Parallel Evaluation of a Spatial Traversability Cost Function on GPU for Efficient Path Planning

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ABSTRACT

A parallel version of the traditional grid based cost-to-go function generation algorithm used in robot path planning is introduced. The process takes advantage of the spatial layout of an occupancy grid by concurrently calculating the next wave front of grid cells usually evaluated sequentially in traditional dynamic programming algorithms. The algorithm offers an order of magnitude increase in run time for highly obstacle dense worst-case environments. Efficient path planning of real world agents can greatly increase their accuracy and responsiveness. The process and theoretical analysis are covered before the results of practical testing are discussed.

Keywords: GPGPU, Dynamic Programming, Concurrent Programming, Path Planning

1. Introduction

Global path planning is an important part of robot navigation through both static and dynamic environments. Many simple techniques exist for immediate obstacle avoidance [1], but without a high level path to follow to a desired destination an agent cannot guarantee near optimal travel. A current popular technique is to generate a two-dimensional occupancy grid based on the robot’s on-board sensing capabilities, such as laser range scanners [2-4]. On this occupancy grid a cost-to-go function is generated from a single destination point to any number of locations. A global holonomic path can be generated from this function by travelling from grid cell to grid cell by choosing the neighbour with the lowest global cost.

Generation of the cost-to-go function from an occupancy grid is an inherently sequential algorithm as the global cost of a cell depends on the calculation of a neighbour closer to the destination point. This paper presents a modified version of the cost-to-go generation algorithm that makes use of the parallel hardware found on modern graphics cards, while maintaining numerically identical results to traditional sequential dynamic programming algorithms. This paper begins by giving a background on developing concurrent algorithms on modern graphics hardware, followed by a literature review of existing sequential techniques. The paper then demonstrates how the proposed method achieves an order of magnitude increase in efficiency before showing experimental results and their practical benefit.

2. Background

Current forefront central processing unit (CPU) technology is making use of multiple cores to improve performance and as such many software developers are attempting to find new methods to parallelise existing algorithms for the new paradigm [5,6]. Running along side this CPU evolution in recent years is the growth of graphics hardware in capability and performance. This growth has been driven by market forces such as consumer demand for more realistic games and interactive media, but it has enabled a new research area known as General Purpose Computing on Graphics Processing Units (GPGPU) [7]. This section will give a brief outline of some foundation ideas of the field of GPGPU.

2.1. General Purpose Computing on Graphics Processing Units

Graphical processing units (GPUs) have evolved over the past two decades to have one to two orders of magnitude more cores than currently found in multi-core CPU technology [8]. Rather than using a multiple instruction multiple data (MIMD) paradigm like multi-core CPUs, GPU manufacturers have gone for a single instruction multiple data (SIMD) paradigm. That is, in SIMD, in each cycle, the same set of instructions is executed over a range of data. Graphical calculations have involved performing
the same transformation, colouring, texture interpolation and lighting calculations on many different vertices and fragments in a three-dimensional scene. From one point of view, this can be seen as limiting the flexibility of the hardware. On the other hand, it relieves the programmer from having to deal with a core problem associated with concurrent programming.

The two main problems faced by concurrent programmers are concurrent access of critical resources, such as common data, and synchronisation between the independent threads of execution. Graphics hardware removes the burden of concurrent data access by using the SIMD paradigm and enforces synchronisation at a regular interval via the rendering of a single frame to screen or other texture buffer.

Some common programming constructs are known under different terminology in a GPGPU context. Memory is usually referred to as either a texture or a frame buffer as the programmer usually accesses data in this memory via pixel or texel lookups.\(^1\) A single program run on each pixel individually is known as a shader program from a traditional graphics programming perspective, or more recently a kernel in GPGPU research.

A common example demonstrating the power of concurrent programming on GPU is the grey scaling of an image. For an image that is \(n \times n\) pixels a sequential algorithm will have to process each pixel individually and average the red, green and blue channels to calculate the grey scale intensity. Therefore, the sequential algorithm runs in \(O(n^2)\) time. On a GPU, since each pixel’s calculation is independent this kernel can be run on each pixel concurrently, therefore running in \(O(1)\) time.\(^2\) Analysis of similar computer vision methods ported to GPU are referred to in [9].

Another textbook example is matrix-matrix multiplication [10,11]. Usually a sequential algorithm will run in \(O(n^3)\) as each of the \(n \times n\) cells in the solution matrix requires a process running in \(O(n)\) time (the sum of \(n\) multiplications). Again, this kernel can be run on each cell in the solution matrix independently, so a parallel implementation of this process will run in \(O(n)\) time. Of course, in practice, this linear algorithm would run slightly slower depending on the number of processors available. For example, many current consumer graphics cards have approximately one hundred thread processors, whereas a high end General Purpose GPU such as the nVidia Tesla [12] has around five hundred thread processors. Table 1 shows the increase in cycles based on the number of calculations to be completed spread over the number of processors available.

More generally this can be specified as the complexity of the sequential algorithm divided by the number of processors available. For current consumer graphics hardware this means the matrix-matrix multiplication algorithm could run anywhere between \(O(n^2)\) and \(O(n)\) time depending on the size of \(n\) relative to the number of thread processors. However, with no modification to the algorithm itself, this can reach \(O(n)\) in current forefront graphics hardware and therefore next generation consumer and mobile graphics hardware.

### 2.2. The Ping-Pong Method

In GPGPU, many simple algorithms require a single iteration or frame to be calculated, like image grey scaling or matrix-matrix multiplication. There are however many algorithms that have to be evaluated through a series of steps.\(^3\) The state of each step must be saved in a frame buffer ready for the next calculation. A textbook method outlined in [13] is the ping-pong method. This method uses the standard two texture/buffer approach of having one input buffer and one output buffer, rather than outputting to screen like normal graphics operations. The difference is that between each step the roles of the two buffers are reversed so that the input from one step becomes the output of the next step. By have one texture as read-only and the other write-only in each iteration, each kernel is allowed to make a clean state change. The example used in [13] is finding the maximum value in an array of values. Sequentially, this algorithm runs in \(O(n)\) time, whereas the parallel version can run in \(O(\log n)\) time. The parallel algorithm partitions a two-dimensional matrix of these values into \(2 \times 2\) grids. A kernel will take these four values, calculate the highest value and place the result in a known cell closer to the origin. This cell becomes part of another \(2 \times 2\) group that comprises of other running local maxima. For example, given 16 numbers in a \(4 \times 4\) grid, each of the four \(2 \times 2\) quadrants

\(^1\)From a low-level hardware point of view graphics card memory is indifferent to traditional CPU accessible memory. However, a programmer using graphics programming languages is given access to this memory more like a two-dimensional array of RGBA values.

\(^2\)Assuming averaging the RGB channels of a pixel is negligible.

**Table 1. Comparison of cycles required for differing number of available processors.**

<table>
<thead>
<tr>
<th>(n)</th>
<th>(n^2)</th>
<th>(\text{cycles pr} = 100)</th>
<th>(\text{cycles pr} = 500)</th>
</tr>
</thead>
<tbody>
<tr>
<td>4</td>
<td>16</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>8</td>
<td>64</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>16</td>
<td>256</td>
<td>3</td>
<td>1</td>
</tr>
<tr>
<td>32</td>
<td>1024</td>
<td>11</td>
<td>3</td>
</tr>
<tr>
<td>64</td>
<td>4096</td>
<td>41</td>
<td>9</td>
</tr>
<tr>
<td>128</td>
<td>16,384</td>
<td>164</td>
<td>33</td>
</tr>
</tbody>
</table>

\(^3\)Some algorithms cannot be completely parallelised as there is some dependence between steps within an algorithm. However, many of these algorithms can have disjoint steps run concurrently.
can have their maximum calculated concurrently and placed in a corresponding cell in the bottom-left quadrant. On the next iteration the maximum of all 16 numbers will be the result of running the kernel on the $2 \times 2$ bottom-left quadrant. This results in the maximum of 16 values being calculated in two steps.

3. Literature Review

Cost-to-go function generation on an occupancy grid is a well established technique used for robot path planning [2] and other non-linear control problems such as Model Predictive Control (MPC) [14]. Many techniques for generating this cost, given obstacles and a goal, initially relied on a potential function where obstacles have a repulsive “charge” and the goal state(s) have an attractive “charge” [15,16]. The advantages of this method allowed a robot to not only plan a path towards a goal state by avoiding obstacles but by staying a distance from obstacles if the environment allowed for it. Some publications, such as [16], have shown that for some state spaces this method can produce local minima which do not involve the goal state(s).

Another method widely used that removes the local minima weakness of the potential field method above is the Laplacian method [17]. Given a single destination location on an occupancy grid, with a cost of zero, the cost from any point on the grid to that destination can be calculated using the Laplacian method by treating the destination point as a point charge. As a result, the cost-to-go evaluation runs as an increasing wave front emanating from the destination point. As with other graph based path planning algorithms [18,19] the cost of a node or cell cannot be globally determined until one of its immediate neighbours’ cost has been determined.

In practice, when working with multiple agents it may be desirable for more than one agent to travel to a given common location. As a result, an exhaustive cost-to-go function can benefit all agents rather than a path planning algorithm that terminates when a single path is found for one agent. If the environment doesn’t change then this cost-to-go function can be used continuously without change until both agents reach the desired destination location.

There is also an emerging direction in robot path planning of calculating a path in a non-discretised domain [20]. Instead of the previously mentioned technique of using an occupancy grid to plan on, this area of path planning decided on a path in continuous space. This can be desirable as the path generated is smoother. However, these techniques are inefficient for practical use with current hardware capabilities.

Even though the method proposed in this paper relies upon a discretised space, if the algorithm does not have an overly strict time requirement then it can be given an occupancy grid with higher resolution. This may allow the solution to be close enough to continuous for all practical means without reducing the coverage or responsiveness the pre-existing algorithm provided.

Similar path planning techniques have also been applied to multi-core architectures such as the GPU for considerable increase in efficiency. Bayesian estimation over a grid based representation of an environment is proposed in [21]. The technique makes use of matrix and vector calculations that are inherently efficient on GPU, for the prediction and update steps of maintaining a probabilistic belief of an agent’s location, for example.

4. Method

The generation of a cost-to-go function cannot be purely concurrent for every cell as there is a dependency between one cell’s global cost and its neighbour’s cost. However, as a cost generation function expands away from the zero cost destination cell, many of the cell calculations are spatially disjoint from one another. This section outlines how these disjoint cell cost calculations can be done concurrently.

4.1. Parallel Cost Function Generation

Existing algorithms for cost function generation go through every cell that is on the edge of the classified/non-classified border and calculate a value for that cell $x$:

$$C^*(x) = \min [C^*(y) + \delta(x,y)] \forall \delta(x,y) < \infty$$

This equation is the basis of existing path planning algorithms as it maintains the Principle of Optimality [22] associated with dynamic programming.

Figure 1 shows the definition of the terminology used in this paper. In particular, the wavefront is defined as the cells that have not been given a global cost value, but have at least one neighbour with a global cost. This algorithm assumes each cell has eight neighbours and hence includes the diagonal neighbours in each calculation. This means on the subsequent iteration of the algorithm every cell in the wavefront will have a global cost calculated for it and become “classified”.

For a grid containing $n \times n$ cells this algorithm will need to run a calculation for each cell, meaning an $O(n^2)$ run time from a traditional sequential point of view, let alone the overhead of maintaining an ordered priority queue of yet to be visited paths. The concurrent algorithm does not require any supporting data structures.

As a simplified example, the very first step of generating the cost function is to assign the destination cell a value of zero. After this you would logically go to each of the neighbouring cells and calculate the cost of each neighbour.
Figure 1. Definition of different classification of cells. Black represents a non-traversable grid square (obstacle), blue represents a classified traversable region, green represents the wave front about to be evaluated/classified, and white represents the unknown/unclassified region. The figure shows the progression of the wave front through the first five iterations of the algorithm.

If the cost of traversal between cells is a uniform cost then the optimal cost value for each of these first generation neighbouring cells can equally be based on the value of the original destination cell. In a two dimensional grid this means eight separate disjoint calculation steps. In a GPU these calculations can be done concurrently, thus using one iteration of the process rather than eight. In the subsequent wave front a sequential algorithm would require sixteen cell calculations, which can also be done concurrently on a GPU. Table 2 shows how this number of “iterations” increases at different rates for sequential and parallel solutions.

This reinforces the fact that from a theoretical point of view the traditional sequential algorithm runs in \( O(n^2) \) time whereas the proposed parallel algorithm runs in \( O(n) \) time.\(^4\)

In theory, for uniform local cost environments it would be optimal if the cost calculation for a cell could only be run on a cell once. In practice, this is not possible without having to do some per-cell calculation in the kernel to determine whether the calculation should be done, which can be inefficient for large \( n \).

The kernel makes the most significant progress when it is applied to cells in the wavefront. Therefore, a larger wave front will enable more cells to be calculated concurrently in one iteration. In practice wide open spaces in the occupancy grid promote larger wave fronts, while corridors limit wavefront growth. In an optimal best case where there are no obstacles an \( n \times n \) grid can be completely evaluated in \( \frac{n}{2} \) steps, that is, \( O(n) \) time. In the worst case of a single cell wide corridor, the algorithm runs equally as slow as the sequential algorithm. As seen in the Experimental Results section the algorithm runs closer to \( O(n) \) time than \( O(n^2) \) in practice for environments a field robot would be expected to traverse.

### 4.2. Implementation

The parallel algorithm was implemented using Open GL’s Shading Language (GLSL) [23]. Other mainstream programming languages such as nVidia’s CUDA [24], ATI’s Stream [25] and the generic successor OpenCL [26] were considered, but the grid based nature of an occupancy grid intuitively maps to the texture-kernel paradigm associated with traditional graphical applications.

In most path planning algorithms mentioned in the Literature Review section, unclassified cells are given the value of infinity to allow the algorithm to update the cell’s value given any initial finite calculation. In practice, this algorithm defines unclassified cells as having a value of \(-1\). This value was chosen to represent unclassified cells for possible forward compatibility with stencil buffer operations (see the Future Work section). Obstacle cells are stored as \(-0.5\). Negative numbers are used for determining special cells as normal classified cells will have a global finite cost greater than or equal to zero.

### 4.3. Multiple Destination Point Optimisation

As mentioned earlier in this section, the progression rate of the algorithm is related to the size of the wave front. That is, wave fronts that take up more grid cells can

| #   | cells | \( \Sigma \) cycles (seq) | \( \Sigma \) cycles (||) |
|-----|------|--------------------------|-------------------------|
| 1   | 8    | 8                        | 1                       |
| 2   | 16   | 24                       | 2                       |
| 3   | 24   | 48                       | 3                       |
| 4   | 32   | 80                       | 4                       |
| 5   | 40   | 120                      | 5                       |
| 6   | 48   | 168                      | 6                       |
| ... | ...  | ...                      | ...                     |

\(^4\)As stated in the Background section, in practice this may be hindered by the number of processors available on the hardware. As graphics card manufacturers continue to increase the number of processors in GPUs this algorithm will tend towards linear execution time in practice.
make use of more processors in a single algorithm iteration. When working with multiple agents, there may be a situation where one or more agents are required to travel to more than one destination, where each destination has equal weight of importance. This scenario greatly benefits from the parallel implementation by providing multiple wave fronts and hence more grid cells situated in the wave front state every iteration.

4.4. Optimisation of Calculation using Expanding Texture

In the textbook definition of the ping-pong method the entire texture is requested to be rendered/evaluated every iteration. In the early stages of this algorithm there are many unclassified cells that are a large distance from the wave front that can be given to the shader pipeline if the entire texture (occupancy grid) is rendered every frame to iterate the algorithm.

One rough method to speed up the process is to render a sub-area of the texture as large as the wave front could possibly be at a given iteration of the calculation. Under this method, the sub-texture requested to be rendered gradually increases in size for each iteration of the algorithm. This technique is shown in Figure 2. Any texture cell not rendered will be left in its prior state, and hence will not change. It should be noted that this only applies for single destination cases.

As an example, in the very first iteration all eight of the destination’s neighbours will be evaluated. Therefore, the wave front will never be further than one cell from the destination and therefore no other cells are required to be given to a shader program for evaluation for this iteration. In short, at each stage, the area of the texture that is requested to be rendered gradually grows with the wavefront, so that (in the early generations) many of the redundant calculations of cells away from the wave front are not even considered. This is not perfect, as seen in the fourth iteration in Figure 2, as cells immediately behind an obstacle are within the expanding texture’s bounds, but are still run through a shader program. Figure 3 shows the execution time difference with and without expanding textures for an indoor environment of similar complexity to the L205 Office experiment detailed later in this paper.

Using the expanding textures method of evaluation cuts the total number of calculations down by an order of magnitude. For environments with uniform local cost of traversability between cells, this method still has a high level of redundancy, as cells closer to the middle of the expanding texture will be evaluated to the same value many times. Using a stencil buffer to limit evaluation to only wave front cells is mentioned in Future Work.

However, for a non-uniform environment, revaluation of cells not on the edge of the expanding texture is required for correcting cells with differing traversal costs. In particular, this applies for paths that are spatially longer (take up more cells) but have a lower cost due to efficient traversal between cells. Figure 4 shows a contrived example of this situation.

Figure 2. The first four iterations of the algorithm. The red outline represents the size of the expanding texture requested to be evaluated/rendered by the GPU each iteration.

Figure 3. Comparison of execution time with and without expanding textures.

Figure 4. Example of a non-uniform environment. Black cells represent non-traversable cells, blue cells have a traversal cost of one unit and red cells have a traversal cost of two units. Two possible paths from point A to point B are shown.
Figure 4 shows two possible paths from A to B. Given the cost of traversal $C_t$ between two cells, $x$ and $y$, with local costs $C_l(x)$ and $C_l(y)$ can be given by the following equation:

$$C_t = f(C_l(x), C_l(y)) = k \frac{C_l(x) + C_l(y)}{2}$$

where $k = 1$ for horizontal and vertical neighbours and $k = \sqrt{2}$ for diagonal neighbours, then the cost of the upper and lower paths is summarised in Table 3.

If the algorithm starts executing at B and begins to expand, at the seventh iteration it will first give cell A a global cost of 13.000. The wavefront that happens to be gradually evaluating the upper path will have just classified cell b1 in the seventh iteration. Two iterations later, the algorithm will re-evaluate cell a3 and give it the more favourable value of 9.828. Unlike traditional path planning algorithms, which expand based on lowest cost first, this algorithm expands on proximity or closest cell first, regardless of cost.

The inefficient re-evaluation of cell values due to an implementation limitation helps the more efficient wave fronts “wash over” already classified cells to make the resulting cost-to-go function more accurate. In Figure 4 this can also be seen in cell b3. As the algorithm evaluates the lower path, b3 is initially given a global cost of 11.5 from cell c3. When the upper path wave front eventually “washes around” the obstacle from cell a2 it gives cell b3 a better global cost of 10.949. A cost-to-go function is said to be mature if each cell had reached its final optimal value. Until the more efficient, but spatially longer, wave fronts wash over their sub-optimal counterparts the function is considered immature.

4.5. Reading Values from Graphics Hardware

Current consumer graphics card hardware can have a large bottleneck in reading texture data back from graphics memory into the main memory used by the CPU relative to algorithm execution time. The initial intention of this algorithm from a practical context was to make an identical function interface to the existing sequential algorithm so that there would be no modification to the other components of the system. That is, from an outsider’s view, both CPU and GPU based versions of the exhaustive cost-to-go evaluation would take in an occupancy grid, a local traversal cost grid and a destination cell and fill out the traversable regions of the grid with global cost-to-go values. As a result, in the CPU based implementation, the entire grid evaluation for further path planning is provided ready of planning in CPU accessible RAM. Although, internally, the evaluation step on a GPU is at least an order of magnitude better than the CPU version, the upload of values from graphical memory to main memory provided a bottleneck that can, on some systems, nullify the GPU benefit.

In practice, however, only the immediate area around an agent is required at any one time for path planning to occur. That is, even though a complete agent to destination evaluation is required to properly plan on a global scale, the short term values of the evaluated traversability grid can be used to plan a local immediate path. For example, Figure 5 shows how three agents plan a path to a common destination. Although the complete path from each agent to destination is calculated, it is only practical (even more so in dynamic environments) to decide upon a short term path. Therefore, only a small subregion of the cost-to-go function needs to be transferred back from GPU to CPU memory, as shown in the insets.

This removes the burden imposed by the hardware bottleneck while still allowing the GPU to evaluate an exhaustive cost-to-go function efficiently.\textsuperscript{5}

Table 3. Comparison of lower and upper path options with differing traversal costs.

<table>
<thead>
<tr>
<th>Path</th>
<th>Cost (3d.p.)</th>
<th>Number of cells</th>
</tr>
</thead>
<tbody>
<tr>
<td>upper</td>
<td>9.828</td>
<td>9</td>
</tr>
<tr>
<td>lower</td>
<td>13.000</td>
<td>7</td>
</tr>
</tbody>
</table>

Figure 5. Example of multiple agents planning paths to a common destination on campus. Even though the cost-to-go function needs to be globally evaluated from the destination, only a small immediate region around the agent is required at any instant to do short term path planning.

\textsuperscript{5}Of course, as graphics and motherboard hardware improves in terms of bus transfer speeds, a larger local area can be requested as required. A local cost-to-go is adequate for current field robotics applications.
4.6. Algorithm Termination

Depending on the density of obstacles in the environment and layout of traversable paths for the evaluation wave front to travel through, the algorithm can take a different number of iterations before it generates a complete cost-to-go of the entire grid, or at least a required sub grid. For example, an environment with a low proportion of obstacles will allow the algorithm to spread quickly. On the other hand, a spiral or snaking corridor arrangement will force more iterations as less cells are being evaluated each iteration.

Knowing when to terminate the algorithm is important for practical applications as you may want to have an exhaustive solution without having to do more calculations than required. The easiest approach is to look at the state space being evaluated and arbitrarily decide on a fixed number of iterations that would guarantee exhaustive evaluation. For example, for an $n \times n$ grid, choosing an iteration count of a nominal value like $5n$ would work, but will result in redundant calculations being done on environments with open spaces. The advantage of this method, however, is that there is no overhead in determining algorithm termination.

An alternate method of measuring the progression of the algorithm is to calculate either the maximum or sum of the differences of all cells between step $k-1$ and $k$. That is, if

$$\exists x \text{ where } C_{k-1}(x) \neq C_k(x)$$

then the wavefront is still progressing into previously unclassified cells. From an implementation point of view, this condition can be applied to the entire grid via either of the following conditions of termination:

$$\sum \| C_{k-1}(x_i) - C_k(x_i) \| = 0$$

$$\max \| C_{k-1}(x_i) - C_k(x_i) \| = 0$$

Both sum and maximum methods can theoretically run in $O(1) + O(\log n)$ time, although the maximum method provides more numerical stability on hardware that does not meet the full 32-bit floating point specification. Like the minimum cell value method mentioned above, this can also be run periodically to reduce overhead on execution time.

For environments that are explored and developed at a slow rate a hybrid method can be utilised. Such a method would periodically use a maximum of differences algorithm to check on the state for one complete evaluation. The number of algorithm iterations to complete this evaluation can then be stored and assumed as a set value for a number of cycles, as previously discussed in the arbitrary value method.

The algorithm could also potentially be terminated early by monitoring the cell an agent resides in or monitoring a user defined set of cells or subregions, which include an agent’s cell. When the subregion’s cells all contain non-negative values, a path exists from the subregion to the destination cell. This path may not be the optimal path, but under some circumstances it could be close enough to optimal for practical purposes. Of course, if the agent does not have a valid path to the destination then the algorithm would still need to terminate based on a secondary condition.

5. Experimental Results

The concurrent algorithm was applied to both simulated and real world environments to test both extreme cases and average operational cases, respectively. To test true practical benefit against an existing sequential algorithm, the execution or completion time of each test includes sending the occupancy grid data from CPU memory to graphical memory, performing the algorithm and copying a small portion of the texture back from graphical memory to perform CPU based path planning on. Images of the cost-to-go in the following experiments were generated via a CPU based simulator. Real GPU benchmarks were done in two stages. The first stage used an on board laptop graphics card (ATI Radeon HD 2400 XT) for more realistic test of practical applications. The second stage used a high end desktop graphics card (nVidia GeForce GTX 480) to show future potential of on board systems as well as potential live off board execution time.

The bench marked times are an approximation of the execution of one completion of the algorithm for a single occupancy grid as mentioned above. There is an inherent setup\(^5\) and shutdown time involved in executing the program, which should not be included in the timing of the completion of the algorithm. Therefore, the following formula was used to estimate the single completion of the algorithm.

$$T = \frac{\text{setup} + 512\text{completions} + \text{shutdown}}{512}$$

\(^5\)Path planning was not included in the experiment.
\(^6\)Setup time includes acquiring a render context, compiling and linking the kernel program and requesting handles to the textures and texture buffers. On the on board laptop mentioned in this section, this equated to approximately 0:21 seconds.
Each two-dimensional occupancy grid is generated from a three-dimensional laser range scan, fused with pose data derived from other sensor devices. Details of this configuration can be found in [2]. Figure 6 shows the three-dimensional point cloud data overlaid on the subsequently real-time generated black and white occupancy grid. A video of this process running in real time can be found in [28]. The following sections outline a sample of the experiments run.

5.1. UNSW Middle Campus

This experiment was initially run under a CPU based simulator, where exhaustive evaluation was achieved in 370 wavefront progressions. Figure 7 shows the original occupancy grid generated from point cloud data. Figure 8 shows the mature cost-to-go function. Table 4 shows the bench marked times of the algorithm calculating a single exhaustive mature cost-to-go function on both the laptop and desktop machines. A video of the algorithm running in the CPU based simulator can be found with [28].

5.2. L205 Office

Under the CPU based simulator this occupancy grid achieved exhaustive evaluation in 1118 iterations. Figure 9 shows the mature cost-to-go function from the simulator output. Table 5 shows the bench marked times of the algorithm calculating a mature cost-to-go function. The grid labelled normal sized denotes the original level of detail available from sensor processing to generate an occupancy grid. However, this would provide grid cells between 2.5 cm and 3.0 cm, which is more than enough for our UGVs. A quarter sized resolution occupancy grid was produced for benchmarking as it better matches the occupancy grid granularity required by our robots for successful navigation.10 The algorithm can be seen applied to the normal sized grid in [28].

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10The normal sized grid was used in the CPU simulation.
Figure 9. Cost-to-go function applied to a hybrid technical drawing and post-processed occupancy grid of the layout of walls and large obstacles the ME-L205 office at UNSW. The grid represents an area of approximately 33 m × 22 m and uses 1099 × 731 grid cells, resulting in an approximate grid size of 2.5 cm to 3 cm. As before, black pixels represent obstacles, indigo to red pixels represent increasing global cost and the red dot represents the destination location. Grey regions represent areas that are unreachable from the destination location and have hence remained unclassified.

Table 4. Comparison of the execution time of the parallel algorithm on different machines.

<table>
<thead>
<tr>
<th></th>
<th>On Board Laptop</th>
<th>Desktop</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>4.498s</td>
<td>0.0478s</td>
</tr>
</tbody>
</table>

Table 5. Comparison of the execution time of the parallel algorithm on different machines for normal and quarter sized occupancy grids.

<table>
<thead>
<tr>
<th>Size</th>
<th>On Board Laptop</th>
<th>Desktop</th>
</tr>
</thead>
<tbody>
<tr>
<td>Normal Size</td>
<td>11.113s</td>
<td>0.0580s</td>
</tr>
<tr>
<td>Quarter Size</td>
<td>0.714s</td>
<td>0.0362s</td>
</tr>
</tbody>
</table>

6. Future Work

The next step of this research will be to see if the stencil or depth buffer can be used to cull already classified cells within the bounds of the expanding texture at the current iteration. This would then enable the kernel to only be evaluated on cells in the wave front rather than those simply within the expanding texture. The stencil buffer is traditionally used to render pixels to screen that pass a stencil test. The test is based on a per-pixel value in the stencil buffer and is optimised to run outside the kernel execution in hardware.

Another direction this work is heading in is to expand the algorithm to a third dimension so that non-holonomic path planning can be done with, for example, \((x, y, \theta)\) like co-ordinates. It is also under investigation whether higher dimensional problems can be mapped to a two-dimensional equivalent for use in the proposed algorithm.

For this to happen, an efficient algorithm is to be investigated that can map an undirected weighted graph to the grid structure for efficient cost generation. As a preview of the work Figure 10 shows how a given graph can be mapped to “channels” in a grid structure for evaluation as a traditional occupancy grid. The channel approach enforces one path between nodes that are connected in the original graph, but limits the number of paths from each node to four. If a node is more than degree four it can be split into two or more virtual nodes connected by zero cost channels.

A CUDA [24] implementation will also be investigated from an implementation point of view, however the core algorithm will remain the same.

7. Conclusions

This paper presented a parallel implementation of a common sequential algorithm in the field of path planning using dynamic programming techniques. It provides an order of magnitude improvement on computation time for particularly bad cases and many orders of magnitude for low obstacle density environments. In practice the implementation has provided an option that remains real-time responsive while allowing for the additional option of either larger traversability grids, more responsive calculation or more fine grained grid structures tending towards approximating continuous space.

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REFERENCES


