Parallel Search: A* with CUDA

15-418 Final Report

Aditya Kannan Gabriel Lee

Summary

We developed an approach to implement the A^* algorithm in parallel called GA^* . A^* is a classical search algorithm that is used for computing the optimal path between a source and a sink node. We develop several data structures, optimized for use in parallel applications, and designed an algorithm that allows us to run A^* efficiently on a GPU. We implemented a serial version of A^* and sequential version of the GA^* algorithm. While we made many attempts at creating GA^* in CUDA, we were ultimately unsuccessful in this endeavor. We applied the A^* problem to a slider game. We demonstrate the solution generated by our algorithm in a visual demonstration of the solution to the sliding game.

Background

2.1 The A* Algorithm

The A* algorithm is a solution to the single source shortest path problem. It is applied on a graph (directed or undirected) with various costs for all the edges of the graph. A* finds the path of edges from a given source and goal node that minimizes the total sum of the edge costs.

It improves on algorithms like Dijkstra's algorithm by using a heuristic to help guide the search for the end goal. Every node n in the set of vertices V has a distance g(n) from the source node that was discovered by exploring the different paths from the start to n. There is also a heuristic function h such that h(n) represents an estimate of the remaining distance to the goal node. Nodes on the frontier of the search are placed on a priority queue. They are then picked to be explored from the frontier based on a combination of the cost and heuristic functions:

$$f(n) = q(n) + h(n)$$

By using f(n) to prioritize nodes that we believe are more likely to reach the goal faster, we are able to expand fewer nodes and find the least cost path more efficiently then Dijkstra's algorithm.

In order to ensure that we have an accurate estimate of the priority of a node, we need our heuristic to be consistent in order for the A^* algorithm to generate a provably correct output. A heuristic is consistent if for a node s and any of its neighbors s', it satisfies the following inequality:

$$f(s) \le d(s, s') + f(s')$$

where d(s, s') is the distance between the two nodes.

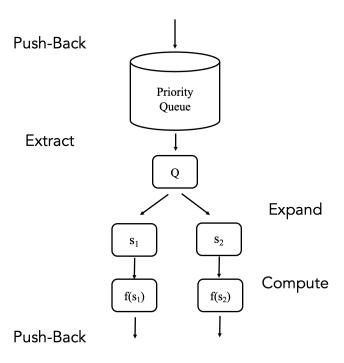


Figure 2.1: A* workflow

2.2 Algorithm $(GA^*)[1]$

The A* algorithm as described above is not amenable to parallelization. The priority queue requires that only the topmost entry in the priority queue be used for exploration. As a result, it is not possible to create a parallel version of the algorithm described above without changing the data structures.

GA* is a general version of A* that can be run in a parallel setting. We present the pseudocode of our algorithm below:

```
// every thread expands one state
        let si = qi.pop()
        // update goal state if found state is shorter path to goal
        if si = t then
            if m = None or si.total_cost < m.total_cost then</pre>
                m <- si
            break
        // add neighbors
        S.extend(s' for s' in s.next)
    done
    // check if found goal state is closer than all other states
    if m != None and m.total_cost <= min si.total_cost then return m.path
    // remove expanded states that have already been seen cheaper
    parallel for i in [0, s.length] do
        if si in H and H[si] < si.path_cost then s.pop(i)
    done
    parallel for i in [0, s.length] do
        // push si to some q in Q evenly
        Q.push(si)
        // store seen expanded paths
        H[si] <- si.path_cost</pre>
    done
done
return m
```

At a high level, this algorithm uses multiple priority queues to generate a path between the start state and goal state. An explicit graph data structure is unnecessary in this scenario as we can determine neighbor states from the current state (and anyways, an explicit graph structure would be very much too large in scenarios that would motivate parallelization on this scale).

The pseudocode is commented at each step, but the high level idea is that each thread has its own priority queue of states. Each thread expands the top node of its own priority queue, checks if it has reached a goal state and if it has been visited already and expands the neighbors of this state.

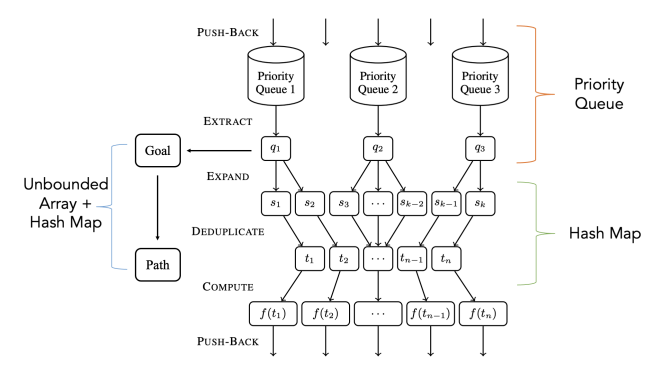


Figure 2.2: GA* workflow with relevant data structures (based on [1])

When states are expanded, the number of states grows exponentially, resulting in a smoothing of the states distribution across all of the threads. This results in state duplication among independent threads that may separately want to push the same state onto the frontier. To handle this, we need to keep track of which nodes have been explored and/or will be on the frontier. We utilize the global hash table (described below) for this purpose.

2.3 Data Structures

2.3.1 Serial

Standard libraries cannot be used in CUDA code due to GPU memory working differently than CPU memory. Thus, certain serial data structures need to be re-implemented to accommodate for this.

Unbounded Array

This structure is used to generate the path to the goal state once the algorithm terminates. Thus, unlike a standard unbounded array, it only requires a push operation. It has a growth factor of 2 (resize to double its original size).

Priority Queue (Heap)

Every thread contains their own priority queue for expanding states. These are min-heaps (for retrieving closest perceived state) that operate on top of an unbounded array structure (same as above, but with a shrinkage factor of 2 as well). In accordance with the above algorithm, they have pop, push, and is_empty operations.

2.3.2 Parallel

Hash Table (Parallel Hashing with Replacement)[1]

This structure is used to keep a global collection of visited states and their path costs. Because of this global attribute, this is the bottleneck in the algorithm. A standard hash table contains an unbounded array of linked lists, but the structure's dynamic nature (on item inserts) makes it extremely difficult to parallelize. To tackle this problem, we implement parallel hashing with replacement[1] which uses a static list and multiple hash functions. Details are explored in the Approach chapter. As in a standard hash table, this has assignment, existence, and modification operations. These operations are allowed to run in parallel with one another and as a result must be safe for parallelism.

2.4 Workload

In the sequential algorithm for A*, a single priority queue is used to determine the next state to extract. Since node neighbors are fully dependent on the nodes within the priority queue, this data structure becomes the bottleneck if we were to parallelize the algorithm. If instead (as demonstrated in the pseudocode), we allow for states to be expanded independently of one another using multiple independent priority queues, we will allow for duplicate expansions. Thus, especially for large problem spaces, the approach is massively parallel.

Since each thread operates independently in the sense that they expand nodes without dependencies on nodes held by other threads, this program is task parallel. Locality and vector instructions are not as relevant due to this.

Approach

3.1 Setup & Hardware

All of our code was written from scratch.

The goal of our project was to create a working GA* implementation on GPU. To this end, we used C++ for serial implementations and CUDA programming to develop our GPU code. We evaluated our code on the GHC machines.

3.2 Work Distribution

The parallelism of our work comes from how we explore the frontier of the A* search more efficiently. The original A* algorithm as designed for sequential execution is not amenable to parallel speedup, so we used the GA* algorithm instead to judge parallel execution.

Each of our threads corresponded to one priority queue. It was responsible for popping the top node in the priority queue, and evaluating it for if it is the goal state and for finding its neighbors. With this approach, we could have thousands of priority queues, which is especially useful in problems where the number of states increases exponentially, such as the slider problem we describe in the Results. In the rest of our algorithm, we encounter a lot of communication and synchronization.

We allocate a global array S for storing all the neighbors that are being added to the frontier due to the expansion of the priority queue nodes. This array is accessible to all the threads. Once each thread generates the neighbors it expands in the current iteration, it adds those neighbors to S.

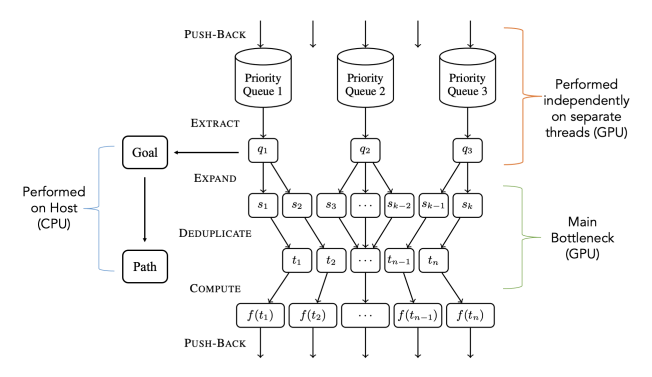


Figure 3.1: GA* workflow with relevant data structures (based on [1])

We also allocate the hash table as a global variable. Once the threads populate the array S, each thread works on deduplicating nodes in the array S (not necessarily the nodes it added to S!). This is done by looking at the global hash map for the same node and replacing it if there's a node in S that has lower cost. If not, we discard this node from S.

Finally, we redistribute the remaining nodes evenly to the many different priority queues. We do this by rearranging the indices in S that correspond to the thread.

3.3 Iterations of our Approach

We began by implementing and verifying a sequential version of the original A* algorithm. These results are presented in the Results below. We then created a sequential version of the GA* algorithm that worked (i.e. it could run on a single CPU thread, not on CUDA. Nodes were mapped to a random priority queue so the structure imitated the final result we were hoping for). Finally, we made numerous attempts at a CUDA implementation for the GA* algorithm, but we were ultimately unsuccessful.

Along the way, we made a couple optimizations to our approach. We decided to allocate nodes in our frontier in S in one ordering, and then we redistribute the nodes to threads in a different order. This way, by having nodes in different parts of the graph, the work done

by the threads is evenly spread out. That is, we are less likely to run into situations where many priority queues are empty and do not have nodes to expand or add to S.

Another aspect we found was that we could optimize our deduplication to haves less synchronization and therefore potentially save time. There were two methods that [1] suggested for a parallel implementation of a hash table. The first one, Cuckoo Hashing, is a more complicated method that involved greater synchronization between threads. We opted to go with the second approach, Hashing with replacement, which required no synchronization at the cost of missing some duplicate nodes. We decided this tradeoff was acceptable because this approach would not affect the final path returned.

Results

4.1 Sliding Game

Our metric for algorithmic performance involves running A* on the sliding game (see the Youtube video for a demonstration). The sliding game is a particularly convenient search problem for two reasons: state representation is small and the state space is very large (blows up exponentially with respect to the puzzle dimensions). The latter is particularly important since our goal was to measure the effectiveness of a parallel A* algorithm, and the expansive state space would decrease potential overhead.

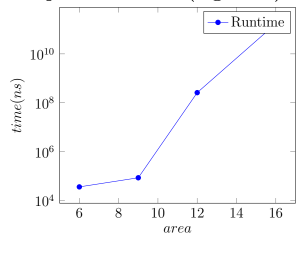
Sliding Game Sequential Times				
Board Size	Runtime (ns)	Runtime (ms)	Runtime (s)	
2x3	35880	0	0	
3x3	84711	0	0	
3x4	258638033	259	0	
4x4	171623957677	171624	172	

0			
5	15	4	3
9	14	8	2
1	13	0	12
7	6	11	10

Figure 4.1: The puzzle grid used for the 4x4 measure

The object of the game is to order the tiles in increasing numerical order. The 0 tile represents an empty tile where adjacent tiles can "slide" to. Thus, every turn can result in 4 more possible states (exponential). The graph below is a visual representation of the above runtimes in log scale, supporting that the state space blows up.

Sequential Times (log scale)



Bibliography

[1] Yichao Zhou and Jianyang Zeng. Massively parallel a* search on a gpu. *Proceedings of the AAAI Conference on Artificial Intelligence*, 29, 02 2015. URL https://ojs.aaai.org/index.php/AAAI/article/view/9367.