A Comparative Review of Component Tree Computation Algorithms

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Abstract-Connected operators are morphological tools that have the property of filtering images without creating new contours and without moving the contours that are preserved. Those operators are related to the max-tree and min-tree representations of images, and many algorithms have been proposed to compute those trees. However, no exhaustive comparison of these algorithms has been proposed so far, and the choice of an algorithm over another depends on many parameters. Since the need for fast algorithms is obvious for production code, we present an in-depth comparison of the existing algorithms in a unique framework, as well as variations of some of them that improve their efficiency. This comparison involves both sequential and parallel algorithms, and execution times are given w.r.t. the number of threads, the input image size, and the pixel value quantization. Eventually, a decision tree is given to help the user choose the most appropriate algorithm with respect to the user requirements. To favor reproducible research, an online demo allows the user to upload an image and bench the different algorithms, and the source code of every algorithms has been made available.

Keywords—Component tree, max-tree, min-tree, connected operators, filtering, mathematical morphology, algorithms.

EDICS Categories: ARS-RBS, TEC-PRC, SMR-STM

I. INTRODUCTION

N mathematical morphology, connected filters are those that modify an original image by only merging flat zones, hence those that preserve some of the original image contours. Originally, they were mostly used for image filtering [1, 2]. Major advances came from max- and min-tree as hierarchical representations of connected components and from an efficient algorithm able to compute them [3]. Since then, usage of these trees has soared for more advanced forms of filtering: based on attributes [4, 5], using new filtering strategies [3, 6], allowing new types of connectivity [7]. They are also a base for other image representations. In [8] a tree of shapes is computed from a merge of the min- and max- trees. In [9] a component tree is computed over the attributes values of the max-tree. Maxtrees have been involved in many applications: computer vision through motion extraction [3], features extraction with MSER [10], segmentation, 3D visualization [11]. With the increase of applications comes an increase of data type to process: 12bit images in medical imagery [11], 16-bit or float images in astronomical imagery [12], and even multivariate data with special ordering relation [13]. With the improvement of optical sensors, images are getting bigger (so do image data sets) which argues for the need for fast algorithms. Many algorithms have been proposed to compute the max-tree efficiently but only partial comparisons have been proposed. Moreover, some of them are dedicated to a particular task (e.g., filtering) and are unusable for other purposes.

In a short paper [14], we have presented a first comparison of many state-of-the-art max-tree algorithms in a unique framework, i.e., same architecture, same language (C++) and same outputs. Yet this comparison was performed on a single image, the pseudo-code of all the algorithms were not listed, and the description of those algorithms and their comparison were short. This paper aims at correcting those three drawbacks, so it presents a full and exhaustive comparative review of the state-of-art component tree computation algorithms.

The paper is organized as follows. Section II recalls basic notions and manipulations of max-tree. Section III describes the algorithms and implementations used in this study; in particular, a new technique that improves the efficiency of union-find-based algorithms is presented in Section III-A4. Section IV is dedicated to the comparison of those algorithms both in terms of complexity and running times through experimentations. Last we conclude in Section V.

II. A TOUR OF MAX-TREE: DEFINITION, REPRESENTATION AND ALGORITHMS

A. Basic notions for max-tree

Let $f : \Omega \to V$ be an image on a regular domain Ω , having values on a totally preordered set (V, \leq) and let \mathcal{N} be a neighborhood on Ω . Let $\lambda \in V$, we note $[f \leq \lambda]$ the set $\{p \in \Omega, f(p) \leq \lambda\}$. Let $X \subset \Omega$, we note $CC(X) \subset \mathcal{P}(\Omega)$ the set of connected components of X w.r.t. the neighborhood \mathcal{N} ; $\mathcal{P}(\Omega)$ being the power set of all the possible subsets of Ω . $\{CC([f = \lambda]), \lambda \in V\}$ are *level components* and $\Psi = \{CC([f \geq \lambda]), \lambda \in V\}$ (resp. \leq) is the set of upper components (resp. lower components). The latter endowed with the inclusion relation form a tree called the max-tree (resp. min-tree). Since min- and max-trees are dual, this study obviously holds for min-tree as well. Finally, the peak component of p at level λ noted P_{p}^{λ} is the upper component $X \in CC([f \geq \lambda])$ such that $p \in X$.

B. Max-tree representation

Berger et al. [12], and Najman and Couprie [15] rely on a simple and effective encoding of component-trees using an

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image that stores the *parent* relationship. The latter exists between two components A and B whenever A is *directly* included in B (*parent* is actually the covering relation of (Ψ, \subseteq)). An upper component is represented by a single point called the *canonical element* [12, 15] or *level root*. Let two points $p, q \in \Omega$, and p_r be the root of the tree. We say that p is canonical if $p = p_r$ or f(parent(p)) < f(p). A *parent* image shall satisfy the following three properties: 1) *parent*(p) = $p \Rightarrow p = p_r$ - the root points to itself and it is the only point verifying this property - 2) $f(parent(p)) \le f(p)$ and 3) *parent*(p) is canonical.

Furthermore, having just the *parent* image is an incomplete representation since it is not sufficient to easily perform classical tree traversals. For that, we need an extra array of points, $S : \mathbb{N} \to \Omega$, where points are stored so that $\forall i, j \in \mathbb{N} \ i < j \Rightarrow S[j] \neq parent(S[i])$. Thus browsing S elements allows to traverse the tree downwards i.e from the top (the root) to the bottom of the tree (the leaves). On the contrary, a reverse browsing of S is an upward tree traversal. Note that having both S and *parent* thus makes it useless to store the children of each node. Fig. 1 shows an example of such a representation of a max-tree. This representation only requires 2nI bytes memory space where n is the number of pixels and I the size in bytes of an integer, since points stored in S and *parent* are actually positive offsets in a pixel buffer. The algorithms we compare have all been modified to output the same tree encoding, that is, the couple (*parent*, S).

C. Attribute filtering and reconstruction

A classical approach for object detection and filtering is to compute some features called attributes on max-tree nodes. A usual attribute is the number of pixels in components. Followed by a filtering, it leads to the well-known area opening. More advanced attributes have been used like elongation, moment of inertia [16] or even mumford-shah like energy [9]. Some maxtree algorithms [17, 10] construct the *parent* image only; they do not compute S. As a consequence, they do not provide a "versatile" tree, i.e., a tree that can be easily traversed upwards and downwards, that allows attribute computation and nontrivial filtering. Here we require every algorithms to output a "complete" tree representation (*parent and S*) so that it can be multi-purposedly usable. The rationale behind this requirement is that, for some applications, filtering parameters are not known yet at the time the tree is built (e.g., for interactive visualization [11]). In the algorithms we compare in this paper,

 $\begin{array}{l|l} \textbf{function COMPUTE-ATTRIBUTE}(S, parent, f) \\ & p_{root} \leftarrow S[0] \\ \textbf{for all } p \in S \ \textbf{do} \ attr(p) \leftarrow \hat{a}(p, f(p)) \\ & \textbf{for all } p \in S \ backward, p \neq p_{root} \ \textbf{do} \\ & | \ q \leftarrow parent(p) \\ & | \ attr(q) \leftarrow attr(q) \hat{+} attr(p) \\ & \textbf{return } attr \end{array}$

function DIRECT-FILTER(S , parent, f , attr)	
$p_{root} \leftarrow S[0]$	
if $attr(p_{root}) < \lambda$ then $out(p_{root}) \leftarrow 0$	
else $out(p_{root}) \leftarrow f(p_{root})$	
for all $p \in S$ forward do	
$ q \leftarrow parent(p)$	
if $f(q) = f(p)$ then $out(p) \leftarrow out(q)$	\triangleright ⁽¹⁾
else if $attr(p) < \lambda$ then $out(p) \leftarrow out(q)$	⊳ (2)
else $out(p) \leftarrow f(p)$	⊳ (3)
return out	

⁽¹⁾ p not canonical, ⁽²⁾ Criterion failed, ⁽³⁾ Criterion passed

Fig. 2. Computation of attributes and filtering.

no attribute computation nor filtering are performed during tree construction for clarity reasons; yet they can be augmented to compute attribute and filtering at the same time. Fig. 2 provides an implementation of attribute computation and direct-filtering with the representation. $\hat{a}: \Omega \times V \to \mathcal{A}$ is an application that projects a pixel p and its value f(p) in the attribute space \mathcal{A} . $\hat{+}: \mathcal{A} \times \mathcal{A} \to \mathcal{A}$ is an associative operator used to merge attributes of different nodes. COMPUTE-ATTRIBUTE starts with computing attributes of each singleton node and merges them from leaves toward root. Note that this simple code relies on the fact that a node receives all information from its children before passing its attribute to the parent. Without any ordering on S, it would not have been possible. DIRECT-FILTER is an implementation of direct filtering as explained in [3] that keeps all nodes passing a criterion λ and lowers nodes that fail to the last ancestor "alive". This implementation has to be compared with the one in [17] that only uses *parent*. This one is shorter, faster and clearer above all.

III. MAX-TREE ALGORITHMS

Max-tree algorithms can be classified in three classes: **Immersion algorithms.** It starts with building N disjoints singleton for each pixel and sort them according to their gray value. Then, disjoint sets merge to form a tree using the unionfind algorithm [18, 19].

Flooding algorithms. A first scan allows to retrieve the root which is a pixel at lowest level in the image. Then, it performs a propagation by flooding first the neighbor at highest level i.e. a depth first propagation [3, 20].

Merge-based algorithms. They divide an image in blocks and compute the max-tree on each sub-image using another max-tree algorithm. Sub max-trees are then merged to form the tree of the whole image. Those algorithms are well-suited for parallelism using a *map-reduce* (or *divide-and-conquer*) approach [21, 17]. When blocks are image lines, dedicated 1D max-tree algorithms can be used [22, 23, 24].

A. Immersion algorithms

1) Basic principle: Berger et al. [12], Najman and Couprie [15] proposed two algorithms based on Tarjan's union-find. They consist in tracking disjoints connected components and merge them in a bottom-up fashion. First, pixels are sorted in an array S where each pixel p represent the singleton set $\{p\}$. Then, we process pixels of S in backward order. When a pixel p is processed, it looks for already processed neighbors $(\mathcal{N}(p))$ and merges with neighboring connected components to form a new connected set rooted in p. The merging process consists in updating the *parent* pointer of neighboring component roots toward p. Thus, the union-find relies on three processes: make-set(parent, x) that builds the singleton set $\{x\}$, find-root (parent, x) that finds the root of the component that contains x, and merge-set(parent, x, y) that merges components rooted in x and y and set x as the new root. Based on the above functions, a simple max-tree algorithm is given below:

procedure $MAXTREE(f)$
$S \leftarrow$ sort pixels increasing
for all $p \in S$ backward do
make-set $(parent, p)$
for all $n \in \mathcal{N}_p$ processed do
$r \leftarrow \text{find-root}(parent, n)$
if $r \neq p$ then merge-set(parent, p, r)

find-root is a O(n) function that makes the above procedure a $O(n^2)$ algorithm. Tarjan [19] discussed two important optimizations to avoid a quadratic complexity: root path compression and union-by-rank.

When *parent* is traversed to get the root of the component, points of the path used to find the root collapse to the root the component. However, path compression should not be applied on *parent* image because it removes the hierarchical structure of the tree. As consequence, we apply path compression on an intermediate image *zpar* that stores the root of disjoints components. Path compression bounds union-find complexity to $O(n \log n)$ and has been applied in [12] and [15].

When merging two components A and B, we have to select one of the roots to represent the newly created component. If A has a rank greater than B then $root_A$ is selected as the new root, $root_B$ otherwise. When rank matches the depth of trees, it enables tree balancing and guaranties a $O(n \log n)$ complexity for union-find. When used with path compression, it allows to compute the max-tree in quasi-linear time $(O(n.\alpha(n)))$ where $\alpha(n)$ is the inverse of Ackermann function which is very lowgrowing). Union-by-rank has been applied in [15].

Note that *parent* and *zpar* encode two different things, *parent* encodes the max tree while *zpar* tracks disjoints set of points and also uses a tree. Thus, union-by-rank and root path compression shall be applied on *zpar* but never on *parent*.

The algorithm given in Fig. 3 is the union-find-based maxtree algorithm as proposed by [12]. It starts with sorting pixels that can be done with a counting sort algorithm for lowquantized data or with a radix sort-based algorithm for high quantized data [25]. Then it annotates all pixels as *unprocessed* with -1 (in common implementations, pixels are positive offsets in a pixel buffer). Later in the algorithm, when a pixel p is processed it becomes the root of the component

```
 \begin{array}{l} \text{function Find-root}(par, \ p) \\ \mid \ \text{if } par(p) \neq p \ \text{then } par(p) \leftarrow \text{Find-root}(par, \ par(p)) \\ \mid \ \text{return } par(p) \end{array}
```

 $\begin{vmatrix} \text{ for all } p \text{ in } S \text{ forward } \mathbf{do} \\ | q \leftarrow parent(p) \\ \text{ if } f(q) = f(parent(q)) \text{ then } parent(p) \leftarrow parent(q) \end{vmatrix}$

Fig. 3. Union-find without union-by-rank.

i.e parent(p) = p with $p \neq -1$, thus testing $parent(p) \neq -1$ stands for *is p* already processed. Since *S* is processed in reverse order and merge-set sets the root of the tree to the current pixel *p* ($parent(r) \leftarrow p$), it ensures that the parent *p* will be seen before its child *r* when traversing *S* in the direct order.

```
procedure MAXTREE(f)
for all p do parent(p) \leftarrow -1
 S \leftarrow sort pixels increasing
 for all p \in S backward do
     parent(p) \leftarrow p; zpar(p) \leftarrow p
                                                                      ⊳ make-set
     rank(p) \leftarrow 0; repr(p) \leftarrow p
     z_p \leftarrow p
     for all n \in \mathcal{N}_p s.t. parent(n) \neq -1 do
         z_n \leftarrow \text{FIND-ROOT}(zpar, n)
         if z_n \neq z_p then
             parent(repr(z_n)) \leftarrow p
             if rank(z_p) < rank(z_n) then swap(z_p, z_n)
             zpar(z_n) \leftarrow z_p
                                                                     ⊳ merge-set
             repr(z_p) \leftarrow p
             if rank(z_p) = rank(z_n) then rank(z_p) \leftarrow rank(z_p) + 1
 CANONICALIZE(parent, S)
return (parent, S)
```

Fig. 4. Union-find with union-by-rank

2) Union-by-rank: The algorithm given in Fig. 4 is similar to the one in Fig. 3 but augmented with union-by-rank. It first introduces a new image rank. The make-set step creates a tree with a single node, thus with a rank set to 0. The rank image is then used when merging two connected sets in zpar. Let z_p be the root of the connected component of p, and z_n be the root of connected component of $n \in \mathcal{N}(p)$. When merging two components, we have to decide which of z_p or z_n becomes the new root w.r.t their rank. If $rank(z_p) < rank(z_n), z_p$ becomes the root, z_n otherwise. If both z_p and z_n have the same rank then we can choose either z_p or z_n as the new root, but the rank should be incremented by one. On the other

hand, the relation *parent* is unaffected by the union-by-rank, p becomes the new root whatever the rank of z_p and z_n . Whereas without balancing the root of any point p in *zpar* matches the root of p in parent, this is not the case anymore. For every connected components we have to keep a connection between the root of the component in *zpar* and the root of max-tree in *parent*. Thus, we introduce an new image *repr* that keeps this connection updated.



Fig. 5. Union-by-rank. (a) State of the algorithm before processing the neighbor H from E. (b) State of the algorithm after processing.

The union-by-rank technique and structure update are illustrated in Fig. 5. The algorithm has been running until processing E at level 12, the first neighbor B has already been treated and neighbors D and F are skipped because not yet processed. Thus, the algorithm is going to process the last neighbor H. z_p is the root of p in zpar and we retrieve the root z_n of n with find-root procedure. Using *repr* mapping, we look up the corresponding point r of z_n in *parent*. The tree rooted in r is then merged to the tree rooted in p (*parent* merge). Back in zpar, the components rooted in z_p and z_n merge. Since they have the same rank, we choose arbitrary z_p to be the new root.

The algorithm in Fig. 4 is slightly different from the one of [15]. They use two union-find structures, one to build the tree, the other to handle flat zones. In their paper, lowernode $[z_p]$ is an array that maps the root of a component z_p in *zpar* to a point of current level component in *parent* (just like repr (z_p) in our algorithm). Thus, they apply a second union-find to retrieve the canonical element. This extra union-find can be avoided because lowernode [x] is already a canonical element, thus *findoot* on *lowernode* (z_p) is useless and so does *parent* balancing on flat zones.

3) Canonicalization: Both algorithms call the CANONICAL-IZE procedure to ensure that any node's parent is a canonical node. In Fig. 3, canonical property is propagated downward. Sis traversed in direct order such that when processing a pixel p, its parent q has the canonical property that is parent(q) is a canonical element. Hence, if q and parent(q) belongs to the same node i.e f(q) = f(parent(q)), the parent of p is set to the component's canonical element: parent(q).

4) Level compression: Union-by-rank provides time complexity guaranties at the price of an extra memory requirement. When dealing with huge images, it results in a significant drawback (e.g. RAM overflow...). Since the last point processed always becomes the root, union-find without rank

```
function MAXTREE(f)
 for all p do parent(p) \leftarrow -1
 S \leftarrow sort pixels increasing
 i = N - 1
 for all p \in S backward do
     parent(p) \leftarrow p; zpar(p) \leftarrow p
                                                                       ▷ make-set
     z_n = p
     for all n \in \mathcal{N}_p s.t. parent(n) \neq -1 do
          z_n \leftarrow \text{FIND-ROOT}(zpar, n)
         if z_p \neq z_n then
             if f(z_p) = f(z_n) then SWAP(z_p, z_n)
              zpar(z_n) \leftarrow z_p
                                                                       ⊳ merge-set
             parent(z_n) \leftarrow z_p
             S[j] \leftarrow z_n; j \leftarrow j-1
 S[0] \leftarrow parent[S[0]]
 CANONICALIZE(parent, S)
 return (parent, S)
```

Fig. 6. Union-find with level compression.

technique tends to create a degenerated tree in flat zones. Level compression avoids this behavior by a special handling of flat zones. In Fig. 6, p is the point in process at level $\lambda = f(p)$, n a neighbor of p already processed, z_p the root of P_p^{λ} (at first $z_p = p$), z_n the root of P_n^{λ} . We suppose $f(z_p) = f(z_n)$, thus z_p and z_n belong to the same node and we can choose any of them as a canonical element. Normally p should become the root with child z_n but level compression inverts the relation, z_n is kept as the root and z_p becomes a child. Since parent may be inverted, S array is not valid anymore. Hence S is reconstructed, as soon as a point p gets attached to a root node, p will not be processed anymore so it is inserted in back of S. At the end S only misses the tree root which is parent[S[0]].

B. Flooding algorithms

A second class of algorithms, based on flooding, contrasts with the immersion-based algorithms described in the previous section III-A. Salembier et al. [3] proposed the first efficient algorithm to compute the max-tree. A propagation starts from the root that is the pixel at lowest level l_{min} . Pixels in the propagation front are stored in a hierarchical queue composed by as many FIFO queues as the number of levels. It allows to access directly any pixel in the FIFO queue at a given level. Fig. 7 shows a slightly modified version of Salembier's original algorithm where the original STATUS image is replaced by the *parent* image having the same role. The flood (λ, r) procedure is in charge of flooding the peak component P_r^{λ} and building the corresponding sub max-tree rooted in r. It proceeds as follows: first pixels at level λ are retrieved from the queue, their *parent* pointer is set to the canonical element rand their neighbors n are analyzed. If n is not in the queue and has not yet been processed, then n is pushed in the queue for further processing and n is marked as processed (parent(n))is set to INQUEUE which is any value different from -1). If the level l of n is higher than λ then n is in the childhood of the current node, thus flooding is halted at the current level and a recursive call to flood initiates the process for the peak component P_n^l rooted in *n*. During the recursive flooding, some points can be pushed in the queue between level λ and l. Hence, when flood ends, it returns the level l' of n's parent.

function FLOOD(λ , r) while $hqueue[\lambda]$ not empty do $p \leftarrow \text{POP}(hqueue[\lambda])$ $parent(p) \gets r$ if $p \neq r$ then INSERT_FRONT(S, p) for all $n \in \mathcal{N}(p)$ s.t. parent(p) = -1 do $l \leftarrow f(n)$ if levroot[l] = -1 then $levroot[l] \leftarrow n$ PUSH(hqueue[l], n) $parent(n) \leftarrow INQUEUE$ while $l > \lambda$ do $| l \leftarrow flood(l, levroot[l])$ \triangleright Attach to parent 12: $levroot[\lambda] \leftarrow -1$ $lpar \leftarrow \lambda - 1$ while $lpar \ge 0$ and levroot[lpar] = -1 do $| lpar \leftarrow lpar - 1$ if $lpar \neq -1$ then $| parent(r) \leftarrow levroot[lpar]$ INSERT_FRONT(S, r)return lpar function MAX-TREE(f) for all h do $levroot[h] \leftarrow -1$ for all p do $parent(p) \leftarrow -1$ $l_{min} \leftarrow \min_p f(p)$ $p_{min} \leftarrow \arg\min_p f(p)$ $PUSH(hqueue[l_{min}], p_{min})$ $levroot[lmin] \leftarrow p_{min}$ $FLOOD(l_{min}, p_{min})$ Fig. 7. Salembier et al. [3] max-tree algorithm.

1: **procedure** PROCESSSTACK(r, q)2. $\lambda \leftarrow f(q)$ 3: POP(levroot) 4: while *levroot* not empty and $\lambda < f(\text{TOP}(levroot))$ do INSERT_FRONT(S, r) 5: $r \leftarrow parent(r) \leftarrow POP(levroot))$ 6: 7: if *levroot* empty or $f(\text{TOP}(levroot)) \neq \lambda$ then PUSH(levroot, q)8: $parent(r) \leftarrow TOP(levroot)$ 9: INSERT_FRONT(S, r) 10: function MAX-TREE(f) for all p do $parent(p) \leftarrow -1$ 11: $p_{start} \leftarrow any point in \Omega$ 13: $PUSH(pqueue, p_{start}); PUSH(levroot, p_{start})$ 14: $parent(p_{start}) \leftarrow INQUEUE$ 15: loop $p \leftarrow \text{TOP}(pqueue); r \leftarrow \text{TOP}(levroot)$ 16. 17: for all $n \in \mathcal{N}(p)$ such that parent(p) = -1 do 18: PUSH(pqueue, n) $parent(n) \leftarrow \text{INQUEUE}$ 19. if f(p) < f(n) then 20: 21: PUSH(levroot, n)22: goto 16 23: $\{ p \text{ is done } \}$ 24: POP(pqueue) 25: $parent(p) \leftarrow r$ if $p \neq r$ then $\text{INSERT}_{\text{FRONT}}(S, p)$ 26. 27: while pqueue not empty do; 28: { all points at current level done ? } 29: $q \leftarrow \text{TOP}(pqueue)$ 30: if $f(q) \neq f(r)$ then \triangleright Attach r to its parent 31: PROCESSSTACK(r, q)32: repeat 33: $root \leftarrow POP(levroot)$

```
INSERT_FRONT(S, root)
```

34:

Fig. 8. Non-recursive max-tree algorithm [26, 20].

If $l' > \lambda$, we need to flood levels l' until $l' \leq \lambda$ i.e. until there are no more points in the queue above λ . Once all pixels at level λ have been processed, we need to retrieve the level lpar of the parent component and attach r to its canonical element. A levroot array stores canonical element of each level component and -1 if the component is empty. Thus we just have to traverse *levroot* looking for $lpar = max\{h < h\}$ λ , $levroot[h] \neq -1$ and set the parent of r to levroot[lpar]. Since the construction of *parent* is bottom-up, we can safely insert p in front of the S array each time parent(p) is set. For a level component, the canonical element is the last element inserted ensuring a correct ordering of S. Note that the pass which gets the minimum level of the image is not necessary. Instead, we could have called flood in Max-tree procedure until the parent level returned by the function was -1, i.e the last flood call was processing the root. Anyway, this pass has other advantages for optimization that will be discussed in the implementation details section.

Salembier et al. [3]'s algorithm was rewritten in a nonrecursive implementation in [27] and later by [26] and [20]. These algorithms differ in only two points. First, [20] uses a pass to retrieve the root before flooding to mimics the original recursive version while [26] does not. Second, priority queues in [26] use an unacknowledged implementation of heap based on hierarchical queues while in [20] they are implemented using a standard heap (based on comparisons). The algorithm given in Fig. 8 is a code transcription of the method described in [26]. The array levroot in the recursive version is replaced by a stack with the same purpose: storing the canonical element of level components. The hierarchical queue hqueue is replaced by a priority queue pqueue that stores the propagation front. The algorithm starts with some initialization and choose a random point p_{start} as the flooding point. p_{start} is enqueued and pushed on *levroot* as a canonical element. During the flooding, the algorithm picks the point pat highest level (with the highest priority) in the queue, and the canonical element r of its component which is the top of *levroot* (p is not removed from the queue). Like in the recursive version, we look for neighbors n of p and enqueue those that have not yet been seen. If f(n) > f(p), n is pushed on the stack and we immediately flood n (a goto that mimics the recursive call). On the other hand, if all neighbors are in the queue or already processed then p is *done*, it is removed from the queue, parent(p) is set its the canonical element r and if $r \neq p$, p is added to S (we have to ensure that the canonical element will be inserted last). Once p removed from the queue, we have to check if the level component has been fully processed in order to attach the canonical element r to its parent. If the next pixel q has a different level than p, we call the procedure ProcessStack that pops the stack, sets parent relationship between canonical elements and inserts them in Suntil the top component has a level no greater than f(q). If the stack top's level matches q's level, q extends the component so that no more processing is needed. On the other hand, if the stack gets empty or the top level is lesser than f(q), then q is pushed on the stack as the canonical element of a new component. The algorithm ends when all points in queue have been processed, then S only misses the root of the tree which is the single element that remains on the stack.

C. Merge-based algorithms and parallelism

Whereas the algorithms of the two first classes (Sections III-A and III-B) are sequential, this section is dedicated to parallel algorithms. Merge-based algorithms consist in computing max-trees on sub-parts of images and merging back trees to get the max-tree of the whole image [21, 17, 22]. Those algorithms are typically well-suited for parallelism since they adopt a map-reduce idiom. Computation of sub maxtrees (map step), done by any sequential method and merge (reduce-step) are executed in parallel by several threads. In order to improve cache coherence, images should be split in contiguous memory blocks that is, splitting along the first dimension if images are row-major. Fig. 9 shows an example of parallel processing using a map-reduce idiom. The domain has been split into five sub-domains $\{D_1, D_2, \ldots, D_5\}$, we thus have 5 map operations which run a sequential algorithm and 4 joins that merge the sub-trees. Figs. 9b and 9c show a possible distribution of the tasks on 3 threads. Note that map-steps and reduce-steps may occur in parallel, but a single thread may also be in charge of several sub-tree construction. For instance, the first thread is in charge of computing the sub-trees T_1 and T_2 for D_1 and D_2 , merging them into a tree T_{12} and then merging it with the tree computed by the second thread. Choosing the right number of splits and jobs distribution between threads is a difficult topic that depends on the architecture (number of threads available, power frequency of each core). If the domain is not split enough (a number of chunks no greater than the number of threads) the parallelism is not maximal, some threads become idle once they have done their jobs, or wait for other thread to merge. On the other hand, if the number of split gets too large, merging and thread synchronization cause significant overheads. Since work balancing and thread management are outside the current topic, they are delegated to high level parallelism libraries such as Intel Threading Building Blocks (TBB).

The procedure in charge of merging sub-trees T_i and T_j of two adjacent domains D_i and D_j is given in Fig. 10. For two neighbors p and q in the junction of D_i , D_j , it connects components of p's branch in T_i to components of q's branch in T_j until a common ancestor is found. Let x and y be the canonical elements of the components to merge with $f(x) \ge$ f(y) (x is in the childhood to y) and z be the canonical element of the parent component of x. If x is the root of the sub-tree then it gets attached to y and the procedure ends. Otherwise, we traverse up the branch of x to find the component that will



Fig. 9. Map-reduce idiom for max-tree computation. (a) Sub-domains of f. (b) A possible distribution of jobs by threads. (c) Map-reduce operations. M is the map operator, \oplus the merge operator.

procedure CONNECT(p,q) $x \leftarrow \text{FINDREPR}(parent, p)$ $y \leftarrow \text{FINDREPR}(parent, q)$ if f(x) < f(y) then SWAP(x, y)while $x \neq y$ do ▷ common ancestor found ? $parent(x) \leftarrow FINDREPR(parent, parent(x));$ $z \leftarrow parent(x)$ if x = z then $parent(x) \leftarrow y; y \leftarrow x$ else if $f(z) \ge f(y)$ then $x \leftarrow z$ else $parent(x) \leftarrow y; x \leftarrow y; y \leftarrow z$ function FINDREPR(par, p)if $f(p) \neq f(par(p))$ then return p $par(p) \leftarrow FINDREPR(par, par(p))$ return par(p)**procedure** MERGETREE (D_i, D_j) for all $(p,q) \in D_i \times D_j$ such that $q \in \mathcal{N}(p)$ do CONNECT(p,q)

Fig. 10. Tree merge algorithm.

be attached to y that is the lowest node having a level greater than f(y). Once found, x gets attached to y, and we now have to connect y to x's old parent. Function findrepr(p) is used to get the canonical element of p's component whenever the algorithm needs it.

Once sub-trees have been computed and merged into a single tree, it does not hold canonical property (because noncanonical elements are not updated during merge). Also, the reduction step does not merge the S arrays corresponding to sub-trees (it would imply reordering S which is more costly than just recomputing it at the end). Fig. 11 shows an algorithm that canonicalizes and reconstructs S array from *parent* image. It uses an auxiliary image *dejavu* to track nodes that have already been inserted in S. As opposed to other max-tree algorithms, construction of S and processing of nodes are top-down. For any points p, we traverse in a recursive way its path to the root to process its ancestors. When the recursive call returns, *parent*(p) is already inserted in S and holds the canonical property, thus we can safely insert back p in S and canonicalize p as in Fig. 3.

D. Implementation details

Algorithms have been implemented in pure C++ using STL implementation of some basic data structures (heaps, priority queues), MILENA image processing library to provide fundamental image types and I/O functionality, and Intel TBB [28]

procedure CANONICALIZEREC(p)
$$dejavu(p) = true$$
 $q \leftarrow parent(p)$ if not $dejavu(q)$ then $|$ CANONICALIZEREC(q)if $f(q) = f(parent(q))$ then $|$ parent(p) $\leftarrow parent(q)$ $|$ INSERTBACK(S, p)

 $\begin{array}{c|c|c|c|c|c|} \textbf{procedure POST-PROCESS(parent, f)} & | & \textbf{for all } p \textbf{ do } dejavu(p) \leftarrow False \\ & | & \textbf{for all } p \in \Omega \text{ such that not } dejavu(p) \textbf{ do} \\ & | & \text{CANONICALIZEREC}(p) \\ & | & \textbf{return } (parent, S) \\ \end{array}$

Fig. 11. Canonicalization and S computation algorithm.

for parallelism. Specific implementation optimizations are listed below:

Sort optimization. A counting sort is used when quantization is lower than 18 bits. For large integer of q bits, it switches to 2^{16} -based radix sort requiring q/16 counting sorts.

Pre-allocation. Queues and stacks are pre-allocated to avoid dynamic memory reallocation. Hierarchical queues are also pre-allocated by computing image histogram as a pre-processing.

Priority-queues. An heap is implemented with hierarchical queues when quantization is less than 18 bits. For large integer it switches to the STL standard heap implementation. A "y-fast trie" data structure [29] can be used for large integer ensuring a better complexity (see Section IV-A) but no performance gain has been obtained.

Map-reduce. In the parallel version of the algorithms, all instructions that deal about S construction and canonicalization have been removed since S is reconstructed from scratch and *parent* canonicalized by the procedure in Fig. 11

IV. ALGORITHMS COMPARISON

A. Complexity analysis

Let n = H * W with H the image height, W the image width and n the total number of pixels. Let k be the number of values in V.

1) Immersion algorithms: They require sorting pixels, a process of $\Theta(n + k)$ complexity $(k \ll n)$ for small integers (counting sort), $O(n \log \log n)$ for large integers (hybrid radix sort), and $O(n \log n)$ for generic data types with a more complicated ordering relation (comparison sort). Union-find is $O(n \log n)$ and $O(n\alpha(n))$ when used with union-by-rank¹. Canonicalization is linear and does not use extra memory. Memory-wise, sorting may require an auxiliary buffer depending on the algorithm and histograms for integer sorts thus $\Theta(n + k)$ extra-space. Union without rank requires a *zpar* image for path compression ($\Theta(n)$) and the system stack for recursive calls in findroot which is O(n) (findroot could be non-recursive, but memory space is saved at cost of a higher computational time). Union-by-rank requires two extra images (*rank* and *repr*) of *n* pixels each.

2) Flooding algorithms: They require a priority queue to retrieve the highest point in the propagation front. Each point is inserted and removed once, thus the complexity is $\Theta(np)$ where p is the cost of pushing or popping from the heap. If the priority queue is encoded with a hierarchical queue as in [3, 26], it uses n + 2k memory space, provides constant insertion and constant access to the maximum but popping is O(k). In practice, in images with small integers, gray level difference between neighboring pixels is far to be as large as k. With high dynamic image, a heap can be implemented with a y-fast trie [29], which has insertion and deletion in $O(\log \log k)$ and access to maximum element in O(1). For any other data type, a "standard" heap based on comparisons requires n extra space, allows insertion and deletion in $O(\log n)$ and has a constant access to its maximal element. Those algorithms need an array or a stack of respective size k and n. Salembier's algorithm uses the system stack for a recursion of maximum depth k, hence O(k) extra-space.

3) Merge-based algorithms: The complexity depends on $\mathcal{A}(k,n)$, the complexity of the underlying method used to compute the max-trees of sub-domains. Let $s = 2^h$ the number of sub-domains. The map-reduce algorithms require s mapping operations and s - 1 merges. A good map-reduce algorithm would split the domain to form a full and complete tree so we assume all leaves to be at level h. Merging sub-trees of size n/2 has been analyzed in [17] and is $O(k \log n)$ (we merge nodes of every k levels using union-find without union-by-rank). Thus, the complexity of a single reduction is $O(Wk \log n)$. Assuming s constant and $H = W = \sqrt{n}$ the complexity as a function of n and k of the map-reduce algorithm is $O(\mathcal{A}(k,n)) + O(k\sqrt{n}\log n)$. When there is as many splits as rows, s is now dependent on n. This leads to Matas et al. [22] algorithm whose complexity is O(n) + $O(k\sqrt{n}(\log n)^2)$. Contrary to what they claim, when values are small integers the complexity stays linear and is not dominated by merging operations. Finally, canonicalization and S reconstruction have a linear time complexity (CanonicalizeRec is called only once for each point) and only use an image of *n* elements to track points already processed. The complexity analysis for each algorithm as well as the memory required by any auxiliary data structure (including preallocated stacks and queues) is summerized in Table I.

B. Experiments

Benchmarks were performed on an Intel Core i7 (4 physical cores, 8 logical cores). The programs were compiled with gcc 4.7, optimization flags on (-O3 -march=native). Tests were conducted on a dataset of 8-bit images that were re-sized by cropping or tiling the original image. Over-quantization was performed by shifting the eight bits left and generating missing lower bits at random. Fig. 12 depicts performance of the sequential algorithms w.r.t to the size and the quantization. As a first remark, we notice that all algorithms are linear in practice. On natural images, the $n \log n$ upper bound complexity of the [20] and [12] algorithms is not reached. Algorithms from [12] and [15] have quite the same running time ($\pm 6\%$ on average), however the performance of [15] algorithm drops

TABLE I. COMPLEXITY AND SPACE REQUIREMENTS OF MANY MAX-TREE ALGORITHMS. *n* is the number of pixels and *k* the number of gray levels.

	Time complexity			Auxiliary space requirement		
Algorithm	Small int	Large int	Generic V	Small int	Large int	Generic
Berger [12]	$O(n \log n)$	$O(n \log n)$	$O(n \log n)$	n+k+O(n)	2n + O(n)	n + O(t)
Berger + rank	$O(n \alpha(n))$	$O(n \log \log n)$	$O(n \log n)$	3n+k+O(n)	4n + O(n)	3n + O
Najman and Couprie [15]	$O(n \alpha(n))$	$O(n \log \log n)$	$O(n \log n)$	5n+k+O(n)	6n + O(n)	5n + C
Salembier et al. [3]	O(nk)	$O(nk) \simeq O(n^2)$	N/A	3k+n+O(n)	2k + n + O(n)	N/A
Nistér and Stewénius [26]	O(nk)	$O(nk) \simeq O(n^2)$	N/A	2k+2n	2k + 2n	N/A
Wilkinson [20]	$O(n \log n)$	$O(n \log n)$	$O(n \log n)$	3n	3n	3n
Salembier non-recursive	O(nk)	$O(n \log \log n)$	$O(n \log n)$	2k + 2n	3n	3n
Map-reduce	O(A(k,n))	$O(A(\bar{k},n)) + O$	$(k\sqrt{n}\log n)$	$\ldots + n$	$\ldots + n$	$\ldots + r$
Matas et al. [22]	O(n)	$O(n) + O(k\sqrt{2})$	$\overline{n}(\log n)^2)$	k+n	2n	2n



Fig. 12. (a) Comparison of the algorithms on 8-bit images as a function of the size; (b) Comparison of the algorithms on 8 Mega-pixels images as a function of the quantization.

significantly at 256 Mega-pixels. Indeed, at that size each auxiliary array/image requires 1 GB memory space, thus [15] who use a lot of memory exceed the 6 GB RAM limit and need to swap. Our implementation of union-by-rank uses less memory and is on average 42% faster than [15]. Level compression is an efficient optimization that provides 35% speedup on average on [12]. However, this optimization is only reliable on low quantized data. Fig. 12b shows that it is relevant up to 18 bits. It operates on flat-zones but when quantization gets higher, flatzones are less probable and the tests add worthless overheads (see Fig. 14). Union-find is not affected by the quantization but sorting does, counting sort and radix sort complexities are respectively linear and logarithmic with the number of bits. The break in union-find curves between 18 and 20 bits stands for the switch from counting to radix sort. Flooding-based algorithms using hierarchical queues outperform our unionfind by rank on low quantized image by 41% on average. As expected, [3] and [26] (which is the exact non-recursive version of the former) closely match. However, the exponential cost



(c) Fig. 13. (a,b) Comparison of the parallel algorithms on a 6.8 Mega-pixels 8bits image as a function of number of threads. (a) Wall clock time; (b) speedup w.r.t the sequential version; (c) Comparison of the parallel algorithms using 8 threads on a 6.8 Mega-pixels image as a function of the quantization.

of hierarchical queues w.r.t the number of bits is evident on Fig. 12b. By using a standard heap instead of hierarchical queues, [20] does scale well with the number of bits and outperforms every algorithms except our implementation of union-by-rank. In [20], the algorithm is supposed to match [3]'s method for low quantized images, but in our experiments it remains 4 times slower. Since [15]'s algorithm is always



Fig. 16. Decision tree to choose the appropriate max-tree algorithm.

outperformed by our implementation of union-find by rank, it will not be tested any further. Furthermore, because of the strong similarities of [26] and [20], they are merged in our single implementation (called *Non-recursive Salembier* below) that will use hierarchical queues when quantization is below 18 bits and switches to a standard heap implementation otherwise. Finally, the algorithm *Berger* + *level compression* will enable level compression only when the number of bits is below 18.

Fig. 13 shows the results of the map-reduce idiom applied on many algorithms and their parallel versions. As a first result, we can see that better performance is generally achieved with 8 threads that is when the number of threads matches the number of (logical) cores. However, since there are actually only 4 physical cores, we can expect a $\times 4$ maximum speedup. Some algorithms benefit more from map-reduce than others. Union-find-based algorithms are particularly well-suited for parallelism. Union-find with level compression achieves the best speedup, 3.6 times faster that the sequential version while the union-find by rank, second, performs a $\times 3.1$ speedup. More surprising, the map-reduce pattern achieves significant speedup even when a single thread is used ($\times 1.7$ and $\times 1.4$ for union-find with level compression and union-find by rank respectively). This result is explained by a better cache coherence when working on sub-domains that balances tree merges overheads. On the other hand, flooding algorithms do not scale that well because they are limited by post-processes. Indeed, Fig. 15 shows that 76% of the time of parallelized Salembier's algorithm is spent in post-preprocessing (that is going to happen as well for union-find algorithms on architectures with more cores). In [17] and [22], they obtain a speedup almost linear with the number of threads because only a *parent* image is built. If we remove the canonicalization and the S construction steps, we also get those speedups. Fig. 13cshows the exponential complexity of merging trees as number of bits increases that makes parallel algorithms unsuitable for high quantized data. In light of the previous analysis, Fig. 16 provides some guidelines on how to choose the appropriate max-tree algorithm w.r.t. to image types and architectures.

V. CONCLUSION

In this paper, we tried to lead a fair comparison of maxtree algorithms in a unique framework. We highlighted the fact that there is no such thing as the "best" algorithm that outranks all the others in every case and we provided a decision tree to choose the appropriate algorithm *w.r.t.* to data and hardware. We proposed a max-tree algorithm using union-by-rank that outperforms the existing one from [15]. Furthermore, we proposed a second one that uses a new technique, *level compression*, for systems with strict memory constraints. Extra-materials including the image dataset used for this comparison, and a "reproducible research" code, intensively tested, is available on the Internet at http://www.lrde.epita.fr/Olena/MaxtreeReview.

Actually the union-find algorithm is a versatile tool used in many algorithms. A recent publication [30] shows that the morphological *tree of shapes*, which is a self-dual representation of the image contents, can also be computed using union-find. In [31], a specific binary tree, corresponding to an ordered version of the edges of the minimum spanning tree, is computed thanks to a Kruskal-like algorithm and involves the union-find algorithm. Thus, the results presented in this paper also apply to obtain those trees in the most efficient way.

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Fig. 14. Time distribution of the sequential union-find-based algorithms.



Fig. 15. Time distribution of the parallel versions of the algorithms.

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