Parallel implementation of flow and matching algorithms

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Abstract

In our work we present two parallel algorithms and their lock-free implementations using a popular GPU environment Nvidia CUDA. The first algorithm is the push-relabel method for the flow problem in grid graphs. The second is the cost scaling algorithm for the assignment problem in complete bipartite graphs.

1 Introduction

The maximum flow problem has found many applications in various computer graphics and vision problems. For example, the algorithm for the graph cut problem is an optimization tool for the optimal MAP (Maximum A-Posteriori Probability) estimation of energy functions defined over an MRF (Markov Random Field) [4, 11, 12, 13, 17]. Another new and most interesting for us idea consists in computing optical flow by reducing it to the assignment (weighted matching) problem in bipartite graphs ([18]). Therefore it is important to look for new solutions to improve the execution time of algorithms solving the max flow and related problems.

The new approach to acceleration of algorithms uses the power of GPU (graphics parallel units). This has motivated us to pose the main purpose of this research: find an efficient parallel algorithm for the weighted matching problem and implement it in a popular GPU environment Nvidia CUDA. Naturally, such an algorithm can use max flow computation techniques and the easiest method to compute maxflow in parallel is the push-relabel algorithm. Hence in the first part of our work we develop our own CUDA implementation of Hong’s lock-free push-relabel algorithm. It can be used to find graph cuts in graphs constructed by Kolmogorov et. al. in [12] and is suitable for minimization of any energy function of the class characterized by these authors.

In the second part of the paper we focus on the assignment problem, i.e. the max weight matching problem. The cost scaling algorithm solving this problem has also found applications in many computer vision tasks such as recognition and tracking of images. Finally we present our implementation
of the cost scaling algorithm for the assignment problem, where the core
*refine* procedure is implemented lock-free on CUDA.

This work is organized as follows. Section 2 contains all needed defi-
nitions in our paper. Section 3 introduces the CUDA programming envi-
rонment. Section 4 presents the sequential push-relabel algorithm and its
parallel counterparts: a blocking version of the parallel push-relabel algo-

rithm, presented by Vineet and Narayanan [4], and a lock-free push-relabel
algorithm by Hong [5]. In the end of this section we present our CUDA
implementation of the lock-free push-relabel algorithm for grid graphs. Section
5 starts from the presentation of two sequential algorithms: a scaling
minimum-cost flow method and its counterpart for the assignment problem,
both developed by Goldberg et al. [2,8,9]. Then we present our own par-
allel cost scaling algorithm for the assignment problem using the lock-free
push-relabel algorithm. In the end of section 5 we present our CUDA imple-
mentation of this algorithm for arbitrary graphs. Figure 1 shows the diagram
of the reductions between the main analyzed problems.

2 CUDA Programming

CUDA (Compute Unified Device Architecture) is a parallel computing
architecture for Nvidia GPUs. In the past the GPU was used only as the co-
processor of the CPU to reply on the many graphics tasks in real-time. Now,
thanks to increasing computation power of GPUs, they are also very efficient
on many data-parallel tasks. Therefore GPUs are used also for many non-
graphics applications like the push-relabel max flow algorithm.

We present two algorithms implemented in CUDA 4.0. Both were tested

Figure 1: The algorithms described in Section 5
on Nvidia GTX 560 Ti (i.e. on a device of compute capability 2.1).

The programs operating on the GPU are implemented in the CUDA C language, an extension of C. CUDA C allows to define C functions called kernels that can be executed in parallel. The C program is run on the host (or CPU) by the host thread. The kernels (CUDA programs) are launched by the host thread and run on the device (or GPU) by many CUDA threads. The number of threads executing a kernel is defined by the programmer. The running threads are split into three-dimensional blocks. The set of blocks forms a three-dimensional grid. An example of a kernel declaration is:

```c
__global__ void kernel_function(int* data);
```

which must by called like this:

```c
dim3 gD = 10; /* dimension of grid, unspecified component is initialized to 1*/
dim3 bD = (32, 8, 1); /* dimension of block */
kernel_function <<gD, bD>>(someArray);
```

Each thread executing a kernel is given the following built-in three-dimensional variables:

```c
dim3 threadIdx = (threadIdx.x, threadIdx.y, threadIdx.z) /* unique for each thread in the block */
dim3 blockIdx = (blockIdx.x, blockIdx.y, blockIdx.z) /* unique for each block in grid */
dim3 blockDim = (blockDim.x, blockDim.y, blockDim.z)
dim3 gridDim = (gridDim.x, gridDim.y, gridDim.z)
```

Hence for each thread we can calculate its unique index in grid in the following way:

```c
int threadsInBlock = blockDim.x * blockDim.y;
int numberOfBlockInGrid = (blockIdx.y * gridDim.x) + blockIdx.x;
int numberOfThreadInBlock = (threadIdx.y * blockDim.x) + threadIdx.x;
nthid_id = (threadsInBlock * numberOfBlockInGrid) + numberOfThreadInBlock;
```

The host and the device have separated memory spaces called the host memory and the device memory, both residing in the dynamic random-access memory (DRAM). There are several types of device memory: global, local, shared, constant and texture. There are also registers. The shared memory and the registers are located on the chip of the GPU. The others are located off the chip so they have large access latency. However the sizes of the shared memory and the registers are much smaller than of the others. Note
that since the local memory is located off chip then its access latency is also big.

The scopes and the lifetimes of the local memory and the registers are restricted to one thread. The scope of the shared memory and its lifetime is restricted to all threads of the block. The lifetime and access to other of memory are available for all launched threads and the host.

For devices of compute capability 2.x the local and the global memory are cached. There are two types of cache: an L1 cache for each multiprocessor and an L2 cache shared by all multiprocessors on the GPU. The size of the L2 cache is fixed equal to 768 KB. The L1 cache and the shared memory are stored in the same on-chip memory and their initial sizes are 48 KB of shared memory and 16 KB of L1. These values can be reversed by the function `cudaFuncSetCacheConfig()` (or `cuFuncSetCacheConfig()` for Driver API).

In our implementations the shared memory is useless and its size that we use is not bigger than 16 KB. However, we use the first configuration because it gives better running times.

The threads can be synchronized in the scope of a block by the function `__syncthreads()`. It sets a semaphore which causes that the execution of the further code waits until all parallel threads reach the specified point.

In our implementations we use atomic functions: `atomicAdd()` and `atomicSub()`, available for devices of compute capability 2.x, which perform a read-modify-write operations on 64-bit words residing in the global memory. The atomic operations are slower than their non-atomic counterparts but they allow us to implement the programs without any synchronization of the threads.

A bandwidth is the rate at which data can be transferred. The bandwidth between the global memory on device and the global memory on the host is much smaller than the bandwidth between the global memory on device and the memory space on GPU. Therefore it is important to minimize the data transfer between the device and the host. In our implementation we strived to reduce the copying only to necessary arrays of data.

To allocate and deallocate memory on the device we use the functions `cudaMalloc()` and `cudaFree()` and to copy memory between the device and the host we use the function `cudaMemcpy()`.

In our implementations we use the cutil.h library available in the GPU computing SDK which allows us to detect the errors returned by the device.

### 3 Basic Graph Definitions

Let $G = (V, E)$ be a directed graph and $u : E \rightarrow \mathbb{R}^+$ be an edge capacity function. For formal reasons, if $(x, y) \notin E$ then we set $u(x, y) = 0$. Let $s \neq t$ be two distinguished vertices in $V$, the source and the sink, respectively. Then the triple $F = (V, E, u, s, t)$ is called a flow network.
A pseudoflow is a function \( f : V \times V \rightarrow \mathbb{R} \) such that, for each \( (x, y) \in E \), \( f \) satisfies the capacity constraints: \( f(x, y) \leq u(x, y) \), and the skew symmetry: \( f(x, y) = -f(y, x) \). We say that the pseudoflow \( f \) is a flow if for each node \( x \in V - \{s, t\} \), \( \sum_{(x,y) \in E} f(x, y) = 0 \). We say that an edge \((x, y)\) is in the flow \( f \), if \( f(x, y) > 0 \). The value of the flow \( f \) is \( |f| = \sum_{(s,x) \in E} f(s, x) \).

In the max flow problem we are given a flow network \( F \) with a capacity function \( u \) and distinguished vertices \( s, t \). We must find a flow \( f \) from \( s \) to \( t \), such that \( |f| \) is maximum.

The residual capacity of an edge \((x, y)\) \( \in E \) is \( u_f(x, y) = u(x, y) - f(x, y) \). An edge, for which \( u_f(x, y) > 0 \), is called a residual edge. The set of all residual edges in \( G \) is denoted by \( E_f \). The graph \( G_f = (V_f, E_f) \) is called a residual graph.

It is easy to see that if \( u_f(x, y) > 0 \) then \( u_f(y, x) > 0 \) and it is possible to push more units of flow through \((y, x)\). Let \( u_f(x, y) > 0 \) for \((x, y) \in E \). If \( f(x, y) > 0 \) than \( f(y, x) < 0 \). This implies that \( u_f(y, x) = u(y, x) - f(y, x) > 0 \) because of \( u(y, x) \geq 0 \).

To present the push-relabel algorithm we need to introduce the following definitions. Let \( F \) be a flow network and \( f : V \times V \rightarrow \mathbb{R} \) be a pseudoflow function. For each \( x \in V \) we define \( e(x) \), the excess of the node \( x \), as the sum: \( e(x) = \sum_{(x,y) \in E} f(x, y) - \sum_{(y,x) \in E} f(y, x) \). Note if \( f \) is a flow then \( e(x) = 0 \) for all \( x \in V \). If \( e(x) > 0 \), for some node \( x \in V \), then we say that \( x \) is an active node. The height of a node \( x \), denoted \( h(x) \) is a natural number from \([0, n] \).

Now we give definitions related to the matching problems and max flow min cost algorithms. Let \( G = (V = X \cup Y, E) \) be a bipartite graph, \(|X| = |Y| = n, |E| = m \). The bipartite matching problem is to find a largest cardinality subset \( M \subseteq E \) such that each vertex \( x \) belongs to at most one edge of \( M \). In case \(|M| = n \), \( M \) is called a perfect matching.

If \( w : E \rightarrow \mathbb{R} \) is a weight function for the edges then \( w(M) = \sum_{(x,y) \in M} w(x, y) \) is the weight of the matching \( M \). The assignment problem is to find the largest cardinality matching \( M \subseteq E \) of the maximum weight \( w(M) \).

For any graph \( G = (V, E) \), the cost function is \( c : E \rightarrow \mathbb{R} \). Hence, the cost of a pseudoflow \( f \) is \( c(f) = \sum_{(x,y) \in E} c(x, y) f(x, y) \).

Assume a flow network \( F = (V, E, u, s, t) \) is given with an aditional edge cost function \( c \). The max flow min cost problem consist in finding a maximum flow with the lowest possible cost.

Following [9] we introduce definitions used in the cost scaling algorithm, they extend the notation given above.

Let \( I' = (V, E, u, s, t, c) \) be a instance of the max flow min cost problem. A price of a node is given by a function \( p : V \rightarrow \mathbb{R} \). A reduced cost of an edge \((x, y) \in E \) is \( c_p(x, y) = c(x, y) + p(x) - p(y) \), while a part-reduced cost of an edge \((x, y) \in E \) is \( c'_p(x, y) = c(x, y) - p(y) \).

For a constant \( \epsilon \geq 0 \) a pseudoflow \( f \) is \( \epsilon \)-optimal with respect to a price function \( p \) if, for every residual edge \((x, y) \in E_f \), we have \( c_p(x, y) \geq -\epsilon \).
For a constant $\epsilon \geq 0$ a pseudoflow $f$ is $\epsilon$-optimal if it is $\epsilon$-optimal with respect to some price function $p$.

We say that a residual edge $(x, y) \in E_f$ is admissible if $c_p(x, y) < 0$.

4 Push-Relabel Algorithm

4.1 Sequential Algorithm

In this section we focus on a standard sequential algorithm solving the max flow problem.

Among many sequential algorithms solving the max flow problem, there are three basic methods. The Ford-Fulkerson and the Edmonds-Karp algorithms are the most common and easiest. The Ford-Fulkerson algorithm calculates the flow in time $O(E|f^*|)$, where $|f^*|$ is value of the max flow $f^*$. The running time of the Edmonds-Karp algorithm is $O(VE^2)$. In every step, these two algorithms look for a new augmenting path from the source to the sink. If found, they increase the flow on the edges of the path. Otherwise they stop and return the flow found up to this moment (it is optimal).

The third solution is the push-relabel algorithm. We present its generic version whose time complexity is $O(V^2E)$. Next we discribe two heuristics which significantly improve its execution time. The generic push-relabel version with two heuristics can be effectivly parallalized and provides a basis for our further considerations.

At the beginning of algorithm $h(x) = 0$, for all $x \in V - \{s\}$, and $h(s) = |V|$. We have also $e(x) = 0$, for all $x \in V$, and $e(s) = \infty$ (Algorithm 4.1).

\begin{algorithm}
\caption{Init operation for the push-relabel algorithm}
\begin{algorithmic}
  \For {each $(x, y) \in E$}
    \State $f(x, y) \leftarrow 1$
    \State $f(y, x) \leftarrow 0$
  \EndFor
  \For {each $x \in V - \{s\}$}
    \State $e(x) \leftarrow 0$
    \State $h(x) \leftarrow 1$
    \State $e(s) \leftarrow \infty$
    \State $h(s) \leftarrow |V|$
  \EndFor
\end{algorithmic}
\end{algorithm}

As opposed to the previous two algorithms, push-relabel does not look for augmenting paths but acts locally on the nodes (Algorithm 4.2).

\begin{algorithm}
\caption{The push-relabel algorithm}
\begin{algorithmic}
  Init()
  make set $S$ empty
  \State $S \leftarrow s$
  \While {$(S$ is not empty)}
    \State $x \leftarrow S.pop()$
    \State discharge($x$)
    \If {($x$ is active node)}
      \State $S.push(x)$
    \EndIf
  \EndWhile
\end{algorithmic}
\end{algorithm}
The discharge operation for each active node $x$ with the set $S$ selects either push or relabel operation (Algorithm 4.3).

**Algorithm 4.3.** discharge($x$), push($x$, $y$) and relabel($x$) operations

```
discharge($x$):
if ($\exists (x, y) \in E_f : h(x) = h(y) + 1$) push($x$, $y$)
else relabel($x$)

push($x$, $y$):
$\delta \leftarrow \min \{u_f(x, y), e(x)\}$
$e(x) \leftarrow e(x) - \delta$
$e(y) \leftarrow e(y) + \delta$
$f(x, y) \leftarrow f(x, y) + \delta$
$f(y, x) \leftarrow f(y, x) - \delta$

relabel($x$):
$h(x) \leftarrow \min \{h[y] : (x, y) \in E_f\} + 1$
```

The push operation is performed on an active node $x$, for which there exists an outgoing residual edge $(x, y) \in E_f$ and the node $y$ satisfies the height constraint: $h(x) = h(y) + 1$. If the node $x$ is active and every edge $(x, y) \in E_f$ does not satisfy this constraint then the relabel operation is performed.

It can be shown that the generic push-relabel algorithm is correct, terminates and its running time is $O(V^2E)$. The proofs and a comprehensive discussion about the generic push-relabel algorithm and its improvements, can be found in [1] and [2].

### 4.2 Heuristics: Global and Gap Relabeling

The above version of the push-relabel algorithm has poor performance in practical applications. To improve the running time two heuristics are used: global relabeling and gap relabeling [2]. To get intuition how these heuristics work, we make some observations.

During the execution of the algorithm, both the excess and the height of the source and the sink are not changing. Then, to the end of the algorithm, the source height is $|V|$ and the sink height is 0. Let us define a distance function.

**Definition 4.1 (distance function).** Let $G = (V, E)$ be a network with a flow $f$ and let $E_f$ be the set of its residual edges. The function $h : V \rightarrow \mathbb{N}$ is a distance function if $h(s) = |V|$, $h(t) = 0$, and for each edge $(x, y) \in E_f$, $h(x) \leq h(y) + 1$. 
It can be proved ([1]) that the height function satisfies the properties of a distance function at each step of the algorithm. Hence we can think about the node height as its distance from the sink to the source. The major issue from which the algorithm’s performance suffers is an execution of a lot of unnecessary relabel operations. It can be proved ([1]) that during the execution, a node height can reach a limit of $2|V| - 1$. The global relabeling heuristic (Algorithm 4.4) prevents the heights of the nodes from growing fast and assigns them the smallest admissible heights.

Algorithm 4.4 global relabeling heuristic

make $Q$ empty queue
for each $x \in V$
    $x$ scanned ← false
$Q.enqueue(t)$
$t.scanned$ ← true
while ($Q$ not empty) do
    $x ← Q.dequeue()$
    current ← $h(x)$
    current ← current + 1
    $\forall (y,x) \in E_f$: $y$ not scanned do
        $h(y) ← current$
        $y$ scanned ← true
        $Q.enqueue(y)$

The global relabeling technique consists in performing a breadth-first backwards search (BFS) in the residual graph and assigning the new heights, equal to the level number of a node in the BFS tree. Obviously BFS takes linear time $O(m + n)$. Usually global relabeling is performed once every $n$ relabels. This heuristic significantly improves the performance of the push-relabel method.

The second heuristic is gap relabeling. The gap relabeling "removes" from the residual graph the nodes that will never satisfy the height constraint. This improves the performance of the push-relabel method because of reducing the number of active nodes. However its result is not so significant for the running time as the global relabeling. The gap relabeling heuristic also can be done in linear time.

The gap relabeling can be performed after the BFS loop of the global relabeling. Any non-scanned node $x$ is not reachable from the sink so we can set its height to $|V|$. This makes the pushed flow omit $x$ and go to another node $y$ (from which there is an augmenting path). As a result the pushed flow gets faster to the sink.

We have presented the generic push-relabel algorithm with two additional techniques which will be used next in parallel versions. Further improvements of the sequential push-relabel algorithm can be found e.g. in [1], [2] and [3].
4.3 Parallel Approach

The push-relabel algorithm was parallelized by Anderson and Setubal in 1992 [14]. One of the first CUDA implementations was proposed by Vineet and Narayanan ([4]). They presented the push-relabel algorithm to graph cuts on the GPU, which is a tool to find the optimal MAP estimation of energy functions defined over an MRF [11, 12, 13].

Vineet and Narayanan’s CUDA implementations were tested on the Nvidia 280 and 8800 GTX graphic cards (devices of compute capability at most 1.3). Their algorithm works on grid graphs which arise in MRFs defined over images. The dataset and CUDA implementations are available from [http://cvit.iiit.ac.in/index.php?page=resources](http://cvit.iiit.ac.in/index.php?page=resources).

The authors assumed that each node of a graph is handled by one thread and the number of outgoing edges per node is fixed, equal 4. Authors suggested that the algorithm can be implemented for the expanded 3D graphs, that is with 8 outgoing edges per node. Their algorithm requires a computer architecture that could launch the same number of threads as the number of the graph nodes so possibilities to run this algorithm on a CPU are small.

The authors prepared two implementations of push-relabel algorithm: atomic and non-atomic. The first of them requires two phases: push and relabel. The second implementation, which was designed for devices of compute capability lower than 1.2, additionally requires a pull phase. Further we will describe only the first implementation, more details about the second can be found in [4].

Vineet and Narayanan have used the graph construction of Kolmogorov et. al. ([12]) which maintains the grid structure, suitable for the CUDA architecture. The data of a grid graph are stored in the global and the shared memory of the device. They are in 8 separated tables. Table of the heights is stored in the shared memory and other tables are stored in the global memory. Among them are the excesses, the relabel masks (which say whether the node is active), the residual capacities of edges upwards/downwards nodes, the residual capacities of edges towards the nodes on the left/right and the residual capacities of edges toward the sink. Access to the element in table is by calculating its index.

Before running the algorithm, the host thread copies data to the global memory on device. The main loop of algorithm is executed on CPU and for each phase the host thread calls another kernel. After finishing a push kernel, the control is returned to the host thread and can be launch the next relabel kernel. However, authors suggested that for some grid graphs, running \( m \) push phases before each relabel phase, improved the execution time. Algorithm stops when all excesses stay the same after a few iterations of loop.

In first step of the push kernel, each node saves its height in the shared memory of thread-block. After the saving, each node whose relabel mask is set to 1 pushes the flow toward its neighbors, if they satisfy a height
constraint. In this step the threads read the heights saved in shared memory.

In the relabel phase first, each thread sets a new relabel mask. If excess of the node is positive and is connected with neighbor, which satisfied a height constraint, the mask is set to one (it is active). If node only has positive excess, the mask is set to zero (node is passive). Otherwise, mask is set to two (inactive node) and this node will never be active. After setting all the relabel masks, threads can calculate new heights of the nodes. They read the old heights saved in the shared memory and write the new to the global memory. It is preparation for next a push phase.

To improve an execution time of algorithm, Authors experimented with varying numbers of threads per block. The best result, they obtained for a thread-block of size 32 × 8.

In the Vineet et al. implementation, the synchronization of threads is assured by __syncthreads() CUDA function. This approach to the parallel push-relabel algorithm blocks the threads which are ready to run next operation before finishing other threads. Therefore it causes long execution time of the algorithm.

4.4 Parallel Lock-Free Algorithm

In 2008 Hong presented a lock-free multi-threaded algorithm for the max flow problem ([5]) based on Goldberg’s version ([2]) of the push-relabel algorithm. Implementation of Hong’s algorithm requires a multi-threaded architecture that supports read-modify-write atomic operations. In our implementation we have used a Nvidia CUDA atomicAdd(int*, int, int) and atomicSub(int*, int, int) functions.

Without lost of generality we assume that the number of running threads is |V| and each of them handles exactly one node of the graph, including all push and relabel operations on it. In several, a few nodes can be handled by one thread.

Let x be the running thread representing the node x ∈ V. In Hong’s algorithm (Algorithm 4.5) each of the running threads has the following private attributes. The variable e′ stores the excess of the node x. The variable h′ stores the height of the currently considered neighbour y of x such that (x, y) ∈ E. The variable ˜h stores the height of the lowest neighbour ˜y of x.

Other variables are shared between all the running threads. Among them there are the arrays with excesses and heights of nodes, and residual capacities of edges.

First, the Init operation is performed by the master thread, in CUDA programing it is the host thread. This init code is the same as its counterpart in the sequential push-relabel version. Next, the master thread starts the threads executing in parallel the lock-free push-relabel algorithm (in CUDA the host thread launches kernels).
The basic changes, introduced by Hong, deal with the selection of operation (push or relabel) that should be executed by $x$, and to which of the adjacent nodes $\tilde{y}, (x, \tilde{y}) \in E_f$, a flow must be pushed. In opposite to the push operation of the generic sequence version where any node $y$ connected by a residual edge to $x$ such that $h(x) = h(y) + 1$ could be pushed, it selects the lowest node among all the nodes connected by a residual edges (lines 4-9). Next if the height of $\tilde{y}$ is less than the height of $x$ (line 10), the push operation is performed (lines 11-15). Otherwise, the relabel operation is performed, that is, the height of $x$ is modified to $h(\tilde{y}) + 1$ (line 17). Note that the relabel operation need not be atomic because only the $x$ thread can change the value of the height of $x$. Furthermore, all critical lines in the code where more than two threads execute the write instruction are atomic. Hence it is easy to see that the algorithm is correct in respect to read and write instructions.

Algorithm 4.5. Lock-free multi-threaded push-relabel algorithm by Bo Hong

Init():

\[
\begin{align*}
& h(s) \leftarrow |V| \\
& \text{for each } x \in V - s \\
& & h(x) \leftarrow 0 \\
& \text{for each } (x,y) \in E \\
& & u_f(x,y) \leftarrow u_{xy} \\
& & u_f(y,x) \leftarrow u_{yx} \\
& \text{for each } (s,x) \in E \\
& & u_f(s,x) \leftarrow 0 \\
& & u_f(x,s) \leftarrow u_{xs} + u_{sx} \\
& e(x) \leftarrow u_{sx}
\end{align*}
\]

lock-free push-relabel():

1. /*$x$ - node operated by the $x$ thread*/
2. while ($e(x) > 0$) do
3. \quad $e' \leftarrow e(x)$
4. \quad $\tilde{y} \leftarrow \text{NULL}$
5. \quad for each $(x,y) \in E_f$ do
6. \quad \quad $h' \leftarrow h(y)$
7. \quad \quad if $\tilde{h} > h'$ do
8. \quad \quad \quad $\tilde{h} \leftarrow h'$
9. \quad \quad $\tilde{y} \leftarrow y$
10. \quad \quad if $h(x) > \tilde{h}$ do /*the $x$ thread performs PUSH towards $y$*/
11. \quad \quad \quad $\delta \leftarrow \min\{e', c_f(x, \tilde{y})\}$
12. \quad \quad \quad $c_f(x, \tilde{y}) \leftarrow c_f(x, \tilde{y}) - \delta$
13. \quad \quad \quad $c_f((y),x) \leftarrow c_f((y),x) + \delta$
14. \quad \quad \quad $e(x) \leftarrow e(x) - \delta$
15. \quad \quad \quad $e((y)) \leftarrow e((y)) + \delta$
16. \quad else do /*the $x$ thread performs RELABEL*/
17. \quad \quad \quad $h(x) \leftarrow h(x) + 1$

All running threads have access to a critical variable in the global memory, but indeed their task are performed sequentialy thanks to the atomic access to the data. The order of the operations in this sequence cannot be predicted. Despite this the algorithm can be proved correct.
It can be proved that the lock-free algorithm terminates after at most \(O(V^2E)\) the push/relabel operations. Because it is executed in parallel by many threads, the complexity of algorithm is analyzed in the number of the operations, not in the execution time.

### 4.5 Lock-Free Algorithm and Heuristics

In 2010 Hong and He ([6]) improved the lock-free push-relabel algorithm, by adding to it a sequential global relabel heuristic performed on CPU (Algorithm 4.6). According to the authors and their experimental result, the new CPU-GPU-Hybrid scheme of the lock-free push-relabel algorithm is robust and efficient.

**Algorithm 4.6. CPU-GPU-Hybrid of push-relabel algorithm by Hong and He**

\begin{algorithm}
\begin{algorithmic}
  \State Init():
  \State 1. initialize \(e, h, u_f\) and \(ExcessTotal\)
  \State 2. copy \(e\) and \(u_f\) from the CPU main memory to the CUDA global memory

  \State push-relabel-cpu():
  \State 1. while \((e(s) + e(t)) < ExcessTotal\) do
  \State 2. copy \(h\) from the CPU main memory to the CUDA global memory
  \State 3. call push-relabel-kernel()
  \State 4. copy \(u_f\), \(h\) and \(e\) from CUDA global memory to CPU main memory
  \State 5. call global-relabel-cpu()
\end{algorithmic}
\end{algorithm}

Following the Authors we will talk about the CPU-GPU-Hybrid algorithm in the context of CUDA programming. Similarly as in the previous algorithm we assume that each node is operated by at most one thread. The previous version of the lock-free push-relabel algorithm will be called the generic algorithm, while the CPU-GPU-Hybrid scheme will be named the hybrid algorithm.

The initialization of the hybrid algorithm is the same as its counterpart in the generic algorithm. The hybrid algorithm maintains 3 arrays with excesses, heights and residual capacities of the nodes and the edges and keeps the global variable \(ExcessTotal\), equal to the value of the flow pushed from the source. \(ExcessTotal\) resides in the global memory on the host and can be changed during the global relabeling.

In opposite to the generic algorithm, the main body of the hybrid algorithm is controlled by the host thread on CPU. The host thread executes the while loop until the cumulative value of the excesses stored in the source and the sink achieves the value of \(ExcessTotal\). In this moment all the valid flow gets to the sink, and the rest of a flow returns to the source. Then, the excess at the sink equals the value of the maximum flow.

In the first step of the while loop, the host thread copies the heights of the nodes to device and launches the push-relabel-kernel. When the control is returned back to the host thread, the calculated pseudoflow and the heights
of the nodes are copied to the CPU memory and the *global-relabel-cup* is performed.

**Algorithm 4.7. Initialization for CPU-GPU-Hybrid**

Init():
1. \(h(s) \leftarrow |V|\)
2. \(e(s) \leftarrow 0\)
3. for each \(x \in V - s\)
   4. \(h(x) \leftarrow 0\)
   5. \(e(x) \leftarrow 0\)
6. for each \((x, y) \in E\)
   7. \(u_f(x, y) \leftarrow u_{xy}\)
   8. \(u_f(y, x) \leftarrow u_{yx}\)
9. for each \((s, x) \in E\)
   10. \(u_f(s, x) \leftarrow 0\)
   11. \(u_f(x, s) \leftarrow u_{sx} + u_{xs}\)
12. \(e(x) \leftarrow u_{sx}\)
13. \(ExcessTotal \leftarrow ExcessTotal + u_{sx}\)

**Algorithm 4.8. Lock-free push-relabel and global relabel for CPU-GPU-Hybrid**

lock-free push-relabel():
1. /*x - node operated by the x thread*/
2. while (\(CYCLE > 0\)) do
3.  if \((e(x) > 0 \text{ and } h(x) < |V|)\) do
4.   \(c' \leftarrow c(x)\)
5.   \(y \leftarrow NULL\)
6.   for each \((x, y) \in E_f\) do
7.     \(h' \leftarrow h(y)\)
8.     if \(h > h'\) do
9.       \(h \leftarrow h'\)
10. \(y \leftarrow y\)
11. if \(h(x) > h\) do /* then the x thread perform PUSH towards y*/
12.   \(\delta \leftarrow \min\{c', c_f(x, y)\}\)
13.   \(c_f(x, y) \leftarrow c_f(x, y) - \delta\)
14.   \(c_f(y, x) \leftarrow c_f((y), x) + \delta\)
15.   \(e(x) \leftarrow e(x) - \delta\)
16.   \(e(y) \leftarrow e(y) + \delta\)
17. else do /* then the x thread perform RELABEL*/
18.   \(h(x) \leftarrow h + 1\)
19. \(CYCLE \leftarrow CYCLE - 1\)

global relabeling heuristic():
1. for all \((x, y) \in E\) do
2.  if \((h(x) > h(y) + 1)\) then
3.   \(e(x) \leftarrow e(x) - u_f(x, y)\)
4.   \(e(y) \leftarrow e(y) + u_f(x, y)\)
5.   \(u_f(y, x) \leftarrow u_f(y, x) + u_f(x, y)\)
6.   \(u_f(x, y) \leftarrow 0\)
7. do a backwards BFS from the sink and assign the height function
8. with each node’s BFS tree level
9. if (not all the nodes are relabeled) then
10.  \(\forall x \in V\) do
11.   if \((x \text{ is not relabeled and marked})\) then
12.     mark \(x\)
13. \(ExcessTotal \leftarrow ExcessTotal - e(x)\)

13
The push-relabel-kernel algorithm differs from the generic algorithm in the timing of executing the kernel. The thread stops the while loop after \( CYCLE \) iterations (where \( CYCLE \) is an integer constant defined by the user) and not when its node becomes inactive. After stopping the loop the heuristic is called, and then the loop is initialized again.

Since the while loop can terminate at any moment (randomly in respect to the original sequential flow computation) it may occur that the property of some residual edge \((x, y) \in E_f\) is violated, i.e. \( h(x) > h(y) + 1 \). Then, before computing the new heights of nodes, all the violating edges must be canceled by pushing the flow. It is made in lines 1-6 of the global relabeling heuristic. Next, the nodes are assigned new heights by performing in the residual graph a backwards BFS from the sink towards the source. The new heights are equal to the shortest distances towards the sink in the residual graph. The excesses of nodes, which are not available from the sink in backwards BFS tree, must be subtracted from \( \text{ExcessTotal} \), because it is a stored excess which will never reach the sink.

In 2011 Hong and He improved both heuristics to a new Asynchronous Global Relabeling (ARG) method ([7]). So far, the global and gap relabeling heuristics were run independent from the lock-free algorithm (in CUDA implementation, to run the heuristics the control was returned to the CPU). The main reason for this was that the push and the relabel operations are mutually exclusive with the global and gap relabeling heuristics (a critical moment is when both the heuristics and the relabel operation want to set a new height for the same vertex). However, this problem does not occur for the non-lock-free versions of the push-relabel algorithms (which is contrary to our expectations). In the new approach, the ARG heuristic is executed by a distinguished thread which corresponds to any vertex and runs periodically, while the other threads asynchronously run push or relabel operations. It significantly improves the execution time of the algorithm. The only problem is that ARG needs to maintain a queue of the unvisited vertices whose size is \( O(V) \). In CUDA programming, a queue of that size can be maintained only in the global memory the access to which is very slow. Perhaps this is why the implementation presented in [7] uses C and the \texttt{pthread} library for multi-threaded constructions.

### 4.6 Our Implementation

In our implementation of the lock-free push-relabel algorithm we have tried to use the ARG heuristic but from the reasons mentioned above this algorithm turns out to be slower than the algorithm using a heuristic launched on CPU. Therefore we use the approach presented in Algorithm 4.8 with
an additional improvement. In the end of the global relabeling phase we add the gap relabeling heuristic which for each unvisited node in the BFS tree sets its height to $|V|$. We implement the procedure of the lock-free push-relabel algorithm as a CUDA kernel. It is executed by $|V|+2$ threads. After $CYCLE$ iterations of the while loop in the kernel, the control is returned to the Host thread. The constant $CYCLE$ is set to 7000 by a preprocessor macro (in our tests this value yielded best results). Next the host thread calls the C procedure running on CPU which performs a global relabeling. The algorithm runs until all the flow of the value $ExcessTotal$ gets to the sink, then the excess of the sink is equal $ExcessTotal$. Note that during the execution of the heuristic, the value of $ExcessTotal$ can be decreased.

In our implementation a vertex is a structure $node$ holding four pointers: $excess$, $height$, $toSourceOrSink$ and $firstOutgoing$. The pointer $toSourceOrSink$ points to the edge leading directly towards the source or the sink. This makes the access to the source and the sink faster. The pointer $firstOutgoing$ points to the first edge on the adjacency list of the vertex. The structure of an edge (named $adj$) also holds four attributes. The pointer $vertex$ points to the neighbor to which the edge leads. The pointer $flow$ points to the location in the global memory where the residual capacity of the edge is stored. The attribute $mate$ is a pointer to the backward edge in the residual graph and $next$ is a pointer to the next edge on the adjacency list.

During the push operation performing on the edge $(x,y)$ both the residual capacities of edges $(x,y)$ and $(y,x)$ are changed. Then to improve the cache utilization, the residual capacities of edges $(x,y)$ and $(y,x)$ are stored one after another.

Throughout the algorithm only flows, excesses and heights need to be copied between the device and host. Therefore to minimize the data transfer between the host and the device we separate arrays containing these data from the structures. Since global relabel heuristic gets all these arrays at the beginning than they are copied onto the host. On the other hand after performing the heuristic only the heights need to be copied back to the device. Therefore we store the excesses and the residual capacities in a single array and the heights are stored in a separate array.

Before starting the algorithm the two arrays are allocated on the device containing nodes and edges respectively. The first keeps the structures of type $node$ and the second contains the structures of type $adj$. During the execution the algorithm arrays of nodes and edges are not copied to the host.

5 Cost Scaling Algorithm

It is known that the non-weighted matching problem can be easily reduced to the max flow problem (for more details see [1, paragraph "Maximum
bipartite matching"). In [9] Goldberg and Kennedy present a way how to efficiently solve the weighted matching problem with a cost scaling algorithm. In their work they reduce the assignment problem to the transporation problem and present their implementation of the algorithm.

We present an analogous reduction from the assignment problem to the max flow min cost problem. In [9], for a given graph $G' = (V' = X' \cup Y', E')$, they additionally define a supply $d(x)$, $x \in V$ that $\forall x \in X$, $d(x) = 1$ and $\forall y \in Y$, $d(y) = -1$. It stimulates preflow pushed from the source to every node of $X$ and preflow pushed from every node of $Y$ to the sink. Hence the push relabel algorithm starts execution with $e(x) = d(x)$. In our work, instead of defining the supply and the transporation problem, we initialize the push-relabel algorithm with $e(x) = 1$, $x \in X$ and $e(x) = -1$, $y \in Y$.

Let $I$ be an instance of the assignment problem: $I = (G, w)$, $G = (V = X \cup Y, E)$ where $|X| = |Y|$ and $w$ is a weight function for edges. We construct an instance $I' = (G', u, c)$ of the max flow min cost problem as follows. For each edge $(x, y) \in E$ we add $(x, y)$ and $(y, x)$ to $E'$. For each $(x, y) \in X \times Y$ define capacities: $u(x, y) = 1$ and $u(y, x) = 0$, and costs: $c(x, y) = w(x, y)$ and $c(y, x) = -w(x, y)$. The graph $G'$ is still bipartite.

Figure 2: Cost scaling algorithm.
5.1 Sequential Cost Scaling Algorithm

The two cost scaling algorithms with efficient implementations which are described by Goldberg et al. in [8,9], are slightly different. The first of them [8] is the generic cost scaling algorithm and was proposed by Goldberg in [10]. The second of them [9] was applied to the assignment problem.

Now we present the first version (Algorithm 5.0), next we point out differences from the second which requires some changes in the definitions of the \( \epsilon \)-optimal pseudoflow and the admissible edge. Our version of the algorithm uses the unmodified definitions and will be presented in subsection 5.3.

Let \( C \) be the largest cost of an edge in \( G \).

**Algorithm 5.0. The cost scaling algorithm**

1. Min-Cost():
   1. \( e \leftarrow C \)
   2. for each \( x \in V \)
      1. \( p(x) \leftarrow 0 \)
   3. while \( (e \geq 1/n) \)
      1. \( (e, f, p) \leftarrow \text{Refine}(e, p) \)

1. Refine\((e, p)\):
   1. \( e \leftarrow e/\alpha \)
   2. for each \( (x, y) \in E : c_p(x, y) < 0 \)
      1. \( f(x, y) \leftarrow 1 \)
   3. while (\( f \) is not a flow) do
      1. apply a push or a relabel operation
   4. return \( (e, f, p) \)

1. push\((x, y)\):
   1. \( \delta \leftarrow \min \{e(x), u_f(x, y)\} \)
   2. send \( \delta \) units flow from \( x \) to \( y \)

1. relabel\((x)\):
   1. \( p(x) \leftarrow \max_{(x, z) \in E} \{p(z) - c(x, z) - e\} \)

The main procedure of the algorithm is a Min-Cost method which maintains a variable \( e \), a flow \( f \), and a price function \( p \). These variables are changed in the while loop by the procedure Refine. On the beginning of each step of the loop the flow \( f \) is \( \epsilon \)-optimal with respect to \( p \). The while loop stops when \( e < 1/n \). It was proved that if Refine reduces the parameter \( \epsilon \) by a constant factor, the total number of iterations is \( O(m \log n) \) (paper [10], Theorem 4.5).

The procedure Refine starts with decreasing \( \epsilon \) to \( \epsilon/\alpha \) and saturating every admissible edge \( (x, y) \), ie. \( c_p(x, y) < 0 \). This spoils the initial flow \( f \) such that \( f \) becomes an \( \epsilon \)-optimal pseudoflow, for \( \epsilon = 0 \) (because \( \forall (x, y) \in E, c_p(x, y) \geq 0 \)). This makes also some nodes active and some with a negative excess. Next the pseudoflow \( f \) become an \( \epsilon \)-optimal flow by making a se-
ries of the flow and the price update operations, each of which preserves $\epsilon$-optimality. There are two kinds of the update operations: push and relabel.

The push$(x, y)$ operation is applied to an active node $x$ and a residual edge $(x, y)$ that is admissible: $c_p(x, y) < 0$. Then $\delta$ units of the flow is put to the node $y$: decreasing $e(x)$ and $f(x, y)$ by $\delta$ and increasing $e(y)$ and $f(y, x)$ by $\delta$.

The relabel$(x)$ operation is applied to an active node $x$ such that $(x, y) \in E_f$ and $(x, y)$ does not satisfy the admissible constraints, i.e. $c_p(x, y) \geq 0$. The new value of $p(x)$ is the smallest value allowed by the $\epsilon$-optimality constraints, i.e. $\max_{(x, z) \in E_f} \{p(z) - c(x, z) - \epsilon\}$.

Goldberg proved that the Refine procedure is correct, that is, if it terminates, the pseudoflow $f$ is an $\epsilon$-optimal flow. Hence the min-cost algorithm is also correct (see [10]).

The procedure Refine maintains a set $S$ containing all the active nodes. The loop terminates when $S$ becomes empty. The generic implementations of the procedure Refine runs in $O(n^2 m)$ time, giving an $O(n^2 m \min\{\log(nC), m \log n\})$ time bound for computing a minimum-cost flow.

The differences between the algorithms mentioned above occur in a new definition of an admissible edge, in the initialization stage of the refine procedure, and in the relabel operation.

Let $\epsilon > 0$. A residual edge $(x, y) \in E_f$ is admissible, if $(x, y) \in (X \times Y) \cap E_f$ and $c_p(x, y) < \frac{1}{2} \epsilon$ or $(x, y) \in (Y \times X) \cap E_f$ and $c_p(x, y) < -\frac{1}{2} \epsilon$. Then we have different conditions for the two types of edges.

The $\epsilon$-optimality notation is closely related to the admissible edge definition. Therefore after changing it we must later update the former, as well.

For a constant $\epsilon \geq 0$ a pseudoflow $f$ is $\epsilon$-optimal with respect to a price function $p$ if, for every residual edge $(x, y) \in (X \times Y) \cap E_f$, we have $c_p(x, y) \geq 0$, and for every residual edge $(y, x) \in (Y \times X) \cap E_f$, we have $c_p(y, x) \geq -\epsilon$. For a constant $\epsilon \geq 0$ a pseudoflow $f$ is $\epsilon$-optimal if it is $\epsilon$-optimal with respect to some price function $p$. 
Algorithm 5.1. The cost scaling algorithm, version 2

1. Refine$(\epsilon, p)$:
   2. $\epsilon \leftarrow \epsilon / \alpha$
   3. for each $(x, y) \in E$
      4. $f(x, y) \leftarrow 0$ /* it can make some node active */
   5. for each $x \in X$
      6. $p(x) \leftarrow -\min_{x, y \in E} \{c(x, y) \}^p$
   7. while (f is not a flow) do
      8. apply a push or a relabel operation
   9. return $(\epsilon, f, p)$

1. relabel$(x)$:
   2. if $x \in X$ then
      3. $p(x) \leftarrow \max_{x, y \in E} \{p(y) - c(x, y)\}$
   4. else if $x \in Y$ then
      5. $p(x) \leftarrow \max_{z, x \in E} \{p(z) + c(z, x) - \epsilon\}$

On the start of the procedure refine, $\epsilon$ and an $\epsilon$-optimal flow $f$ are given. Similarly as in the first algorithm, $\epsilon$ is decreased to $\epsilon / \alpha$. Next, for each saturated edge $(x, y)$, the flow is removed from $(x, y)$ back to its node $x$. This makes some nodes active and some nodes obtain negative excess: $e(x) < 0$. Before the loop starts, for each node $x \in X$, $p(x)$ is set to $-\min_{x, y \in E} c(x, y)$. Then, we get an $\epsilon$-optimal pseudoflow $f$, for $\epsilon = 0$ (because of $\forall (x, y) \in E_f \cap (X \times Y)$ $c_p(x, y) \geq 0$ and $\forall (y, x) \in E_f \cap (Y \times X)$ $c_p(y, x) \geq -\epsilon$). Further, the push and the relabel operations are performed to make $f$ an $\epsilon$-optimal flow.

It is easy to see that the differences between the algorithms do not result in any change in the returned output but they have impact on the efficiency. Therefore, our idea is a combination these two codes (Algorithm 5.2.). We assume the first version of the definitions of admissible edges and $\epsilon$-optimal pseudoflows.

Algorithm 5.2. The cost scaling algorithm, our approach

1. Refine$(\epsilon, p)$:
   2. $\epsilon \leftarrow \epsilon / \alpha$
   3. for each $(x, y) \in E$
      4. $f(x, y) \leftarrow 0$ /* it can make some node active */
   5. for each $x \in X$
      6. $p(x) \leftarrow -\min_{(x, y) \in E} \{c_p(x, y) + \epsilon\}$ /* it makes pseudoflow $f$ $\epsilon$-optimal, (also 0-optimal) */
   7. while (f is not a flow) do
      8. apply a push or a relabel operation
   9. return $(\epsilon, f, p)$

1. relabel$(x)$:
   2. $p(x) \leftarrow -\min_{(x, z) \in E} \{c_p(x, z) + \epsilon\}$
The initialization of the procedure refine (lines 2-6) is the same as in Algorithm 5.1 except line 6, where we must adapt it to the appropriate definition the admissible edge. The relabel operation is the same as in Algorithm 5.0.

5.2 Heuristics: Price Updates and Arc Fixing

The papers [8] and [9] provide a lot of improvements to the cost scaling algorithm for its sequential version. We focus our attention on the price updates and arc fixing heuristics. The idea of the price updates heuristic (introduced in [10] and described also in [15]) is similar to Dijkstra’s shortest path algorithm, implemented using buckets as in Dial’s implementation [16] of the shortest path algorithm.

Algorithm 5.3. Price updates heuristic

```plaintext
price-updates-heuristic():
make empty sets B[i] /* buckets of unscanned nodes*/
make empty set S /* set of active nodes*/
make array l /* constisted the labels of nodes*/
for each x ∈ V
  x.scanned ← false
  l(x) ← ∞
  if e(x) < 0 then
    B[0].push(x) /* bucket 0 is consisted of the x which excess is negative*/
    x.bucket ← 0
  else
    x.bucket ← ∞
  if (e(x) > 0) then
    S.push(x) /* active node x is pushed to S*/
i ← 0
while (S is not empty) do
  while (B[i] is not empty) do
    x ← B[i].pop()
    for each (y, x) ∈ E
      if (y.scanned = false and ⌊cp(y, x)/ε⌋ + 1 < y.bucket) do
        old ← y.bucket
        new ← ⌊cp(y, x)/ε⌋ + 1
        y.bucket ← new
        B[old].pop(y)
        B[new].push(y)
    x.scanned ← true
    l(x) ← i
    if (e(x) > 0) then
      S.pop(x)
      last ← i
      i ← i + 1
for each x ∈ V
  if l(x) < ∞
    p(x) ← p(x) − ε l(x)
  else
    p(x) ← p(x) − ε (last + 1)
```

In our implementation, the price update heuristic maintains the set of buck-
ets $B[i]$. Each node belongs to at most one bucket. Any node with a negative excess is in $B[0]$ and initially buckets with the number larger than 0 are empty. Therefore, each node with a non-negative excess belongs to number $\infty$. In general, for any node the number of its bucket equals its distance from any node residing in the 0-bucket. Additionally, the heuristic maintains an array $l$ containing the labels, which are distances to some node with negative excess i.e. a node residing in the 0-bucket. The distances are multiplies of $\epsilon$.

In each iteration of algorithm, a node with the lowest distance (i.e. a node residing in a nonempty bucket with the lowest number) is scanned. After the scanning, the distance of that node is set to the number of its bucket and the node is marked as scanned.

During the scanning, residual edges entering the scanned node are tested. If a neighbor was not scanned yet and its new calculated distance is smaller than the present, then its distance is updated to a new and this neighbor is removed from the old bucket and put into a new bucket.

In our implementation we also use the arc heuristic described in [8]. It deletes some edges from a residual graph thus decreasing the running time. For the $\epsilon$-optimal flow $f$ and edge $e$, if $c_p(e) > 2n\epsilon$ then the flow of $e$ will never be changed. Therefore this edge can be permanently omitted.

5.3 Applying Lock-Free Push-Relabel Method to Cost Scaling Algorithm

To get better running time of the Refine procedure than in a sequential algorithm, we improve it by applying the lock-free push-relabel algorithm (Algorithm 5.4).
Algorithm 5.4. Push-relabel, our approach

refine\((G, e, p)\):
1. /* x - node operated by the x thread */
2. while \((e(x) > 0)\) do
3. \(e' \leftarrow e(x)\)
4. \(y \leftarrow \text{NULL}\)
5. \(\text{min}_c'p \leftarrow \infty\)
6. for each \((x, y) \in E_f\) do
7. \(\text{tmp}_c'p \leftarrow c'_p(x, y)\)
8. if \(\text{min}_c'p > \text{tmp}_c'p\) do
9. \(\text{min}_c'p \leftarrow \text{tmp}_c'p\)
10. \(y \leftarrow y\)
11. if \((\text{min}_c'p < -p(x))\) do /* that is: \(c_p(x, y) < 0\) - edge is admissible */
12. /* then the x thread performs PUSH towards \(y\) */
13. \(u_f(x, y) \leftarrow u_f(x, y) - 1\)
14. \(u_f(y, x) \leftarrow u_f(y, x) + 1\)
15. \(e(x) \leftarrow e(x) - 1\)
16. \(e(y) \leftarrow e(y) + 1\)
17. else do /* then the x thread performs RELABEL */
18. \(p(x) \leftarrow -(\text{min}_c'p + \epsilon)\)

As in Hong’s algorithm, we assume that one node can be operated by at most one thread. Each thread has the following private attributes. The variable \(e'\) stores the excess of the node \(x\). The variable \(y\) stores the node with the lowest reduced cost. The variable \(\text{min}_c'p\) stores the lowest partially reduced cost of the edge \((x, y)\). The variable \(\text{tmp}_c'p\) stores the temporarily reduced cost of the edge \((x, y)\). The arrays with excesses \(e\) and prices \(p\) of nodes, and costs \(c\) and residual capacity \(u_f\) of edges are shared between all the running threads.

Let \(x\) be an active node operated by thread \(x\). Before performing push or relabel operation, \(x\) select the residual edge \((x, y) \in E_f\), whose reduced cost is the lowest among all residual edges outgoing from \(x\) (lines 6-10). Let \((x, y)\) be the edge with the reduced cost \(\text{min}_c'p\). The thread \(x\) verifies whether \((x, y)\) is admissible (line 11). If so, \(x\) puts the unit of flow towards \(y\): decreasing excess of \(x\) and residual capacity of \((x, y)\), and increasing excess of \(y\) and residual capacity of \((y, x)\) (lines 13-16). Otherwise, if \((x, y)\) is not admissible, the relabel operation is performed on \(x\) and new price of \(x\) is \(p(x) \leftarrow -(\text{min}_c'p + \epsilon)\) (line 18).

Obviously, the decreasing and increasing variables in the push operation must be atomic because of the write conflicts, which may occur when two threads will want to change excess of the same node.

5.4 Correctness of Algorithm

The correctness of the algorithm follows from the correctness of the Refine procedure.

The main difference between our algorithm and the push-relabel algorithm for the max flow problem is that in the second method for each node,
each of its neighbors sees the same height of that node. In our algorithm for each node, each of its neighbors sees the same price of that node but to determine the $\epsilon$-optimality edge, the reduced cost of an edge (not the price of the node) must be considered, which can be different for each neighbors.

Our proof is based on Hong’s proof of the correctness of the lock-free push-relabel algorithm [5] and Goldberg and Tarjan’s proof of the correctness of the generic refine subroutine [2].

**Lemma 5.1.** During the execution of the refine operation, for any active node one of two operations: push or relabel can be applied.

*Proof.* Let $x$ be an active node. Then there must be an edge $(x, y) \in E_f$ because a push operation has occurred which increased the value of $c(x)$ and $u_f(x, y)$. Let $c_p(x, y) >= 0$. Hence $(x, y)$ is not admissible and a push operation is impossible. Therefore a relabel operation can be applied.

**Lemma 5.2.** During the execution of the refine operation the price of a node never increases.

*Proof.* Let $x$ be an active node. If there is a residual edge $(x, y)$ such that $c_p(x, y) < 0$ then the push operation can be applied to it and the price of $x$ does not increase.

Otherwise, if for each residual edge $(x, y)$, $c_p(x, y) >= 0$ then the relabel operation can be applied to $x$. Let $(x, \tilde{y}) \in E_f$ have the smallest reduced cost among all the residual edges outgoing from $x$. Since $c_p(x, \tilde{y}) >= 0$ then $c(x, \tilde{y}) + p(x) - p(\tilde{y}) >= 0$ and $p(x) >= p(\tilde{y}) - c(x, \tilde{y}) = -c'_p(x, \tilde{y})$. Additionally, after the relabel operation we have $p(x) = -(c'_p(x, \tilde{y}) + \epsilon)$. Therefore the price of $x$ does not increase.

Now, following Hong’s proof, we define a trace, a preparation stage and a fulfillment stage and two basic types of computation (history) traces consisting of push and/or relabel operations.

The *trace* of the interleaved execution of multiple threads is the order in which instructions from the threads are executed in real time.

Each operation (both push or relabel), can be split into 2 stages: the preparation and the fulfillment.

For the push operation the preparation stage is performed in lines 6-11 of Algorithm 5.4 and the fulfillment stage is lines 12-16. For the relabel operation the preparation stage is in lines 6-10 of Algorithm 5.4 and the fulfillment stage is in line 18.

The preparation stage tests if the operation is applicable. The fulfillment stage finishes the operation. The P and F notations denote the preparation and the fulfillment stage respectively.

A *stage-clean* trace consists of non overlapping operations, for example: (P(push1), F(push1), P(relabel1), F(relabel1), P(push2), F(push2)).
In *stage-stepping* trace all the preparation stages are performed before any fulfillment stage, for example: (P(push1), P(relabel2), P(relabel1), F(relabel2), F(push1), F(relabel1)).

**Lemma 5.3.** Each trace consisting of push and/or relabel operations is semantically equivalent to one of the two basic traces: stage-clean or stage-stepping.

**Proof.** There are 5 nontrivial cases in which the stages of push and relabel operations on common data can interleave:

1. *push(x, y) and push(y, z)*

   The fulfillment stage of *push(x, y)* increases \( e(y) \). However the preparation stage of *push(y, z)* reads \( e(y) \). The reading and the writing operations can occur in the following three scenarios.

   (a) \( P(push(x, y)) - F(push(x, y)) - P(push(y, z)) \).

   This is a stage-clean trace: \( push(x, y) -> push(y, z) \). In the fulfillment of *push(x, y)* the thread \( x \) increases \( e(y) \) before the thread \( y \) reads \( e(y) \) in the preparation stage.

   (b) \( P(push(y, z)) - F(push(y, z)) - F(push(x, y)) \).

   This is a stage-clean trace: \( push(y, z) -> push(x, y) \). In the fulfillment of *push(y, z)* thread \( y \) decreases \( e(y) \) before the thread \( x \) writes to the \( e(y) \) in fulfillment stage.

   (c) \( P(push(y, z)) - F(push(x, y)) - F(push(y, z)) \).

   This is also a stage-clean trace: \( push(y, z) -> push(x, y) \). In the fulfillment of *push(x, y)*, the thread \( x \) increases \( e(y) \), which was earlier remembered by thread \( y \). Next, thread \( y \) pushes the flow, whose value is depended from the old value of \( e(y) \), to the node \( z \). So it is equivalent that the thread \( y \) pushes the flow to \( z \) and then the push operation from \( x \) to \( y \) is performed.

2. *push(x, y) and push(z, y)*

   Both *push(x, y)* and *push(z, y)* increase the value of \( e(y) \) without reading and storing this value before. It may occur in two different scenarios, each of them is equivalent to a stage-clean trace.

3. *push(x, y) and relabel(y)*

   In the preparation of *push(x, y)* the thread \( x \) chooses the edge \( (x, y) \) because it has the lowest partially reduced cost \( c'_p(x, y) \). However, in the fulfillment stage the thread \( y \) updates the price \( p(y) \). We have six different scenarios interleaving the stages of *push(x, y)* and *relabel(y)*.
(a) \( P(push(x, y)) - F(push(x, y)) - P(relabel(y) - F(relabel(y)) \)

Obvious. It is equivalent to the stage-clean trace: \( push(x, y) \rightarrow relabel(y) \).

(b) \( P(relabel(y)) - F(relabel(y)) - P(push(x, y)) - F(push(x, y)) \)

Obvious. It is equivalent to the stage-clean trace: \( relabel(y) \rightarrow push(x, y) \).

(c) \( P(push(x, y)) - P(relabel(y)) - F(push(x, y)) - F(relabel(y)) \)

\( P(push(x, y)) - P(relabel(y)) - F(relabel(y)) - F(push(x, y)) \)

\( P(relabel(y)) - P(push(x, y)) - F(relabel(y)) - F(push(x, y)) \)

\( P(relabel(y)) - P(push(x, y)) - F(push(x, y)) - F(relabel(y)) \)

In this case two preparation stages are performed before any fulfillment stage one. Therefore they are equivalent to the stage-stepping trace: \( P(push(x, y)) \rightarrow P(relabel(y)) \rightarrow F(push(x, y)) \)

\( \rightarrow F(relabel(y)) \).

4. \( push(x, y) \) and \( relabel(z) \)

Let \( \tilde{p}(z) \) be the new price of \( z \) after the fulfillment stage of \( relabel(z) \).

According to lemma 4.2: \( \tilde{p}(z) \leq p(z) \) must hold.

We have \( c_p(x, y) = \min_{(x, w) \in E_f} c_p(x, w) \leq c_p(x, z) \).

Then \( c_p(x, z) = c(x, z) + p(x) - p(z) \leq c(x, z) + p(x) - \tilde{p}(z) = \tilde{c}_p(x, z) \).

Since \( \tilde{c}_p(x, z) \geq c_p(x, z) \geq c_p(x, y) \geq 0 \), then the operation \( relabel(z) \) does not impact the operation \( push(x, y) \) and this scenario is equivalent to a stage-clean trace.

5. \( relabel(x) \) and \( relabel(y) \)

In the fulfillment stage of \( relabel(x) \) the thread \( x \) updates \( p(x) \) and \( (y, x) \) may be the residual edge read by the thread \( y \) before or after the fulfillment stage of \( relabel(y) \). Then there are six scenarios interleaving the stages of \( relabel(x) \) and \( relabel(y) \). The proof is analogous to the case 3.

Lemma 5.4. For any trace consisting of three or more \( push \) and/or \( relabel \) operations there exists an equivalent sequence consisting only of stage-clean or stage-stepping traces.

Proof. Similar as above. □

Lemma 5.5. When the algorithm terminates \( f \) is an \( \epsilon \)-optimal flow.

Proof. We show that during the execution of the algorithm any residual edge \( (x, y) \) satisfies the constraint \( c_p(x, y) \geq -\epsilon \) with one exception which is transient. Let \( f \) be an \( \epsilon \)-optimal pseudoflow. There may occur the following situations:
1. applying the push operation
   Obviously.

2. applying the relabel operation
   Let $x$ be the relabeled node and $\tilde{p}(x)$ be the new price of $x$. According to Lemma 5.2, we have $\tilde{p}(x) < p(x)$. Then any edge of the form: $(x, y)$ or $(y, x)$ can spoil the $\epsilon$-optimality of $f$. For any residual edge $(x, y)$ outgoing from $x$, we have $c_p(x, y) = c(x, y) + p(x) - p(y) < c(x, y) + \tilde{p}(x) - p(y) = \tilde{c}_p(x, y)$ Therefore, if $\tilde{c}_p(x, y) \geq c_p(x, y) \geq -\epsilon$ then the relabel operation preserves the $\epsilon$-optimality.

3. applying relabel$(x)$ and relabel$(y)$ operations
   According to Lemma 5.3 this scenario can be reduced to a stage-clean trace or stage-stepping trace. A clean-stage trace can be reduced to cases: 1. and 2. above. A stage-stepping trace has the following cases.

   (a) $(x, y) \in E_f$ and $(y, x) \in E_f$

   In this case, $c_p(x, z) \geq 0$, for all $(x, z) \in E_f$, and $c_p(y, w) \geq 0$, for all $(y, w) \in E_f$. Therefore, since $c_p(x, y) = c(x, y) + p(x) - p(y) = -(c(y, x) + p(y) - p(x)) = -c_p(y, x)$

   then $\tilde{c}_p(x, y) = c_p(x, y) = 0$.

   Furthermore, $c_p(x, y) = \min \{c_p(x, z), (x, z) \in E_f\}$ and $c_p(y, x) = \min \{c_p(y, w), (y, w) \in E_f\}$.

   Therefore, the fulfillment stages of relabel$(x)$ and relabel$(y)$ update the prices of $x$ and $y$ respectively:

   $\tilde{p}(x) \leftarrow -(c_p'(x, y) + \epsilon)$ and $\tilde{p}(y) \leftarrow -(c_p'(y, x) + \epsilon)$.

   Let us check whether the new prices preserve $\epsilon$-optimality of the residual edges $(x, y)$ and $(y, x)$. Since $\tilde{c}_p(x, y) = c(x, y) + \tilde{p}(x) - \tilde{p}(y) = c(x, y) - c_p'(x, y) - \epsilon + c_p'(y, x) + \epsilon = c(x, y) - c(x, y) + p(y) + c(y, x) - p(x) = c(y, x) + p(y) - p(x) = c_p(y, x) = 0 >= -\epsilon$, then $(x, y)$ preserves $\epsilon$-optimality of the pseudoflow $f$. Because of $\tilde{c}_p(x, y) = -\tilde{c}_p'(y, x)$ then also $c_p(y, x) = 0$ and $(y, x)$ preserves $\epsilon$-optimality of $f$.

(b) $(x, y) \in E_f$ and $(y, x) \notin E_f$

   We want to see if the edge $(x, y)$ preserves $\epsilon$-optimality of the pseudoflow $f$ after the fulfillment stages of relabel$(x)$ and relabel$(y)$. Let $(x, z)$ be the edge that affects on the new price $\tilde{p}(x)$ of $x$.

   Since $c_p(x, z) \leq c_p(x, y)$ then $c_p'(x, z) \leq c_p'(x, y)$ and $\tilde{p}(x) = -(c_p'(x, z) + \epsilon) >= -(c_p'(x, y) + \epsilon) = -c(x, y) + p(y) - \epsilon \geq -c(x, y) + \tilde{p}(y) - \epsilon$.

   Therefore $\tilde{c}_p(x, y) = c(x, y) + \tilde{p}(x) - \tilde{p}(y) \geq -\epsilon$. Q.E.D
(c) \((x, y) \notin E_f \) and \((y, x) \in E_f\)

The proof is the same as above.

(d) \((x, y) \notin E_f \) and \((y, x) \notin E_f\)

If \((x, y)\) and \((y, x)\) are not residual then after the relabel operations they will not be either. Therefore, this case is trivial.

4. applying push\((x, y)\) and push\((z, y)\) operations

Following Lemma 5.3, this case is equivalent a stage-clean trace hence it reduces to the cases: 1 and 2.

5. applying push\((x, y)\) and relabel\((y)\) operations

This case is equivalent to either a stage-clean or a stage-stepping trace. The first scenario can be reduced to the cases 1 and 2, while the second has the following cases. Note \((x, y) \in E_f\) because the push operation is applicable.

(a) \((y, x) \in E_f\)

Since the push operation preserves \(\epsilon\)-optimality of the pseudoflow \(f\), then from the inductive assumption, the pseudoflow \(f\) is \(\epsilon\)-optimal after the push operation.

The fulfillment stage of push\((x, y)\) does not affect the value of \(c_p(y, x)\). If \(0 \leq c_p(y, x) \leq \min \{c_p(y, z), (y, z) \in E_f\}\) before the fulfillment stage of relabel\((y)\) then the new price of the node \(y\) satisfies \(\hat{p}(y) = -(\min_{(y,z) \in E_f} c'_p(y,z) + \epsilon) \geq -(c'_p(y,x) + \epsilon)\). Therefore, after the fulfillment stage of relabel\((y)\) we obtain: \(\hat{c}_p(x) = c(x,y) + \hat{p}(y) - p(x) \geq c(y,x) - c'_p(y,x) - \epsilon - p(x) = c(y,x) - c(y,x) + p(x) - \epsilon - p(x) = -\epsilon\). Then the residual edge \((y, x)\) is \(\epsilon\)-optimal with respect to the new price of the node \(y\).

(b) \((y, x) \notin E_f\)

There are two subscenarios.

i. \((y, x) \in E\)

In the situation that \((y, x) \in E\) and \((y, x) \notin E_f\) we have \(f(y, x) = u_f(y, x)\). However, the fulfillment stage of push\((x, y)\) pushes the flow through the edge \((x, y)\) and hence adds the reverse edge \((y, x)\) to \(E_f\). Otherwise, the preparation stage of relabel\((y)\), which is performed before the fulfillment stage of push\((x, y)\), would not consider the edge \((y, x)\) (because it would be added to \(E_f\) after that stage). Let \((y, z_0)\) be the edge that affects the new price of \(y\) in the preparation stage of relabel\((y)\). After the trace, it may occur that the new residual edge \((y, x)\) has a lower reduced cost than \((y, z_0)\), that is \(c_p(y, x) < c_p(y, z_0)\) and hence \(c'_p(y, x) < c'_p(y, z_0)\).
Therefore \(-c'_p(y, x) + \epsilon > -c'_p(y, z_0) + \epsilon\) what implies that \(-c(y, x) - p(x) + \epsilon > \tilde{p}(y)\) and hence \(c(y, x) + \tilde{p}(y) - p(x) < -\epsilon\) and \(c_p(x, y) < -\epsilon\). Then the residual edge \((y, x)\) spoils the \(\epsilon\)-optimality of the pseudoflow \(f\). Further we show that this situation is temporary and in the next step for the node \(y\) \(f\) becomes an \(\epsilon\)-optimal pseudoflow. For the node \(y\), \(c(y, x) > \epsilon\). Moreover, the residual edge \((y, x)\) has the smallest reduced cost among all the residual edges outgoing from \(y\). Then the push operation towards \(x\) can be performed on \(y\) and \(u_f(y, x) = \min\{u_f(y, x), e(y)\}\). Therefore, in the next step for the node \(y\), the push operation removes the residual edge \((y, x)\) from the residual graph and hence remove the requirement \(c_p(y, x) \geq -\epsilon\).

ii. \((y, x) \notin E\)

In the situation that \((y, x) \notin E\) and \((y, x) \notin E_f\) there must be \(f(x, y) = 0\) (if \(f(x, y) > 0\) then \(u_f(y, x) = u(y, x) - f(y, x) = u(y, x) + f(x, y) > 0\) and then it would be \((y, x) \in E_f\)). Similarly to the previous case, the residual edge \((y, x)\) will be added to \(E_f\) in the fulfillment stage of \(\text{push}(x, y)\) and the \(\epsilon\)-optimality of the flow might be violated because of \((y, x)\).

However, like in the previous case, the residual edge \((y, x)\) will be removed from \(E_f\) because of the push operation from \(y\) towards \(x\) which will push the flow of value \(f(y, x)\).

6. applying \(\text{push}(x, y)\) and \(\text{push}(y, z)\) operations.

According to Lemma 5.3 this trace is equivalent to a stage-clean trace, where \(\text{push}(x, y)\) and \(\text{push}(y, z)\) are performed one by one. Then this case can be reduced to the case 2 above. The \(\epsilon\)-optimality is preserved by this trace.

7. applying \(\text{push}(x, y)\) and \(\text{relabel}(x)\) operations.

According to Lemma 5.2 this two operations cannot be intereleaved because the active node can be dealt with only by one operation per step of the algorithm.

8. applying \(\text{push}(x, y)\) and \(\text{push}(y, x)\) operations.

These two operations cannot be intereleaved because the two conditions \(c_p(x, y) < 0\) and \(c_p(y, x) < 0\) are mutually exclusive.

9. applying more operations than two.

Any trace consisting of more than two operations is interleaved. The proof is similar to the above and is omitted. 

\[\square\]
Lemma 5.6. When the algorithm terminates the pseudoflow $f$ is $\epsilon$-optimal.

Proof. The initial pseudoflow $f$ is $\epsilon$-optimal. According to lemma 5.5, any trace consisting of the flow and/or the price update operations preserve $\epsilon$-optimality of $f$. According to lemma 5.1, the refine procedure can terminate only if there is no active node, which implies that the pseudoflow $f$ is an $\epsilon$-optimal flow.

5.5 Our Implementation

Our implementation of the cost scaling algorithm uses the method presented in Algorithm 5.4, the global price update heuristic in Algorithm 5.3 and the arc-fixing heuristic described above. We also use our implementation of the lock-free push-relabel algorithm changed appropriately for the cost-scaling method.

After loading the input, the graph is constructed in the host global memory. Next, all arrays are copied to the global memory on device. Since the copied addresses point to locations in the host memory and the graph can be arbitrary we use an additional array to store information about adjacency lists of vertices and set valid addresses in the device memory. Further, while running the algorithm only three arrays are copied between host and device: the prices, the excesses and the flows.

In each iteration of the refine loop the host thread calls the push-relabel kernel until the pseudoflow $f$ becomes a flow, that is, until $\text{EXCESS}(\text{source}) = 0$ and $\text{EXCESS}(\text{sink}) = \text{Total}$, where $\text{Total} = |X| = |Y|$. The kernels are launched by $|V| + 2$ threads. The control is returned back to the host in two cases. First, after each $\text{CYCLE}$ iterations, ($\text{CYCLE}$ has been preset to 500000). In the second case, if $\text{EXCESS}(\text{source}) = 0$ and $\text{EXCESS}(\text{sink}) = \text{Total}$. The first case means that the calculation is not finished, but the control is returned to the host. The second means that the pseudoflow $f$ becomes a flow. In the first case, it is common that the running kernel is interrupted and restored without changes in the graph. The purpose of this is to avoid terminating the execution of the kernel by GPU (launch timed out). This often happens when the graph has many edges.

In our implementation only after the first running of the push-relabel kernel the heuristics are performed. They can be executed many times but the way we did it yields the best results. Similarly as in [15] we chose the value of $\text{ALPHA}$ constant equal 10 because in our tests other values much extended the running time of the program. The best results of the algorithm are achieved for the thread block of size $32 \times 16$.

The first heuristic executed is $\text{arc\_fixing}$ kernel then $\text{globalPriceUpdate}$ C procedure. The $\text{arc\_fixing}$ kernel deletes edges from the graph residing in the device memory by removing them from the adjacency lists and sets
the flows on them to $-10$ (any negative constant would do) in order not to free the memory. Then the arrays of flows and prices are copied to the host memory. To make the graphs on the device and on the host equal, the procedure of the global relabeling deletes from the host graph the edges which were previously deleted from the graph on the device memory. They are recognized by the value of a flow equal $-10$. Next the global relabeling is performed and further the new prices are copied back to the device memory.

6 Discussion and Future Work

In this paper we have presented an efficient implementation of Hong’s parallel non-blocking push-relabel algorithm and an implementation of the cost-scaling algorithm for the assignment problem. The Refine procedure of the cost-scaling algorithm uses the parallel push-relabel lock-free algorithm and runs in $O(n^2m)$ time, where the complexity is analyzed in respect to the number of push and relabel operations. The amount of work performed by the program (the total number of instructions executed) is the same as in the sequential Algorithm 5.2, i.e. $O(n^2m \min \{\log(nC), m \log n\})$. Both implementations run on GPU using CUDA architecture.

For our purpose, the cost scaling algorithm for the assignment problem is used for the complete bipartite graphs of size $|X| = |Y| < 30$. The execution time of our implementation for graphs of this size and costs of edges at most 100, is about $1/20s$ which allows for real-time applications.

The maximum efficiency of an algorithm implemented in CUDA architecture is achieved when the number of threads is large and all threads execute the same task. Otherwise the running time can be equal the running time of a sequential algorithm. In the case of the large complete bipartite graphs the presented algorithm is not efficient.

Further research could be made towards the use of the CUDA architecture to finding a maximum cardinality matching of maximum weight for arbitrary (nonbipartite) graphs.
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