Performance of RDF Query Processing on the Intel SCC
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Abstract—Chip makers are envisioning hundreds of cores in future processors for throughput oriented computing. These processors, called manycore processors, require new architectural innovations for scaling to a large number of cores as compared with today’s multicore processors. We report an early study on the performance of RDF query processing on a manycore processor. In our study, we use the Intel SCC, an experimental manycore processor from Intel Labs. This processor has new architectural features, namely, 48 Pentium cores, a high speed, on-chip mesh network to communicate between cores and access memory controllers, on-chip message passing buffers for high speed message passing, and software controlled fine-grained power management. We classify queries based on their I/O footprint and study the impact of two standard models, namely, task and data parallel programming models. Based on our experiments with synthetic and real RDF datasets on the SCC, we conclude that the task parallelism model provides an immediate way to boost the performance of RDF query processing.

I. INTRODUCTION

Chip makers are envisioning hundreds of cores in future processors for throughput oriented computing. In throughput oriented computing, we expect abundant parallelism opportunities in the workload, and aim to achieve high throughput using a large number of simple cores, while compromising the latency on individual cores [1]. A processor with such large number of cores is called a manycore processor. The cores may be homogeneous or heterogeneous. New architectural innovations for faster on-chip communication and efficient power management are necessary to scale to a large number of cores as compared with today’s multicore processors.

In recent years, a few manycore prototypes have emerged (e.g., 80 core processor called Polaris [2], Larrabee [3], Intel Single-chip Cloud Computer (SCC) [4]). Of particular interest to us is the Intel SCC, an experimental manycore processor from Intel Labs. This processor has new architectural features, namely, 48 Pentium cores, a high speed, on-chip mesh network to communicate between cores and access memory controllers, on-chip message passing buffers for high speed message passing, and software controlled fine-grained power management.

In this work, we attempt to understand the benefits and limitations of the SCC for parallel RDF query processing. RDF (Resource Description Framework) is a popular language for representing data on the Web [5]. It enables the interchange and machine processing of data by considering its semantics. The essence of RDF lies in the notion of representing any fact as subject, predicate, and object. Formally, RDF represents resources as a directed, labeled graph where a pair of adjacent nodes denotes two things and the directed, labeled edge represents their relationship. The source node denotes the subject; the sink node denotes the object; and the edge label is the predicate (or property). This “subject-predicate-object” relationship is commonly referred to as an RDF triple. SPARQL is a popular query language for RDF graphs [6]. Using SPARQL, complex graph pattern queries can be expressed on individual RDF graphs as well as across multiple RDF graphs.

In recent years, the RDF data model has become increasingly important in domain-specific applications and the WWW. Through RDF technologies, one can reason over semantic data, which is highly appealing in domains such as healthcare, defense and intelligence, biopharmaceuticals, and so forth. With the rapidly growing size of RDF datasets (e.g., DBPedia [7], Billion Triples Challenge [8]), there is a pressing need for high performance RDF processing tools. With the emergence of manycore processors, it is natural and timely to ask whether a manycore processor can boost the performance of RDF query processing – through parallel processing. To the best of our knowledge, there is no published work in this area. Recent studies on the Intel SCC have focused on low level aspects such as on-chip message passing performance, memory access latency, and power and energy consumption on benchmarks from high performance computing [4], [9].

In our study, we adopt standard task parallel and data parallel programming models for parallel RDF query processing. We categorize RDF queries on real and synthetic RDF datasets into two different query workloads based on their I/O footprint – one with small I/O footprint queries and the other with large I/O footprint queries. We study the effect of inter-query parallelism via the task parallel programming model on these workloads. We also study the effect of intra-query parallelism via the data parallel programming model on these workloads.

The rest of the paper is organized as follows. We present background and related work in Section II; we present the methodology of our study in Section III; we present the empirical findings in Section IV; and we conclude in Section V with a note on future work.

II. PRIOR WORK ON RDF QUERY PROCESSING

Today, there are a number of open-source and commercial tools for storing and querying RDF graphs. These tools either
store and process RDF in main-memory, use an RDBMS, or a native RDF database. The popular approach has been to use relational database systems for storing, indexing, and querying RDF [10], [11], [12], [13], [14], [15]. Abadi et al. proposed a vertical partitioning approach and leveraged a column-oriented DBMS for achieving an order of magnitude performance improvement over previous techniques [16]. RDF-3X [17] and Hexastore [18] demonstrated that storing RDF data in a single triples table and building exhaustive indexes on the six permutations of \((s, p, o)\) triples can significantly outperform the vertical partitioning approach [16] and also support a larger class of RDF queries efficiently. Recently, BitMat [19] was proposed to overcome the overhead of large intermediate join results in RDF-3X and Hexastore when queries contain low selectivity triple patterns. (Low selectivity implies large result set size.)

There are some RDF stores that operate in shared-nothing clusters (e.g., YARS2 [20], 4store [21], Clustered TBD [22]) by hashing triples/quadruples and distributing them on different nodes in the cluster. Parallel query processing is performed. The scalability of these approaches has been demonstrated on small sized clusters. Weaver et al. [23] have studied RDF query processing on supercomputers. More recently, tools for data intensive computing such as Apache Hadoop and Pig have been used for query processing and analytics over RDF data [24], [25], [26]. These approaches are more suitable for batch processing of queries. A few researchers have focused on parallel RDF reasoning [27], [28]. More recently, Huang et al., developed parallel RDF query processing techniques for large RDF graphs [29].

On the Intel SCC, Vidal et al., studied the parallelization of an AI automated planner using a hash-based distribution of tasks [30]. Petrides et al., studied the performance of relational decision support queries on the SCC [31]. However, none of the previous work has studied the performance impact of parallel RDF query processing on the Intel SCC.

III. OUR METHODOLOGY

In this section, we introduce our methodology for parallel RDF query processing on the SCC. Our first approach is to express inter-query parallelism via the task parallel programming model. Our second approach is to express intra-query parallelism via the data parallel programming model. While we adopt standard techniques for task and data parallelism, these techniques provide good insights into the benefits and limitations of the Intel SCC for RDF query processing. The query workloads we study are I/O bound in nature, unlike prior work on SCC [4], [9], which focused on high performance computing benchmarks. We consider two different types of query workloads: one that has relatively smaller I/O footprint and the other that has relatively larger I/O footprint.

A. Message Passing Interface

We use the popular Message Passing Interface (MPI) for writing parallel programs. MPI contains a standard library of routines for writing portable message-passing based programs. The MPI routines that we used for the task parallel and data parallel programming models are listed in Table I. MPI programs essentially create a collection of processes. MPI_Send and MPI_Recv allow a process to exchange messages with another process (point-to-point communication); MPI_Barrier enables processes to synchronize at certain points during execution; and MPI_Bcast, MPI_Scatter, and MPI_Gather are collective communication operations, which allow a process to communicate with a group of other processes.

B. Impact of Granularity

In parallel computing, granularity denotes the ratio between the amount of computation to the amount of communication. In fine-grained parallelism, we break a problem into relatively smaller sized computation tasks and therefore, may require more frequent communication between processors. In coarse-grained parallelism, we break a problem into relatively larger sized computation tasks and therefore reduce the frequency of communication between processors. However, fine-grained parallelism enables better load balancing than coarse-grained parallelism. But it may increase communication cost and synchronization overhead. By design, Intel SCC provides a high speed, on-chip network to enable fast communication between cores. Therefore, we attempt to partition the tasks as fine-grained as possible in our experiments. Because the query workloads we study are I/O bound, we use the I/O footprint to characterize the granularity of a task.

<table>
<thead>
<tr>
<th>MPI routines</th>
<th>Usage</th>
</tr>
</thead>
<tbody>
<tr>
<td>MPI_Send</td>
<td>Is called when a process wants to send a message in its local buffer to another process</td>
</tr>
<tr>
<td>MPI_Recv</td>
<td>Is called when a process wants to receive a message from another process</td>
</tr>
<tr>
<td>MPI_Barrier</td>
<td>Is called by a process to enter a barrier</td>
</tr>
<tr>
<td>MPI_Bcast</td>
<td>Is called by a process to broadcast the message to all processes in the group</td>
</tr>
<tr>
<td>MPI_Scatter</td>
<td>Is called by a process to scatter an array of data items to other processes</td>
</tr>
<tr>
<td>MPI_Gather</td>
<td>Is called by all processes in the group (one receiver, multiple senders) so that the receiver can collect different sized messages from the senders synchronously</td>
</tr>
</tbody>
</table>

TABLE I
MPI ROUTINES USED

C. Task Parallel Programming Model

We express inter-query parallelism via a straightforward task parallel programming model. Each query is regarded as a task. Our model is as follows. On one core, we run the master and on the other cores, we run workers. Algorithm 1 describes the set of actions performed by the master and workers. Lines 1-1 denote the actions taken by the master. Lines 1-1 denote the actions taken by a worker. The master maintains a single task pool. Once the master and workers have started (as MPI processes), each worker sends a message to the master. The master responds to a worker with a query from the task pool. The worker then executes the query locally on the index. (The index is constructed over the entire dataset and is shared by the workers.) Once completed, the worker...
IV. PERFORMANCE EVALUATION

We used RDF-3X [17], a state-of-the-art RDF query processing engine in our evaluation. RDF-3X was implemented in C++ and was compiled to run on the SCC using a 32 bit GCC compiler. The SCC cores ran Linux and had a NFS mounted file system where the indexes were stored. We did not modify the memory organization/configuration of the SCC and used the default setting.

We implemented the task and data parallel programming models described in Algorithms 1 and 2 using RCKMPI, a modified MPICH2 for the Intel SCC [33]. RCKMPI uses the message passing buffers (MPBs) in the SCC to allow low latency high bandwidth message passing. The SCC platform was initialized to run with tile frequency of 800 MHz, mesh frequency of 800 MHz, and memory controller frequency of 800 MHz.

A. Dataset and Queries

We used two real datasets, namely, YAGO2 [34] and Uniprot [35]. YAGO2 is a semantic knowledge base derived from Wikipedia, WordNet, and Geonames. Uniprot is a comprehensive resource for protein sequence and annotation data. We also generated a synthetic dataset using the Lehigh University Benchmark (LUBM) [36]. The ontology for this dataset is based on a university domain.

Note that the SCC cores generate 32 bit addresses. RDF-3X leverages memory mapping of index files and therefore, recommends 64 bit processors for indexing and querying large RDF datasets. To cope with the 32 bit addressing on the SCC, we indexed a set of triples in each dataset such that the index size was at most 2GB in size, a limit set by the underlying OS. This ensured that RDF-3X successfully ran the queries on the SCC. For YAGO, we indexed 27,331,797 triples; for Uniprot, we indexed 46,972,851 triples; and for LUBM, we...
indexed 35,612,176 triples. (The SPARQL queries used for the experiments are listed in a technical report [37].)

B. Query Workload Classification

The queries used in our evaluation are I/O bound in nature. Using the iostat command, we measured the I/O footprint of each query. (We dropped the file system buffer cache before running each query by issuing echo 3 > /proc/sys/vm/drop_caches.) Based on the I/O footprint, we classified the queries into two categories, small and large. Queries that were classified small had relatively smaller I/O footprint. Queries that were classified large had relatively larger I/O footprint. Table II shows the queries and their classification after running each query serially. (The block size used by the filesystem was 4096 bytes.) In addition, the serial time (on a single core) and the % CPU utilization for each query is shown. Queries that had higher CPU utilization (e.g., QU2), typically returned more results. Note that internally RDF-3X stores long string literals in a mapping dictionary. For YAGO, it maps back these ids to literals using the dictionary. For Uniprot, we had one separate RDF files, we grouped the triples from one file and and placed it in a bucket. All the files were distributed across the buckets in a round-robin fashion. For LUBM, as the generator produced separate RDF files, we grouped the triples from one file and placed it in a bucket. All the files were distributed across the buckets in a round-robin fashion.

Figure 1(b) shows the efficiency. For all three datasets, the ST model was obtained for YAGO, LUBM, and Uniprot, respectively. 47 workers), a promising speedup of 34.92, 32.74, and 32.27 for the STAR model was obtained for YAGO, LUBM, and Uniprot, respectively. Figure 1(a) shows the speedup obtained for parallel RDF query processing using Algorithm 1. On 48 cores (1 master + 47 workers), a promising speedup of 34.92, 32.74, and 32.27 was obtained for YAGO, LUBM, and Uniprot, respectively. Figure 1(b) shows the efficiency. For all three datasets, the efficiency reached close to 70% on 48 cores. The tasks were relatively fine-grained due to their small I/O footprints and were well distributed across the workers. There was effective load balancing of tasks across the workers resulting in good speedup and efficiency. (This is evident from the mean and standard deviation of the number of tasks processed by each worker.)

1) The ST Model: The query workload for each dataset consisted of queries marked small in Table II. The task pool consisted of these queries put in order and scaled by a factor of 100. (For example, the task pool for YAGO consisted of queries QU1, QU2, QU3, QU4, ..., QU1, QU2, QU3, QU4, ...). Figure 1(a) shows the speedup obtained for parallel RDF query processing using Algorithm 1. On 48 cores (1 master + 47 workers), a promising speedup of 34.92, 32.74, and 32.27 was obtained for YAGO, LUBM, and Uniprot, respectively. Figure 1(b) shows the efficiency. For all three datasets, the efficiency reached close to 70% on 48 cores. The tasks were relatively fine-grained due to their small I/O footprints and were well distributed across the workers. There was effective load balancing of tasks across the workers resulting in good speedup and efficiency. (This is evident from the mean and standard deviation of the number of tasks processed by each worker.)

TABLE II

<table>
<thead>
<tr>
<th>Query</th>
<th>Dataset</th>
<th>I/O footprint</th>
<th>Type</th>
<th>% CPU</th>
<th>Serial time</th>
</tr>
</thead>
<tbody>
<tr>
<td>QU1</td>
<td>YAGO</td>
<td>14,756 KB</td>
<td>small</td>
<td>29</td>
<td>4.73 secs</td>
</tr>
<tr>
<td>QU2</td>
<td>YAGO</td>
<td>15,004 KB</td>
<td>small</td>
<td>40</td>
<td>9.23 secs</td>
</tr>
<tr>
<td>QU3</td>
<td>YAGO</td>
<td>22,832 KB</td>
<td>small</td>
<td>29</td>
<td>6.51 secs</td>
</tr>
<tr>
<td>QU4</td>
<td>YAGO</td>
<td>35,492 KB</td>
<td>small</td>
<td>21</td>
<td>9.27 secs</td>
</tr>
<tr>
<td>QU5</td>
<td>YAGO</td>
<td>216,564 KB</td>
<td>large</td>
<td>42</td>
<td>74.55 secs</td>
</tr>
<tr>
<td>QU6</td>
<td>YAGO</td>
<td>272,848 KB</td>
<td>large</td>
<td>30</td>
<td>120.08 secs</td>
</tr>
<tr>
<td>QU7</td>
<td>YAGO</td>
<td>332,944 KB</td>
<td>large</td>
<td>43</td>
<td>218.83 secs</td>
</tr>
<tr>
<td>QL1</td>
<td>LUBM</td>
<td>2,668 KB</td>
<td>small</td>
<td>25</td>
<td>1.4 secs</td>
</tr>
<tr>
<td>QL2</td>
<td>LUBM</td>
<td>3,132 KB</td>
<td>small</td>
<td>35</td>
<td>1.47 secs</td>
</tr>
<tr>
<td>QL3</td>
<td>LUBM</td>
<td>9,804 KB</td>
<td>small</td>
<td>19</td>
<td>5.5 secs</td>
</tr>
<tr>
<td>QL4</td>
<td>LUBM</td>
<td>636,204 KB</td>
<td>large</td>
<td>32</td>
<td>299.99 secs</td>
</tr>
<tr>
<td>QL5</td>
<td>LUBM</td>
<td>67,392 KB</td>
<td>large</td>
<td>29</td>
<td>286.58 secs</td>
</tr>
<tr>
<td>QU1</td>
<td>Uniprot</td>
<td>4,468 KB</td>
<td>small</td>
<td>39</td>
<td>2.98 secs</td>
</tr>
<tr>
<td>QU2</td>
<td>Uniprot</td>
<td>10,344 KB</td>
<td>small</td>
<td>39</td>
<td>6.46 secs</td>
</tr>
<tr>
<td>QU3</td>
<td>Uniprot</td>
<td>48,020 KB</td>
<td>large</td>
<td>31</td>
<td>19.39 secs</td>
</tr>
<tr>
<td>QU4</td>
<td>Uniprot</td>
<td>62,188 KB</td>
<td>large</td>
<td>19</td>
<td>15.48 secs</td>
</tr>
<tr>
<td>QU5</td>
<td>Uniprot</td>
<td>166,808 KB</td>
<td>large</td>
<td>17</td>
<td>43.51 secs</td>
</tr>
</tbody>
</table>

D. Data Partitioning Approach

For the data parallel programming model, we partitioned a dataset depending on how many cores were available to run the workers. (Note that partitioning was done once before executing all the queries.) Each worker was assigned one partition and used the index for that partition during query processing. Different approaches were followed for each of the three datasets. The main goal was to assign the triples corresponding to weakly connected directed subgraphs in the RDF graph into buckets. For LUBM, as the generator produced separate RDF files, we grouped the triples from one file and placed it in a bucket. All the files were distributed across the buckets in a round-robin fashion. For Uniprot, we had one single XML/RDF file, and we created fragments of this XML file at points where a new protein was described. The triples from each fragment were stored together in a bucket. All the fragments were distributed across the buckets in a round-robin fashion.

The YAGO2 dataset was available in N-Triples format. First, we extracted graphs of a particular type from the dataset, which we call star-shaped graphs. A star-shaped graph is a weakly connected directed graph, where the degree of all vertices except one is exactly 1. All the triples from a star-shaped graph were put into a bucket. On the remaining non-star graphs, we ran the METIS [32] algorithm to partition the graphs. After obtaining n partitions, we assigned the triples for each partition into one bucket. (We replicated the cut edges in each partition.) As mentioned earlier, our approach may miss results.

E. Results

We focus on four possible combinations of workload and parallel programming models, namely, ST (small I/O footprint, task parallelism), LT (large I/O footprint, task parallelism), SD (small I/O footprint, data parallelism), and LD (large I/O footprint, data parallelism). We will refer to these as the ST, LT, SD, and LD models in subsequent discussions. Note that all I/O requests go through the MCPC connected to the SCC platform via the PCIe bus. We measured wall clock time by ensuring a cold cache scenario. (We dropped the file system buffer cache before a query was executed on a core.)

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worker as shown in Figures 2(a) and 2(b).) As shown in Figure 4, the average CPU utilization varied marginally (from 2 to 48 cores), indicating negligible I/O contention in ST.

2) The LT Model: The query workload for each dataset consisted of queries marked large in Table II. Similar to ST, the task pool consisted of these queries put in order and scaled by a factor of 33, 50, and 33 for YAGO, LUBM, and Uniprot, respectively. Figures 1(c) and 1(d) show the speedup and efficiency for parallel RDF query processing using Algorithm 1. On 48 cores, the speedup ranged between 25 to 30 for the three datasets. This is promising given that the queries had larger I/O footprint than those used in the ST model. The load was fairly well distributed across the workers. (See Figures 3(a) and 3(b).) As shown in Figure 4, the drop in the average CPU utilization (from 2 to 48 cores) was higher for LUBM and Uniprot as compared to YAGO, indicating higher I/O contention for these datasets.

3) The SD Model: The query workload for each dataset consisted of queries marked small in Table II. Each query was run multiple times using Algorithm 2. Although the data parallel approach created fine-grained tasks for a query with increasing number of cores, there was load imbalance as many of the workers returned no results on their partitions. This resulted in poor speedup and efficiency as the number of cores was increased. We show the plots in Figures 5(a) and 5(b).

4) The LD Model: The query workload for each dataset consisted of queries marked large in Table II. Each query was run multiple times using Algorithm 2. As more cores were used to process a query, I/O contention became an issue. This is evident from the fact that the average CPU utilization for LD was lower than that for LT on all datasets. As a result, poor speedup and efficiency were obtained. We show the plots in Figures 5(c) and 5(d).

F. Summary of Results on the SCC

- The task parallel programming model yielded good speedup and efficiency for parallel RDF query processing. This was true for both small I/O and large I/O footprint queries. The ST model, however, gave better results than the LT model.
- Although the data parallel programming model created fine-grained tasks, the speedup and efficiency for both the SD and LD models were poor due to either load imbalance or I/O contention. Further research is necessary to address these issues.

V. CONCLUSIONS AND FUTURE WORK

We have presented an early study of the performance of parallel RDF query processing on the Intel SCC, an experimental manycore processor. Using real and synthetic RDF datasets, we studied how inter-query parallelism (via
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