D-iteration: Evaluation of a Dynamic Partition Strategy

Dohy Hong
Alcatel-Lucent Bell Labs
Route de Villejust
91620 Nozay, France
dohy.hong@alcatel-lucent.com

ABSTRACT

The aim of this paper is to present a first evaluation of a dynamic partition strategy associated to the recently proposed asynchronous distributed computation scheme based on the D-iteration approach. The D-iteration is a fluid diffusion point of view based iteration method to solve numerically linear equations. Using a simple static partition strategy, it has been shown that, when the computation is distributed over K virtual machines (PIDs), the memory size to be handled by each virtual machine decreases linearly with K and the computation speed increases almost linearly with K with a slope becoming closer to one when the number N of linear equations to be solved increments. Here, we want to evaluate how further those results can be improved when a simple dynamic partition strategy is deployed and to show that the dynamic partition strategy allows one to control and equalize the computation load between PIDs without any deep analysis of the matrix or of the underlying graph structure.

Categories and Subject Descriptors
G.1.0 [Mathematics of Computing]: Numerical Analysis—Parallel algorithms; G.1.3 [Mathematics of Computing]: Numerical Analysis—Numerical Linear Algebra; C.2.4 [Computer Systems Organization]: Computer-Communication Networks—Distributed Systems

General Terms
Algorithms, Performance

Keywords
Distributed computation, Iteration, Fixed point, Eigenvector.

1. INTRODUCTION

Solving efficiently a very large linear equation systems (and the related initial problems) is a very classical problem and challenge for the algorithm design. The complexity of the problem to solve numerically a very large linear systems may increase rapidly with the dimension of the vector space. There are many known approaches to solve such a class of problems: Gauss elimination, Jacobi iteration, Gauss-Seidel iteration, SOR (successive over-relaxation), Richardson, Krylov, Gradient method, power iteration, QR algorithm etc [11, 31, 4, 10, 25, 83]. And there are more specific approaches in more particular cases when the linear equations are associated to a sparse matrix (in particular, in the context of PageRank equation [26, 9, 7, 3]: power method [22] with adaptation [20] or extrapolation [12, 21, 8], iterative aggregation/disaggregation method [27, 15, 29], adaptive on-line method [2, etc]. The case of the symmetric and diagonally dominant (SDD) systems [9, 22, 24] is also a very interesting case that was deeply investigated. In parallel, there have been a lot of researches concerning the distributed computation of the linear equations [4, 19, 23, 26, 29, 14, 28], with a particular interest on asynchronous iteration scheme.

The algorithm proposed here is a new solution for a class of problem we could call diagonally dominant (DD) systems based on the recent research results on the D-iteration. The D-iteration method was initially introduced in [17] to solve numerically the eigenvector of the PageRank equation (the eigenvector defining the score of the page importance). Its applicability in a general linear equation has been described in [16]. The distributed architecture based on this algorithm was first proposed in [15] and then evaluated through simulations in [14] when static partition strategies are applied. It has been shown in [14] that, when the computation is distributed over K virtual machines (PIDs), the memory size to be handled by each virtual machine decreases linearly with K and the computation speed increases almost linearly with K with a slope becoming closer to one when the number N of linear equations to be solved increases. However, those results were obtained under the assumption that the information diffusion cost can be neglected, in particular the computation cost of the fluid quantities to be diffused were neglected. Such an assumption is not realistic when K becomes larger or more precisely when N/K becomes smaller.

Refining and redesigning the algorithms that were proposed in [16, 14, 15], we propose here to revisit the results in [14] and evaluate the benefit of a simple and natural dynamic partition strategy in order to control and equalize the work load of each virtual machine when parallel computation is used. Such a dynamic scheme may be also required when we assume that the underlying graph structure is evolving continuously in time and updates are applied continuously (cf. [15]).

In Section 2, we describe the distributed architecture that is considered in this paper. Section 3 presents the evaluation analysis based on synthetic data and dataset of web graph.

2. DISTRIBUTED ARCHITECTURE

In this paper, we will evaluate the performance of the proposed distributed algorithm focusing to the eigenvector problem associated to PageRank type equation. However,
the algorithm is described here in a more general case. We assume given a square matrix $P$ of size $N \times N$ and an initial condition $B$ (a vector of size $N$). The D-iteration applied on $(P, B)$ solves $X$ (a vector of size $N$) satisfying:

$$X = PX + B.$$ 

The approach proposed here should work as soon as the spectral radius of $P$ is strictly less than 1 (this is what we could call a diagonally dominant system that was mentioned in the introduction). In particular, the entries of $P$ or $B$ may be positive or negative (cf. [16]). However, for a better intuitive understanding, we chose here to focus on the case where all entries of $P$ are non-negative and implicitly associated to a transition matrix.

### 2.1 D-iteration: diffusion approach

We recall that the D-iteration is based on the fluid diffusion approach where one step of the iteration consists in choosing a node $i_n$ ($n$-th step) and diffusing all fluid at node $i_n$ to its children nodes (non zero entries of the $i_n$-th column of $P$): at each step of the iteration, we keep two state vectors: the current residual/transient fluids are described by the vector $F_n$ and the history (counting the amount of diffused fluid by each node) of the fluid diffusion by the vector $H_n$. 

Below, an adaptation of the pseudo-code in [14] for the general case:

**Initialization:**
For $i=1..N$:
1. $H[i] := 0$; // History (counter)
2. $F[i] := B_i$; // Fluid

**Iteration:**
While $(r > \text{Target\_Error})$:
1. Choose $i$; // node selection
2. $sent := F[i]$;
3. $H[i]$ += $sent$;
4. $F[i] := 0$;
5. For $(j \text{ such that } p(j,i) != 0)$:
   - $F[j] += sent \times p(j,i)$;
   - $r := |F| = \text{sum}_{j} |F[j]|$;

When the above scheme converges (DD system), we have asymptotically (when $\text{Target\_Error} \rightarrow$ zero) $X = H$.

### 2.2 Distributed algorithm

We assume that the set $\Omega = \{1,...,N\}$ is partitioned in $K$ sets $\Omega_k, k = 1,...,K$ (static or dynamic, see Section 2.5). We set $L$ the number of non zero entries of the matrix $P$ (total number of links).

#### 2.2.1 Local information and diffusion

We distribute the computation tasks of the D-iteration scheme between $K$ virtual machines (we call PIDs) as follows (cf. [14]):

- each PID$_k$ keeps information on:
  - the set of nodes it is responsible for: $\Omega_k$;
  - the extracted matrix $C_k(P) = (p_{ij})_{i \in \Omega_k, j \in \Omega_k}$, the column vectors of $P$ corresponding to $\Omega_k$;
  - the marginal fluid vector $[F]_k = ((F_i))_i \in \Omega_k$;
  - the marginal history vector $[H]_k = ((H_i))_i \in \Omega_k$;
  - the previous history vector $[H_{old}]_k = ((H_i)_i \in \Omega_k$,
    - the history vector value at the moment of the last fluid transmission (to other PIDs);
  - its activity state: active or idle state;
  - the target error value: $\text{Target\_Error}$;

- each PID$_k$ maintains two local variables (evaluated periodically):
  - the local residual fluid: $r_k = |[F]_k|$;
  - the fluid to be transmitted: $s_k = |C_k(P)([H]_k - [H_{old}]_k)|$;

- each PID$_k$ applies the local diffusion algorithm (*) below (when not in idle state);

- activity state:
  - initialized to active;
  - PID$_k$’s state is set to idle when
    $$r_k < \max(s_k/10.0, \text{Target\_Error} \times \epsilon/K/10),$$
    where $\epsilon$ is a factor depending on $P$: for PageRank equation, $\epsilon = 1 - \text{damping\_factor}$;

- each PID$_k$ select the node to be diffused by a cyclic check-up of elements of $\Omega_k$ of the condition:
  $$(F)_i \times w_i > T_k,$$
  where $T_k$ is a threshold value initialized to an arbitrary value larger than $\max_{i \in \Omega_k} (F)_i \times w_i$ and $w_i$ the weight we associate to the node $i$; the greedy approach would set $w_i = 1$; other candidates are: $w_i = 1/(\#\text{out}_i)$ or $w_i = 1/(\#\text{out}_i \times \#\text{in}_i)$, where $\#\text{out}_i$ and $\#\text{in}_i$ are respectively the number of of the outgoing links from (number of non zero entries of i-th column of $P$) and the incoming links to node $i$ (number of non zero entries of i-th line of $P$). By default, we choose in this paper $w_i = 1/(\#\text{out}_i)$. When for all $i$, the condition $(F)_i \times w_i > T_k$ is not satisfied, we apply: $T_k := T_k/\gamma$ (by default, $\gamma = 1.2$).

**Local diffusion for PID(k): (*)**

Choose $i$ in Omega$_k$;
1. $sent := F[i]$;
2. $H[i]$ += $sent$;
3. $F[i] := 0$;
4. For $(j \text{ in Omega}_k \text{ such that } p(j,i) != 0)$:
   - $F[j] += sent \times p(j,i)$;

#### 2.2.2 Fluid exchange

The transmission of fluid from PID$_k$ to other PIDs is done when:

$$s_k > r_k/2. \quad (1)$$

The idea is just to anticipate a bit the moment when $s_k$ and $r_k$ becomes equal. The PIDs (PID$_{k'}$) receiving $\text{received} = |C_k(P)([H]_k - [H_{old}]_k)|$ fluids reinitialize $T_{k'}$ to min($T_{k'} \times (r_{k'} + \text{received})/r_{k'}, \text{received}$).
2.3 PID modelling

As in [14], we consider a time stepped approximation for the simulation of the distributed computation cost (for now running on a single PC); during each time step, each PID can execute \( PID_{speed_k} \) operations. By default, we set: \( PID_{speed_k} = PID_{speed} = N/K \) (by default, PIDs are assumed to compute at the same speed).

When a PID is active, it increments \( count_{active_k} \) each time an elementary operation (a diffusion from one node to another node in the same \( \Omega \) set, which roughly corresponds to a product of one entry \( (F)_i \) with one entry of the matrix \( (P)_{ij} \) and the addition of the product to \( (F)_j \)) is done.

Every time step, we set a local counter that counts the number of elementary operations that are not consumed (because entering in the idle state). When a PID is idle, the wasted operations are then added to \( count_{idle_k} \).

In the following, the number of iterations is defined as the normalized quantity:

\[
\frac{count_{active_k} + count_{idle_k}}{L}
\]

so that it can be easily compared to the cost of one matrix-vector product, or one iteration in power iteration.

2.4 Computation cost

The computation effort of \( PID_k \) is indirectly estimated through \( count_{active_k} \). This counter is incremented:

- by one each time there is a local diffusion from one node to another;
- by one to the receiver for each diffusion to one node (during fluid exchange) managed by the receiver; for the sender, we increment by one for each diffusion coming from \( C_k(P)([H]_k - [H_{idle}]_k) \); this is the quantity that was underestimated in [14];
- by the number of nodes re-affected for the partition set adaptation.

2.5 Partition sets

2.5.1 Static partition sets

As in [14], we consider two simple \( K \) partition sets for comparison purpose:

- Uniform partition: \( \Omega_1 = \{1, 2, ..., N/K\} \), \( \Omega_2 = \{N/K + 1, 2, ..., 2 \times N/K\} \), etc
- Cost Balanced (CB) partition: \( \Omega_k = \{\omega_1, \omega_2, 1, ..., \omega_k+1-1\} \) such that \( \sum_{r=1}^{\omega_k+1} (#out_{\omega_k}) = L/K \),

such that \( \{1, ..., N\} = \Omega = \cup_{k=1}^{\omega_k} \Omega_k \). The intuition of the cost balanced partition is that when we apply the diffusion iteration on all nodes of each \( \Omega_k \), the diffusion cost is constant. The main reason why we chose this is the simplicity of its computation [14].

2.5.2 Dynamic partition sets

In the initial state, we start with the uniform or CB partition sets. Then, we update the following quantity every time step (\( PID_{speed} \) operations in active or idle state):

\[
slope_k := slope_k \times (1 - \eta) - log(r_k + s_k + \varepsilon)/log(10.0) \times \eta
\]

where \( \varepsilon = \text{target}\_\text{error}/K/1000 \) is added to avoid undefined value of \( \text{slope}_k \). The quantity \( -\text{slope}_k \) measures the moving averaged value of the exponent (base 10) of \( r_k + s_k \): if we plot the curve \( r_k + s_k \) as a function of the number of iteration (normalized) in logscale on y-axis, the exponent represents the slope of the curve. By default, we used \( \eta = 0.5 \).

Then, every time step, we compute \( k \) which maximize and minimize \( \text{slope}_k \) (resp. \( \text{max}, \text{min} \)). If the difference is more than 50%:

\[
\text{if } (\text{slope}_\text{min} < \text{slope}_\text{max} + log(0.5)/log(10.0))
\]

then, we reaffect:

\[
\Omega_{\text{min}} \times \min \left( \frac{\text{slope}_\text{min} + 1}{\text{slope}_\text{max} + 1}, 0.1 \right)
\]

nodes from \( \Omega_{\text{min}} \) to \( \Omega_{\text{max}} \) (\( \text{min} \) identifies the slowest PID).

To minimize the oscillation behaviour, the sets that are just re-affected (decreased or increased) can not be re-affected during the next \( Z \) steps (by default \( Z = 10 \)).

When the \( \Omega_k \) set is re-affected, we increment its operation cost counter \( count_{active_k} \) (by the number of nodes modified for \( \Omega_{\text{min}} \) and \( \Omega_{\text{max}} \)).

3. EXPERIMENTS AND EVALUATION

3.1 Synthetic data

We first used a synthetic data generated as follows: assuming a power-law \( 1/k^\alpha \) (\( \alpha = 1.5 \) used here) for the in-degree and the out-degree distribution, we generated random links between pair of nodes (see [17] for more details).

3.1.1 Analysis of \( K = 2 \): \( N = 1000 \)

Let us start with 2 PIDs case for an easier illustration of the problem. Figure 1 shows the plots of the convergence speed (given by the ratio of \( r_k + s_k \) and the number of iterations) in logscale on y-axis, when starting with a static partition sets of 250 + 750, 500 + 500 and 750 + 250 (in this case, \( L = 9543 \)).
can improve by factor above 2. Figure 2 shows the plots of the convergence speed integrating the fluid exchange cost: the relative gain to the single PID case is much less important than previous results, illustrating the importance of this factor even when $K$ is small.

![Figure 2: Illustration of convergence speed: fluid exchange cost integrated.](image1)

Figure 2: Illustration of convergence speed: fluid exchange cost integrated.

Figure 3 illustrates the impact of the partition adaptation on the convergence speeds that are made closer: in this case, we took initial partition sets of 750-250 and let the system self adapts.

![Figure 3: Illustration of the impact of the dynamic partition.](image2)

Figure 3: Illustration of the impact of the dynamic partition.

Figure 4 illustrates the evolution of the partition sets when starting from 750-250 (here, we took $Z = 1$ for a quicker adaptation).

![Figure 4: Illustration of the evolution of the dynamic partition.](image3)

Figure 4: Illustration of the evolution of the dynamic partition.

Table 1 gives a comparative computation time (number of iterations of the slowest PID) of different approaches for $K = 1$ to 128 ($Z = 10$): by construction, this can be considered as the most favourable situation for the uniform partition (links are independently and identically distributed to all nodes). We see that the dynamic strategy can still improve in almost all situations (but not too much).

To further illustrate the advantage of the dynamic adaptation, we biased the nodes ordering replacing the complete random one (previous) by the number of outgoing links (cf.

| $K$ | From Uniform Partition | From CB Partition |
|-----|------------------------|-------------------|
|     | Static | Dynamic | Static | Dynamic |
| 1   | 2.39   | 2.39    | 2.39   | 2.39    |
| 2   | 1.39   | 1.25    | 1.31   | 1.38    |
| 4   | 0.85   | 0.85    | 0.81   | 0.80    |
| 8   | 0.56   | 0.53    | 0.49   | 0.47    |
| 16  | 0.37   | 0.35    | 0.43   | 0.38    |
| 32  | 0.29   | 0.27    | 0.31   | 0.26    |
| 64  | 0.26   | 0.22    | 0.30   | 0.24    |
| 128 | 0.26   | 0.26    | 0.35   | 0.29    |

Table 1: Illustration of the computation time for a target error of $1/N$: $N = 1000$. 

To further illustrate the advantage of the dynamic adaptation, we biased the nodes ordering replacing the complete random one (previous) by the number of outgoing links (cf.
| K  | From Uniform partition Static | From Uniform partition Dynamic | From CB partition Static | From CB partition Dynamic |
|----|-------------------------------|--------------------------------|--------------------------|----------------------------|
| 1  | 3.79                         | 3.79                          | 3.79                     | 3.79                      |
| 2  | 3.07                         | 2.96                          | 2.83                     | 2.33                      |
| 4  | 2.48                         | 2.16                          | 3.42                     | 2.68                      |
| 8  | 1.97                         | 1.53                          | 5.09                     | 2.63                      |
| 16 | 1.57                         | 1.02                          | 6.01                     | 2.40                      |

Table 2: Illustration of the computation time for a target error of $1/N$: $N = 1000$. Nodes are ordered by the number of outgoing links.

| K  | From Uniform partition Static | From Uniform partition Dynamic | From CB partition Static | From CB partition Dynamic |
|----|-------------------------------|--------------------------------|--------------------------|----------------------------|
| 1  | 4.96                         | 4.96                          | 4.96                     | 4.96                      |
| 2  | 3.65                         | 3.48                          | 3.55                     | 3.02                      |
| 4  | 2.97                         | 2.03                          | 2.57                     | 1.91                      |
| 8  | 2.93                         | 1.69                          | 2.48                     | 1.62                      |
| 16 | 3.14                         | 1.35                          | 2.28                     | 1.25                      |

Table 3: Illustration of the computation time for a target error of $1/N$: $N = 1000$. Nodes are ordered by the number of incoming links.

Table 2: we see that the CB static strategy is not good and when $K \geq 4$ its performance is even degraded.

The results of the case when the nodes are ordered by the number of incoming links are shown in Table 3: here the uniform partition is the worst one.

Globally, what we observe is that when $N/K$ becomes too small, the gain is limited or the performance may be even degraded due to the fluid exchange cost. Finally, we observe a very good stability/performance of the dynamic partition strategy in all situations.

### 3.2 Web graph datasets

For the evaluation purpose, we experimented the dynamic partition strategy on a web graph imported from the dataset uk-2007-05@1000000 (available on [1]) which has 41,247,159 links on 1,000,000 nodes (45,766 dangling nodes).

Below we vary $N$ from 1,000 to 100,000 extracting from the dataset the information on the first $N$ nodes.

Figure 5 shows the summarized results on the convergence speeds (normalized to the convergence cost for $K = 1$) for $N = 1000$, 10000, 100000 starting from the uniform partition (unfortunately, we could not yet handle $N = 1000000$ case because of the memory limitation on a single PC). We clearly see that because of the fluid exchange cost, the convergence becomes slower when $K$ is too large compared to $N$ and that for larger $N$ the optimal $K$ value is larger.

Figure 6 shows the summarized results on the convergence speeds starting from the CB partition. Figure 5 and Figure 6 correct results reported in [14] (fluid exchange cost was underestimated) when $N/K$ is small. However, the main conjecture/result which states that, when the computation is distributed over $K$ virtual machines, the computation speed increases almost linearly with $K$ with a slope becoming closer to one when the number $N$ of linear equations to be solved increases is still true: we conjecture that the slope goes to one asymptotically for large $N/K$ and this is very clearly visible in the curves of the dynamic partition based approaches (Unif+DYN or CB+DYN) when $N$ is increased.

Figure 7 shows the consequence on the proportion of the PIDs’ idle state

$$\sum_k \text{count}_{\text{idle}}_k$$

when different approaches are applied ($N = 10000$). We see a clear reduction of the idle state with the dynamic strategy when the fluid exchange is not dominant.

Figure 8 shows the typical result of two different convergence speeds: in this case PID2 is the slowest one. The fluid exchange is done every about 1.2 iterations which is clearly visible here. We can see that PID1 can enter in the

![Figure 5: Convergence speed-up factor: starting from uniform partition.](image-url)

![Figure 6: Convergence speed-up factor: starting from CB partition.](image-url)

Table 4: Extracted graph: $N = 1000$ to 100000.

| $N$  | $L$ (nb links) | $L/N$ | $D$ (Nb dangling nodes) |
|------|----------------|-------|--------------------------|
| 1000 | 12,935         | 12.9  | 41 (4.1%)                |
| 10000| 125,439        | 12.5  | 80 (0.8%)                |
| 100000| 3,141,476     | 31.4  | 2729 (2.7%)              |

![Table 4](image-url)
idle state because it is waiting for inputs from PID2 (for $x$ between 4 and 5, between 6.5 and 7.5 etc) when it reaches the target value $\max(s_k/10.0, \text{target error} \times \epsilon/K/10)$: this is globally not optimal in terms of the PID’s computation capacity utilization.

Figure 7: Proportion of the idle state: $N = 10000$.  

Figure 8: Evolution of convergence: $K = 2$, $N = 100000$ with static uniform partition.

Figure 9: Evolution of convergence: $K = 2$, $N = 100000$. Dynamic partition from the uniform partition.

Figure 10: Evolution of partition sets: $K = 2$, $N = 100000$.

Figure 11: Evolution of convergence: $K = 2$, $N = 100000$ with static CB partition.
strategy is above 2.

Figure 12: Evolution of convergence: $K = 4$, $N = 100000$. Comparison of static unif., static CB and dynamic from unif. and CB.

Figure 13: Evolution of convergence: $K = 8$, $N = 100000$. Comparison of static unif., static CB and dynamic from unif.

Figure 14 shows the result of 128 different convergence speeds with $K = 128$: in this case, we can identify 2 slowest PIDs. The computation capacities of 126 other PIDs are likely to be wasted.

Figure 15 shows the impact of the dynamic partition starting from the uniform partition for the same case than Figure 14. In this case, the speed-up factor is about 4 thanks to a better computation effort redistribution between PIDs.

Figure 16 and Figure 17 show the global convergence (an upper bound on the $L_1$ norm to the distance) for different approaches ($N = 10000$): the benefit of the dynamic adaptation is more visible for $K \geq 8$. Note that those curves must be strictly decreasing function: we observe here some local fluctuation due to the artefact of the time stepped approximation (linked to the fluid exchange cost): when the fluid exchange cost becomes important, the concerned PID is likely to be frozen during that time.

Figure 18 and Figure 19 show the global convergence for $N = 100000$: when $K$ and $N$ are larger, the analysis be-
comes much more complex: we can observe significant and sudden slope modification during the iteration. See for instance Unif-DYN or CB curves for $K = 512$ in Figure 19.

One of very visible effect is the impact of the fluid exchange cost which is increased for larger value of $K$.

4. CONCLUSION

In this paper, we presented an adaptive dynamic partition strategy applied to a distributed computation architecture of the D-iteration method. Through experiments on synthetic data and real dataset, we showed that a dynamic partition strategy brings a robustness and a better efficiency guarantee compared to the static partition strategy, especially when $N$ is large. We believe that, even though this is preliminary results that need to be confirmed by a real deployment of a distributed system with possibly further adaptation/modification of the algorithm design, we showed here the potential of a new promising distributed computation architecture to solve a very large diagonal dominant class of linear systems.

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