \varphi \in \psi
\]  

(28)
where \( c(t) \) is the specific heat, \( k(t) \) is the thermal conductivity for isotropic conduction, and \( r^t \) a heat source related to the inelastic deformations, defined at time \( t \). We have made the assumption that heat generated by elastic deformation is negligible with respect to heat generated by plastic deformation.

In the following, we assume \( c \) and \( k \) constant in time. \( \varphi \) is the virtual temperatures, \( \psi \) is the space of virtual temperatures, and \( \bar{q} \) represents the heat transfer at the boundary \( \partial_2 \Omega \). Following similar arguments as in the mechanical problem (26), the weak form of the heat balance results, with the CNEM discretization, in the ODE system:

\[
CT(t) + KT(t) = Q(t)
\]

which in the context of an explicit scheme can be written as

\[
CT^{n+1} = [C + \Delta t K] T^n + \Delta t Q^n
\]

The matrix \( C \) becomes diagonal in the context of CNEM when SCNI is used. The heat source resulting from the inelastic deformations is given by

\[
r^t = \chi \sigma^t : \dot{D}^p_t, \quad \dot{D}^p_t = \text{sym} \left( \left( \dot{F}^p_t \right) (F^p_t)^{-1} \right)
\]

where \( \chi \) is the Taylor-Quinney parameter [56] representing the fraction of plastic work converted into heat. In the present work, we have used \( \chi = 0.9 \) [57].

When mechanical and thermal problems are coupled, a staggered solution approach is usually adopted, which solves the mechanical and the thermal problems in an uncoupled manner with data exchange performed at the end of each time step or increment. In particular, the nodal temperatures are transferred to the mechanical procedure, while plastic work is communicated to the thermal solver.

6.5. Constitutive law

For the examples below (section 7), we use the Johnson-Cook hardening law (31)[58]. This law allows, in a simple way, to take into account a thermomechanical coupling: effects of inelastic deformations and softening effects due to temperature.

\[
\sigma_y(\bar{\varepsilon}^p) = [A + B(\bar{\varepsilon}^p)^n] \left[ 1 + Cln \left( \frac{\bar{\varepsilon}^p}{\bar{\varepsilon}_0} \right) \right] \left[ 1 - \left( \frac{T - T_0}{T_f - T_0} \right)^m \right]
\]

33
$A$, $B$, $C$, $m$, $n$ and $\dot{\varepsilon}_0^p$ are material parameters, $\varepsilon^p$ represents the equivalent plastic strain and $\dot{\varepsilon}^p$ represents the rate of equivalent plastic strain. $T$ and $T_0$ correspond to the current and initial temperature respectively.

7. Simulation of manufacturing processes

The objective of this section is to show the ability of the CNEM to simulate problems where very large strains (around 10) are present. We chose an example of high speed blanking and two examples of machining (orthogonal cutting and oblique cutting). For these examples we give theses following data:

- Nodes: the number of nodes at the beginning / end of the simulation,
- $T_{tot}$: total duration of the simulation,
- $N_I$: number of corresponding increments,
- $N_U$: number of updates of the updated Lagrangian procedure,
- $T_U$: average time for one update,
- $R_U$: percentage of total time taken for all the updates versus $T_{tot}$.

For simplicity reason we took the same material for all these examples (table 2). All the simulations were launched on the same computer:

- OS: Linux RedHat 64 bits
- Compiler: Intel c++ compiler 9.1
- Processor: 4 x Intel Itanium 2 Monticito - 1,6GHz - 8 cores used
- RAM: 32G - DDR2 - 667MHz

SCNI and constitutive law integration are parallelized by using TBB (Threading Building Blocks) task primitive.

<table>
<thead>
<tr>
<th>$A$(MPa)</th>
<th>$B$(MPa)</th>
<th>$C$</th>
<th>$n$</th>
<th>$m$</th>
<th>$\dot{\varepsilon}_0^p$(s$^{-1}$)</th>
<th>$T_i$(C)</th>
<th>$T_f$(C)</th>
</tr>
</thead>
<tbody>
<tr>
<td>866</td>
<td>318</td>
<td>0.008</td>
<td>0.25</td>
<td>1.055</td>
<td>5.77 x 10$^{-2}$</td>
<td>20</td>
<td>1670</td>
</tr>
</tbody>
</table>

Table 2: TA6V – Johnson-Cook coefficients (from [59])

Figures 27, 29 and 30 are structured in the same way for the three examples. On the top is represented the equivalent plastic strain (for three different instants), and on the bottom the equivalent Von Mises stress (for the same instants).
7.1. High speed blanking

Figure 26 gives the diagram of the studied example: the punch is blue, the sample is grey/red (due to symmetry reason only a quarter is modeled), and the die is yellow. The table 3 gives the corresponding data. \(v\) is the relative speed between the punch and the die.

\[
\begin{array}{cccccc}
v & e_e & l_e & h_e & r_p/r_m & l_p & j \\
10 \text{ m/s} & 2 \text{ mm} & 17 \text{ mm} & 32 \text{ mm} & 0.5 \text{ mm} & 16 \text{ mm} & 0.1 \text{ mm}
\end{array}
\]

Table 3: High speed blanking – Geometric and process data

7.2. Orthogonal and oblique cutting

Figure 28 gives the diagram of the studied cutting examples. \(\gamma\) is equal to 0 for orthogonal cutting, otherwise we are in oblique cutting. Table 4 gives the corresponding data. \(v\) is the relative speed between the tool and the workpiece.
Figure 27: High speed blanking – Equivalent plastic strain and stress contours

Figure 28: Machining – Diagram of the apparatus
Figure 29: Orthogonal cutting – Equivalent plastic strain and stress contours

Figure 30: Oblique cutting – Equivalent plastic strain and stress contours
7.3. Results

The computational data are given in table 5. One can see that the needed time for all the updates of configuration is non-negligible with respect to the total computational time. In general the ratio is near 30%. It is interesting to see that the number of updates remains relatively low with regard to the number of increments: \( \frac{N_I}{N_U} \) is always greater than 300.

<p>| | | | | | | |</p>
<table>
<thead>
<tr>
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</tr>
</thead>
<tbody>
<tr>
<td>Orthogonal cutting</td>
<td>20 m/s</td>
<td>0.2 mm</td>
<td>0.5 mm</td>
<td>0.05 mm</td>
<td>5°</td>
<td>30°</td>
</tr>
<tr>
<td>Oblique cutting</td>
<td>20 m/s</td>
<td>0.2 mm</td>
<td>0.5 mm</td>
<td>0.05 mm</td>
<td>5°</td>
<td>30°</td>
</tr>
</tbody>
</table>

Table 4: Machining – Geometric and process data

<p>| | | | | | | |</p>
<table>
<thead>
<tr>
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<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Nodes</td>
<td>( T_{tot} )</td>
<td>( N_I )</td>
<td>( N_U )</td>
<td>( T_U )</td>
<td>( R_U )</td>
<td></td>
</tr>
<tr>
<td>Blanking</td>
<td>73385/147974</td>
<td>147 h</td>
<td>305944</td>
<td>153</td>
<td>6.82 min</td>
<td>12%</td>
</tr>
<tr>
<td>Oth. cutting</td>
<td>14386/78092</td>
<td>32.2 h</td>
<td>148651</td>
<td>449</td>
<td>2.3 min</td>
<td>54%</td>
</tr>
<tr>
<td>Obl. cutting</td>
<td>14386/20699</td>
<td>10.55 h</td>
<td>149093</td>
<td>254</td>
<td>0.75 min</td>
<td>30%</td>
</tr>
</tbody>
</table>

Table 5: Computational data

8. Conclusion

We can say that up to now the main difficulties inherent to the implementation of the CNEM 3d are overcome: construction of the constrained Voronoï diagram in 3d, optimization of the calculation of Sibson shape functions, domain discretization in nodal elementary volumes for the stabilized nodal integration. The elastostatic tests are successful. Although having the same rate of convergence of linear finite elements, the CNEM divides the global error by 3. In finite transformations, for thermo-elasto-viscoplastic behavior, the proposed approach also gives very promising results.

However, the CNEM requires a correct description of the border (no autointersection) for the construction of the constrained Voronoï diagram. This involves a possible update of this last one: a remeshing of the border in the area where there are surface folds, excessive elongations or where nodes become to close. For high speed processes (cutting or blanking for example) it is still necessary, to realize more representative simulations, to take into account dynamic effects of the tools and the machine (die, punch, ...). To simulate a process like blanking, it is also necessary to be able to take into
account matter separation. A simple manner to introduce such a separation may consist in using a “kill cell” approach.

If numerical simulation is a key issue for the development of efficient cutting or forming processes, the definition, and the identification, of constitutive laws adapted to high speeds and high temperatures remain still a major difficulty.

9. Acknowledgement

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Appendix A. Study the CNN shape functions near highly non-convex borders

For non-convex domains, there exist locations for which the insertion of a point in the CVD creates a non-convex Voronoï cell. In order to illustrate that, let take the domain $\Omega$ depicted in Figure A.31. The insertion of the point $x$ (coordinates $[-0.1, 0, 0]$), in the CVD of $\Omega$, leads to a non-convex cell (see Figure A.32). We can also see in this Figure, that the cell associated with the neighbor node $v$ ($n_0$ or $n_1$) is also non-convex. When $x$ moves the topology of its associated cell evolves continuously from a non-convex topology to a convex topology. In order to highlights this phenomenon, the point $x$ moves gradually, in Figure A.33, from the position $[-0.25, 0, 0]$ to $[0, 0, 0]$ (displacement along the $X$ axis).

Having noticed this non-convex property, it is interesting to study the evolution of the shape functions according to different positions of a point $x$. The graph A.34 gives, according to the point $x$ coordinate along the $X$ axis, the volume measures of:
- the point $x$ cell, $(\text{Vol}(c_x))$,
- the polyhedron intersections associated with each neighbor of $x$.

The graph A.35 gives, again according to the the point $x$ coordinate along the $X$ axis, the value of the shape functions associated with each neighbor of $x$. All shape function properties stated in 3.4 remain verified. We can however notice that there is a position $x_0$ for which the volume of the cell of the point $x$ cancels. This gives place to an indetermination of the Sibson shape function (division by 0). Such position is not unique. Indeed, if we
calculate the volume measure of the point $x$ cell in a vicinity of $x_0$, the location where the volume measure cancel forms a surface (Figure A.36).

It should be noted that even for Laplace CNN shape functions, places for which they are unspecified also exist, as the graph in the Figure A.37 shows it.

Figure A.31: Non-convex domain – Location of the point $x$
Figure A.32: Non-convex domain – Cell $c'_x$ – Intersection with the cell $c_v$ of a neighbor node
Figure A.33: From a convex cell to a non-convex cell according to the position of \( x \).

Figure A.34: Volume measurement – Cell of node \( x \) – Intersection with neighboring cells
Figure A.35: Sibson shape functions at $x$ for different neighbors

Figure A.36: Space location where the volume of $x$ cell is null
Figure A.37: Laplace shape functions at $x$ for different neighbors
References


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