# GPU implementation of a region based algorithm for large images segmentation

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Abstract—Image segmentation is one of the most challenging issues in image computing. In this work, we focus on region-based active contour techniques (snakes) as they seem to achieve a high level of robustness and fit with a large range of applications. Some algorithmic optimizations provide significant speedups, but even so, execution times are still non-neglectable with the continuing increase of image sizes. Moreover, these algorithms are not well suited for running on multi-core CPU's. At the same time, recent developments of Graphical Processing Units (GPU) suggest that higher speedups could be obtained by use of their specific design. We have managed to adapt a specially efficient snake algorithm that fits recent Nvidia GPU architecture and takes advantage of its massive multithreaded execution capabilities. The speedup obtained is most often around 7.

## Keywords-GPU; segmentation; snake;

#### I. INTRODUCTION

Segmentation and shape detection are still key issues in image computing. These techniques are used in numerous fields ranging from medical imaging to video tracking, shape recognition or localization. Since 1988, the active contours (snakes) introduced by and Kass et al. [1], have proved to be efficient and robust, especially against noise, for a wide range of image types.

The main shortcoming of these algorithms is often their high dependence on the initial contour, though several contributions have lowered this dependency and also brought more accurate segmentation of non convex shape [4].

The information that drives a contour model comes either from the contour itself or from the characteristics of the regions it defines. For noisy images, the second option is often more suitable as it takes into account the statistical fluctuations of the pixels. One approach [4] proposes a geometric (polygonal) region-based snake driven by the minimization of the likelihood (ML).

An important issue of image processing, especially segmentation, has always been the computation time of most algorithms. Over the years, the increase of CPU computing capabilities, although quite impressive, has not been able to fulfill the combined needs of growing resolution and real-time computation. Since having been introduced in the early 1980's, the capabilities and speed of graphics accelerators have always been increasing. So much so that the recent GPGPU (General Purpose Graphic Processing Units) currently benefit by a massively parallel architecture for general purpose programming, especially when dealing with large matrices or vectors. On the other hand, their specific design obviously imposes a number of limitations and constraints. Some implementations of parametric snakes have already been tested, such as [6]. However, a similar solution (computation per small tile) is not suited for the algorithm we have implemented.

Our goal was to propose a way to fit such a snake algorithm to the Nvidia<sup> $\bigcirc$ </sup> Tesla GPU architecture. The remainder of this paper presents the principles of the algorithm and notations in section II. In section III, the details of the sequential CPU implementation are explained. Section IV summarizes Nvidia's GPU important characteristics and how to deal with them efficiently. Then sections V and VI detail our GPU implementation and timing results. Finally, the conclusion of section VII evaluates the pros and cons of this implementation and then gives a few direction to be followed in future works.

#### II. SEQUENTIAL ALGORITHM : OUTLINES

The goal of the active contour segmentation method (snake) we studied [4] is to distinguish, inside an image I, a target region T from the background region B. The size of Iis L x H pixels of coordinates (i, j) and gray level z(i, j). Zrepresents the gray levels data of I. We assume that the gray levels of T and B are vectors of independent and identically distributed values, each with a probability density function (PDF)  $p^{\Omega}$  ( $\Omega \in \{T; B\}$ ). The present implementation uses a Gaussian PDF, but another one can easily be used as Gamma, Poisson,...(Cf. [4])...

The *active contour* S, which defines the shape of T is chosen as polygonal. The purpose of the segmentation is then to determine the shape that optimizes a generalized log-likelihood-based criterion (GL). This is done by an iterative

process which is initialized with an arbitrary shape, then at each step :

- 1) it modifies the shape
- 2) it estimates the parameters of the Gaussian functions for the two regions and evaluates the criterion.
- 3) it validates the new shape if the criterion has a better value.

A simplified description of it is given in *Algorithm 1* which features two nested loops : the main one, on iteration level, is responsible for tuning the number of nodes ; the inner one, on step level, takes care of finding the best shape for a given number of nodes. *Figure 1* shows intermediate results at iteration level. Sub-figure *1a* shows the initial rectangular shape, *1b* shows the best four-node shape that ends the first iteration. Sub-figures *1c* and *1d* show the best shape for an eight-node contour (resp. 29-node) which occurs at the end of the second iteration (resp. fourth).

Algorithm 1: Sequential algorithm : outlines								
1: begin with a rectangular 4 nodes contour;								
2: repeat /* iteration leve								
3:	repeat	/* step level */						
4:	Test some other positions for each node, near							
	its current position;							
5:	Find the best GL and adjust the node's position;							
6:	until no more node can be moved;							
7:	Add a node in the middle of each long enough							
	segment;							

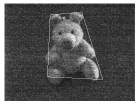
s: until no more node can be added;



(a) Initial contour



(c) End of second iteration (8 nodes)



(b) End of first iteration (4 nodes)



(d) End of fourth iteration (29 nodes)

Figure 1. segmentation of a noisy image

## **III. SEQUENTIAL ALGORITHM : DETAILS**

A. Criterion

Let  $p^{\Omega}$  be a Gaussian PDF. Its vector of parameters  $\Theta_{\Omega}$  $(\Omega \in \{T; B\})$  has two components, the average value  $\mu$  and the standard deviation  $\sigma$ . The likelihood for the regions  $\Omega$  $(\Omega \in \{T; B\})$  is given by

$$P[Z|T, B, \Theta_T, \Theta_B] = P(Z|T, \Theta_T)P(Z|B, \Theta_B)$$

where

$$P(Z|\Omega,\Theta_{\Omega}) = \prod_{(i,j)\in\Omega} p^{\Omega}[z(i,j),\Theta_{\Omega}] \quad (\Omega \in \{T;B\})$$

The log-likelihood of the region  $\Omega$  is then

$$-N_{\Omega} \log \left(\sqrt{2\pi}\right) - N_{\Omega} \log \left(\sigma\right) - \frac{1}{2\sigma^2} \sum_{(i,j)\in\Omega} \left(z(i,j) - \mu\right)^2$$

inside which the vectors of parameters  $\Theta_\Omega$  are determined by ML estimation

$$\widehat{\Theta_{\Omega}} \left( \begin{array}{c} \widehat{\mu_{\Omega}} = \frac{1}{N_{\Omega}} \sum_{\substack{(i,j) \in \Omega \\ \sigma_{\Omega}^2}} z(i,j) \\ \widehat{\sigma_{\Omega}^2} = \frac{1}{N_{\Omega}} \sum_{\substack{(i,j) \in \Omega \\ (i,j) \in \Omega}} (z(i,j) - \widehat{\mu_{\Omega}})^2 \end{array} \right)$$

Considering the two regions, the criterion to be optimized is then, up to a constant, the Generalized Likelihood (GL):

$$GL = \frac{1}{2} \left( N_B \log \left( \widehat{\sigma_B}^2 \right) + N_T \log \left( \widehat{\sigma_T}^2 \right) \right)$$

## B. CPU implementation

Let  $S_{n,l}$  be the polygonal contour state at step l of iteration n, and  $S_{n,l}^i$  the node i of  $S_{n,l}$  ( $i \in [0, N_n]$ ).  $S_{n,l}^{i,w}$  is the neighbor of index  $\omega$  of the node  $S_{n,l}^{i}$  in a 8connexity meaning with d pixels scope. Each segment of  $S_{n,l}$  is considered as an oriented list of discrete points. Chesnaud & Réfrégier, based on the Green-Ostogradski theorem, have shown how to replace the 2 dimensions (2D) sums needed to estimate  $\Theta_{\Omega}$  by 1 dimension sums along  $S_{n,l}$ [4]. This approach leads to compute a pair of transformed images, called cumulated images, at the very beginning of the process, which are then used as lookup tables. It also involves weighting coefficients for pixels and segments of the contour. See [4] for details. Therefore, beyond this point, we will talk about the contribution of each point to the 1D sums. By extension, we also talk about the contribution of each segment to the 1D sums.

A more detailed description of the sequential algorithm is given by *Algorithm 2*. The process starts with the computation of cumulated images ; an initialization stage takes place from line 3 to line 9. Then we recognize the two nested loops (line 10 and line 11) and finally the heart of the algorithm stands on line 15 which represents the main part of the calculations to be done :

- 1) compute the various sums without the contributions of both segments connected to current node  $S_{n,l}^i$ .
- 2) compute the contributions of both segments, which requires :
  - To determine the coordinates of every discrete pixel of both segments connected to  $S_{n,l}^{i,w}$ .
  - To compute every pixel contribution.
  - To sum pixel contributions to obtain segment contributions.
- 3) compute the GL given the contribution of each segment of the tested contour.

## Algorithm 2: Sequential simplified algorithm

1: read image from HDD; 2: compute\_cumulated\_images(); 3: iteration  $n \leftarrow 0$ ; 4:  $N_0 \leftarrow 4$ ; 5:  $S_{n,l} \leftarrow S_{0,0};$ 6: step  $d \leftarrow d_{max}$  an arbitrary power of 2 value; 7: current node  $S_{0,0}^i \leftarrow S_{0,0}^0$ ; **8**:  $l \leftarrow 0$ ; 9: compute  $GL_{ref}$ , the GL of  $S_{n,0}$ ; 10: repeat /\* iteration level, n index \*/ /\* step level, l index \*/ repeat 11: for i = 0 to  $N_n$  do 12:  $S_{n,l}^{i,w}$  ( $w \in [0;7]$ ) are the neighbors of  $S_{n,l}^{i}$ 13: by d pixels; for w = 0 to 7 do 14: compute  $GL_w$  for  $S_{n,l}$  when  $S_{n,l}^{i,w}$ 15: replaces  $S_{n,l}^i$ ; if  $GL_w$  is better than  $GL_{ref}$  then 16:  $\begin{aligned} GL_{ref} \leftarrow GL_w; \\ \text{move node } S^i_{n,l} \leftarrow S^{i,w}_{n,l}; \end{aligned}$ 17: end 18: 19: end  $l \leftarrow l + 1;$ 20: until no node move occured; 21: add new nodes,  $N_n \leftarrow N_n + N_{newnodes}$ ; 22: if d > 1 then  $d \leftarrow d/2$  else d = 1; 23:  $n \leftarrow n+1$ ; 24: compute  $GL_{ref}$ , the GL of  $S_{n,0}$ ; 25: 26: until no new node added;

The profiling results of the CPU implementation shown in *Figure 2* display the relative costs of the most timeconsumming functions. It appears that more than 80% of the total execution time is always spent by only three functions :

- compute\_segment\_contribution() which is responsible for point 2 above,
- compute\_cumulated\_images() which computes the 3 lookup tables at the very beginning,

 compute\_pixels\_coordinate() which is called by compute\_segment\_contribution().

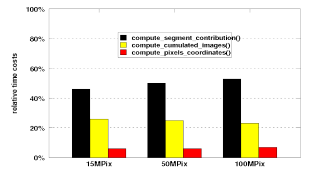


Figure 2. the three most-consumming functions for various image sizes

Measurements have been performed for several image sizes from 15 MPixels (about 3900 x 3900) to 144 MPixels (about 12000 x 12000). On the one hand, we can notice that function compute\_segment\_contribution() always lasts more than 45% of the total running time, and even more when the image gets larger. On the other hand, the function compute\_cumulated\_images() costs more than 23%, decreasing with image size, while function compute\_pixels\_coordinate() always takes around 6%. It confirms that the need for parallelization resides in line 15 and line 2 of Algorithm 2 as they contain every call to those three functions.

The following sections detail how we managed to implement these time-consumming functions in parallel, but a brief reminder on GPU's recent architecture is presented first.

# IV. NVIDIA'S GPU ARCHITECTURE

GPUs are multi-core, multi-threaded processors, optimized for highly parallel computation. Their design focuses on a Single Instruction Multiple Threads (SIMT) model by devoting more transistors to data processing rather than datacaching and flow control [7].

For example, Figure 3 shows a Tesla C1060 with its 4GB of global memory and 30 SM processors, each including :

- 8 Scalar Processors (SP)
- a Floating Point Unit (FPU)
- a parallel execution unit (SIMT) that runs threads by warps of 32.
- 16KB of shared memory, organized in 16 banks of 32 bits words

Nvidia uses a parameter called the *compute capability* of each GPU model. Its value is composed of a major number and a minor number ; for example the C1060 is a sm13 GPU (major=1 minor=3) and C2050 is a sm20 GPU.

The recent Fermi cards (eg. C2050,) have improved performances by supplying more shared memory in a 32 banks array, a second execution unit and several managing

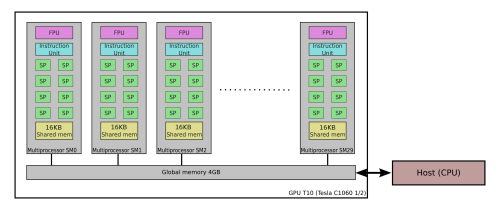


Figure 3. schematic diagram of GPU's internal architecture

capabilities on both the shared memory and level 1 cache memory ([7], [8], [9]. However, writing efficient code for such architectures is not obvious, as re-serialization must be avoided as much as possible. Thus, when designing, one must keep a few key points in mind :

- the CUDA model organizes threads by a) threads blocks in which synchronization is possible, b) a grid of blocks with no possible synchronization between blocks.
- there is no way to know in what order the blocks are to be scheduled during one single kernel execution.
- data must be kept in GPU memory, to reduce the overhead due to copying between CPU and GPU memories.
- the total amount of threads running the same computation must be maximized.
- the number of execution branches inside a block should be reduced as much as possible.
- global memory accesses should be coalescent, *ie*. memory accesses done by physically parallel threads (16 at a time) must be consecutive and contained in a 128 Bytes range.
- shared memory is organized by 16 x 32 bits wide banks. To avoid bank conflicts, each parallel thread (16 at a time) must access a different bank.

All the above charasteristics make it always a quite constrained problem to solve when designing a GPU code. Moreover, a non suited code would probably run even slower on GPU than on CPU due to the automatic serialization which would be done at run time.

## V. GPU IMPLEMENTATION

In the implementation described below, pre-computations and proper segmentation are discussed separately. To keep data in GPU memory, the whole computation is assigned to the GPU. CPU still hosts :

- data reading from HDD
- data writing on HDD if needed
- main loops control (corresponding to lines 10 and 11 of Algorithm 2)

It must be noticed that controlling these loops is achieved with only a very small amount of data being transferred between host (CPU) and device (GPU), which does not produce high overhead.

Morever, the structures described below need 20 Bytes per pixel of the image to process (plus an offset of about 50 MByte). It defines the maximum image size we can accept : approximately 150 M Pixels.

## A. Pre-computations

To replace 2D sums by 1D sums, Chesnaud *et al.* [4] have shown that the three matrices below should be computed :

$$C_1(i,j) = \sum_{k=0}^{k=j} (1+k)$$
$$C_2(i,j) = \sum_{k=0}^{k=j} z(i,k)$$

and

$$C_{z^2}(i,j) = \sum_{k=0}^{k=j} z^2(i,k)$$

Where z(i, k) is the gray level of pixel of coordinate (i, j), so that  $C_1$ ,  $C_z$  and  $C_{z^2}$  are the same size as image I.

First, we chose not to generate  $C_1(i, j)$ , which requires that values should be computed when needed, but saves global memory and does not lead to any overhead. The computation of  $C_z$  and  $C_{z^2}$  easily decomposes into series of *inclusive prefixsums* [10]. However, by keeping the *1 thread per pixel* rule, as the total number of threads that can be run in a grid cannot exceed  $2^{25}$  (Cf. [7]), slicing is necessary for images exceeding a size threshold which can vary according to the GPU model (e.g. 33 MPix for sm13 GPU, eg. C1060). It's quite easy to do, but it leads to a small overhead as the process requires multiple calls to one kernel. Slicing can be done in two ways :

• all slices are of the same size (balanced)

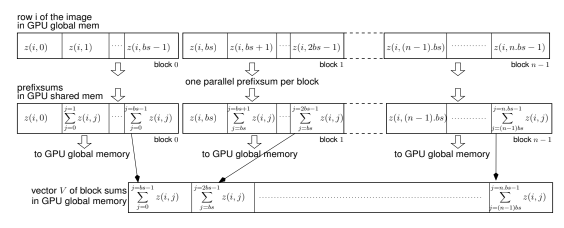


Figure 4. compute\_blocks\_prefixes() details.

• slices fit the maximum size allowed by the GPU, leaving one smaller slice at the end of the process (full-sized).

The balanced slice option has proved to run faster.

For example : if a given image has 9000 lines and the GPU can process up to 4000 lines at a time, it's faster to run 3 times with 3000 lines rather than twice with 4000 and once with 1000.

As the sums in  $C_z$  and  $C_{z^2}$  are row-wide, it is easy to see that every block-wide sum will be needed before being able to use it in the global sum. But as mentioned earlier, the scheduling of blocks must be considered as random. So, in order to ensure synchronizations, each row of the original image is then treated by three different kernels :

- compute\_blocks\_prefixes().
- scan\_blocksums().
- add\_sums2prefixes().

Figures 4, 5 and 6 show relevant data structures for a given row i of I. We assume that each thread block runs bs threads in parallel and each row of  $C_z$  needs n blocks to cover its L pixels.

Figure 4 shows the details of the process for row i of the original image I, already stored in GPU global memory. Operands are first copied into GPU shared memory for efficiency reasons. An inclusive prefixsum is then performed inside each independant thread block. At this point, only the first shared memory block contains the final values. Its last element contains the sum of all elements in the corresponding block of I. In order to obtain the right values for the row i of  $C_z$ , every element value in the other blocks must then be summed with an offset value. This offset value is the sum of all element values in every corresponding previous block of row i.

As the scheduling of blocks is fully unpredictable, the necessary intermediate results have to be stored in GPU global memory before exiting from kernel. Each element of the prefixsums in GPU shared memory has been stored in its corresponding position in  $C_z$  (GPU global mem), along with the vector of block sums which will be passed later to the next kernel scan\_blocksums().

The kernel scan\_blocksums () (Figure 5) only makes an exclusive prefixsum on the vector of block sums described above. The result is a vector containing, at index x, the value to be added to every element of block x in each line of  $C_z$ .

This summing is done in shared memory by kernel add\_sums2prefixes() as described by Figure 6.

The values of  $C_{z^2}$  are obtained together with those of  $C_z$  and in exactly the same way. For publishing reasons, figures do not show the  $C_{z^2}$  part of structures.

With this implementation, speedups are quite significant (Table I). Moreover, the larger the image, the higher the speedup is, as the step-complexity of the sequential algorithm is of  $O(N^2)$  and  $O(N \log(N))$  for the parallel version. Even higher speedups are achieved by adapting the code to specific-size images, especially when the number of columns is a power of 2. This avoids inactive threads in the grid, and thus improves efficiency. However, since the use of 64-bit sums is imposed by image sizes (up to 12000 pixel wide) and 16-bit pixel coding, computations are made with a 2-way bank conflict as sums are based on 64-bit words, thus creating overhead.

#### B. Segment contributions

The choice made for this implementation has been to keep the *1 thread per pixel* rule for the main kernels. Of course, some reduction stages need to override this principle and will be pointed out.

As each of the  $N_n$  nodes of the contour  $S_{n,l}$  may move to one of the eight neighbor positions as shown in *Figure* 7, there is  $16N_n$  segments whose contribution has to be estimated. The best combination is then chosen to obtain  $S_{n,l+1}$ (Figure 7). Segment contributions are computed in parallel by kernel GPU\_compute\_segments\_contrib().

The grid parameters for this kernel are determined according to the size of the longest segment  $npix_{max}$ . If  $bs_{max}$  is

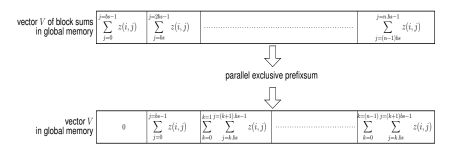


Figure 5. scan\_blocksums() details.

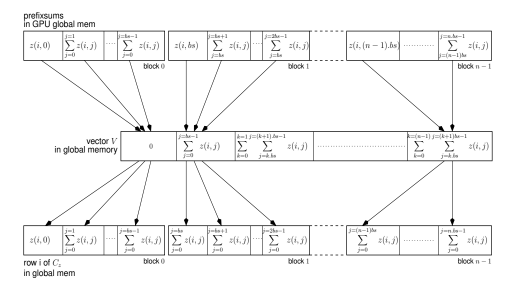


Figure 6. add\_sums2prefixes() details.

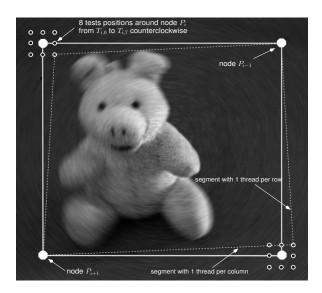


Figure 7. Optimization of node locations using 8 position tests around each node.

the maximum theoritical blocksize that a GPU can accept,

• the block size bs is taken as

- $npix_{max}$ 's next power of two if
- $npix_{max} \in [33; bs_{max}]$
- 32 if  $npix_{max} < 32$
- $bs_{max}$  if  $npix_{max} > 256$
- the number of threads blocks assigned to each segment,  $N_{TB} = \frac{npix_{max} + bs - 1}{bs}$

Our implementation makes intensive use of shared memory and does not allow the use of the maximum theoritical blocksizes (512 for sm13, 1024 for sm20, see [8] and [7]). Instead we set  $bs_{max}^{sm13} = 256$  and  $bs_{max}^{sm20} = 512$ . Anyway, testing has shown that most often, the best value is 256 for both *sm13* and *sm20* GPU's.

Then GPU\_compute\_segments\_contrib() computes in parallel :

- each pixel coordinates for all  $16N_n$  segments. Since the contour is only read in one direction, we have been able to use a very simple parallel algorithm instead of Bresenham's. It is based on the slope k of each segment : one pixel per row if |k| > 1, one pixel per column otherwise.
- each pixel contribution by reading the corresponding values in the lookup tables.

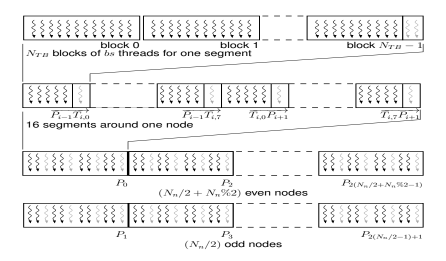


Figure 8. structure for segments contributions computation. Gray symbols help to locate inactive threads as opposed to black ones that figure active threads.

• each thread-block sum of individual pixel contributions by running a *reduction* stage for each block.

The top line of Figure 8 shows the base data structure in GPU shared memory which is relative to one segment. We concatenate the single segment structure as much as necessary to create a large vector representing every pixel of every test segment. As each segment has a different size (most often different from any power of two), there is a non-neglectable number of inactive threads scattered in the whole structure. Two stages are processed separately : one for all even nodes and another one for odd nodes, as shown in the two bottom lines of Figure 8.

The process is entirely done in shared memory ; only a small amount of data needs to be stored in global memory for each segment :

- the coordinates of its middle point, in order to be able to add nodes easily if needed.
- the coordinates of its first and last two points, to compute the slope at each end of the segment.

The five values above are part of the weighting coefficients determination for each segment and node.

The GPU\_sum\_contribs() takes the blocks sums obtained by GPU\_compute\_segments\_contrib() and computes a second stage parallel summing to provide the  $16N_n$  segment contributions.

# C. Segments with a slope k such as $|k| \leq 1$

Such a segment is treated with 1 thread per column and consequently, it often has more than one pixel per row as shown by Figure 9. In an image row, consecutive pixels which belong to the target define an interval which can only have one low and one high ends. That's why, on each row, we choose to consider only the contributions of the innermost pixels. This selection is also done inside GPU\_compute\_segments\_contrib() when reading the lookup tables for each pixel contribution. We simply set a null contribution for pixels that need to be ignored.

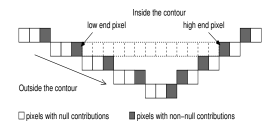


Figure 9. Zoom on part a of segment with |k| < 1, at pixel level.

#### D. Parameters estimation

A GPU\_compute\_GL() kernel computes in parallel :

- every  $8N_n$  vector of parameters values corresponding to each possible next state of the contour. Summing is done in shared memory but relevant data for these operations are stored in global memory.
- every associated pseudo likelihood value.
- every node substitution when better GL have been found and if it does not lead to segments crossing.

#### E. End of segmentation

Segmentation is considered achieved out when no other node can be added to the contour (Algorithm 3). A very simple GPU kernel adds every possible node and returns the number it added.

#### VI. SPEEDUPS

Results are given in Table I. CPU timings were measured on an Intel Xeon E5530-2.4GHz with 12Go RAM (LIFC cluster). GPU timings were obtained on a C2050 GPU with 3GB RAM (adonis-11.grenoble.grid5000.fr).

Execution times reported are means on ten executions. The

A	gorithm 3: Par	rralel GPU algorithm : outlines.						
<<	<<<>>> indicates a GPU kernel parallel process.							
1: le	oad images;							
2: t1	2: transfer image from CPU to GPU;							
3: <	3: << <compute 2="" cumulated="" images="" the="">&gt;&gt;;</compute>							
4: << <i contour="" the="">&gt;&gt;;</i>								
5: r	repeat /* iteration level */							
6:	repeat	/* step level */						
7:	<< <find be<="" td=""><td colspan="5">&lt;&lt;<find best="" contour="" neighbor="">&gt;&gt;;</find></td></find>	<< <find best="" contour="" neighbor="">&gt;&gt;;</find>						
8:	<< <adjust i<="" td=""><td colspan="5">&lt;&lt;<adjust node's="" positions="">&gt;&gt;;</adjust></td></adjust>	<< <adjust node's="" positions="">&gt;&gt;;</adjust>						
9:	transfer the	transfer the number of moves achieved from						
	GPU memory to CPU memory.							
10:	<b>until</b> no more node can be moved;							
11:	<< <add nodes="">&gt;&gt;;</add>							
12:	transfert the number of nodes added from GPU							
	memory to CPU memory.							

13: **until** no more node can be added;

image of figure 1a (scaled down for printing reasons) is based on a real noisy image ( $800 \times 800$ ), 16-bit gray level. Contrast has been enhanced for better viewing ; its various sizes have been obtained by interpolation and addition of gaussian noise.

We separately give the timings of pre-computations as they are a very general purpose piece of code. Segmentations have been performed with strictly the same parameters (initial shape, threshold length). The neighborhood distance for the first iteration is 32 pixels. It has a slight influence on the time process, but it leads to similar speedups values of approximately 7 times faster than CPU.

Though it does not appear in Table I, we observed that during segmentation stage, higher speedups are obtained in the very first iterations, when segments are made of a lot of pixels, leading to a higher parallelism ratio.

Several parameters prevent from achieving higher speedups :

- accesses in the lookup tables in global memory cannot be coalescent. It would imply that the pixel contributions of a segment are stored in consecutive spaces in  $C_z$  and  $C_{z^2}$ . This is only the case for horizontal segments.
- the use of 64-bit words for computations in shared memory often leads to 2-way bank conflicts.
- the level of parallelism is not so high, ie. the total number of pixel is not large enough to achieve impressive speedups. For example, on C2050 GPU, a grid can run about 66 million of threads, but a contour in a 10000 x 10000 image would be less than 0.1 million pixel long.

		CPU	GPU	Speedup
Image 15MP	total	0.51 s	0.06 s	x8.5
	pre-comp.	0.13 s	0.02 s	x6.5
	segment.	0.46 s	0.04 s	x11.5
Image 100MP	total	4.08 s	0.59 s	x6.9
	pre-comp.	0.91 s	0.13 s	x6.9
	segment.	3.17 s	0.46 s	x6.9
Image 150Mp	total	5.7 s	0.79 s	x7.2
	pre-comp.	1.4 s	0.20 s	x7.0
	segment.	4.3 s	0.59 s	x7.3

Table I GPU (C2050, sm20) vs CPU timings.

## VII. CONCLUSION

The algorithm we have focused on is not easy to adapt for high speedups on GPGPU, though we managed to make it work faster than on CPU. The main drawback is clearly its relative low level of parallelism. Nevertheless, we proposed different kernels that allowed us to take advantage of the computation power of GPUs. In future works, we plan to try and manage to benefit from larger computing grids of thread blocks. Among the possible solutions, we plan to work on:

- slicing the image and proceeding the parts in parallel. This is made possible since sm20 GPU provide multi kernel capabilities.
- slicing the image and proceeding the parts on two different GPUs, hosted by the same CPU.

To extend the scope of this work beyond our present hypothesis (based on *single* target segmentation), we are also going to investigate achieving speedups in *multiple* target segmentation of large images. This might be useful in a wide range of applications.

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