Pipelined Workflow in Hybrid MPI/Pthread runtime for External Memory Graph Construction

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Abstract—Graph construction from a given set of edges is a data-intensive operator that appears in social network analysis, ontology enabled databases, and, other analytics processing. The operator represents an edge list to what is called compressed sparse row (CSR) representation (or sometimes in adjacency list, or as clustered B-Tree storage). In this work, we show how to scale CSR construction to massive scale on SSD-enabled supercomputers such as Gordon using pipelined processing. We develop several abstraction and operations for external memory and parallel edge list and integer array processing that are utilized towards building a scalable algorithm for creating CSR representation.

Our experiments demonstrate that this scheme is four to six times faster than currently available implementation. Moreover, our scheme can handle up to 8 billion edges (128GB) by using external memory as compared to prior schemes where performance degrades considerably for edge list size 26 million and beyond.

I. INTRODUCTION

Web click stream, online social networks, semantic web etc. are some of the domains where inherently graph structured data is produced as a stream of edges or triples. Consider, for example, a set of tuples (product, customer) produced from a high traffic e-commerce website indicating the products viewed by its customers in last hour. Collection of pairs (source, destination) generated within a Twitter network that indicate message exchanges between its users is another scenario where edge list arises. Many large data stores are maintained in Resource Description Framework (RDF) format which represents data as collection of edges. Such data stores are often very large. As an example DBpedia consisting of approximately 2 billion RDF triples returns a large collection of triples.

In all these scenarios the edge or triples data set is best converted into graph representation, i.e., as an adjacency list, compressed sparse row format (CSR), or clustered B-Tree indexed storage. This conversion allows for efficient graph analysis and navigation over the received data. For example, a market analyst would be interested in finding the most connected bipartite graph of product and customers. In Twitter dataset we may want to know the most influential twitterer (a person whose tweet diffused the most) in last one hour. An “informatican” (domain knowledge engineer) would like to perform guided navigation on the graph representation of the returned RDF triples. Such a navigation can either be performed using variations of Sparql such as nSparql, or pSparql [12] (and references therein) or through hard coded algorithms for finding graphs that are homeomorphic to a given pattern [3], [14].

One of the most important use of conversion from edge list to graph representation arises in the world of Ontology-enabled databases[2], [4], [19]. Such databases are popular in medical and life-sciences domains. They have two distinct but connected parts: the ontology and annotation component and instance and experiment data component. The ontologies and annotations describe the metadata (and their relationships) while instance data consists of entities, experimental readings, and their relationships. A typical query in such databases selects a subgraph from the metadata component. The returned sub-graph is the set of nodes from the instance domain which have links from the selected metadata subgraphs. Further processing on the returned sub-graph is a very common step. For example, a user may select all “Gaba-ergic neurons” and its sub-type in the metadata component. The system then returns all the publications, experimental findings, biological entities that are annotated/have connection from at least one of the metadata nodes. Following this, a user may request to find the most “central” publications or perform path pattern query on the returned node. Again, it is efficient to build the graph representation over the selected sub-graph to answer such queries efficiently.

The problem of scalable algorithms for edge list to graph construction (either in external memory or main memory) has been studied less frequently in literature. Most RDF processing toolkits primarily provide sequential, main memory or relational table based storage and retrieval of RDF triples [24], [13], [17], [8]. To the best of our knowledge, we do not know of any parallel (shared memory) implementation of RDF-toolkit. Some of the RDF libraries provide techniques to build the graph on external memory (disk). In most work the edges are stored as a relational table. A B+-tree index structure, primarily over the source field, is constructed as well. Among the set of existing tools, B+-tree is the most efficient method for external memory graph representation of a given edgelist. In [23] showed that B-Tree index does not work well for graph style workload.

Customized storage mechanisms have also been proposed for RDF triples [23]. These mechanisms primarily focus on optimizing queries expressed using Sparql language. Such
queries are conjunctive selection and join predicates and do not express navigation or higher order graph operations such as BFS, centrality etc. Distributed RDF processing, especially using the map-reduce framework, has been an active area of research [7, 16, 13, 21]. Here again the focus has been on expediting queries expressed under the restricted framework of Sparql. In addition to these, several distributed general purpose graph processing engines have been proposed recently [9], [11]. Although each of these scheme has some mechanism for graph ingestion, the focus is primarily on optimizations for processing of graph operators such as BFS, pagerank, centrality etc and not on edge list to graph construction.

The Parallel Boost Graph Library (PBGL) [6] addresses edge list to CSR construction in distributed settings. The library provides several distributed mechanisms for constructing the CSR representation of graph from edgetlist. However, with the exception of one, all the rest of proposed mechanisms require that edges be present on all compute nodes and require additional temporary main memory storage. A mechanism is also provided that does not have the requirement and can handle edges being distributed across the compute-nodes. This scheme performs in-place construction of graph representation (CSR) if the edges are represented with two vectors  and . i.e., pair  is an edge for any index . This scheme can handle much larger edge lists than prior scheme but it induces too many memory swaps thereby exponentially increasing the processing time. A shared memory parallel graph library is presented in [10]. However, it does not address the problem of graph construction in parallel. Once the graph is constructed it provides routines for several graph operations such as BFS, connected component etc.

In this paper, we consider a hybrid distributed+shared memory out-of-core approach for building graph representation from a collection of edge lists. Specifically, we consider the problem of building distributed graph representation of a given set of edges (or triples) in the MPI/Pthread framework when the edge list and graph representations are maintained in external memory (preferably in SSDs). To the best of our knowledge this variation of problem has not been studied before. On a single laptop we can process 32GB graph in less than 1400 secs and 128GB graph in 2400 secs. The primary contribution of our approach is that it alleviates the limitation on main memory and can scale to massively large number triples (2 billion edges) and beyond. With our scheme the only true limitations are the number of cores available per CPU, the bandwidth and IOPS of the external memory, and, the communication cost associated with the interconnect fabric.

All these aspects are growing very fast in the industry. Multi-core cpus with 32-128 cores and potentially 1000s of cores would be possible on modest enterprise class CPUs.
while the bandwidth and IOPS of modern SSDs and phase change memory devices are upwards of 5600 MB/s and 1.2M IOPS (assuming 8K block size) respectively. The interconnects costs too has driven down significantly. These trends together with our distributed algorithm imply that in near future massive scale graph (64 billion edges) processing would be feasible even on modest size computers (32–64 compute nodes).

**Contribution:** Our contribution in this paper are as follows:

- We propose a set of generic abstractions and operators for data manipulation and movement over out-of-core (external, SSD) storage and network subsystem.
- We provide efficient implementation of these abstractions within our framework which consists of clusters of solid state enabled machines connected over high bandwidth low-latency network.
- We propose a buffered network reader, a novel scheme which allows us to pipeline operators in the hybrid MPI/Pthread communication framework without any additional synchronization overhead. Pipelining operators is crucial because it avoids the need for reading and writing massive amounts of data in external memory. An additional benefit of pipelining is that it can produce results early (and possibly in sorted order) which in same cases has been shown to aid further optimization.
- We design a scheme for building distributed CSR representation. The scheme exploits pipelining, SSDs, multicores, and, fast interconnect to scale this operation to billions of edges. Figure 1 shows the pipeline stages and the communication channels between them for constructing distributed CSR representation. Figure 2 shows the flow of messages across the stages in the pipeline. We see a workflow with smoothed movement of data.
- Our experiments show that the scheme is 4 –6 times faster than standard implementation and that it can handle 10 time bigger size edge list. Our experiments identify limitations with the current hybrid MPI/Pthread runtime that limit the scalability to two compute nodes.

The rest of the paper is organized as follows: We begin with definitions and then formally setup the problem and our objective. We present all the abstractions, operators, and their implementation in the hybrid MPI/Pthread framework. We then describe our procedure for building distributed CSR representation using these operators. Finally, we present experiments that study the performance trade-offs and scalability of the proposed approach.

### A. Setup and Problem Definition

A cluster of nb compute nodes, each with nc cores, are connected via high bandwidth network, possibly an interconnect. A few notations used to setup the problem of building CSR representation is as follows:

- **Label:** A character string used as an identifier to an object.
- **Vertex:** A node in the graph and represented as \( u, v, w \) etc. The vertex is either identified through a label or a numeric identifier.
- **Edge:** A pair \((u, v)\).
- **Adj\((u)\):** List of vertices adjacent to \( u \).
- **Graph:** \( G = (V, E) \) represents a graph with \( n = |V| \) vertices and \( m = |E| \) edges.
- **InducedEdgelist\((V')\) :** Given a subset \( V' \subset V \), its induced edge list is the subset of edges \( E' \) whose source and target belong to \( V' \).
- **CSR\((G)\):** Compressed sparse row representation (CSR) of any graph \( G \) consists of two vectors, namely, an offset vector \((offv)\) and an adjacency vector \((adjv)\). The \( offv \) indexes into the \( adjv \) vector. The adjacency (edge) information is stored in \( adjv \) vector. The neighbors of node \( nid \) are stored as entries in \( adjv \) vector from range \([offv[nid], offv[nid + 1])\).
- **box :** A compute node in the cluster is termed as a box.
- **nb :** Number of compute nodes (boxes) in the cluster. Symbols \( box_0, box_1, \ldots, box_{nb-1} \) identify box numbered as \( 0,1, \ldots \), and \( nb - 1 \) respectively.

**Problem Definition:** We are given a collection of pair of labels (representing edge list) \( E_l = \{e_1 = (u_1, v_1), e_2 = (u_2, v_2), e_3 = (u_3, v_3) \ldots \} \). A label in this setup uniquely identifies a vertex and \( E_l \) represents an edge list that induces a graph. Our objective is to find a compressed sparse row representation of \( E_l \), i.e., first, each node \( u \) is assigned a unique numeric identifier \( nid \) from range \([0 : t] \), where \( t \) is the total number of unique labels. We call this operation as *relabeling*. Second, build two arrays, the offset vector \( offv \) and adjacency vector \( adjv \), such that for any label (or vertex) \( u \), its adjacency entries are in \( adjv \) from index \( offv[nid] \) to \( offv[nid + 1] \), if \( nid \) is the numeric identifier for node \( u \).

Furthermore, we require the CSR representation be distributed across the \( nb \) boxes. In order to achieve this, each label is mapped to a unique box. Let \( L_b \) be the set of labels at box \( b \). Each label is assigned a unique local identifier between 0 to \(|L_b|\). The global identifier of a label is a pair consisting of the box id the label is mapped to and its local identifier at that box.

The edges in \( E_l \) are distributed in accordance with the label partitioning, i.e., edges are placed on the same box on which its source is mapped. Let, \( E_{lb} = InducedEdgelist(L_b) \) denote the set of edges placed at box \( b \). We also say that edges in \( E_{lb} \) are *owned* by box \( b \). In the distributed CSR representation for the edge list \( E_l \), each box \( b \) has an \( offv \) and \( adjv \) vectors that define the CSR representation \( E_{lb} \). The entries in \( adjv \) are the global identifier of the adjacent node. Figure 3 shows relabeling and distribution performed for a sample edge list over two boxes assuming odd/even number based mapping from label to box. The local identifiers are assigned based up on the lexicographic ordering of the labels at the box. The final column shows the relabeled edges.

Finally, we observe that after relabeling, if the edge list \( E_l \) (or \( E_{lb} \), after relabeling, is sorted based upon source field then building CSR representation is straightforward as shown in algorithm [4].

Hence, we can reformulate the problem of building CSR representation as that of (a) distributing the labels, (b) assigning unique local identifiers to the labels (we term the mapping from labels to identifiers as *identifier map* (c) partitioning
Algorithm 1 build_csr(edge list el)

Require: edge list el be sorted
1: aidx = 0, offv[0] = aidx, elidx = 0, csr = 0
2: while 1 do
3:   if csr ≠ el[elidx].src then
4:     ++csr
5:   offv[csr] = elidx
6:   continue
7: end if
8: adjv[elidx] = el[elidx].des, ++elidx
9: end while

and distributing $E_i$ based upon the source field (d) relabeling destination field with the global numeric identifier e) relabeling source field local numeric identifier, and, finally (d) sorting $E_{i_b}$ at each box based upon the relabeled source identifier.

II. OUR APPROACH FOR BUILDING EXTERNAL MEMORY BASED DISTRIBUTED CSR GRAPH USING HYBRID MPI/PTHREAD PARALLEL PROGRAMMING MODEL

We now describe our approach for building the distributed CSR representation assuming an unordered collection of edge list. We begin with few definitions followed by functions (utilities) needed from operating system/runtime, such as file I/O, mmap, communication primitives etc. We then define high order operators and primitive and provide implementation using the system provided calls and utilities. Finally, the complete procedure for building distributed CSR is put together using these operators.

a) System Utilities and Functions: The list of systems calls and runtime utilities that are used by our algorithm for building distributed CSR:

- $S(\tau)$: Gives the number of bytes required to store a particular data object of type $\tau$. For example $S(int)$ is 8 bytes in our setup. Similarly, an edge requires 16 bytes as it a pair.
- $mmap(fn, offset, sz)$ : map the data in file fn starting from offset and of sz bytes in the virtual address space of the calling process.
- msg : A message, msg, is simply a sequence of blk_sz number of bytes. It is read as an array of type $\tau$. The capacity of the array depends up on the storage requirements of the element from $\tau$ i.e the capacity $C_\tau = blk_sz/S(\tau)$. Following operations are defined on msg array:
  - size(msg) : number of elements stored in the message
  - full(msg) : check if size(msg) == $C_\tau$
  - add(msg) : add an element to the message msg, if there is room
- channel : Channel identifies a single session of communications between sender and receiver. Usually, a session between sender and receiver consists of transferring a complete stream by breaking it into a sequence of messages and transferring those one by one. Each message exchange between sender and receiver is tagged with the channel identifier to uniquely qualify this session.
- send(msg, receiver, channel) : A blocking call that sends the bits stored in msg to receiver. The communication between the sender and receiver is identified by the channel variable. The receiver should post a receive command (shown next) with the same channel number and sender field set to either ANY i.e. receive from any sender, or to the specific sender in order to receive the message from this sending box.
- msg $\leftarrow$ recv(sender, channel) : The corresponding receive blocking call to receive message in this channel. In general, there can multiple simultaneously active channels between the same sender, receiver pair.
- reader : A reader is a simple encapsulation over the receive operation. We need this concept to contrast the buffered reader approach, presented in section III-B with the one without it. The encapsulation allows us to keep the design and algorithmic description of build CSR unchanged. Following operations are associated with reader object:
  - reader $\leftarrow$ init(channel) : initialize a reader to read messages on a particular channel
  - read(reader, sender) : read data from sender on the channel with which the reader is initialized with.

A. A note of presentation of abstractions and algorithms

Before we proceed to a detailed description we briefly visit the presentation style followed in the rest of the paper. Essentially, we have opted for a functional programming style for description of all our concepts and algorithms as opposed to the more frequently used imperative style of programming. The functional style is a better fit as it naturally allows us to express generic (type agnostic) operations and pipelining. Both these are crucial for our work. Furthermore, it makes the presentation of algorithms terse and highlights core computation by reducing the amount of unnecessary syntactic expressions.

However, in order to not let the functional practices come in the way of presentation we make use of functional expressions in a restricted setting. To begin with, a generic type is represented using symbol $\tau$. All the functors have arity at most three. Moreover, we only use two functional programming
specific constructs, viz., bind and apply. Bind takes a functor and an object (which can either be another functor or an object instance) and returns a new functor of lesser arity by binding the corresponding argument to the instance. This operation is denoted using symbol \( \beta \). For example, \( \text{sort}(\text{cmp}, \text{iter}) \) takes two arguments: one, a comparison function and, the other, an iterator. The expression \( \beta(\text{sort}, \text{int\_cmp}, _1) \) returns a new sorting function that has arity one, i.e., it only takes \text{iter} as an argument. The sorting (of the collection represented by \text{iter}) by this new function is dictated by the \text{int\_cmp} function. The function \( \text{apply}(F, C) \) (denoted by \( \Gamma \)) takes a single arity functor \( F \) and an iterator. It returns a new collection whose elements are derived from applying \( F \) to elements in the collection.

A few additional terms used here are as follows:

\textit{tuples:} A tuple, as usual, represents an ordered sequence of values. The function \( \text{first}, \text{second}, \text{third}, \ldots \) extract the first, second, third and so on field of the tuple.

\textit{iterators:} The collection of elements of basic data types or of tuples is represented using iterators. The interface of iterators consists of:

- \textit{init}: Takes a collection or stream and initializes it to the corresponding iterator (integer or edges)
- \textit{eos}: Tells if the iterator has reached the end
- \textit{next}: advance the iterator to next element (int or edge)
- \textit{get}: returns the current element or object
- \textit{xget}: Is derived from the get function. In returns tuple \( \langle \text{iter}, \text{get(\text{iter})} \rangle \).

Any operator, using these functions, can access every single element of the collection represented using the \text{iter}. For example to simply scan the elements the following code can be used:

\begin{verbatim}
scan(\text{iter}):  
1. while(!\text{eos(\text{iter})})  
2. \hspace{1em} x \leftarrow \text{get(\text{iter})}  
3. \hspace{1em} do some work using x  
4. \hspace{1em} next(\text{iter})
\end{verbatim}

\textit{Functors:} Following functors are used in this paper

- \textit{filter(\text{p, \text{iter})}: \text{p} is a predicate and \text{iter} is an iterator representing a collection. \text{filter} represents a new iterator consisting of elements from the collection for which the predicate holds true.
- \textit{enum(\text{iter})}: enumerates the elements present in the \text{iter} collection.
- \textit{uniq(\text{iter})}: Assuming \text{iter} represents a sorted sequence of elements, unique creates a new iterator where the duplicates are removed from the input sequence.
- \textit{seq\_nb}: An iterator that represents sequence of integer \([0 : nb - 1]\).

\textbf{B. Streams and Iterators}

We now begin with definitions of streams and their iterators designed specifically to address the problem of building scalable CSR representation on SSD-enabled cluster of compute nodes such as Gordon \cite{15}. 

\( \) (Byte) Stream: The most basic, non-trivial, type is a (byte) stream object. A stream object is a sequence of bytes and can be of two types:

- \textit{Transient}: A transient stream arrives over the network and can be read only once.
- \textit{Persistent}: A persistent stream is stored on physical media and can be read several times.

A transient (or in-network) stream is identified by \( \langle \text{source, destination, channel\_id} \rangle \). A persistent stream is addressed using attributes set \( \langle \text{file\_name, sz, off} \rangle \). Variable \text{file\_name} is the name of the file that store the stream which is of size \text{sz} bytes and starts at offset \text{off}. We define following operations on a stream:

- \textit{store}: Store a persistent or transient stream to a new location on the physical media
- \textit{load}: load the stream into the main memory (assuming sufficient memory)
- \textit{split}: split a stream into several small streams of a fixed given size

We describe several iterators and discuss their implementation in our framework. Each iterators also encapsulates an operation; for example, the \textit{in\_network} iterator encapsulates the operation of transferring a persistent stream from one box to another. Similarly, the \textit{sort\_merge\_join} iterator encapsulates a join operation over two streams. The instantiation of an iterator gives the program a handle to its execution. However, instantiation does not execute the operation. The execution of the operation happens when a scan is performed on the iterator.

In this respect the iterators can seen as non-blocking operators. Moreover, they can be pipelined which in our case turn out to be crucial for overcoming main memory constraints.

\textit{Formatted Iterators:} Formatted iterators, such as integer iterator \( \langle \text{int\_iter} \rangle \) or edge iterator \( \langle \text{edge\_iter} \rangle \), defines iterator object over the byte stream. A specialization of formatted iterators are the random access iterators. Random access iterators support two additional functions: size and get\_at. Size returns the number of elements in the iterator while get\_at returns the element at the required index. Random access iterators provide the vector/array functionality. In order to derive a random access iterator over the stream it is necessary (for performance reasons) to load the stream in memory.

- \textit{em\_stream\_iter (\text{iter\_esi})}: This iterator is used to scan elements of type \( \tau \) stored as stream on external memory. In this work, \( \tau \) can be integer or edges. The state of the iterator is defined by variables \text{cursor}, \text{curr\_blk}, \text{ptr}, \text{sz}. The entire stream is broken into blocks. Variable \text{curr\_blk} points to currently active block. A block is made active by memory mapping its content. \text{ptr} points to this memory mapped region of curr\_blk portion of stream and read as an array of elements of type \( \tau \). Variable \text{cursor} indexes into the array starting at \text{ptr} while \text{sz} represents the total size of the stream in bytes.

The working of iterator functions \text{init}, \text{eos}, \text{get}, and \text{clean} are evident from the code description. The function \text{next} advances the cursor. If it is the end of the block, then the current block is unmapped and the next block is mapped into the memory. The cursor is reset back is zero (line 2–6).

\text{init}(fn, sz, off):
1. \( \text{iter}_{\text{esi}} \leftarrow \{ \text{cursor} = 0, \text{curr}_{\text{blk}} = 0, \text{ptr} = \text{munmap}(\text{fn}, \text{ofset}, \text{blk}_{\text{sz}}) \} \)

\( \text{next}(\text{iter}_{\text{esi}}) \):
1. \( \text{cursor} = \text{cursor} + 1 \)
2. if \( (\text{cursor} \times S(\text{\tau}) == \text{blk}_{\text{sz}}) \)
3. \( \text{munmap}(\text{ptr}, \text{blk}_{\text{sz}}) \)
4. \( \text{curr}_{\text{blk}} = \text{curr}_{\text{blk}} + 1 \)
5. \( \text{ptr} = \text{munmap}(\text{fn}, \text{curr}_{\text{blk}} \times \text{blk}_{\text{sz}}, \text{blk}_{\text{sz}}) \)
6. \( \text{cursor} = 0 \)

\( \text{eos}(\text{iter}_{\text{esi}}) \):
1. if \( (\text{curr}_{\text{blk}} \times \text{blk}_{\text{sz}} + \text{cursor} \times S(\text{\tau}) < \text{sz}) \)
2. return false
3. return true

\( \text{get}(\text{iter}_{\text{esi}}) \):
1. return \( \text{ptr}[\text{cursor}] \)

\( \text{clean}(\text{iter}_{\text{esi}}) \):
1. \( \text{munmap}(\text{ptr}, \text{sz}) \)

\( \text{c) Sorted merge iterator:} \) Sorted merge iterators represent a sorted sequence of elements obtained from merging an input collection of iterators (of the same type). It assumes that the input iterators are also representative sorted sequences. A scan operation over this iterator performs a sorted merge of the input sequence.

The set \( \{ H, \text{in} \text{_iters} \} \) encapsulates the state of this iterator. \( H \) represents a heap object while \( \text{in}_\text{iters} \) is a collection of iterators over which the sorted merge operation is performed. The heap \( H \) maintains a collection of \( (\text{iter}, \text{value}) \) tuples, where the \( \text{value} \) is the current value accessed from the iterator, i.e., \( \text{value} = \text{get}(\text{iter}) \). We assume following operations on the heap object:

- \( \text{top} \) returns the top element of the heap
- \( \text{pop} \): removes the top elements and returns its copy
- \( \text{insert} \): insert a \( (\text{key}, \text{value}) \) pair in the heap

The algorithm implemented here is the canonical sort-merge, i.e., retrieve the element from the stream that has smallest elements among all the current elements of the stream. We refer to a standard textbook for further details [20].

\( \text{init}(\text{in}_\text{iters}) \):
1. \( \text{iter}_{\text{smi}} \leftarrow \{ H, \text{in}_\text{iters} \} \)
2. \( \Sigma(\beta(\text{insert}, H, \_1), \Gamma(\text{get}, \Gamma(\beta(\text{filter}, \text{eos}, \_1), \text{in}_\text{iters}))) \)

\( \text{next}(\text{iter}_{\text{smi}}) \):
1. if \( (\text{eos}(\text{first}(\text{top}(H))) \)
2. \( \text{insert}(H, \text{get}(\text{first}(\text{pop}(H)))) \)
3. return
4. \( \text{pop}(H) \)

\( \text{eos}(\text{iter}_{\text{smi}}) \):
1. if \( (\text{empty}(H)) \)
2. return true
3. return false

\( \text{get}(\text{iter}_{\text{smi}}) \):
1. return \( \text{second}(\text{top}(H)) \)

\( \text{clean}(\text{iter}_{\text{smi}}) \):
1. while \( (\text{eos}(\text{iter}_{\text{smi}})) \)
2. next(\( \text{iter}_{\text{smi}} \))

\( \text{d) in-network stream iter:} \) In-network stream iterators are used to scan sequences of elements sent by another thread running within the same or remote box. The set \( \{ \text{sender}, \text{cursor}, \text{msg}, \text{reader}(\text{channel}) \} \) encapsulate the state of this iterator. The variable \( \text{channel} \) identifies this communication, i.e., sender and receiver issue send and receive command parametrized with the \( \text{channel} \) value.

\( \text{init}(\text{channel}, \text{sender}) \):
1. \( \text{iter}_{\text{nsi}} \leftarrow \{ \text{reader}(\text{channel}), \text{cursor} = 0, \text{msg} = \text{buf}(\text{blk}_{\text{sz}}) \} \)
2. \( \text{msg} = \text{read}(\text{reader}, \text{sender}) \)

\( \text{next}(\text{iter}_{\text{nsi}}) \):
1. \( \text{cursor} = \text{cursor} + 1 \)
2. if \( (\text{cursor} == c_e) \)
3. \( \text{msg} = \text{read}(\text{reader}, \text{sender}) \)
4. \( \text{cursor} = 0 \)

\( \text{eos}(\text{iter}_{\text{nsi}}) \):
1. if \( (\text{size}(\text{msg}) < c_e) \)
2. if \( (\text{cursor} == \text{size}(\text{msg})) \)
3. return false
4. return false

\( \text{get}(\text{iter}_{\text{nsi}}) \):
1. return \( \\text{second}(\text{top}(H)) \)

\( \text{clean}(\text{iter}_{\text{nsi}}) \):
1. while \( (\text{eos}(\text{iter}_{\text{nsi}})) \)
2. next(\( \text{iter}_{\text{nsi}} \))

\( \text{e) sort-merge-join iter:} \) The join operator is required to perform the relabeling of source and destination. In this paper, we adhere to a restricted notion of join: a join between streams \( S \) and \( J \) given functions \( \text{fields}_S, \text{field}_J, \) and \( \text{join}_{SJ} \), results in a new stream whose elements are derived as follows:

\[ \text{Join}(S, J) = \{ \text{join}_{SJ}(s, j) \mid \text{field}_S(s) == \text{field}_J(j), s \in S, j \in J \} \]

Furthermore, we assume that the streams are sorted based on the join attributes, i.e., elements in stream \( S \) are ordered according to values in \( \text{field}_S \) of each element. Similarly, for stream \( J \). We now describe the sort-merge-join implementation for the resulting stream assuming this condition holds. The by-product of a sort-merge-join operation is that the resulting stream is sorted based on join field (the \( \text{fields}_S, \text{field}_J \) for stream \( S \) and \( J \), or the field corresponding to these in new stream).

The set \( \{ \text{in}_\text{iter}, \text{out}_\text{iter}, \text{join}_\text{fn}, \text{in}_\text{join}_\text{field}, \text{out}_\text{join}_\text{field} \} \) encapsulates the state of this iterator. \( \text{in}_\text{iter} \) and \( \text{out}_\text{iter} \) represents the inner and outer input (sorted) streams (or relations), i.e., streams \( S \) and \( J \). The function \( \text{join}_\text{fn} \) takes two tuples, one from each relation, to produce an output relation. In addition, comparison operators are required over the join fields. In our case it is the canonical integer or integer derived comparison operator. The following algorithm describes the textbook implementation of sort-merge-join using in our proposed framework. The two streams are read simultaneously. When the join criteria is satisfied the \( \text{join}_\text{fn} \) is called with arguments set as the current
elements of the streams. The result of join_fn produces the current element for the resulting iterator.

\[ \text{join}(\text{in}_\text{iter}, \text{out}_\text{iter}, \text{join}_\text{fn}, \text{in}_\text{join}_\text{field}, \text{out}_\text{join}_\text{field}) ; \]

1. \( \text{iter}_\text{smji} \leftarrow \{\text{in}_\text{iter}, \text{out}_\text{iter}, \text{join}_\text{fn}, \text{in}_\text{join}_\text{field}, \text{out}_\text{join}_\text{field}\} \)
2. if(!\( \text{eos}(\text{out}_\text{iter}) \))
3. while(\( \text{in}_\text{join}_\text{field}(\text{in}_\text{iter}) < \text{out}_\text{join}_\text{field}(\text{out}_\text{iter}) \))
   \( \text{next}(\text{in}_\text{iter}) \)
4. \( \text{next}(\text{iter}_\text{smji}) \): 1. \( \text{next}(\text{out}_\text{iter}) \)
2. if(!\( \text{eos}(\text{out}_\text{iter}) \))
3. if(\( \text{out}_\text{join}_\text{field}(\text{get}(\text{out}_\text{iter})) == \text{in}_\text{join}_\text{field}(\text{get}(\text{in}_\text{iter})) \))
   \( \text{return} \)
4. while(\( \text{in}_\text{join}_\text{field}(\text{in}_\text{iter}) < \text{out}_\text{join}_\text{field}(\text{out}_\text{iter}) \))
5. \( \text{next}(\text{in}_\text{iter}) \)
6. \( \text{eos}(\text{iter}_\text{smji}) \): 1. return \( \text{eos}(\text{out}_\text{iter}) \)
get(\( \text{iter}_\text{smji} \)):
1. return \( \text{join}_\text{fn}(\text{get}(\text{in}_\text{iter}), \text{get}(\text{out}_\text{iter})) \)
\( \text{clean}(\text{iter}_\text{smji}) \): 1. \( \text{clean}(\text{in}_\text{iter}) \)
2. \( \text{clean}(\text{out}_\text{iter}) \)

The generic implementation of sort-merge-join allows us to perform join over any kind of streams: in-network, external memory, sorted-merge, or, even a stream obtained from sort-merge-join operation. This generality and the iterator based implementation would be crucial towards developing efficient and pipelined algorithm for building CSR. We shall join an in-network stream (identifier map stream) with an external memory edge stream in order to relabel the edges.

C. Blocking Operators

Here, we define two communication operations over the streams. These operators behave differently than the iterator based operators where the instantiation is decoupled from execution, and therefore allows for pipelining, i.e., several iterators can be instantiated at once and chained together so that all of them execute in pipelined fashion. The blocking operators, on the other hand, are executed upon call and the calling process is blocked until the operation is complete.

\text{broadcast\ stream}: The broadcast operation takes as input a stream and sends a copy to every other box. In order to do so the stream is partitioned into blk_sz messages. Each message is broadcasted over the network. The ordering of messages is maintained due to the use of blocking compunction operation, viz. MPI_Send and MPI_Recv, from the runtime.

\( \text{broadcast\ stream}(\text{iter}) ; \)
1. \( \text{msg} = \text{buf}(\text{blk}\_sz) \)
2. while(\( \text{eos}(\text{iter}) \))
3. \( \text{add}(\text{msg}, \text{get}(\text{iter})) \)
4. if(\( \text{full}(\text{msg}) \))
5. \( \Gamma(\beta(\text{MPI}_\text{Send}, \text{msg}, _1), \text{seq}_\text{nb}) \)

\text{scatter\ stream}: The scatter stream takes an additional mapping function which maps elements of the stream to a unique box. The input stream is partitioned into nb sub streams based upon the mapping function. The sub streams are send to their respective boxes.

\( \text{scatter\ stream}(\text{iter}, \text{map}_\text{fn}, \text{channel}\_\text{id}) ; \)
1. \( M \leftarrow \{\text{msg}_1, \ldots, \text{msg}_\text{nb}\} \) where \( \text{msg}_\alpha = \text{buf}(\text{blk}\_sz) \)
2. while(\( \text{eos}(\text{iter}) \))
3. \( \text{o} \leftarrow \text{map}_\text{fn}(\text{get}(\text{iter})) \)
4. \( \text{add}(\text{msg}_\text{o}, \text{get}(\text{iter})) \)
5. \( \text{MPI}_\text{Send}(\text{msg}_\text{o}, \text{o}, \text{channel}\_\text{id}) \)
6. \( \text{reset}(\text{msg}_\text{o}) \)
7. \( \text{next}(\text{iter}) \)
8. \( \Gamma(\beta(\text{MPI}_\text{Send}, _1, 2, \text{channel}\_\text{id}), \text{seq}_\text{nb}) \)
\( \Gamma(\beta(\text{MPI}_\text{Send}, _1, 2, \text{channel}\_\text{id}), M) \)

These operators require a thread at the receiving box to collect the incoming messages via the MPI_Recv primitive. This is achieved by invoking a new thread which scans a in-network stream iterator (described earlier in section [I-B0d]) whose state variables (sender and channel) are set accordingly i.e. sender is set to the rank of the box broadcasting (or scattering) the stream and channel is set to a value that uniquely identifies this communication.

III. EDGE LIST TO DISTRIBUTED CSR : THE ALGORITHMS

We now describe the approach for building the distributed CSR representation using the proposed iterators and operators. The algorithm ties the various iterators and operators in a way that allows for pipelined distributed processing and shared memory/parallel processing.

Our approach consists of five phases: setup, assign unique local identifiers, relabel edges (destination), relabel edges (source), and finally build CSR.

1) The driver routine: The buildCSR procedure is invoked on the console with requisite parameters such as number of boxes, number of cores per box, the channel identifier for input edgelist. As with any MPI-based program, the runtime launches a process on each of the box with the same set of parameters. Each box is also assigned a unique id from the range \([0: nb]\) which is known as the rank of the process.

Each MPI process executes the phases, in that order, and in sync with other processes. It may also launch nc pthreads either for parallel processing or for pipelined processing.

2) Setup: This step collects the edges generated by external processes and sets all the system parameters accordingly. The external processes generates the edges as a transient stream. This stream is split into nc stream, one per core worker. The core worker transforms the received stream into a persistent stream by storing in a physical location.
3) Assign unique local identifier: In this phase each label is mapped to a unique box where it is assigned a unique numeric identifier. In order to do so all the labels mapped to a box are maintained at that box in sorted order.

The steps to distribute the labels across boxes and build this sorted ordering is as follows: Each box partitions the edge stream (maintained in external memory) into sub streams of size \( mnc \) bytes. Each of these sub streams are converted into integer random access iterator by loading the sub stream into the main memory and then casting into an integer iterator. The stream is then sorted in the memory and then written back to the external storage in a new location. The computational expression being:

1. \( op = \beta(\text{save}, \beta(\text{sort}, \beta(\text{int}_\text{iter}, \beta(\text{load}, \_1)))) \)
2. \( S1 \leftarrow \Gamma(op, \text{split}(S, mnc)) \)

where \( S \) is the input edge stream on the box.

Next we scatter the identifiers based upon a mapping function. The scattering is performed in a manner such that they arrive at the destination in sorted order. To achieve this, each box performs a sorted merge over streams in \( S1 \) and then calls the blocking scatter operation:

1. \( \text{scatter}(\text{sorted_merge}(S1), \text{LABEL\_SCATTER\_CHANNEL}) \)

The scatter operation at a box produces a sub-stream for every other box. Hence each box \( b \) receives \( nb \) streams, one from each scatter operation at a box (see figure 5). These streams contain labels (in sorted order) that are mapped to box \( b \). A collector thread, at each box, opens \( nb \) in-network stream iterator, to receive the in-coming streams. The collector also performs a sorted merge over these streams to arrive at a single stream of sorted labels. The pseudo-code describe the procedure for the collector thread:

1. \( R1 \leftarrow \text{sorted_merge}(\Gamma(\beta(\text{in-network_stream} \_1)), \text{seq}_{bn}) \)

Assigning unique identifier from range \([0, t]\), where \( t \) is the total number of unique labels in the resulting stream can be done efficiently. Specifically, the computational expression is \( \text{enumerate}(\text{uniq}(R1)) \). We observe that with initializing \( R1 \) we have only so far instantiated a handle on the entire step and at that we haven’t yet executed it. A scan over \( R1 \) i.e. \( \text{scan}(R1) \) will execute the step and produce a sorted stream of labels on the box.

We call the stream \( R1 \) as identifier map as it contains the mapping from labels to new numeric identifiers.

4) Relabel edges (destination field): We now proceed to relabel the destination with the new identifier i.e. the numeric identifiers for the labels determined in the previous step. If \((u, v)\) is an labeled-edge s.t. \( v \) is assigned (hashed) to box \( b \) and has numeric identifier \( nid \) then, in this step, we relabel these edges as \((u, < b, nid >)\).

We achieve this by performing a join over two streams, \( E \) and \( I \), where \( E \) represents an stream of edges ordered by destination and \( I \) represents a stream of all (identifier, label) tuple sorted by labels. The result of this join operation is a new stream of edges with destination relabeled to new identifier i.e

\[
\text{join}(E, I) = \{(u, \langle \text{bid}, \text{nid} \rangle) | (u, v) \in E, \quad (v, \langle \text{bin}, \text{nid} \rangle) \in I \}
\] (1)

Since both the streams are sorted on join field we can obtain \( \text{join}(E, I) \) by perform sort-merge-join over the streams \( E \) and \( I \) i.e.

\[
\text{join}(E, I) = \text{sort_merge_join}(E, I, \text{relabel_des}, \text{second}, \text{first}) \quad (2)
\]

where relabel_des is a function with following signature:

\[
\text{relabel \_des}(e, < l, \text{id} >) :
1. \text{return} (e.\text{src}, \text{id}) \text{ if} e.\text{des} == l
\]

The algorithm below describe the procedure of building stream \( E \). The local edgelist is partitioned into a collection of streams each of size \( mnc \) (represented by function chunk_partition). This collection is further divided among the participating cores. Each core then sorts the streams assigned to it and materializes at a new location. A sorted merge over these streams results in \( E \) i.e. a stream of edges sorted by the destination field.

1. \( C \leftarrow \text{chunk\_partition}(\text{split}(S, mnc), \text{nc}) \)
2. \( op = \beta(\text{save}, \beta(\beta(\text{sort}, \text{des\_comp} \_1), \beta(\text{edge\_iter}, \beta(\text{load}, \_1)))) \)
3. for each thread, \( \text{tid}, \text{where} \text{tid} \in [0 : \text{nc}] \)
4. \( S2 \leftarrow \Gamma(op, C[\text{tid}]) \)
5. wait for all threads
6. \( E \leftarrow \text{sorted\_merge}(S2) \)

We observe that stream \( I \) can be obtain by performing sorted merge over the identifier maps derived in the previous “assigned unique identifier” phase. However, we cannot perform sorted merge directly over the streams since they are distributed across the boxes. In order to bring all the identifier map stream together (on every box) each box broadcasts its identifier map stream to all other boxes. Boxes also launch another thread to receive the identifier maps. It then performs a sorted merge over these streams to arrive at stream \( I \). The computational expression to derive \( I \) is

1. \( I \leftarrow \text{sorted\_merge}(\Gamma(\beta(\text{in-network_stream} \_1), \text{seq}_{nb})) \)

Hence the computation of this step on each box is distributed across two threads: first thread receives the sorted labels and performs sorted merge to build identifier map. It also broadcast the identifier maps. The second thread performs the sort-merge join over stream \( E \) and \( I \) which in turn receives all the identifier maps. Figure 2 illustrate this execution on a box. The first thread performs sorted merge and broadcasts the identifier map (inside boxes shown in top right corner). The
second thread’s operation is illustrated in detail. The sorted-
merge edge stream (from below) and sorted-merge identifier 
map stream (from right) are joined together to produce a 
relabeled edge.

5) Relabel edges (source field): We now relabel the source 
field of the edges. This step is similar to the step of relabeling 
destination with minor differences: sort the edges (this time 
based on source field), launch thread to broadcast identifier 
maps, perform sort merge join over two streams: One obtained 
by sorted merge of edge stream and another obtained by sorted 
merge of identifier stream. As opposed to previous step, in this 
step we only derive the handle for sort_merge_join style algorithm, 
i.e., derive the iterator but do not perform the scan operation 
over this iterator. The scan operation is performed in the next 
sort edge phase.

A. Scatter relabeled edge and build CSR

In this step, we redistribute the relabeled edges obtained in 
the previous step so that each edge is stored at its owner box. 
Additionally, the edges received at a box during the scatter 
operation are sorted based upon the new identifiers. This is 
the because the new identifiers are assigned in the ascending 
order of the labels and that we use sort-merge-join style algorithm 
for relabeling. This expedites building the CSR representation.

Each box performs scatter(S) operation. This operation 
breaks the relabeled edge (source field) stream into sub-
streams, one for each compute box. Edges arriving at a box 
are owned by that box. Furthermore, as explained earlier they 
are also sorted based upon the new identifier. Hence, in order 
to derive the desired edge stream on each box we only need 
to additionally perform a sorted merge over the incoming 
edge stream i.e. in a separate thread we execute the following

\[
R \leftarrow \text{sorted_merge}(\Gamma(\beta(\text{in\_network\_stream,} \\
\text{edge\_scatter\_channel,}_, 1), \text{seq}_m)) 
\]

B. Avoiding communication deadlocks

The above description although algorithmically correct does 
have the problem of reaching a deadlock state, i.e, threads may 
form a circular dependency of waits using the blocking MPI 
communication primitives MPI_Send and MPI_Recv.

We show this through a simple setup consisting of two boxes 
each having two threads one for sending data and the other for 
receiving. All the threads are communicating over the same 
channel. Figure 5 illustrates this dependency cycle. The arcs 
of this dependency cycle are:

1) Receiver on box 1 is blocked after it has issued an 
   \text{MPI\_Recv}(\text{sender} = 1) command, i.e, it is waiting 
   for sender on box 1 to respond. Hence, we draw an arc 
   from the receiver (on box 1) to the sender (on box 1).

2) The sender on box 1 is blocked after it has issued an 
   \text{MPI\_Send}(\text{receiver} = 0) command, i.e., it is waiting 
   for receiver on box 0 to receive the message. Hence, we 
   have an arc from sender on box 1 to receiver on box 0.

3) The receiver on box 0 has posted a 
   \text{MPI\_Recv}(\text{sender} = 0) command, and therefore blocked. 
   Hence, we have an arc from receiver on box 0 to sender 
   on box 0.

4) The sender on box 0 is waiting for reviver on 
   box 1 in order to complete the communication call 
   \text{MPI\_Send}(\text{receive} = 1). Hence, there is an arc from 
   sender on box 0 to receiver on box 1.

These dependencies together represent a deadlock situation 
where the algorithm makes no further progress.

We present an enhancement to our scheme to break the 
circular dependency without changing its architecture and 
methodology. We introduce a buffered reader object that reads 
messages on behalf of in network streams. This reader is 
shared by all the in_network streams on a box having the 
same channel.

Below we show the complete implementation of the 
buffered reader. The state of buffered reader is determined 
by variables channel, msg_queues, msg_free_pool, 
and msg_allocated. The reader serves the request to read 
the next message from stream originating from sender s as
follows: it maintains a first in first out (FIFO) queue of messages per sender. If the queue for sender $s$ is not empty then it simply pops a message from the sender’s queue and returns it. However, if the queue is empty then then the reader issues repeated $MPI_{\text{Recv}}(sender = \text{ANY})$ calls until it receives a message from the sender $s$. This message is returned to the caller. The rest of the received messages are placed in the queues corresponding to its sender.

$$init(channel, msg\_queues, msg\_free\_pool, msg\_allocated);$$

1. reader $\leftarrow \{channel, msg\_queues,$
   $msg\_free\_pool, msg\_allocated\}$

2. $msg\_allocated[seq, nb] = \text{NULL}$

$read\_msg(reader, sender);$:

1. if($msg\_allocated[sender])
2. push(pool, msg\_allocated[sender])
3. $msg\_allocated[sender] = \text{NULL}$
4. if($msg\_queues[sender].empty())
5. $msg \leftarrow msg\_queue[sender].pop()$
6. $msg\_allocated[sender] = msg$
7. return $msg$
8. while(1)
9. if($pool.empty())
10. $msg = pool.pop()$
11. else
12. $msg = init()$
13. source $\leftarrow MPI_{\text{Recv}}(msg,$
   $MPI_{\text{ANY\_SOURCE}}, channel)$
14. if(source $== sender$)break
15. $msg\_queues[source].push(msg)$
16. return $msg$

IV. Operation Pipeline in MPI/Pthread Primitive

Pipelined processing is an important optimization principle in any data processing system. Our choice of data types (streams and its variants), operators (sorted-merge, sort-merge-join, broadcast, and, scatter) and the algorithms for various phases of building CSR are designed to make pipeline processing feasible. We study our proposed approach as a data workflow and show how MPI multi-threaded interface can be used towards pipelined processing without incurring any additional synchronization overheads.

The entire computation for building CSR on a box can be decomposed into stages. A stage is defined by a group of threads distributed across the boxes performing the same step of computation. All the stages/threads are active simultaneously and communicate with each other through the data distribution operator (scatter and broadcast) and buffered reader operator on different channels.

Figure 1 shows the various stages and the channels over which the distribution and reader operators communicate in our proposed approach. The pipeline stages (in that order) are merge and redistributed labels, build-and-broadcast idmap, relabel and scatter edges, and, merge-and-build-csr. The threads in the first and second stages communicate over the $\text{LABEL\_SCATTER\_CHANNEL}$, threads in second and third stage communicate over $\text{IDMAP\_BROADCAST\_CHANNEL}$ and those in third and last stage communicate over $\text{EDGEL\_SCATTER\_CHANNEL}$.

All the data-distribution threads interfacing a channel communicate with all the readers plugged to the channel. Hence the communication pattern at any channel is a complete bipartite graph $K_{nb,nb}$ as shown in figure 6.

A general method for pipelining operations emerges from this discussion. If the operation can be decomposed into sequence of stages such that the input iterator of a stage comes from the output iterator of the previous stage then it can be implemented in the MPI/Pthread framework using the distributed operators and buffered readers such that it allows for pipelined distributed computing.

The added advantage of the pipelining using our scheme is that it eliminates the need for synchronization between threads executing the stages. Pipelining in multi-threaded environment invariably require synchronization since the data is produced (output as result) by one thread and is consumed (as input) by another thread. Hence, these two threads are in producer consumer relationship and need to synchronize the access to the concerned data [5]. In our approach the threads performing each stage of the computation do not need perform any explicit synchronization. All synchronization needs are handled transparently through the MPI multi-threaded environment and the blocking send/recv calls.

V. Experiments

We conduct experiments over the proposed edgelist to CSR converter to study the impact of pipelining and overheads due to shared memory and distributed computing on its performance. In addition, we also compare our algorithm with that of the standard distributed edgelist to CSR routine provided in the parallel boost graph library (PBGL). PBGL is distributed graph library build on the Boost framework which is a collection of data structures, algorithms, system utilities, and advanced language primitives. Since PBGL is a well used library and has an (unoptimized) standard implementation of edgelist to CSR implementation we consider it to be a good reference point for comparison.

All experiments are conducted on Gordon supercomputer [15] Each compute node contains two 8-core 2.6 GHz Intel EM64T Xeon E5 (Sandy Bridge) processors and 64 GB of DDR3-1333 memory. The compute nodes mount a single 300 GB SSD (280 GB usable space). The latency to the SSDs
is several orders of magnitude lower than that for spinning disk (\(<\) 100 microseconds vs. milliseconds).

In our experiments we used synthetic random and scale-free graph edgelist generator. We also varied various algorithm parameters such as scale, edge factor, block size, \(mnc\), and the number of boxes. The algorithm description so far had assumed one MPI process per box. However, it is possible to launch several MPI process on a single box. We compare performance in both the settings: multiple MPI process on a single box vs. multiple boxes each with one MPI process.

In all the experiments we use the random edge generator unless otherwise specified. The edge factor is set to 8. Figure 7 shows performance for a single mpi process for various scale edge list varying the block size. The edge factor is fixed to 8. We see that the system can gracefully handle a scale 28 edge list. The total storage cost of this graph 32 GB. The total communication cost is determined by both the size of edge list and the blk_sz. At larger block size fewer messages are exchange but each message exchange is of longer duration. We see that the system favors smaller blk_sz and that at large blk_sz the system performance deteriorates considerably.

The reason why higher blk_sz is disadvantageous in our algorithm lies in the implementation of the hybrid MPI/pthread scheme. Currently, this scheme is implemented using global lock which serializes all communication, i.e., only one MPI call can be active at any given instance. When the blk_sz is large a single MPI call continues for a long duration. The rest of threads, after finishing their local work, block and wait for this MPI thread to complete. This leads to a scenario where the rest of the threads stall for longer period of time and therefore suffer decreased performance. This aspect hinders the performance and scalability of our system considerably. However, we conjecture that the current implementation of hybrid MPI/pthread is sub-optimal and in future implementations in which a single active MPI call does not block rest of the calls the algorithm will scale with respect to large blk_sz as well.

In figure 8(a) we study the strong scaling aspect of the algorithm on a single box, i.e., we build CSR representation for scale 28 with varying number of processes on a single box. The algorithm achieves scalability until 4 MPI process and performance drops when the number of processes is increased to 8. This is because at 8 MPI process amounts to 32 active threads on a single node. In this configuration we have over subscribed the number of threads since the compute node only has 16 cores and thus reducing the performance.

We also conduct experiment varying the number of boxes for a fixed scale graph. In this setup, there is one MPI process per box. We observe a small improvement with two boxes. As mentioned earlier, due to high overhead associated with communication in MPI/pthread runtime we do not see strong scaling across the number of boxes. Our hope is that with future improvements to MPI/pthread better scalability is possible.

Although due to communication costs our scheme currently does not scale beyond two nodes or two MPI processes, we show that it has good scalability with respect to scale parameter when compared with the standard algorithm present in PBGL. First, we note that irrespective of the number of nodes used the boost graph algorithm cannot scale beyond scale 26 graph. On the other hand our scheme can generate scale 30 graph on single node and even larger using multiple nodes if communication overheads can be tolerated. We see that PBGL’s time graph quadratically with the graph size due to the use sort function on the entire edge list and non-pipelined implementation. Our scheme on the other hand shows much better scalability with respect to increase in scale. We attribute this primarily to sort-merge-join and pipelined implementation.

Figure 10 compares performance for the first phase of the buildCSR routine, i.e., the sort labels and sort edges, which do not require communication across MPI process. It compares the performance for multiple MPI process on single box vs. multiple boxes, one MPI process per node. As we increase the
number of processes on a single node due to limited resources
(cores, bandwidth on SSDs etc) the performance flattens out.
However, as there is proportionally more resources on latter
setup we see increased scalability.

Figure 2 illustrates the pipeline processing of the algorithm.
We ran a single MPI processing for scale 20 edge list. In
this run every send and receive message is an event that is
depicted as upper or lower triangle respectively. Color denote
the channel of communication. We see that the messages
across channel are interleaved together implying a smooth
pipelined processing.

VI. Conclusion and Future Work

We have shown a pipelined algorithm for constructing
distributed graph representation from a given collection of
dependent list. Our implementation is based up on the hybrid
MPI/pthread runtime that allows for multiple threads to simul-
taneously invoke communication calls. Through experiments we
demonstrate good scalability and 4-6 times speedup on
single and two compute nodes when compared with standard
PBGL implementation. We highlight the deficiency of the
current implementation of the MPI/pthread runtime which hin-
ders scalability beyond two compute nodes. We are currently
investigating the use of asynchronous I/O and other runtimes to
bypass this limitation. Our scheme has implication on general
data-intensive computing that we can currently investigating.

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