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Domain decomposition with discrete element simulations using shared-memory parallel computing for railways applications

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ABSTRACT. Numerical simulation with discrete elements leads to several issues for large scale problems and long loading times, as for the granular dynamic simulations of the ballasted railway behavior. To reduce computational costs, we study the use of two strategies: domain decomposition methods and shared-memory parallelization with OpenMP. An example of a maintenance process, the tamping, on a portion of railway track with 7 sleepers, is simulated.

RÉSUMÉ. La simulation numérique par éléments discrets présente des difficultés pour l’étude de problèmes de grande taille et en temps de sollicitation long, comme la dynamique des milieux granulaires pour le ballast ferroviaire. Afin de résoudre ce problème à moindre coût, on propose d’allier deux stratégies : la décomposition de domaine (DDM) et le calcul parallèle (en mémoire partagée avec OpenMP). Un exemple traitant d’un procédé de maintenance ferroviaire, le bourrage, sur une portion de voie ballastée de 7 blocs de long est étudié.

KEYWORDS: Non Smooth Contact Dynamics, parallelization, LMGC90, ballast, maintenance.
MOTS-CLÉS: Non Smooth Contact Dynamics, parallélisation, LMGC90, ballast, maintenance.
1. Motivations: tamping process optimization

The degradation of ballasted railway under commercial exploitation requires frequent and costly maintenance operations. One of these operations is the tamping process (Paderno, 2010; Azéma, 2007; Azéma et al., 2009; Solomon, 2001) which is intended to correct the railway geometry that have been modified by an inhomogeneous vertical settlement. It is performed by a ballast tamper or a tamping machine which works by vibrating the ballast and pushing it under the sleepers. The study of the mechanical behavior of the ballast during tamping is necessary to propose optimizations of this maintenance process (Perales et al., 2009).

The numerical simulation of a portion of a railway submitted to the tamping process is useful to master the physical phenomenons involved in the ballast (Saussine, 2004). However, the numerical complexity of a simulation at the grain scale requires special treatments to get an affordable simulation.

Within this article, the Non Smooth Contact Dynamics (NSCD) model is used for the simulation of the ballasted railway, with a discrete element model (DEM). A domain decomposition method (DDM) coupled with a shared-memory parallelization technique (using OpenMP) are used to improve the computational efficiency and to reduce the cost and the run time of the simulation. The domain is geometrically split into several subdomains, allowing their simultaneous treatment on several processors or computer cores.

The main results are related to a representative simulation of a portion of a railway track with 7 sleepers, submitted to successive tamping processes. The structure is composed of up to 90,000 polyhedral grains, and at each time step, about 300,000 frictional contacts are involved. The post-treatment concerns the influence of the process under each sleeper, in terms of compactness, and several indicators are used to assess the quality of the numerical simulation, and of the physical quality of the obtained railway.

2. Numerical method

2.1. Granular dynamics

The Non Smooth Contact Dynamics (NSCD) approach developed by Moreau and Jean (Moreau, 1999; Jean, 1999; Cambou et al., 2001) is a discrete element method that may be applied to a large range of discrete systems involving various interactions between elements and various regimes (quasi-static, slow and fast dynamics, dense and diluted granulates, rigid and deformable bodies). In the present framework the NCSD is applied to a collection of rigid polyhedral grains with frictional contact interactions. The main features of the approach are the reduced dynamics and the nonsmooth formulation of contact laws.
2.1.1. Dynamic equation

For each grain, according to the NSCD approach, the discrete dynamic equation over a time step \([t_i, t_{i+1}]\) may be written,

\[
M(V - V^i) = R^d + R
\]

where \(M\) denotes the inertia matrix, \(V^i\) and \(V\) are the generalized velocities of the grains at the instants \(t_i\) and \(t_{i+1}\) (the subscript \(i + 1\) is omitted for the following). \(R^d\) represents the external impulsions and \(R\) are the impulsions due to all contacts on the grain over the time step. All the equations for all the grains may be concatenated formally leading to the same expression. \(V\) and \(R\) have to be determined.

The previous equation \([1]\) may be reduced to the local variables of the contacts, the relative velocity \(v\) and the contact impulsion \(r\). The local variables are defined according to a local-to-global (or contact-to-grain) mapping \(H\) and its transpose \(H^T\):

\[
\begin{align*}
  v &= H^T V \\
  r &= H r
\end{align*}
\]

The dynamics is then given by \(v = v^d + W r\), where \(v^d = H^T (M^{-1} R^d + V^i)\), and \(W = H^T M^{-1} H\) is the so-called Delassus operator.

2.1.2. Contact law

According to the viability lemma (Moreau, 1999), for each contact, the unilateral contact may be formulated with a complementarity condition between the normal relative velocity \(v^\alpha_n\) and the normal contact impulsion \(r^\alpha_n\), once the penetration tested with a predicted gap \(g^\alpha\), for the contact \(\alpha\). The frictional contact law is then,

\[
\begin{align*}
  &\text{if } g^\alpha > 0, \quad r^\alpha = 0 \\
  &\text{if } g^\alpha = 0, \quad 0 \leq v^\alpha_n \perp r^\alpha_n \geq 0 \quad \text{and} \quad \begin{cases} \text{if } \|v^\alpha_t\| = 0, \quad \|r^\alpha_t\| \leq \mu r^\alpha_n \\
\text{if } \|v^\alpha_t\| \neq 0, \quad v^\alpha_t = -\mu r^\alpha_n v^\alpha_t / \|v^\alpha_t\| \end{cases}
\end{align*}
\]

This law may be written formally : \(R(r, v) = 0\).

The reference problem consists then in determining the contact impulsion and the relative velocity of each contact satisfying the following system,

\[
\begin{align*}
  v &= v^d + W r \\
  R(r, v) &= 0
\end{align*}
\]

2.1.3. Non Linear Gauss Seidel solver (NLGS)

A Non Linear Gauss-Seidel type algorithm based on a classical block partitioning of the \(W\) matrix amounts to find for each contact \(\alpha\) and for each iteration \(k\) the pair \((v^{\alpha, k+1}_\alpha, r^{\alpha, k+1}_\alpha)\) verifying the local nonsmooth system,

\[
\begin{align*}
  &W_{\alpha\alpha} v^{\alpha, k+1}_\alpha - v^{\alpha, k+1}_\alpha = -v^{\alpha, k}_\alpha - \sum_{\beta < \alpha} W_{\alpha\beta} v^{\beta, k+1}_\beta - \sum_{\beta > \alpha} W_{\alpha\beta} r^{\beta, k}_\beta \\
  &R(v^{k+1}_\alpha, v^{\alpha, k+1}_\alpha) = 0
\end{align*}
\]
2.2. Domain decomposition and parallel technique

2.2.1. Geometrical partitioning

A dense collection of rigid bodies may be viewed as a set of nodes (grains) and a set of links (contacts) shown on Figure 1. Then two partitioning techniques may be performed (Champaney et al., 2007).

- The primal approach distributes the grains into the subdomains (Figure 2 left). A grain belongs to a subdomain if the coordinates of its mass center is inside a geometrical box previously defined by a regular grid. The interface consists then of links between grains of two neighboring subdomains (Hoang et al., 2011b; Hoang et al., 2011a).

- The dual approach distributes the links into the subdomains (Figure 2 right). A link belongs to a subdomain if the barycenter of the two contacting grains is inside the box. The interface consists then of grains with contacts belonging at least to two subdomains (Iceta, 2010; Iceta et al., 2009).

In this study the primal approach is chosen because it is less intrusive in an industrial software than the dual one. The interface is constituted with contacts only, while the subdomains contain grains and contacts. Therefore the interface is numerically dealt as one subdomain with a nonsmooth solver. This appears clearly when ones examines the algebraic partitioning deriving from the geometric substructuring.
2.2.2. Algebraic partitioning

According to the formulation developed in (Champaney et al., 2007; Nineb et al., 2007) the subscripts $E$, $E'$, $\Gamma$ denote in the following the variables of the subdomain $E$, respectively $E'$ and of the interface $\Gamma$. For sake of simplicity two subdomains are only considered. The capital letters refer to variables associated with grains whereas the lowercase letters refer to variables of the contacts. So the velocities of the grains and the impulsions applied to the grains are defined for all the subdomains as follows,

$$V = \begin{bmatrix} V_E \\ V_{E'} \end{bmatrix} \quad \text{and} \quad R = \begin{bmatrix} R_E \\ R_{E'} \end{bmatrix}$$

On the contrary the relative velocities at contacts and the contact impulsions are written for all the subdomains and for the interface,

$$v = \begin{bmatrix} v_E \\ v_{E'} \\ v_{\Gamma} \end{bmatrix} \quad \text{and} \quad r = \begin{bmatrix} r_E \\ r_{E'} \\ r_{\Gamma} \end{bmatrix}$$

The dynamic equation of the grains belonging to the subdomain $E$ is given as,

$$M_E V_E = M_E V^i_E + R^d_E + R_E + R_{ET}$$

where $R_{ET}$ are the impulsions of the interface $\Gamma$ applied to the subdomain $E$,

$$R_E = H_E r_E \quad \text{and} \quad R_{ET} = H_{ET} r_{\Gamma}$$

The dual relations provide the relative velocities starting from the generalized velocities of the grains,

$$v_E = H^T_E V_E \quad \text{and} \quad v_{\Gamma} = \sum_E H^T_{ET} V_E$$

The reduction of the dynamics to the contacts leads to a partitionned system,

$$\begin{bmatrix} v^d_E \\ v^d_{E'} \\ v^d_{\Gamma} \end{bmatrix} = \begin{bmatrix} 0 & W_{ET} & 0 \\ 0 & 0 & W_{ET'} \\ W_{TE} & W_{TE'} & W_{\Gamma} \end{bmatrix} \begin{bmatrix} r_E \\ r_{E'} \\ r_{\Gamma} \end{bmatrix}$$

with $v^d_E = H^T_E (V_E^i + M^{-1}_E R^d_E)$, $v^d_{E'} = \sum_E H^T_{ET} (V_E^i + M^{-1}_E R^d_E)$, $W_E = H^T_E M^{-1}_E H_E$, $W_{ET} = W^T_{TE} = H^T_E M^{-1}_E H_{ET}$, and $W_{\Gamma} = \sum_E H^T_{ET} M^{-1}_E H_{ET}$.

The parallel process is performed with a shared memory architecture and multi-threading OpenMP directives inserted in the program. Such a strategy is the simplest one to implement in a pre-existing industrial software. The generic solver for the Non-Smooth Contact Dynamics is a NonLinear Gauss Seidel algorithm. In a first version it is applied simultaneously to all the $n_{SD}$ subdomains before solving the global interface. This specific treatment of the coupling interface provides then a synchronous algorithm.
The final algorithm described in Algorithm 1 associates a NonLinear Gauss Seidel method, a Domain Decomposition method and Open-MP directives (DDM–OpenMP–NLGS). The algorithmic parameters are: the number of subdomains $n_{SD}$, the number of DDM iterations $n_{DDM}$, the number of NLGS iterations in the subdomains $n$, the number of NLGS iterations in the interface $m$. The emphasized parts are additional tasks to allow the DDM and parallel treatment.

**Algorithm 1 DDM–OpenMP–NLGS**

Loop on time steps

```
for $i = 1, 2 \ldots$ do
  Gap prediction and contact detection
  Domain partitioning
  Parallel loop on subdomains
  for $E = 1, 2 \ldots n_{SD}$ do
    Compute ‘free’ velocity at each grain of the subdomain $E$
  end for
end for
```

Loop on DDM iterations

```
for $j = 1, 2 \ldots n_{DDM}$ do
  Parallel loop on subdomains
  for $E = 1, 2 \ldots n_{SD}$ do
    NLGS solve for each subdomain $E$, with $n$ iterations
    Compute numerical residual for each subdomain $E$ if required
  end for
  NLGS solve for ‘subdomain’ $\Gamma$, with $m$ iterations
  Compute numerical residual for ‘subdomain’ $\Gamma$ if required
  Convergence test
end for
```

Parallel loop on subdomains

```
for $E = 1, 2 \ldots n_{SD}$ do
  Compute nodal quantities (update velocities, positions)
end for
```

end for

3. Numerical results

The targeted application concerns the behavior of a large slice of the ballasted railway, submitted to the tamping process. The numerical simulation of such a problem requires an unaffordable computational cost. The proposed test in this section relies on the previous solving strategy to reduce the simulation cost, to be able to access extensive parametrical studies of the tamping process. A 3D specimen of size $3.6 \times 2 \times 0.56$ $m$ is modeled as a slice of a track with 7 sleepers. It is submitted to a tamping cycle on sleeper # 4. Figure 3 depicts the geometry of this specimen.
The specimen is made of 88 100 polyhedral grains, and approximatively 310 000 frictional contacts. Friction coefficient is $\mu = 1$ between grains, between grains and sleepers, and between grains and tamping tools; it is selected to $\mu = 0.8$ between grains and planes that define boundary conditions. The numerical parameters are selected as follows:

- The time interval $[0, T]$ with $T = 1.764$ s, is discretized with 8 820 time steps,
- $n_{SD} = 7$ subdomains correspond to the areas under each sleeper,
- $n_{DDM} = 740$,
- $n = m = 1$.

The implementation is performed within the LMGC90 platform (Dubois et al., 2007), and a 8 Gb RAM 2 Dual-Core PC is used (2 processors with 2 cores each). The detailed physical response of the problem is studied with several global quantities such as the compactness in (Hoang et al., 2011b). We are more concerned herein with the numerical indicators: computing time, speedup and efficiency, interpenetrations as residuals.

To estimate the parallel efficiency of the OpenMP implementation, the speedup is defined as: $S_p = T_1/T_p$, where $T_1$ ($T_p$) denotes the run time with the parallel algorithm executed on 1 processor (respectively, $p$ processors). The efficiency is defined as $E_p = S_p/p$ and is expected to be in the interval $[0, 1]$.

Table 1 reports the restitution time of each run, as well as the time profile on different parts of the simulation. The numerical solve with NLGS represents on the average 75% of total time. The contact detection phase is approximatively 22%, and the remaining parts 3%. Up to now, only the resolution phase if parallelized. The 4 processor run is roughly twice as fast as the sequential run.

**Figure 3. Railway slice with 7 sleepers. The tamping cycle is operated on the sleeper # 4 (numbered from left to right)**
Table 1. Profiling computation time, using 1 and 4 processors

<table>
<thead>
<tr>
<th>Number of processors</th>
<th>1</th>
<th>4</th>
</tr>
</thead>
<tbody>
<tr>
<td>restitution time / min</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Total time</td>
<td>14019</td>
<td>7860</td>
</tr>
<tr>
<td>Contact detection</td>
<td>2473</td>
<td>2458</td>
</tr>
<tr>
<td>NLGS solve</td>
<td>11290</td>
<td>5086</td>
</tr>
<tr>
<td>Other parts*</td>
<td>256</td>
<td>316</td>
</tr>
</tbody>
</table>

NLGS solve details
- Treatment of 7 subdomains (in parallel) 10250 4048
- Treatment of the global interface (sequential) 1040 1038

* predictions, updates...

The speedup and efficiency plots on Figure 4 illustrates the better performances of the ‘subdomain’ part (OpenMP part) when compared to the total restitution time (respectively 2.53 and 1.8 for the speedup, 63.3% and 45% for the efficiency). The sequential treatment of the interface, as well as the other parts of the code, impairs the parallel performances. The second source for this loss of efficiency is the load unbalance of the processors (7 subdomains of equivalent sizes for 4 processors).

Figure 4. Parallel implementation performance with 4 processors and 7 subdomains

4. Asynchronous algorithm

In the previous algorithm, the sequential treatment of the global interface allows to synchronize all the subdomain information. In such a case, the numerical results are always identical for a given splitting of the domain, whatever the number of processors is. For a parallelization point of view, this synchronization increases idle time for the processors, and reduces the efficiency, especially when one wishes to use a large number of processors and subdomains.
Asynchronous algorithms would therefore lead to a higher parallel efficiency. They are usually iterative algorithms where the update of the components of the iterated vector solution is performed on-the-fly as soon as computed.

4.1. **Principles of the algorithm**

The OpenMP implementation of the asynchronous version is very close to the previous one: the only modification lies in the global interface which is dealt with as any subdomain in the DDM loop, and has no more a specific treatment. From an algebraic point of view, this is the transformation of a block-Gauss-Seidel splitting of the reference problem [2] of the form:

\[
\begin{bmatrix}
  v_E \\
v_{E'} \\
v_T
\end{bmatrix} - \begin{bmatrix}
  W_L^E & 0 & 0 \\
  0 & W_E^L & 0 \\
  W_T & W_{TE'} & W_T^L
\end{bmatrix} \begin{bmatrix}
  r_E \\
r_{E'} \\
r_T
\end{bmatrix} = \begin{bmatrix}
  v_E^d \\
v_{E'}^d \\
v_T^d
\end{bmatrix} + \begin{bmatrix}
  W_E - W_L^E & 0 & W_{ET} \\
  0 & W_{E'} - W_L^E & W_{ET} \\
  0 & 0 & W_T - W_T^L
\end{bmatrix} \begin{bmatrix}
  r_E \\
r_{E'} \\
r_T
\end{bmatrix}
\]

(where superscript \(L\) denotes the diagonal and lower part) for the synchronous algorithm, into a block-Jacobi (for the subdomains) / Gauss-Seidel (for the inner subdomain problems) splitting:

\[
\begin{bmatrix}
  v_E \\
v_{E'} \\
v_T
\end{bmatrix} - \begin{bmatrix}
  W_E^L & 0 & 0 \\
  0 & W_{E'}^L & 0 \\
  0 & 0 & W_T^L
\end{bmatrix} \begin{bmatrix}
  r_E \\
r_{E'} \\
r_T
\end{bmatrix} = \begin{bmatrix}
  v_E^d \\
v_{E'}^d \\
v_T^d
\end{bmatrix} + \begin{bmatrix}
  W_E - W_E^L & 0 & W_{ET} \\
  0 & W_{E'} - W_{E'}^L & W_{ET} \\
  W_T & W_{TE'} & W_T - W_T^L
\end{bmatrix} \begin{bmatrix}
  r_E \\
r_{E'} \\
r_T
\end{bmatrix}
\]

Following (Renouf, 2004; Chau, 2005), this type of algorithm may lead to several issues:

- Since the order of processing on each contact is continuously changing, the produced admissible solution changes for each run on the same problem. Comparison of two solutions is therefore somehow difficult. Nevertheless the macroscopic global behavior should be similar ;

- The asynchronous writing on the solution storage vector may lead to conflicts when simultaneous writings occur. In such a case, one or more computed values are discarded.
4.2. Algorithms comparison

The same ballasted track testbed is used and a solution is produced with 3 versions of the code: (i) version 1 is the standard code, without domain decomposition and without parallelization; (ii) version 2 is the synchronous algorithm parallelized with OpenMP; (iii) version 3 is the asynchronous version with the same domain decomposition, and parallelized with OpenMP. Simulations are performed on a 8 Gb Dual-Core station with 4 processors.

The corresponding test case is a smaller version of the previous one. Namely, it involves 2272 time steps for a physical duration of \( T = 1.8175 \text{ s} \), 3 subdomains and a global interface are used, the average number of contacts are 9 800 and 22 800 for the two subdomains and 3 000 for the interface. For the iteration counts, \( n_{\text{DDM}} = 500 \) DDM iterations and \( n_{\text{NLGS}} = n = m = 1 \) NLGS iteration are prescribed.

The obtained solutions have been compared with their compactness and their inertial parameter evolutions. Though slightly different, their trends are similar. The asynchronous solution does exhibit a small overshoot for the compactness, which does not exceed 1.7\% the other ones. This allows to conclude that sound solutions are obtained in each case. Indeed, the residual for the non penetration constraint is similar, and sometimes lower, for the asynchronous case when compared to the synchronous algorithm.

Simulation costs are reported in Table 2. Even on a small number of processors, the asynchronous algorithm is more efficient than the synchronous version. The overall efficiency is nevertheless still small (1.71/3) due to a relatively small problem size. This efficiency, as well as the improvement of the asynchronous version, are expected to be improved with the simultaneous increase of the number of subdomains and problem size.

Table 2. Simulation time for the 3 implementation versions

<table>
<thead>
<tr>
<th></th>
<th>Version 1</th>
<th>Version 2</th>
<th>Version 3</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number of processors</td>
<td>Reference</td>
<td>Synchronous alg.</td>
<td>Asynchronous alg.</td>
</tr>
<tr>
<td>Total time</td>
<td>923</td>
<td>621</td>
<td>607</td>
</tr>
<tr>
<td>Contact detection</td>
<td>193</td>
<td>175</td>
<td>172</td>
</tr>
<tr>
<td>NLGS solve</td>
<td>708</td>
<td>424</td>
<td>414</td>
</tr>
<tr>
<td>Other parts*</td>
<td>22</td>
<td>22</td>
<td>21</td>
</tr>
</tbody>
</table>

* predictions, updates...

5. Conclusions

The proposed approach combines a domain decomposition method with a shared-memory parallelization technique. Such a strategy allows to reduce significantly the
computational time for solving industrial problems. The use of 4 processors leads to a gain of 1659 minutes (about 4.3 days). However the parallel efficiency is weak with a speedup equal to 1.8 in comparison with the optimal speedup.

In order to optimize the parallel performance some propositions may be formulated. The balance between the processors may be easily improved by choosing a number of subdomains equal to a multiple of the number of processors. Likewise the geometrical partitioning has to be optimized to get a nearly constant number of contacts per subdomain. The size of the interface may be reduced in choosing compact subdomains. Anyway the number of subdomains is limited both by the use of a shared-memory architecture and the need to restrict the size of the interface to the size of one subdomain. Finally a substantial gain may be expected from the parallelization of the contact detection as suggested by the analysis of the computational time spent in the sequential part compared to the parallel part (Figure 5).

In spite of its simplicity, the present strategy is relevant to deal with large-scale strongly nonlinear industrial problems requiring robustness and a multiparametric study for improving maintenance processes.

![Figure 5](image)

**Figure 5.** Evolution of the elapsed time for the full computation and for the parallel part with respect the number of subdomains

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6. References


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