ABSTRACT

Simultaneous BVH traversal, as a dynamic task of pair-wise proximity tests, poses several challenges in terms of parallelization using GPUs. It is a highly dynamic and data-dependent problem which can induce control-flow divergence and inefficient data-access patterns. We present a simple solution using the bulk-synchronous parallel model to ensure a uniform mode of execution, and balanced workloads across GPU threads. The method is easy to implement, fast and operates entirely on the GPU by relying on a topology-centred work expansion scheme to ensure large concurrent workloads. We demonstrate speedups of up to 7.1× over the widely used "streams" model for GPU based parallel collision detection.

CCS CONCEPTS
• Computing methodologies → Collision detection; Massively parallel algorithms; Shared memory algorithms;

KEYWORDS
collision detection, parallel computing, GPU, BVH, BSP

ACM Reference Format:

1 INTRODUCTION

Collision detection (CD) has a wide-spectrum of practical applications including physics based simulations, robotic motion planning, virtual disassembly, haptic rendering and ray-tracing. It is a well known and long studied problem of finding a number of interactions at low computational cost. As a result, collision detection is at the core of many applications in computer science and engineering today.

However, CD can be computationally expensive due to its potential for having vast workloads. A simple approach exhaustively testing for pair-wise intersection between geometry will not scale optimally due to the inherent $O(N^2)$ complexity. This particular constraint has lead to the common solution of using acceleration data structures such as bounding volume hierarchies (BVHs) [Ericson 2005]. BVHs attempt to reach an optimal case of $O(\log N)$ by quickly culling the search-space of potential collisions.

In spite of this potential benefit, geometry may reach scales of tens- to hundreds-of-thousands of triangles or more which makes the prospect of employing BVHs alone insufficient. Furthermore, BVHs can also degenerate if the enclosed geometry is relatively small, such that traversing entire BVHs becomes a layer of overhead.

Previous methods tackling the problem of optimizing BVH based CD on the GPU [Du et al. 2015; Lauterbach et al. 2010; Tang et al. 2011, 2016] offer in-part successful but also complex solutions which can suffer from GPU under-utilisation. They emulate the logic of conventional single-threaded CPU traversal by relying on thread-level private work-stacks and temporal coherence [Li and Chen 1998]. Such heuristics-based optimisations can serve to complicate traversal logic and thus may constrain GPU performance. Work-stacks serve to reduce memory access and synchronisation costs. However, they are a source of control-flow divergence, and load-imbalance which is managed by a separate GPU task in the execution pipeline. Moreover, pre-existing solutions have used work-stacks to effectively mimic recursion on the GPU because threads perform traversal in-place. Evaluation of pair-wise tests in BVH sub-trees is computed independently as threads push and pop intermediate BVH node-pairs to-and-from work-stack memory which creates the divergence in control-flow as a side-effect. Temporal coherence on the other hand, has a high memory footprint since it is based on explicitly storing the BVH node-pairs where traversal terminates. It is also a potential source of work-flow divergence because simply checking when and how to store such node-pairs contributes to the overhead of branching on GPUs.

We present a simple approach to simultaneously traverse a large number of BVHs for CD in parallel and entirely on the GPU. The method is based on the Bulk-Synchronous Parallel (BSP) model [Valiant 1990] were traversal is reformulated as an iterative “fork and join” scheme to: (1) mitigate explicit load-balancing that requires using separate work-rebalancing tasks on the GPU, (2) minimise control-flow divergence by reducing the amount of work mapped to each thread and performing full-restarts from a user-specified entry level such as the root-level, and (3) allow for efficient memory access patterns that may be coalesced while seamlessly unifying synchronisation, communication and storage by relying on the BSP model. Our tests, which are performed on three UNC dynamic scene benchmarks (see Figure 6), also reveal up to 7.1× speedup over the “streams” model for GPU based CD from Tang et al. [2011], which is currently the standard model employed by others [Du et al. 2017, 2015; Tang et al. 2013, 2016].

Contributions. The contributions form a simple solution, from using the topological structure of BVHs and simplified thread-level operations for reducing control-flow divergence, to efficiently traverse multiple BVHs in parallel on the GPU.
• We present a novel algorithm (section 5) as alternative re-formulation of simultaneous and parallel traversal of multiple BVHs for pair-wise CD on the GPU.

• Parametric workload expansion (subsection 5.2): Adaptive depth-stepping and static workload expansion are introduced as key features for ensuring large concurrent workloads and controlling the rate of traversal, using the topological structure inherent in the traversed BVHs.

• A lock-free scheme to write intermediate BVH node-pairs to global memory using iterative buffered-writes (subsection 5.3), which can be controlled based on the topological properties of BVHs and available hardware resources.

2 RELATED WORK

Collision Detection in Physics-based Animation. Collision detection lends itself well to physics-based simulation problems for real-time and off-line use-cases [Ericson 2005]. It has been particularly useful for large scale problems involving complex non-rigid objects such as cloth [Bridson et al. 2002; Brochu et al. 2012] were the complexity of interactions (including self-collisions) places emphasis on the need for efficient culling of triangle intersection tests which have a high computational cost. BVHs are a common data structure in many such works with their ability to quickly cull of the search space of potential interactions [Teschner et al. 2005]. Numerous approaches including axis-aligned bounding boxes (AABB) [Bergen 1997], oriented bounding boxes (OBB) [Gottschalk et al. 1996], discrete oriented polytopes (k-DOP) [Klosowski et al. 1998] have been introduced for this purpose, which function as approximations to the underlying geometric primitives that they enclose in the form of coarse bounding volumes.

Parallel Collision Detection. Methods to accelerate CD through parallelism on GPUs have been investigated for over a decade now. Early pioneering works such as that of Knott and Pai [2003] made use of the parallel rasterization capabilities of GPUs. Recent methods including Tang et al. [2016] and others [Tang et al. 2011, 2013; Weller et al. 2017; Wong et al. 2014] utilize the general purpose computational capabilities of modern GPUs to accelerate computation following the advent of parallel programming frameworks such as CUDA and OpenCL. Wong et al. [2014] present a parallel adaptive scheme combining octrees and hierarchical grid structures for broad-phase CD with deformable objects. Weller et al. [2017] recently introduce a CUDA based scheme, kDet, which is based on a hierarchical grid structures to find the set of potentially colliding pairs using polygon sizes.

In general, mapping BVH traversal to GPUs is recognised as a challenging task as demonstrated by prior efforts that have advocated for the use of more parallelism through many-core GPUs and multi-core CPUs [Lauterbach et al. 2010; Tang et al. 2010, 2016]. Since naive approaches can easily result in hardware under-utilisation due to low workloads, the most influential methods such as Lauterbach et al. [2010] and other variants [Du et al. 2015; Tang et al. 2011, 2013, 2016] have relied on front tracking [Tang et al. 2010] for sustaining high workloads which is ideal for GPUs. In this approach, the bounding volume test tree (BVTT) [Gottschalk 2000] of BVH node-pairs where traversal terminates is explicitly cached and then used as input for next time. In addition, thread-level private work-stacks are another common feature in these methods, to improve memory access costs and minimise inter-thread synchronisation. However, work-stacks can lead to work-flow divergence and load-imbalance that require a separate GPU task to perform work redistribution between threads (see Lauterbach et al. [2009] and [2010] for details). Similar approaches have also been used in robotic motion planning [Pan et al. 2010; Pan and Manocha 2011]. The related work of Hermann et al. [2013] performs CD for motion planning using voxel maps maintained in GPU global memory.

Our method shares some similarities with these approaches but does not rely on work-stacks nor front-tracking. We adopt the BVTT as the primary input but distinctively express traversal as an iterative one-to-one mapping between threads and evaluated node-pairs. Further, we focus on the specific problem of pair-wise CD between BVH-nodes, whereas many of these approaches are focused on parallelizing the entire CD pipeline. Another distinction is that these previous methods have not considered a case for the ability to use the topological information of BVHs to increase workloads at faster rates, since the maximum number of BVH-node pairs created when two nodes intersect is constrained by the number of children per-node. In order to increase workloads at faster rates, these methods are required to change their BVH construction scheme and thus, traversal logic, in order to incorporate having a larger set of children per-node to speed up traversal rates.

Stackless Traversal. We also note that the presented method is not the first to adopt stack-less traversal since we share a similar design premise to Hapala et al. [2013]. In contrast, Hapala et al. have presented an iterative method for ray-tracing on CPUs and GPUs with backtracking and a state-machine to infer which nodes to process next. Barringer and Akenine-Möller [2013] present a similar stack-less approach with full restarts, while Laine [2010] encodes the traversal trial using bit information. In this paper, we instead propose an approach that is entirely GPU based and strictly forward stepping with no notion of backtracking nor state-machines that are used during traversal. We use the topological information of our BVHs, which is encoded as memory locations and offsets, to infer the BVH nodes to traverse next and additionally use this information to increase workloads at faster rates.

Data Parallel Models for Graph Processing. Parallel traversal shares many challenges with large scale graph problems on GPUs, where issues of load-imbalance (irregularity), control-flow divergence, non-coalesced memory access patterns are most common [Lenharth et al. 2016; Merrill et al. 2012]. Harish et al. [2007] present one of the earliest solutions to solve breadth first search (BFS), single source shortest path, and all-pairs shortest path, while later works such as Cederman et al. [2008] and Tzeng et al. [2010] also address issues of load imbalance at the thread-level. Aila et al. [2010] investigated the related difficulties of divergence on GPUs in context in ray-tracing. Recent work focuses on the design of general frameworks for different kinds of large graph structures on GPUs such as the scheduling model for irregular inhomogeneous workloads proposed by Steinberger et al. [2014]. Khorasani et al. [2014] present a CUDA based model focusing on minimising warp divergence by coarsening parallelism to CUDA warps. Other works have also investigated
languages and frameworks for expressing such large-scale computations. Hou et al. [Hou et al. 2008] previously present a programming language for expressing the BSP model [Valiant 1990] on GPUs by addressing the challenge of producing efficient stream code and barrier synchronization. The recent Enterprise [Liu et al. 2016] and Gunrock [Wang et al. 2016] frameworks define an iterative BFS traversal of large graphs using the BSP model similar to the influential work of Merrill et al. [2012]. In contrast, our inspired work focuses on the specific problem domain of simultaneous BVH traversal for pair-wise CD but also borrows key ideas such as GPU based parallel BFS as a building block. Further, these methods are optimised in large part for massive load imbalance across vertices (as seen in scale-free graphs), but BVHs/BVTTs do not have that kind of imbalance. So different optimisation decisions may be appropriate.

3 METHOD OUTLINE

In what follows, we refer to a pair of BVH-nodes tested for intersection as a BVTT-node and additionally refer to each such BVH-node as an entry-node. During traversal, a BVTT-node is discarded after a bounding volume (BV) intersection test, such that if the result is true, the BVTT-node is expanded by replacing it with a new subset of BVTT-nodes. This new subset is constructed by pairing the descendants of one respective entry-node with those of the other. Alternatively, pairings may be produced between either entry-node of large graphs using the BSP model similar to the influential work of Merrill et al. [2012]. In contrast, our inspired work focuses on the specific problem domain of simultaneous BVH traversal for pair-wise CD but also borrows key ideas such as GPU based parallel BFS as a building block. Further, these methods are optimised in large part for massive load imbalance across vertices (as seen in scale-free graphs), but BVHs/BVTTs do not have that kind of imbalance. So different optimisation decisions may be appropriate.

4 DATA STORAGE AND REPRESENTATION

This section describes the employed BVTT and BVH node representations which enable efficient storage and runtime access for our topologically driven workload expansion scheme described in subsection 5.2.

Figure 1: BVH nodes are stored compactly in one memory buffer (BVH storage array) with addition set of of small arrays holding metadata about each BVH which we use to infer node descendants at runtime.

**BVH Storage and Representation.** A BVH is represented as a simple index-pair where each index is a location of a BVH-node in the global memory. We refer to each index as an entry. The BVH is stored as a large contiguous array in order for threads to access GPU global memory in contiguous and aligned memory blocks [Fauzia et al. 2015] (see Figure 3). The availability of vector load/store instructions on certain GPU architectures allows for efficient bandwidth utilisation which can be beneficial since address accesses of each thread can be combined with single memory transaction issued due to the one-to-one sequential and aligned access to memory [Cook 2013; Luitjens 2013].

**BVH Storage and Representation.** We propose a novel representation and indexing scheme for BVH nodes that enables instant computation of the BVH that a node belongs to, and the descendants of this node. All BVHs are assumed to be stored compactly in a contiguous array at known offsets with the first at the zeroth offset, whereby the employed hierarchy representation is an implicit binary-tree that is full and complete with nodes stored in a Pre-order Traversal manner as shown in Figure 1. We pad each BVH by rounding the number of leaf-nodes to the nearest power-of-two to enable implicit indexing of the descendants of any node. Though padding can potentially result in a higher memory footprint, the additional storage cost is relatively low compared to, for example, the BVH memory itself, since only bounding volume information is stored per-node (its "payload"). Information referencing geometry that is associated with each leaf-node can be stored separately. Given an arbitrary entry-node i, its j-th descendant that is δ levels (δ ≥ 0) deeper than i can be inferred by

\[
c_j = \left(2^\delta n_i + 2^\delta - 1\right) + j, \quad j \in \left[0 \ldots 2^\delta\right]
\]  

where \(n_i\), \(0 \leq n_i \leq N-1\) is the position of node i relative to the root node of a BVH with N nodes, and \(c_j\) is the relative positions of the descendants with respect to node i. This representation is strictly forward-stepping and infers the descendants of a node using statically known formulae and index information (see subsection 5.2).
Layout Arrays. An additional set of small arrays, termed layout-arrays, is also maintained which hold BVH metadata used for computing positional offsets of nodes relative to the root of their BVH, and their descendants at runtime. Layout arrays have the same capacity as the number of BVHs being evaluated, and store low-cost information such as offsets and depths (see Figure 1). Layout arrays can be pre-computed once on the host, during initialization, and then uploaded to the GPU since all information about each BVH may be known at this time.

Inferring BVH Information at Runtime. Since an entry of a BVTT-node does not encode information about the BVH containing its respective entry-node, a unique ID corresponding to each BVH is required to compute the descendants of the entry-node. We refer to this ID of each BVH as the layout ID. The layout ID is used to access layout arrays for the information belonging to the BVH containing a given entry-node. We compute the layout ID of a given entry-node by performing a modified lower-bound binary search [Cormen et al. 2009] over the layout array of BVH offsets, using the entry’s value (memory index) as the search target (see Figure 2). The layout ID gotten from this binary search is then used to read layout arrays for information (e.g. the depth) corresponding to the BVH containing the respective entry-node. Note that the overhead of performing this search operation is negligible since it is done in fast GPU local memory with $O(\log_2 N)$ complexity. Further, our method can handle both cases of static and dynamically changing BVHs since we only require that BVHs follow our storage representation. In addition, the implicit representation of BVHs greatly simplifies the construction process which is ideal and would lend itself well to parallel construction methods on GPUs [Lauterbach et al. 2009].

5 ALGORITHM

This section provides the details on how simultaneous BVH traversal is implemented on the GPU using the BSP model. We first describe the general steps to perform GPU traversal in subsection 5.1 and then describe our topologically-driven workload expansion scheme in subsection 5.2. Finally, we describe how intermediate BVTT-nodes are written to global memory at the end of each iteration on the GPU in subsection 5.3.

### 5.1 Parallel Traversal

The presented method evaluates the intersection of BVHs by iteratively expanding the BVTT using the breadth-first search (BFS) as a the core parallel primitive for traversal. The steps of algorithm 1 outline the pseudo-code of our method. The host (e.g. CPU thread) will invoke the GPU by calling `gpu_traversal()` in an iterative loop that will terminate when the traversal operation is complete. After invoking the GPU, the host must wait for the current iteration to complete which is represented by a call to `synchronise()`. Once the GPU has finished, the host will then read the new number of BVTT-nodes from the GPU using the function `dstFrontierSzRequest()` which is a GPU-to-Host memory copy command for a single integer value. The value read by the host determines the workload size for the next iteration and will be used to check if the traversal operation has completed.

The contents of `srcFrontier` and `dstFrontier` in algorithm 1 are distinguished to be read- and write-only, respectively, in order to implement double buffering, which is used to alias the output of one iteration as input for the next (see Figure 3). Swapping will occur at the end of each iteration on the host with all data remaining on the GPU.

**Algorithm 1: Iterative Bulk-Synchronous Traversal**

```c
// Arguments
1 // [input] srcFrontierDef: i3D iteration
2 // [input] srcFrontier
3 // [input] dstFrontier
4 HOST traversal(srcFrontierDef, srcFrontier, dstFrontier,...)
5 converged ← False
6 src ← srcFrontierDef // stores initial BVTT nodes
7
dst ← dstFrontier
8 do
9   [GPU] gpu_traversal(srcFrontier, dstFrontier,...)
10   if src == srcFrontierDef() then
11     src ← srcFrontier
12     synchronise() // stores new BVTT nodes
13     count ← dstFrontierSzRequest()
14   if count == 0 then
15     converged ← True
16   else
17     dstFrontierSzReset()
18     synchronise()
19   while converged ≠ True
20 return

1 GPU_gpu_traversal(srcFrontier, dstFrontier,...)
2 // Phase 1: read
3 data ← malloc(global, al[srcFrontier,...])
4 // Phase 2: traversal
5 if intersection() then
6   expandBVTT(...)
7 // Phase 3: write
8 write(dstFrontier,...)
9 return
```

**GPU Thread Operations.** To start traversal on the GPU, the host will launch approximately as many GPU threads as there are BVTT-nodes in `srcFrontier` (see Figure 3). This will be either the starting amount of default BVTT nodes if it is the first iteration, or resulting amount of the last iteration returned by `dstFrontierSzRequest()`. In phase 1 of algorithm 1, each thread will read a BVTT-node from `srcFrontier` into private register memory and then subsequently read the bounding volume information of each entry-node to perform intersection tests. Phase 2 defines the main body of computation performed by a thread since it is where the intersection test function is applied followed by BVTT expansion. Using the
we maintain all traversal data on the GPU. The output of one iteration becomes the input of the next as (i.e. c5.2 Work Expansion

in the first iteration of traversal is then obtained by pairing every and account for the actual memory locations of each such node heirarchies.

that are constructed from their descendants at lower levels. Figure 4 provides

BVTT-nodes, where

BVTT-node information that is now in private register memory, a thread will then proceed to evaluate it for intersection followed by expansion of the BVTT with new BVTT-nodes if the entry-nodes are found to intersect. Finally, in phase 3, threads collectively copy the new BVTT-nodes to dstFrontier for the next iteration.

5.2 Work Expansion

In this section, we describe our topologically driven workload expansion scheme. We introduce the concepts of static workload expansion and adaptive depth-stepping which are used to overcome GPU under-utilisation resulting from the small workloads of testing higher levels of BVHs, and to control the rate of traversal.

Static Workload Expansion. Evaluating the levels closest to the root nodes can yield small workloads compared to what is expected by GPUs to reach high throughputs. Therefore, in order to increase workloads for the initial iteration(s), evaluation of BVTT-nodes that are constructed from the root nodes is deferred to those constructed from their descendants at lower levels. Figure 4 provides an illustrative example of deferring the entry-level of three implicit hierarchies.

Given an entry-level \( l_e \), \( 0 \leq l_e \leq d - 1 \) of a BVH with depth \( d \), we compute its nodes using Equation 1 with \( \delta = l_e \) and \( n_1 = 0 \) and account for the actual memory locations of each such node (i.e. \( c_j \)) by adding the storage offset of the BVH. Once the entry-level of each BVH is computed, the set of BVTT-nodes evaluated in the first iteration of traversal is then obtained by pairing every node in the entry-level of one BVH to those of another. Deferring the entry-level yields approximately \( 2^{Nl_e} E \) BVTT-nodes, where \( E = \frac{N(N-1)}{2} + S \) is the number of collision checks between \( N \) BVHs, with \( S \) representing the number of self-collision checks. Such an increment can average-out the workloads over multiple iterations while also reducing total number of iterations since entry-level BVTT-nodes can be pre-computed on the host and uploaded once to the GPU as srcFrontierDef in algorithm 1.

Adaptive Depth-Stepping. Recall that expanding the BVTT is the process of creating new BVTT-nodes from the descendants of every pair of entry-nodes that are found to intersect; We introduce the concept of adaptive depth-stepping to infer the by-level distance to such descendants while accounting for any differences between the depths of tested BVHs. In what follows, the term depth-step is used to denote the by-level (jumping) distance that is computed at runtime, from an entry-node to its descendants: This allows us to (1) continue sprouting the descendants in one BVH while reaching the leaves of another, (2) further increase workloads at faster rates while reducing the number of iterations to complete traversal, and (3) tune for performance when writing to global memory.

We compute the depth-step by:

\[
\Delta d = \min(\mu, \Delta l), \quad 0 \leq \Delta d \leq d - 1
\]

(2)

where \( \Delta l = (d - 1) - \left\lfloor \log_2(n_1 + 1) \right\rfloor \) is the by-level distance to the leaf-level of the BVH containing an entry-node \( n_1 \), and \( d \) assumes the depth of the same BVH. The variable \( \mu, 1 \leq \mu \leq d - 1 \) is the user-specified parameter of expansion, which is used to control the maximum possible depth-step threads are permitted to use. (Note that \( \Delta d \) is zero if an entry-node is a leaf). Once \( \Delta d \) is known, the descendants of an entry-node are then determined by using Equation 1 with \( \delta = \Delta d \), which is then followed by BVTT-expansion.

5.3 Writing Traversal Output

We now describe our lock-free scheme for writing BVTT-nodes to dstFrontier, which is designed upon the BSP philosophy for fully utilizing the massive parallelism of modern GPUs.

All new BVTT nodes written to dstFrontier will be first accumulated in local shared memory and then flushed in coars-grained chunks to global memory to prevent individual thread access to global memory as shown in Figure 5. Traversal can potentially induce non-coalesced access to dstFrontier as a result of control-flow divergence which may be an additional source latency overhead. Thread-groups are used to achieve this by using an iterative write-wait-flush memory update scheme. Algorithm 2 outlines the steps of how the threads \( T \) that performed expansion as part of a group \( G \) copy their collective subset of BVTT-nodes to...


\begin{algorithm}
\caption{Lock-free synchronised write-access}
\begin{algorithmic}[1]
\State \texttt{global\_dst\_offset} \Comment{size of dst\_frontier}
\State \texttt{local\_base\_offset}
\State \texttt{i = 0}
\While{\texttt{(i × k) < n}}
\If{\texttt{num\_data > 0}}
\State \texttt{\_copy\_data = atomic\_add(num\_data, checkpoint)}
\If{\texttt{i < k}}
\State \texttt{r = r - e}
\State \texttt{w = max(num\_data, r)} \Comment{amount written}
\EndIf
\State \texttt{\_write\_fn(c, data, w)}
\State \texttt{\_num\_data = num\_data - w}
\EndIf
\State \texttt{\_synchronise\_group()}
\If{\texttt{i > 0}}
\State \texttt{\_checkpoint = – C}
\If{\texttt{local\_id == 0}}
\State \texttt{\_base\_offset = atomic\_add(dst\_offset, c)}
\EndIf
\State \texttt{\_synchronise\_group()}
\EndIf
\Else
\State \texttt{break}
\EndIf
\State \texttt{i = i + 1}
\EndWhile
\end{algorithmic}
\end{algorithm}

\[ I = \begin{cases} 
1 & \text{if } M \leq \kappa \\
\frac{1}{2} & \text{if } M > \kappa 
\end{cases} \]  
(3)

where \(\kappa\), \(1 \leq \kappa\), is the user-specified capacity of \(Q\) and \(M = 2^{\mu} \times |G|\) is the maximum possible output size of a group assuming all threads in the group performed expansion, where \(|G|\) is the user-specified thread group size.

**Lock-free Synchronisation During Shared Write-Access.** Writing to the local memory region \(Q\) can have serious impact on performance since it is a shared resource. For this reason, a shared variable \(C\) is used, which is a counter allocated per thread-group and is used to atomically compute a writing offset to the shared fixed-size region \(Q\) for each thread. At each iteration, group threads \(T\) compete for write-access to \(Q\) by atomically adding to the counter \(C\) (line 5). Each successful thread reserves a region to write its BVTT nodes such that those obtaining a valid offset is that which is within the bounds of \(\kappa\) will then asynchronously write to \(Q\) (lines 6-10). In essence, the threads that have data to write in the current iteration simultaneously contribute toward computing the offset of their collective output relative to a common base address in dst\_frontier. This base address is computed by the first thread of the group as a final step before flushing, which is done by using a single global atomic add after \(Q\) is filled (lines 12-20). Since \(Q\) is the sole interface to global memory, lone-thread accesses to global memory is reduced significantly, which can be more expensive to synchronise as workloads increase. We note that this scheme is in fact similar to Garanžha at

\[ T \leq \log_4 \left( \frac{\kappa}{\alpha} \right), \quad 4 \leq \kappa \]  
(4)

where \(\alpha\), \(1 \leq \alpha \leq \sqrt{\kappa}\) is a user-specified value for the minimum number of group threads \(T\) guaranteed to write all their BVTT-nodes in single iteration. Thus, the guaranteed threads will collectively write \(\alpha \times \beta\) BVTT-nodes to \(Q\) in the current iteration, such that the \(n^{th}\) thread to atomically offset \(C\), where \(n = \lceil \frac{\kappa - 1}{\beta} \rceil\), will write at-most \(\kappa \mod \beta\) BVTT-nodes to \(Q\) and the rest will be written in the next iteration.

6  RESULTS

We evaluate our method using OpenCL 1.2 on the AMD Radeon R9 280X (3GB VRAM, 32KB Local memory) and Nvidia Geforce GTX 960 (4GB VRAM, 49KB Local memory) GPUs. Three benchmarks (Figure 6) from the UNC Dynamic Scene Benchmarks dataset [Curtis et al. 2017] are used for evaluation purposes: \textit{NBody} (6a) has the largest number of objects at 305 with a total of 146K triangles, it is a rigid-body simulation involving many interacting objects without self collisions. \textit{Funnel} (6b) is a soft+rigid body simulation and is the smallest benchmark made up of four low-resolution meshes (total of 18.5K triangles). In this benchmark, the primary interactions occur between the cloth and funnel. \textit{Cloth-Ball} (6c) is another soft+rigid simulation with two objects that have 92K triangles in total. We use simple axis-aligned bounding boxes (AABB) storing one triangle per leaf-node and our BVHs constructed in bottom-up fashion.
Table 1: Our performance results for simultaneous parallel BVH traversal involving inter- and intra-object collisions.

<table>
<thead>
<tr>
<th>Benchmark</th>
<th>GPU</th>
<th>Geforce GTX 960</th>
<th>Radeon R9 280</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cloth-Ball</td>
<td></td>
<td>6.43</td>
<td>3.08</td>
</tr>
<tr>
<td>Funnel</td>
<td></td>
<td>0.99</td>
<td>0.57</td>
</tr>
<tr>
<td>NBody</td>
<td></td>
<td>2.42</td>
<td>1.16</td>
</tr>
</tbody>
</table>

6.1 Performance

Table 1 summarises the performance of our algorithm when we consider both inter- and intra-object tests for the case of Cloth-Ball and Funnel. The presented results are based on the heaviest workloads (colliding leaf-nodes pair) experienced at the most demanding time-step in each benchmark (Cloth-Ball 3.1mil, Funnel 314K, Nbody 117.6K). During experiments, our GPU kernels are executed at-least eight times to reduce potential noise in time measurements because of system warm-up overhead. However, no significant differences were observed between test runs.

The presented method is able to perform parallel simultaneous queries in real-time. Execution time is fastest on Funnel with query time under 1ms. Cloth-Ball takes the longest time (6.43ms on the GTX 960). This benchmark has the largest workloads with over 3.1million overlapping leaf-node pairs due to the self-collisions induced by the cloth’s motion. For this benchmark, our method is able to complete traversal within 6.5ms. NBody has the lowest number of leaf-node overlaps because is a rigid body simulation. Its BVHs are approximately twice as much slower to evaluate than Funnel due to the larger number of objects (305), and hence, the resulting BVTT. The results reveal our method’s strong ability to exploit large scale parallelism on GPUs to quickly evaluate a large number of BVTT nodes for pair-wise CD.

**Speedup.** We compared the performance against our implementation of the "streams" model by Tang et al. [2011]. Comparisons are made using the time-step/frame with the highest workloads on each benchmark. We did not include intra-object collisions for Cloth-Ball and Funnel to ensure that workloads fit in our global memory buffers for the streams model. A reduced implementation was used with only pair-wise collision queries to ensure a fair comparison. The implementation also used explicit (not deferred) front-tracking with stream registration based on segmented locking mechanism (see Tang et al. for details). According to Tang et al., deferred front-tracking simply trades memory overhead for additional runtime computations.

To emulate the BVTT-node cache (moving front) used by the streams model, we setup the benchmarks as follows: For each benchmark, we extract a pair of keyframes \((k_t, k_{t+\Delta t})\) which are consecutive in time, with each keyframe \(k\) representing the geometry of a particular time-step \(t\). Next, we then build the BVH of each mesh in the benchmark for \(k_t\) and \(k_{t+\Delta t}\). The BVTT-node cache is created by traversing the BVHs of the meshes of \(k_t\) until completion and saving the BVTT nodes at which traversal terminates as described by Tang et al. [2011]. Our traversal tests and comparisons are performed using BVHs constructed from \(k_{t+\Delta t}\) since we can use the BVTT-node cache built from \(k_t\) as input for traversal at \(t + \Delta t\), thereby allowing the streams model to have a valid cached input set from a "previous" time-step. We have not included the cost of work redistribution for the streams model in our evaluation.

Figure 7 shows speedup were comparisons are based on BVH traversal times to find the set of potentially colliding triangles pairs. Performance of our method on all benchmarks is faster with an average speedup of 4.4x on the R9 280X and 4.3x on the GTX 960. The highest speedup is on NBody at 7.1x for R9 280X (6.2x on the GTX 960) which has the largest workloads in our comparison setup. In general, we found that adapting the streams model on arbitrary GPU architectures is non-trivial due to its dependence on the available amount of local memory for the work-stacks and exploiting L1 caches. Our method is an efficient and a more simpler option for mapping traversal to GPUs.

6.2 Parameter Effects and Trade-Offs

The explorable nature of our exposed parameters can make finding correlations between their configurations and the resulting performance unintuitive with no obvious settings. Figure 8 shows the results illustrating the effects of the entry level \(l_e\) and the parameter of expansion \(\mu\) on execution time for each benchmark (with intra-object collision tests for Cloth-Ball and Funnel). We have found that although increasing \(\mu\) reduces the number of iterations in our method, care must be taken when making the choice of value. For our evaluated range \((1 \leq \mu \leq 4)\), making further increments beyond \(\mu = 3\) produces a drastic slow-down where the execution-time is on average 3 to 5 times slower than choosing a value between 1 and 3. The observed spikes in the data of Figure 8 , which are observed for \(\mu = 3\) and \(\mu = 4\), are due to our BVH padding scheme. Some levels in our hierarchies contain more inactive BVH nodes than others, hence the periodic spikes subject to \(l_e\). Note that in our implementation, we filtered out the any BVTT-nodes containing padded BVH-nodes during entry-level construction). Generally, a choice of smaller values of \(\mu\) e.g. 2, is suitable for the case of reducing execution time, even though this choice is at the behest of more iterations to complete traversal. We found \(l_e\) to serve our method well for statically reducing the number of iterations to complete traversal while providing the large workloads that we need to
utilise the GPU. There are some limitations on the exploitation of $l_e$ however, since its chosen value must correlate with the number of BVHs tested to control the input size. On the Nbody simulation, we see a more rapid (exponential) performance drop with $l_e$ compared to the other benchmarks due to the faster rate of increase in the initial input size. For example, we observe that setting $\mu = 2$ and making increments on $l_e$ from 1 to 5 results in a sharp change in execution time from 3ms to 16ms respectively on the GTX 960.

Local memory and thread-group sizes. The allocated local memory size $\kappa$ of the fixed-size region $Q$ and thread group size $|G|$ also have an effect on performance and its scaling properties due to their influence on scheduling. Figure 9 provides our findings regarding the change of execution time relative to $\kappa$ and $|G|$, respectively. Setting either parameter to the highest tested value (e.g. $\kappa = 2^9$ and $|G| = 2^8$ on the R9 280X) while maintaining the other at a minimum (e.g. $2^1$) showed slower performance in most cases with the exception of the NBody simulation on the R9 280X. More generally, we observe similar behavioural patterns on both GPUs with the GTX 960 appearing a little more constrained in terms of the optimal choices of $\kappa$ and $|G|$. In our results we have found that configurations that use mid-range values are sufficient to obtain good performance relative to the worst case for each benchmark. We observe that our method favours medium-to-large thread groups ($|G| \geq 2^5$) and allocated local memory size ($\kappa \geq 2^6$) for good performance. The results of Figure 9 are a demonstration of the importance of the trade-offs to be made through our parameters which is crucial for portability.

7 CONCLUSION

We have presented a simple alternative solution for simultaneously traversing a large number of BVHs for CD on GPUs. Our method utilizes the BSP model to overcome the irregular and data-dependent nature traversal. The simplicity of our approach stems from the use of topological properties inherent within an implicit hierarchical representation to harness the parallelism of GPUs. From this, we have presented our topologically-driven workload expansion scheme which provides fine control over the rate of traversal while also increasing workloads for the first iteration(s). In addition, we have described a simple lock-free global memory updating method that can be controlled to adapt algorithm performance based on the available hardware resources. This can likewise be extended with more complex lock-free synchronisation mechanisms using scan primitives such as prefix-sum [Sengupta et al. 2007]. Our method can evaluate complex hierarchies in real-time, and with a speedup of up to 7.1x over the widely used “streams” model.

Limitations and Future Work. The presented algorithm faces a number of limitations which affect performance. Our solution minimizes the compute workload per thread while increasing the DRAM traffic as a side-effect. This is because threads perform just one intersection test, such that in order to perform it, they need to stream data from global memory. Also, our BVH node array is sparsely populated due to padding, which can easily cause excessive L2 and global memory traffic. Such padding can, in the worst-case, also double the storage requirements per BVH subject to the number of leaf nodes. We also note that it is a possibility that our approach of using a one-to-one mapping between threads and BVTT-nodes may not utilise the benefits of GPU caches because there is no opportunity for the reusing BVTT-nodes from srcFrontier: The initial read operation of phase 1 (see algorithm 1) is effectively a cold start with no opportunity for explicit data reuse since little temporal locality exists when reading BVTT-nodes and BVH node data.

In future work, we plan to extend support for BVH compression by eliminating inactive BVH nodes. This would serve as a solution to the highlighted limitation that the currently employed padding scheme is likely to have limited exploitation of GPU caches leading to excessive global memory traffic alongside the higher memory footprint. Support for moving fronts without complex memory
management would also benefit our method well since it is strictly forward stepping with no notion of “node-collapse” to backtrack up hierarchies.

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REFERENCES


