Complex Grid Computing

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Received: date / Revised version: date

Abstract. This article investigates the functional properties of complex networks used as grid computing systems. Complex networks following the Erdős-Rényi model and other models with a preferential attachment rule (with and without growth) or priority to the connection of isolated nodes are studied. Regular networks are also considered for comparison. The processing load of the parallel program executed on the grid is assigned to the nodes on demand, and the efficiency of the overall computation is quantified in terms of the parallel speedup. It is found that networks with preferential attachment allow lower computing efficiency than networks with uniform link attachment. At the same time, considering only node clusters of the same size, preferential attachment networks display better efficiencies. The regular networks, on the other hand, display a poor efficiency, due to their implied larger internode distances. A correlation is observed between the topological properties of the network, specially average cluster size, and their respective computing efficiency.

PACS. 89.75.Fb Structures and organization in complex systems – 89.20.Ff Computer science and technology – 89.20.Hh World Wide Web, Internet – 02.10.Ox Combinatorics; graph theory – 89.75.Hc Networks and genealogical trees

1 Introduction

Among the many implications of the scientific and technological advances in microelectronics along the last decades, the availability of microprocessors characterized by ever diminishing size, cost and power consumption (per operation), together with increasing computing power, has led to an unprecedented opportunity for parallel computing, allowing simulations of non-linear and complex physical systems. More recently, the advent of the Internet paved the way for using a network of computers to obtain
a very large and powerful computing system, defining the new research areas of grid computing \cite{2,3,4} and parasitic computing \cite{5}.

A parallel computing system consists of a set of processing elements connected by some kind of communication network. A parallel program runs on the system by partitioning the work to be done in several pieces that are executed on the available processing elements. To collaborate in the execution of the program, each piece must, as a rule, communicate with others through the communication network. To show a good performance in the execution of a program, a parallel system must then: (i) make a large number of processing elements available to the application; and (ii) enable fast communications between these processing elements.

For grid computing, the first condition can be met by the large amount of computers available in the Internet, as long as their owners agree to make their processing power available. But the second condition is difficult to meet: even if high bandwidth networks are common, the latency to deliver a message to a site geographically far from the origin is large. The increased processing power of microprocessors due to Moore’s law (i.e. the number of transistors in a chip doubles each 1.5 years) contrasts with the typically slow interconnection between processing elements, thus undermining the performance of parallel systems due to the relatively large time spent to send data for processing elsewhere in comparison with the time taken to process that data. A program will only have good performance on the grid if the amount of computations done is large in comparison with the time needed to send a message between two processors. Thus, the effective use of grid computing remains a challenge, demanding a delicate balance between computation and interprocess communication workloads. As a rule, the weaker the coupling (i.e. the amount of communication needed) between different processing tasks, the higher the overall efficiency in a given parallel system.

Generally, a more densely connected processing network favors faster data transmission, as the mean distance between nodes tends to decrease, but at the expense of additional communication resources. Moreover, the specific kind of processing to be performed and the availability of the computer resources for collaborative (or parasitic) computing also play an important role in defining the overall grid execution performance.

The novel area of complex networks (e.g. \cite{6,7,8}) has drawn increasing interest of the physics community. A large part of the success of such an approach derives from the fact that such networks have been found to adhere, at varying degrees, with important real phenomena such as transmission of infectious diseases, social and ecological interactions, and the Internet. The fact that today most computers are interconnected through the Internet has contributed further to promote the systematic investigation of the Internet characteristics, a task that can greatly benefit from physical modeling approaches.

As several features are shared by grid computing systems and complex networks, much can be gained through integrative and comparative approaches, allowing cross-
fertilization between those two important areas. The underlying idea in the current work is to study the efficiency of parallel/distributed architectures whose interconnections are defined in terms of complex network models. Such an investigation therefore focuses on the integration between topology and function of the networks, an important aspect of complex network research [8]. Regular networks, which are often used in parallel computing, are also considered as a reference for comparison. It is particularly interesting to verify how the specific properties of these interconnecting schemes—such as the average vertex-vertex distance and cluster size—affect the processing time and efficiency for different network configurations. Such a possible dependence between the topological and functional properties of the networks is backed by recent works which verified that the emergent features of complex networks, such as associative memory recall in neuronal networks, can be strongly affected by the network interconnecting scheme and phase transitions [9, 10].

In related works, the complex network paradigm has also been explored from the perspective of search algorithms [11, 12, 13] and information transfer in graphs [14, 15, 16].

2 Network Models Used

The main purpose of this article is to study the influence of some network topological features in the efficiency of a grid system for the execution of a suitable parallel program. Therefore, we will not try to use network models that reproduce the characteristics of the Internet; we instead restrict ourselves to some simple models which are described below. The models are undirected, reflecting the bidirectional transfer of packets on the Internet.

Let a complex network be represented as a graph with \( n \) nodes, identified as \( i, i = 0, \ldots, n - 1 \), and unweighted, undirected edges represented as \((i, j)\). The first model considered is the Erdős-Rényi (ER) model with a fixed number \( c \) of edges. In this model, for each connection, two nodes \( i \) and \( j \) are chosen uniformly among all the nodes to establish the connection \((i, j)\). Self-connections (connections of a node with itself) and duplicate connections (connections between already connected nodes) are avoided in this and all the following models.

ER graphs have a fast decaying degree distribution, with very small probability for nodes with high degree (also called hubs). To widen the degree distribution and increase the probability of high degree nodes, a preferential attachment (PA) model is used. Networks in this model are generated as described in [9]: starting with all nodes without connections and choosing two nodes to connect by drawing nodes from a list of node numbers represented in amount proportional to their respective number of connections (plus one, to account for the unconnected nodes). Note that this network model, although having preferential attachment, has no growth and so does not lead to scale-free networks (see Section 3.1).

Hubs of larger degrees can be found in scale-free networks. A simple model for scale-free networks was proposed by Barabási and Albert [17]. In their model, the network starts with \( m_0 \) connected nodes and grows by the addition of one node at a time. When a node is added,
m new connections from the new node to already existing nodes are made, and each already existing node can be chosen to receive a connection with probability proportional to its degree \(k\). In this model, all nodes in the network form a single large connected component; as we are interested in studying the influence of percolation with growing connectivity, their model is not adequate due to the nonexistence of a percolation transition and to the impossibility of specifying an average connectivity that is not an even integer (the average degree is always \(2m\)). For this reason, their model is here generalized as described in the following. In the model used in this work, which we call scale-free (SF) model, instead of having a fixed number of connections for each new node \(i\), a random number \(m_i\) is chosen using a Poisson distribution with mean \(m\), and \(m_i\) connections from this node to the already existing nodes are made. As some nodes may have \(m_i = 0\), the network will have unconnected nodes; to enable these nodes to receive connections with the addition of new nodes, each already existing node is chosen with probability proportional do \(k+1\) instead of \(k\). In this model, as \(m\) is only a mean value, it can be any real number (instead of only an integer number, as in the Barabási-Albert model); also, as new nodes are not necessarily connected to the already existing nodes, the network consists of many connected components.

For grid computing, as for all kinds of collaborative work between the agents represented by a network, the binding of the agents to other agents of the network is a necessity. There is, therefore, a tendency to bind nodes to the network as new communication resources are available, instead of using them to bind already connected nodes. This suggests a different kind of random network construction: connecting new nodes to the network should be given preference while there are still isolated nodes. For that purpose, we introduce here two new models of random networks. The parameters for their construction are: the number of nodes \(n\) and the number of connections \(c\).

In the first model, one end of each new edge is chosen with uniform distribution among the isolated nodes and the other end with uniform distribution among all nodes (isolated or not). If at some point no more isolated nodes exist, both ends of the remaining edges are drawn with uniform distribution among all nodes.

In the second model, one end of the new edge is drawn randomly from the set of isolated nodes, as for the previous model, but the other end is drawn among all nodes with probability proportional to the node degrees, as in the previously discussed preferential attachment network.

We call these networks insertion networks, because the connections are used to insert the nodes in the network; the first model is called uniform-uniform (UU) insertion network, and the second model uniform-preferential (UP) insertion network.

For comparison, we study also three regular network structures common in parallel systems [18]: the hypercube and the 2D and 3D tori. The construction of an hypercube with \(n\) nodes can be explained as follows. For node \(i\), represent the value of \(i\) in binary using \(\lceil \log_2 n \rceil\) bits; there is a link \((i, j)\) iff the binary representation of
$j$ differ in exactly one bit from that of $i$. For example, in an $n = 8$ network, node 3 (binary 011) is connected to nodes 2 (010), 1 (001) and 7 (111). In the bi-dimensional torus the nodes are distributed in a grid of size $(n_x, n_y)$ (with $n_x n_y = n$), each node receiving a label $(x, y)$, where $x = \lfloor i/n_y \rfloor$ and $y = i \mod n_y$; node $(x, y)$ is connected with node $(x', y')$ iff $x' = x$ and $(y' = y \pm 1) \mod n_y$ or $y' = y$ and $(x' = x \pm 1) \mod n_x$. For example, in a $n = 8$ bi-dimensional torus organized as a $2 \times 4$ grid, node 3 corresponds to coordinates (0, 3) and is connected to nodes 2 (0, 2), 0 (0, 0), and 7 (1, 3). The three-dimensional torus is similarly constructed.

3 Results

The results are divided in two parts. First, some results concerning the properties of the network models described are presented. Then, results obtained by using these network models as communication infrastructure for the simulation of the execution of a parallel program on a grid are shown.

3.1 Network properties

A thorough analysis of the network models described is out of the scope of this paper. Here only some properties of interest to the analysis of the following grid simulation results (Section 3.2) are presented.

The topological properties of those networks are quantified in terms of the following measures: (a) node degree $k$; (b) mean vertex-vertex distance $\ell = \frac{2}{n(n-1)} \sum_{i>j} d_{ij}$ where $d_{ij}$ is the geodesic distance (distance, in number of links, of the shortest path) between nodes $i$ and $j$ and the summation includes only pairs $ij$ that have a path connecting them; and (c) mean cluster size $s$. A cluster, also known as a connected component, is a set of directly or indirectly connected nodes, i.e., nodes that can be reached from all the other nodes on the cluster by a path. To evaluate the mean cluster size, we compute for each node of the network the number of nodes on its cluster and take the average of these values for all nodes on the network. Note that this average includes the largest cluster, and is dominated by it after percolation. To enable the comparison between the different complex networks models with respect to their connectivity, we use the parameter $z$ defined as $z = 2m$ for the SF model and $z = 2c/n$ for the other models. In the limit of large $n$ we have $z = \langle k \rangle$ for all network models.

In the following, network characteristics are quantified as averages for 50 random networks for each set of parameters. Error bars display the 99% confidence interval for the computed average, considering normal distribution of the averaged values.

The degree distributions for each of the considered models for $n = 100000$ and $z = 3$, compared with the ER model, are presented in Figure 1 where $P(k)$ is the cumulative probability distribution (probability of finding a node in the network with degree larger than $k$). The UU model is almost indistinguishable from the ER model in terms of degree distribution. Due to the preferential attachment rule, the PA and UP models have broader de-
gree distributions. For the UP model this effect is not so marked due to the preference given to newly added nodes in new connections. As expected, the SF model follows a power law (with a finite-size cutoff), given rise to a large probability of nodes with high degree.

Fig. 2 gives the average network cluster size \( \langle s \rangle \) normalized by the network size \( n \), and Fig. 3 shows the average node distance \( \langle \ell \rangle \) for the networks. These results are shown as functions of \( z \) comparatively to the ER model.

As Figs. 2(a–d) show, there is an abrupt transition (a percolation transition) from small to large cluster sizes as the connectivity grows. In the small cluster size region, the mean distance tends to grow with the connectivity \( z \) (Fig. 3), as new links result in new connections between previously unconnected nodes; in the large cluster region, increased connectivity reduces mean distance, as most of the nodes are already connected in the largest cluster. A striking feature of the results for the PA model (and to some extent for the SF model) is the small cluster sizes even for high connectivity. The reason is that, as the number of nodes connected in the largest cluster grows, the probability of linking an unconnected node is very small, due to the preferential attachment rule used to choose the ends of new links, resulting in a relatively large number of isolated nodes or small clusters. In the SF model, this problem is minimized by the fact that one end of each new connection always go to a new (previously unconnected) node. The insertion (UU and UP) models show very similar behavior in terms of average distance and cluster size. The formation of a cluster spanning most of the nodes occurs for these models for higher connectivities than for the ER model. The explanation is that links that could be used to connect small clusters to form a larger one are being used to link new nodes. On the other hand, the size of the resulting largest cluster tends to be larger, as isolated nodes are less likely to appear. Distances in these models tend to be higher than in the ER model, because the same number of links is used to connect a larger number of nodes. For sufficiently higher connectivities, this last property is compensated in the UP model by the onset of high degree nodes (hubs), that shorten the mean distance between the nodes of the cluster. The presence of hubs is also the explanation for the smaller distances shown by the PA and SF models; this advantage vanishes as the connectivity grows.

More detail about the size of clusters is given in the left side of Figs. 4(a–e), which display the cumulative probability distribution of cluster sizes (i.e. the probability \( P(s) \) that a randomly selected node is part of a cluster of size greater or equal to \( s \)) against \( s/n \) for some values of \( z \).

These figures show that for small average connectivities the probability of finding a large cluster is negligible. On the other hand, for sufficiently large average connectivities almost all nodes are found in large clusters. The models without preferential attachment (ER, Fig. 4(a), and specially the insertion models, UU Figs. 4(c) and UP Fig. 4(d)) show a sharp transition from a regime with low probability of large clusters and high probability of small clusters to a regime with high probability of large clusters and small probability of small clusters. On the preferen-
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10−7
10−6
10−5
10−4
10−3
10−2
10−1
100
101
102
103
104
105
106

\( P(k) \)

(a) (b) (c) (d)

Fig. 1. Degree distributions for the network models. The ER model results appear in all figures, as a reference for comparison. The networks have \( n = 100000 \) nodes and \( z = 3 \): (a) preferential attachment; (b) insertion with uniform probability; (b) insertion with preferential attachment; and (d) scale-free.

0
0.2
0.4
0.6
0.8
1
0
0.2
0.4
0.6
0.8
1
0
0.2
0.4
0.6
0.8
1
0
0.2
0.4
0.6
0.8
1

\( \langle s \rangle /n \)

(a) (b) (c) (d)

Fig. 2. Normalized average cluster size, for \( n = 1000 \), for the PA (a), UU (b), UP (c) and SF (d) models, compared with the ER model.

tial attachment models (PA, Fig. 4(b), and SF, Fig. 4(e)), this transition is more gradual; they also display a larger probability of medium sized clusters before the percolation.

In the regular networks, all nodes are connected, and so \( \langle s \rangle /n = 1 \). Also, the value of \( z \) is fixed for each network type, given \( n \). The third column of Table 1 shows the values of the average distance for these network types.

3.2 Grid Simulations

The parallel computing systems were obtained by assigning a processing unity to each network node, while mes-
Fig. 3. Average distance between connected pairs, for \( n = 1000 \), for the PA (a), UU (b), UP (c) and SF (d) models, compared with the ER model.

Table 1. The topological and efficiency measurements for the three considered regular topologies \((n = 1000)\).

| Network type   | \( z \) | \( \langle \ell \rangle \) | \( \langle E \rangle \) |
|----------------|--------|----------------|----------------|
| 2D torus       | 4      | 16 ± 7         | 0.664          |
| 3D torus       | 6      | 7 ± 3          | 0.778          |
| Hypercube      | 9.9 ± 0.4 | 5 ± 2       | 0.816          |

messages flow along the network edges. The distributed application considered follows the master/slave paradigm (also known as manager/worker or bag of tasks), where a master delivers processing tasks on demand to slave computers. The computational tasks are assumed to be completely independent, in the sense that each node can proceed without additional communications after receiving the work packet. This arrangement is similar to many grid computing efforts, like SETI@home. The computations are partitioned into \( M \) work packets (tasks), each requiring the same amount \( L \) of computing time. The communication cost is taken to correspond to the minimum number of edges between the master and the slave requesting the data. The edge communication overhead is therefore equal for all edges and adopted as time unit. Taking into account this communication cost model, the very small number of short cycles present in all the considered network models (compared with the Internet) does not represent an additional limitation.

Given a network, each node \( i \) at a time is considered as master. The nodes that are part of the same cluster as the master start requesting tasks. The nodes that are not
Fig. 4. Cumulative probability distribution of cluster size $P(s)$ (probability of a randomly selected node being part of a cluster of size $s' \geq s$) in terms of $s/n$ and parallel efficiency $P(E)$ (probability of a randomly selected node achieving efficiency $E' \geq E$ when chosen as master) for the five models: (a) ER; (b) PA; (c) UU; (d) UP; and (e) SF. Distributions shown for five different average connectivity ($z$) values: $z = 0.4$ (○); $z = 0.8$ (+); $z = 1.2$ (□); $z = 2.0$ (×); and $z = 5.0$ (○). Network size: $n = 1000$; number of work packets $M = 5000$; work packet size $L = 100$. 

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part of the same cluster as the master cannot contribute to the computation because of the lack of connection to the master. After receiving a task from the master, the slave computes the result, taking time $L$, and sends it, together with a request for another task, to the master. When all $M$ tasks have been delivered and their results received, the master terminates the execution and computes its total execution time $T_i$. Isolated nodes cannot take part on a distributed computation and so, when chosen as master nodes, their execution time is considered infinite (they will wait forever to receive a work request from a slave).

To quantify the suitability of the network models for grid computing we compute the average speedup achieved by the execution of the application on the networks. The speedup is defined as the ratio between sequential and parallel execution times. For the problem considered, the parallel execution time for master node $i$ is the value of $T_i$ discussed in the previous paragraph and the sequential execution time is $ML$, so that $S_i = \frac{ML}{T_i}$ is the speedup. The mean speedup of a network is the mean value $\frac{1}{n} \sum S_i$. Note that, for isolated nodes, as discussed, $T_i = \infty$ and so $S_i = 0$; as these nodes are anyway considered in the average, if a network has many isolated nodes its average speedup is low. Another equivalent measure, used in our results, is the normalized speedup $E = S/n$, also known as parallel efficiency. The averages $\langle S \rangle$ and $\langle E \rangle$ are then taken for 50 different random networks for each model and parameter set.

If master $i$ sends a task to slave $j$, the time to complete the task (get the result back) is the sum of the computation time of the task with the time taken to send the task to the slave node and receive it back, that is, $L + 2d_{ij}$. At the same time, the number of nodes computing tasks is $s_i - 1$ ($s_i$ is the size of the cluster of master node $i$ and we subtract one because the master does not compute tasks). This indicates that, for the problem considered, two important network metrics are the average cluster size $\langle s \rangle$ (Fig. 2) and average distance $\ell$ between nodes with a path connecting them (Fig. 3).

The number of tasks $M$ was chosen so that each node has some tasks to process, but avoiding too large number of tasks, because the simulation time is proportional to

![Fig. 5. Parallel efficiency $\langle E \rangle$ as a function of the number $M$ (a) and size $L$ (b) of work packets. Network parameters are $n = 1000$ and $z = 3$. In (a) $L = 100$ and in (b) $M = 5000$.](image)
the size of the tasks $L$ was chosen to achieve computation times larger than the average communication time (two times the average distance; see also Figs. 3(a–d)), but small enough such that the communication time has a detectable effect. If $L$ is too large, the average communication time, and then the network structure, is not important; if it is too small, the grid system is not a good choice for the execution of the application. Fig. 5(a) shows the dependence of average efficiency $\langle E \rangle$ with the number of tasks $M$ (for fixed $L = 100$, $n = 1000$, and $z = 3$) for the five models. If $M$ is small, there will be not enough computational work to distribute evenly among the nodes, resulting in very small efficiencies; after some value of $M$, the efficiency is not much affected by a further increase of $M$. The relation of the efficiency with $L$ is very similar; a larger value of $L$ helps reduce the importance of the communication costs, increasing the efficiency; Fig. 5(b) plots this relation (for $M = 5000$, $n = 1000$, and $z = 3$). It can be seen that after $L \approx 100$ there is no significant increase in efficiency for larger $L$. The following results assume $M = 5000$ and $L = 100$; results for other values show no qualitative differences.

Figs. 6(a–d) present the average parallel efficiency as a function the $z$ for $M = 5000$ and $L = 100$. As can be easily seen by comparing Figs. 2(a–d) and 6(a–d), the average parallel efficiency tends to closely reflect the normalized average cluster size for all considered models.

This is further substantiated by a comparison of the left and right graphs of Figs. 4(a–e). The right graphs show the cumulative probability distribution $P(E)$ of a randomly selected node having efficiency greater or equal to $E$ when chosen as master node. It can be seen that $P(E)$ closely follows $P(s)$. The main differences are that the $P(E)$ curves are smoother and the largest achievable values of $E$ smaller than that of $s/n$. The latter difference is a result of the fact that a cluster with $s$ nodes cannot achieve speedup $S$, because the master node does not compute and the communication costs increase the execution time of each task (in comparison to sequential execution); the former difference is due to the fact that different clusters with the same number of nodes $s$ have different interconnection topology, resulting in different communication costs.

As a consequence, similar conclusions can be drawn for the efficiency of a grid computing network as for the mean cluster sizes of the corresponding network model. Particularly, we note a percolation transition of the efficiency with increasing connectivity from a regime of very small efficiency to a regime of high efficiency. This transition is more abrupt for the ER and particularly for the UU and UP models, and somewhat slower for the PA and SF models. The efficiency of the models with preferential attachment is restricted due to the already discussed higher number of isolated nodes, that contribute to a speedup of zero to the average speedup. The insertion models need a higher connectivity to reach the percolation point, but after that show a higher efficiency, demonstrating the advantage of using network resources to connect new nodes.

The strong correlation between the cluster size and parallel efficiency is further substantiated in Fig. 7 where
Fig. 6. Average parallel efficiency for $n = 1000$, $M = 5000$, $L = 100$, for the PA (a), UU (b), UP (c) and SF (d) models, compared with the ER model.

A scatter plot of efficiency against normalized cluster size is shown, that demonstrates the almost linear correlation between efficiency and normalized cluster sizes. Another interesting result observed from this figure is that the PA and SF networks tend to provide slightly better efficiencies than the other models. In other words, if clusters of similar size are considered, the presence of hubs that “short-circuit” the distances tends to enhance the speed-up of the computations in the networks.

In order to further investigate such a possible effect of the intrinsic connectivity properties of the considered models over the respective performance, the largest clusters of the four models were considered in isolation in the grid simulations. That is, networks with the same connectivity were generated and only their largest clusters were considered. Special care was taken so as to obtain such connected clusters with equivalent number of nodes (about 1000 nodes). The results are shown in Figure 8(a–d), where the efficiency of the parallel execution on the largest clusters, $\langle E_{\text{largest}} \rangle$, are plotted. As we are considering here only the largest clusters, the efficiency is computed with respect to the number of nodes of the cluster,
and not of the whole network. The results indicate a definite tendency of the SF model, and to a lesser extent of the PA model, to outperform the others. Such an effect is possibly a consequence of the shorter average lengths usually observed for this model —see Figure 3(a–d)— and the presence of *hubs* which act as message distribution nodes.

The results obtained for the regular topologies (hypercube and 2D and 3D torus) for $n = 1000$ are shown in Table 1. Interestingly, the 2D and 3D torus regular architectures led to rather low efficiencies, despite their relatively large node degrees, reflecting the respective large average distances, due to the absence of shortcut links in the regular structure of these networks. The hypercube topology allowed efficiencies comparable to the maximum obtained for the network models, but at the expense of almost 10 connections per node, which implies a high network cost.

### 4 Conclusions

While the scale-free and preferential attachment models allowed better efficiency considering only the largest cluster, the Erdős-Rényi model tended to provide better average speedup when all clusters were considered, as a consequence of the smaller number of isolated clusters implied by this type of network. The insertion models resulted in even better efficiencies, due to the inclusion of more nodes in the largest cluster after percolation. The random models had better efficiencies than the regular ones, due to the implied smaller average distances.

The results show that a network is of little use for grid computing before the percolation point is reached, that is, for values of $z$ before the formation of a cluster spanning most of the nodes, because of the very small resulting efficiencies. In other words, the percolation of the network used as a grid computing resource is of fundamental importance to the utility of the grid. Although the Internet already connects a very large number of computers, the use of these computers for grid computing is subject mainly to two limitations: a consent from the part of their owner and the installation of a grid computing software on them. It is therefore appropriate to consider the grid network as distinct from the Internet, and two computers in the Internet as connected in the grid network only if their owners have given permission to use *and* installed an appropriate set of protocols and compatible software. In this aspect, the obtained results motivate the grid community to achieve convergence in protocols and software, as the presence of many incompatible software platforms represent the presence of unconnected clusters of nodes in the network. This conclusion is related with the efficiency of the network as a whole for grid computing; for users in isolated clusters, the execution of an application in these clusters can be nevertheless of interest, if enough speedup is achieved.

Future work should extend the analysis to other types of distributed systems and applications. Network models closer to some real Internet characteristics, as in [20], should be considered. The inclusion of measures to quantify the load of intermediate nodes in packet transmission [21], like betweenness centrality, can improve the results, although centrality is non-trivially related to traffic flow if
congestion is considered [22]. If the assumption of communication through shortest paths (that requires global knowledge) is relaxed, the use of local search algorithms [12] will result in stronger dependence of efficiency with the network structure. A further refinement in this direction is to consider queuing of packets on the nodes and congestion. Recent works [15,16] have shown that transmission times are in this case not directly related to shortest distances, but have a much richer behavior, depending also on the total communication load carried by the network.

Another important generalization is to consider complex networks presenting links with different communication speeds. This can be modeled using weighted networks, in which the weight of the links reflect the bandwidth or inverse latency of the interconnecting links. Also the nodes can display different processing powers. These generalizations make the network models closer to real interconnection networks.

The generalization of the parallel application model to include communications between the tasks is of interest, expanding the classes of applications modeled and due to the importance of the network topology to the efficiency of these communications, resulting in an interesting interplay between network and application characteristics.

We conjecture that even with the generalization of the network, routing and application models, as suggested above, the efficiency will remain strongly related with clus-
ter size, although the correlation with shortest distance may be reduced, and other network features may increase their importance, resulting in stronger influence of the network model used.

L. da F. Costa is grateful to FAPESP (processes 99/12765-2 and 96/05497-3) and CNPq for financial support. G. Travieso was financially supported by FAPESP grant 98/14681-8.

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