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Directed connected operators: Asymmetric hierarchies for image filtering and segmentation

Benjamin Perret, Jean Cousty, Olena Tankyevych, Hugues Talbot, *Member, IEEE*, Nicolas Passat

Abstract—Connected operators provide well-established solutions for digital image processing, typically in conjunction with hierarchical schemes. In graph-based frameworks, such operators basically rely on symmetric adjacency relations between pixels. In this article, we introduce a notion of *directed connected operators* for hierarchical image processing, by also considering non-symmetric adjacency relations. The induced image representation models are no longer partition hierarchies (*i.e.*, trees), but directed acyclic graphs that generalize standard morphological tree structures such as component trees, binary partition trees or hierarchical watersheds. We describe how to efficiently build and handle these richer data structures, and we illustrate the versatility of the proposed framework in image filtering and image segmentation.

Index Terms—Mathematical morphology, connected operators, hierarchical image representation, antiextensive filtering, segmentation.

1 INTRODUCTION

GRAPHS are an effective framework for image processing and analysis. They allow for the representation of various adjacency relations (the edges) between pixels (the vertices). Valuation can appear both on the vertices in order to model some information (*e.g.* luminance) and on the edges as a relationship measure. Following the historical symmetric definition of adjacency [1], [2], most methods rely on undirected graphs. Some recent works have aimed at extending these beyond the symmetry hypothesis in order to improve popular image segmentation algorithms. These works have led to different algorithms based on the directed graph framework, and generally show better performances than their symmetric counterpart. Such works include min-cuts [3], random-walkers [4], and shortest path forests [5]. Following these successful attempts, we propose to explore how directed graphs can enrich and improve another family of graph operators: the connected operators. A preliminary version of this work was presented in [6].

1.1 Connected operators

Connected operators [7], [8], [9] are effective image processing tools set in the framework of mathematical morphology.

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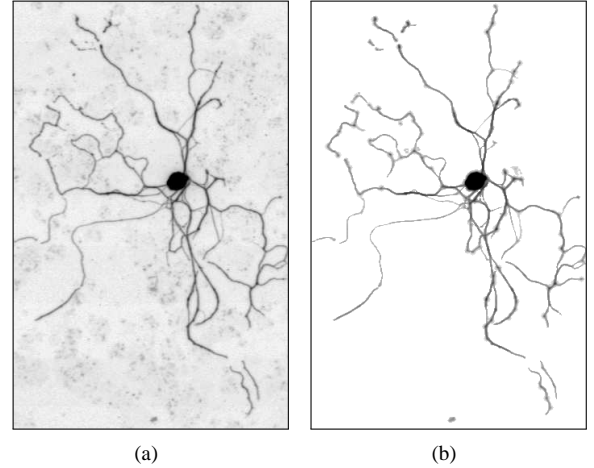


Fig. 1. (a) Neurite image; (b) directed connected filtering.

They have been successful in a wide spectrum of applications (see [10] [11, Ch. 7] for recent surveys). **Connected operators focus on the notion of connected components**, *i.e.*, maximal sets of vertices in which a path exists between any two vertices. Their principle is that the only allowed operation is the deletion of connected components, thus ensuring that they can neither create nor shift contours. The extension of this approach to grayscale images (vertex or edge weighted graphs) leads to the definition of several hierarchical representations: the component tree [12], the binary partition tree [13], or the tree of shapes [14]. Significant effort has been devoted to efficiently construct these hierarchies [12], [15], [16], [17] and to understand the relations that exist between them [18], [19]. A general definition scheme for connected operator consists of four steps: (1) construct the image hierarchical representation; (2) compute attributes at each node of the representation; (3) select relevant nodes according to these attributes; and (4) produce a filtered image or a segmentation map. Connected operators have been used for filtering [12], segmentation [20], interactive segmentation [21], [22], retrieval [23], classification [24], and registration [25]. Applications range from biomedical imaging [26], [27], to astronomy [28], [29], via remote sensing [30], [31] and document analysis [32], [33].

Connected operators face two major issues: (1) making structures of interest appear in the hierarchical representation;

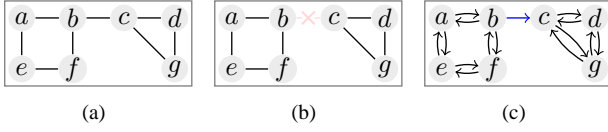


Fig. 2. Undirected and directed graphs (see text).

and (2) discriminating structures of interest in this hierarchy. The first issue has been investigated through the definition of second-generation connections [34], [35], [36], [37], constrained connectivity [38], and hyperconnections [33], [39]. The second issue, *i.e.* selecting relevant nodes of the hierarchy is twofold: (1) defining attributes that provide a suitable feature space able to characterize relevant nodes; and (2) defining robust and accurate node selection processes. Although classical shape attributes (area, elongation, various notions of complexity, ...) are often considered, significant effort has been extended to propose node selection processes. These have evolved from simple global thresholding [9], [12], [20] to energy-minimization strategies [40], [41] and connected filtering in feature spaces [42].

These solutions are effective but are not perfect. We investigate here how the reformulation of connected operators in the context of directed graphs can offer improved practical solutions. Consider the toy example given in Fig. 2(a). The given graph is connected and thus the only two possible results of a connected operator are either the empty graph or the graph itself. To achieve a finer result, for instance knowing that the “rectangle” on the left is only *weakly* connected (perhaps due to noise or some topological considerations) to the “triangle” on the right, one possible solution is simply to remove the edge $\{b, c\}$: this corresponds to second generation connections (Fig. 2(b)). However, by proceeding in this way we lose the information about the initial proximity of the two structures. In the directed graph framework, a **less radical solution is to remove an arc in only one direction**. Then, if we consider the two strongly connected components, we can identify the two parts as separate but still related (Fig. 2(c)). The direction of the remaining arc can also convey some useful information for further processing. An example of this principle is shown in Fig. 1: here the different parts of the neurite are separated using a vesselness [43] prior classification and the directed arcs are constructed so as to always point from least to most reliable structure, as identified in the vesselness: from background, to vessels, to blobs. Filtering based on two attributes that measure the relations (directional information) produces the result shown in Fig. 1(b) (Sec. 6).

1.2 Contributions

In this article, we introduce the new notion of *directed connected component* (*directed component* or *D-component*, for short) which generalizes the notion of connected component to directed graphs (Sec. 2). Furthermore, we establish a bijection theorem (Th. 3) between the D-components and the strongly connected components. In particular, this allows us to rely on well-established tools in graph theory.

We propose the notion of *directed component hierarchy* which extends D-components to weighted graphs (Sec. 2). This structure is a directed acyclic graph, and thus generally is not a tree. However, using a bijection proven in Th. 3, we show that this structure indeed generalizes the standard connected component trees [12], [13].

In Sec. 4 we propose an efficient algorithm for building these hierarchies. The algorithm has a $\mathcal{O}(\ell(n + m))$ time complexity, where m is the number of vertices, n is the number of arcs, and ℓ is the number of weight values.

Then, we present several strategies to select relevant nodes of a D-component hierarchy in order to handle the increased complexity of this structure compared to standard component trees (Sec. 5). These strategies are designed to ensure the consistency of the node selection process in terms of D-components.

Finally, we discuss the methodological and applicative relevance of the D-component hierarchy (Sec. 6). Beyond its obvious relationship with standard symmetric connected operators, we also establish links with non-local paradigms of image processing [44]. In this context, the usefulness of D-component hierarchy is assessed in the challenging case of retinal image segmentation, where it is compared both qualitatively and quantitatively to non-local symmetric morphological approaches as well as gold standard retinal approaches. This analysis is completed by two other application examples in image filtering and segmentation, in order to emphasise the versatility of the proposed framework. In particular, we show how prior information can be injected as a directional information in the graphs and we give examples on how the particular structure of the hierarchy can be used to define new kinds of node attributes.

2 DIRECTED CONNECTEDNESS

The first goal of this article is to extend connected operators from undirected to directed graphs (Sec. 2.1), via employing the *directed connectedness* (or *D-connectedness*) paradigm, which we introduce in Sec. 2.2. Before investigating the differences between D-connectedness and connectedness (defined in the usual frameworks of undirected graphs or connections [45, Ch. 2] [34]), we first discuss the deep links that exist between D-connectedness and the notion of strong connectedness, usually considered on directed graphs (Secs. 2.3, 2.4).

2.1 Graphs

A *directed graph* (or simply, a *graph*) \mathfrak{G} is a pair (V, \mathcal{A}) , where V is a nonempty finite set, and \mathcal{A} is composed of pairs of elements of V , *i.e.*, \mathcal{A} is a subset of $V \times V$. Each element of V is called a *vertex*, a *point*, or a *node* (of \mathfrak{G}) and each element of \mathcal{A} is called an *arc* (of \mathfrak{G}). A *subgraph* of \mathfrak{G} is a graph $\mathfrak{G}_* = (V_*, \mathcal{A}_*)$ such that V_* is a subset of V , and \mathcal{A}_* is a subset of \mathcal{A} . If \mathfrak{G} is a graph, its vertex set is denoted by $V(\mathfrak{G})$ and its arc set by $\mathcal{A}(\mathfrak{G})$.

The *transpose* of a graph \mathfrak{G} is the unique graph with the same vertices as \mathfrak{G} , and such that for any of its arcs (x, y) , the pair (y, x) is an arc of \mathfrak{G} . We say that \mathfrak{G} is *symmetric* if \mathfrak{G} is the same as its transpose. Thus, \mathfrak{G} is symmetric if for

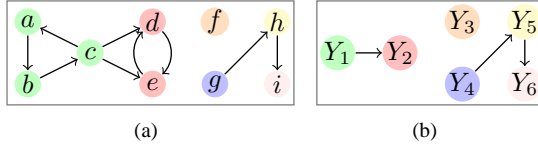


Fig. 3. (a) A directed graph (the vertices and arcs are represented by circles and arrows, respectively) whose D-components are $X_1 = \{a, b, c, d, e\}$, $X_2 = \{d, e\}$, $X_3 = \{f\}$, $X_4 = \{g, h, i\}$, $X_5 = \{h, i\}$ and $X_6 = \{i\}$, and whose S-components are $Y_1 = \{a, b, c\}$, $Y_2 = \{d, e\}$, $Y_3 = \{f\}$, $Y_4 = \{g\}$, $Y_5 = \{h\}$ and $Y_6 = \{i\}$. (b) The DAG $\mathfrak{D}(\mathfrak{G})$ of the S-components of the graph is depicted in (a).

any of its arcs (x, y) , the pair (y, x) is also an arc of \mathfrak{G} . It is well known that any symmetric graph \mathfrak{G} can be associated to a unique undirected graph, and conversely.

Let \mathfrak{G} be a graph, a *path from a vertex x to a vertex y (in \mathfrak{G})* is a sequence (x_0, \dots, x_ℓ) of vertices of \mathfrak{G} such that $x_0 = x$, $x_\ell = y$, and for any i in $\{1, \dots, \ell\}$, the pair (x_{i-1}, x_i) is an arc of \mathfrak{G} . We say that y is a *successor of x (in \mathfrak{G})* and that x is a *predecessor of y (in \mathfrak{G})* if there exists a path from x to y . The singleton (x) is a (trivial) path and therefore x is a successor and a predecessor of itself.

2.2 Directed connected components

In order to take into account “directed subsets” of vertices (i.e., subsets containing some points that play the particular role of “basepoints” or “roots”), we present the notion of a *directed connected component* (or *D-component*).

Definition 1: Let \mathfrak{G} be a graph and let x be a vertex of \mathfrak{G} . The *directed connected component of basepoint x* is the set, denoted by $\text{DCC}_{\mathfrak{G}}(x)$, of all the successors of x in \mathfrak{G} . This set $\text{DCC}_{\mathfrak{G}}(x)$ is also called a *D-component of \mathfrak{G}* , and we denote by $\text{DCC}_{\mathfrak{G}}$ the set of all the D-components of \mathfrak{G} .

For instance, in the graph \mathfrak{G} depicted in Fig. 3(a), the vertices g, h and i are the three successors of g . Thus, the D-component $\text{DCC}_{\mathfrak{G}}(g)$ is the set $\{g, h, i\}$. Observe also that a vertex is a basepoint of a D-component if it is a predecessor of all the vertices in this D-component. For instance, the set $\{a, b, c, d, e\}$ is a D-component and b is a predecessor of all the vertices in this D-component. Therefore, the set $\{a, b, c, d, e\}$ is the D-component of basepoint b . Note that this set is also the D-component of basepoints a and c .

In contrast to connected components, the set of all D-components of a graph is not necessarily a partition of its vertex set. Indeed a vertex may belong to several D-components. For instance (see Fig. 4(a)), let us consider two vertices a and b such that b is a successor of a but a is not a successor of b . Then, the point b is in the D-component of basepoint a and in the D-component of basepoint b . These two D-components are distinct since a belongs to the former one but not to the later. However, these components are linked by inclusion: $\text{DCC}_{\mathfrak{G}}(b) \subseteq \text{DCC}_{\mathfrak{G}}(a)$. More generally, some D-components may intersect without being included in one another. Indeed, let us consider an additional vertex c (see Fig. 4(b)) such that c is a predecessor of b but not a predecessor of a , while c

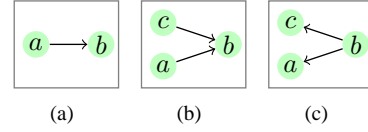


Fig. 4. Some elementary graphs.

is neither a successor of a nor b . Then, the D-components $\text{DCC}_{\mathfrak{G}}(a)$ and $\text{DCC}_{\mathfrak{G}}(c)$ both contain b but are not included in each other since a is in $\text{DCC}_{\mathfrak{G}}(a)$ but not in $\text{DCC}_{\mathfrak{G}}(c)$ and c is in $\text{DCC}_{\mathfrak{G}}(c)$ but not in $\text{DCC}_{\mathfrak{G}}(a)$. However, similarly to the case of connected components, if a vertex x is in a D-component X , then the whole D-component of basepoint x is included in X . In other words, the underlying binary relation “is a successor of” is in general not an equivalence relation but is always reflexive and transitive. We also note that in general the D-components of a graph and of its transpose are not the same. For instance the graphs depicted in Figs. 4(b) and (c) are the transpose of each other and the D-components of the first are $\{c, b\}$, $\{a, b\}$, and $\{b\}$ whereas the D-components of the second are $\{b, a, c\}$, $\{c\}$, and $\{a\}$.

2.3 Strongly connected components

The notion of a strongly connected component is fundamental in graph theory [46, pp. 552–557].

A subset X of the vertex set of a graph \mathfrak{G} is *strongly connected (for \mathfrak{G})* if any two vertices x and y of X are successors of each other, i.e., $x \in \text{DCC}_{\mathfrak{G}}(y)$ and $y \in \text{DCC}_{\mathfrak{G}}(x)$. A *strongly connected component* (or *S-component*) of \mathfrak{G} is a subset X of vertices of \mathfrak{G} that is strongly connected and that is maximal for this property, i.e., any subset of $V(\mathfrak{G})$ which is also a proper superset of X is not strongly connected. We denote by $\text{SCC}_{\mathfrak{G}}$ the set of all S-components of \mathfrak{G} .

This set $\text{SCC}_{\mathfrak{G}}$ of all S-components of a graph \mathfrak{G} – contrarily to the set $\text{DCC}_{\mathfrak{G}}$ of all D-components – is a partition of the vertex set of \mathfrak{G} , i.e., the union of $\text{SCC}_{\mathfrak{G}}$ is $V(\mathfrak{G})$ and the intersection of any two distinct S-components of \mathfrak{G} is empty. In fact, the relation “is in the same S-component as” is an equivalence relation. Thus, for any vertex x of \mathfrak{G} , there is a unique S-component, denoted by $\text{SCC}_{\mathfrak{G}}(x)$, that contains x . For instance, the S-components of the graph depicted in Fig. 3(a) are $\{a, b, c\}$, $\{d, e\}$, $\{f\}$, $\{g\}$, $\{h\}$ and $\{i\}$.

2.4 Links between D- and S-components

Given two vertices x and y of a graph \mathfrak{G} belonging to the same S-component, any successor of x is a successor of y and vice versa. Therefore, the D-components of basepoints x and y are the same. Conversely, if the D-components of basepoints x and y are the same, then x is a successor of y and vice versa, i.e., x and y are in $\text{DCC}_{\mathfrak{G}}(y)$ and in $\text{DCC}_{\mathfrak{G}}(x)$, respectively. In other words x and y are in the same S-component. Hence, S-components and D-components are equivalent according to the following property.

Property 2: Let \mathfrak{G} be a graph. Two vertices x and y of \mathfrak{G} are in the same S-component of \mathfrak{G} if and only if the D-component of basepoint x is equal to the D-component of basepoint y

$$\text{DCC}_{\mathfrak{G}}(x) = \text{DCC}_{\mathfrak{G}}(y) \iff \text{SCC}_{\mathfrak{G}}(x) = \text{SCC}_{\mathfrak{G}}(y) \quad (1)$$

This implies that all the basepoints enabling the definition of a given D-component form a unique S-component. Each D-component is then associated to a unique S-component. In other words, there is a bijection between the set of D-components of \mathfrak{G} and the set of S-components of \mathfrak{G} . This bijection can be expressed based on the *Directed Acyclic Graph* (DAG) of S-components.

We associate to any graph \mathfrak{G} the directed graph $\mathfrak{D}(\mathfrak{G})$, whose vertices are the S-components of \mathfrak{G} and that is such that the pair (X, Y) of S-components of \mathfrak{G} is an arc of $\mathfrak{D}(\mathfrak{G})$ whenever there exists an arc (x, y) of \mathfrak{G} such that x and y are in the S-components X and Y , respectively. This graph has been well studied in graph theory. In particular, it is acyclic, *i.e.*, for any two distinct D-components X and Y , the component X cannot be both a successor and a predecessor of Y in $\mathfrak{D}(\mathfrak{G})$. Therefore, this graph $\mathfrak{D}(\mathfrak{G})$ is called the *DAG of the S-components of \mathfrak{G}* . For instance, the DAG of the S-components of the graph depicted in Fig. 3(a) is depicted in Fig. 3(b). For any S-component X of a graph \mathfrak{G} , we denote by $B_{\mathfrak{G}}(X)$ the union of the successors of X in the graph $\mathfrak{D}(\mathfrak{G})$

$$B_{\mathfrak{G}}(X) = \bigcup \text{DCC}_{\mathfrak{D}(\mathfrak{G})}(X) \quad (2)$$

Theorem 3: Let \mathfrak{G} be a graph.

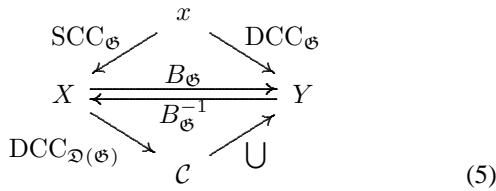
- The map $B_{\mathfrak{G}}$ is a bijection from $\text{SCC}_{\mathfrak{G}}$ to $\text{DCC}_{\mathfrak{G}}$ whose inverse $B_{\mathfrak{G}}^{-1}$ (*i.e.*, $\forall X \in \text{SCC}_{\mathfrak{G}}, B_{\mathfrak{G}}^{-1}(B_{\mathfrak{G}}(X)) = X$) is such that for any D-component X of \mathfrak{G} we have

$$B_{\mathfrak{G}}^{-1}(X) = \{x \in V(\mathfrak{G}) \mid \text{DCC}_{\mathfrak{G}}(x) = X\} \quad (3)$$

- The D-component of any basepoint x is the union of the successors of the S-component $\text{SCC}_{\mathfrak{G}}(x)$ in $\mathfrak{D}(\mathfrak{G})$

$$\text{DCC}_{\mathfrak{G}}(x) = B_{\mathfrak{G}}(\text{SCC}_{\mathfrak{G}}(x)) \quad (4)$$

Th. 3 is illustrated in Diag. (5). In particular, for a given graph \mathfrak{G} , it can be seen that if one knows the S-component X containing a vertex x of \mathfrak{G} , then the D-component Y of basepoint x can be recovered as the direct image of X by the bijection $B_{\mathfrak{G}}$, which can be obtained from the DAG $\mathfrak{D}(\mathfrak{G})$. Conversely, if the directed component Y of basepoint x is known, then one can recover the S-component containing x as the inverse image of Y by $B_{\mathfrak{G}}$. This inverse image $B_{\mathfrak{G}}^{-1}(Y)$ of the D-component Y is called the *root of Y (for \mathfrak{G})*.



In the next sections, we exploit these links between D-components and S-components to design efficient algorithms for image processing.

3 DIRECTED COMPONENT HIERARCHIES

Connected operators act on an image represented as a function through the connected components of its level sets (Sec. 3.1). These connected components are organized – via the inclusion

relationship – in a tree structure, known as the component tree [12]. In this section, we extend this structure from undirected graphs to (directed) graphs. To this end, we present the notions of *strong component tree* (Sec. 3.2) and of *directed component hierarchy* (Sec. 3.3) that encode the inclusion relations of the S-components and D-components of all level sets. The main result of this section is that the directed component hierarchy can be represented as an enriched version of the strong component tree. This enriched version can be further used to define D-connected operators, and to propose efficient algorithms. It is also observed that the directed component hierarchy generalizes the tree structures involved in connected operator definition (Sec. 3.4).

3.1 Stack of graphs

In the framework of undirected graphs, connected operators and component trees have been proposed for the two possible families of weights: those on the vertices and those on the edges. In the first case, a level set is a subset of vertices whose connected components are those of the subgraph induced by these vertices. In the second case, a level set is made of edges, and one considers the subgraphs induced by these edges to obtain connected components. In both cases, the connected components are defined within a series of nested subgraphs induced by the level sets. In order to handle these two cases in a unified and more general setting, one may consider – instead of weights on either edges or vertices – a series of nested subgraphs. Following this approach, we start this section by presenting the notion of a *stack of graphs*.

Definition 4: A *stack (of graphs)* is a finite sequence $\mathcal{S} = (\mathfrak{G}_0, \dots, \mathfrak{G}_\ell)$ of graphs such that, for any i in $\{1, \dots, \ell\}$, the graph \mathfrak{G}_i is a subgraph of \mathfrak{G}_{i-1} . For any i in $\{0, \dots, \ell\}$, we say that \mathfrak{G}_i is a level set of \mathcal{S} (at altitude i). A *S- (resp. D-) component of \mathcal{S}* is a pair (i, X) such that X is a S- (resp. D-) component of the level set of \mathcal{S} at altitude i . The set of all S- (resp. D-) components of the stack \mathcal{S} is denoted by $\text{SCC}_{\mathcal{S}}$ (resp. $\text{DCC}_{\mathcal{S}}$). The stack \mathcal{S} is *connected* whenever \mathfrak{G}_0 is strongly connected.

Fig. 5 first row shows a connected stack composed of five graphs $(\mathfrak{G}_0, \dots, \mathfrak{G}_4)$. In the following, without loss of generality, we assume that the weights of the graphs are positive integers with a maximal value ℓ .

When the domain of an image is considered as the vertex set of a graph \mathfrak{G} , *i.e.*, when the vertices correspond to pixels, the image itself directly leads to a stack of graphs: each level set \mathfrak{G}_i (resp. $\mathfrak{G}_{\ell-i}$) is the subgraph induced by the pixels whose value is greater (resp. lower) than i (*i.e.* the graph whose vertex set the pixels of values greater (resp. lower) than i and whose arc set contains any arc of \mathfrak{G} that links two of these pixels). In this case, the obtained stack is said to be *upper- (resp. lower-) induced by the image*.

For image segmentation tasks, one may also consider similarity measures between pixels that are linked by an arc (for instance, derived from a gradient). Examples of such measures for undirected graphs can be found in [47], [48], [49], [50], [51]. This measure is a function that weights the arcs of the graph \mathfrak{G} . Such an arc-weighted graph also leads to a stack of

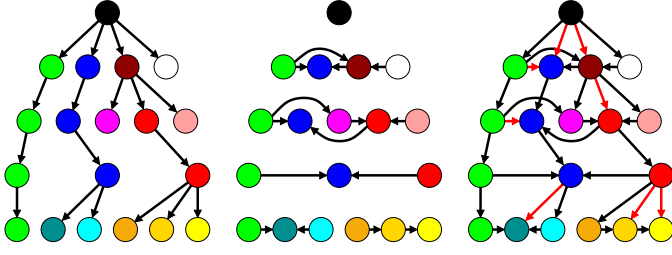


Fig. 6. (a) The S-component tree associated to the stack \mathcal{S} of Fig. 5, first row. (b) The DAGs shown from top to bottom row are the DAGs of S-components of the graphs $\mathfrak{G}_0, \dots, \mathfrak{G}_4$ of Fig. 5, first row. (c) The D-component hierarchy of the stack \mathcal{S} of Fig. 5, second row. This hierarchy is the S-component tree (a) enriched by the relation provided by the DAGs of S-components of all level sets of \mathcal{S} (b). The red arrows are the extra links that are deduced by transitivity.

graphs: each level set \mathfrak{G}_i (resp. $\mathfrak{G}_{\ell-i}$) is the subgraph induced by the arcs of weight greater (resp. lower) than i (i.e., the graph whose arc set contains any arc of weight greater (resp. lower) than i and whose vertex set contains any pixel of \mathfrak{G} linked by one of these arcs). Such a stack is said to be *upper-* (resp. *lower-*) *induced by the similarity measure*. For segmentation methods based on hierarchies of partitions [13], [38], one may want to ensure that all levels in the graph stack remain a partition of the domain by preserving all pixels as vertices of every level set. This can ease further segmentation methods to produce partitions as shown in [18]. A stack obtained by this process is said to be *completed*.

Important notation. In the remaining part of this section, $\mathcal{S} = \{\mathfrak{G}_0, \dots, \mathfrak{G}_\ell\}$ denotes a connected stack.

3.2 Strong component tree

Let X be a S-component of \mathfrak{G}_i , for i in $\{1, \dots, \ell\}$. Since \mathfrak{G}_i is a subgraph of \mathfrak{G}_{i-1} , X is strongly connected in \mathfrak{G}_{i-1} . As the S-components of a graph partition its vertex set, the S-component X of \mathfrak{G}_i is included in a unique S-component of \mathfrak{G}_{i-1} . This unique S-component of \mathfrak{G}_{i-1} that includes X is denoted by $\text{PAR}_{i-1}(X)$ and is called the $(i-1)$ -*parent* of X (in \mathcal{S}). We also say that the S-component $(i-1, \text{PAR}_{i-1}(X))$ of \mathcal{S} is the *parent* of the S-component (i, X) . The set of all S-components of \mathcal{S} equipped with the parent relation is a tree called the *strong component* (or *S-component*) *tree* of \mathcal{S} .

Following the usual terminology on trees, given two S-components (i, X) and (j, Y) of the stack \mathcal{S} , we say that (j, Y) is an *ancestor* of (i, X) and that (i, X) is a *descendant* of (j, Y) if there exists a sequence (C_0, \dots, C_n) of S-components of \mathcal{S} such that $C_0 = (i, X)$, $C_n = (j, Y)$, and C_k is the parent of C_{k-1} for any $k \in \{1, \dots, n\}$. For instance, Fig. 6(a) shows the S-component tree of the stack of Fig. 5.

3.3 Directed component hierarchy

Since distinct D-components of the same graph can be linked by inclusion (see Sec. 2.2), it can be seen that for a given

i in $\{1, \dots, \ell\}$, a D-component of \mathfrak{G}_i can be included in several D-components of \mathfrak{G}_{i-1} . Therefore, contrarily to the case of S-components, the inclusion relations between D-components of successive level sets cannot be directly used for organizing the D-components in a tree structure. Fortunately, as we will see later in this section, the D-components can be arranged as a DAG that is sufficient to recover the inclusion relationship between any two D-components. Furthermore, due to the bijection between S-components and D-components (see Th. 3), this DAG corresponds to an enriched version of the S-component tree. This structure leads to efficient methods, that are described in Secs. 4, 5, for designing D-connected operators. The next theorem is the key result for establishing the properties of this fundamental DAG.

Theorem 5: Let X and Y be two D-components of \mathfrak{G}_i and \mathfrak{G}_{i-1} , respectively, with i in $\{1, \dots, \ell\}$. The D-component X is a subset of the D-component Y if and only if the $(i-1)$ -parent of the root of X is a successor of the root of Y in the DAG of S-components of \mathfrak{G}_i

$$X \subseteq Y \iff \text{PAR}_{i-1}(B_{\mathfrak{G}_i}^{-1}(X)) \in \text{DCC}_{\mathfrak{D}(\mathfrak{G}_{i-1})}(B_{\mathfrak{G}_{i-1}}^{-1}(Y)) \quad (6)$$

More generally, a D-component X of \mathfrak{G}_i is a subset of a D-component Y of \mathfrak{G}_j (with $i \geq j$) if and only if the intersection between the ancestors of the root of X and the successors of the root of Y in $\mathfrak{D}(\mathfrak{G}_j)$ is nonempty. In other words, the set of DAGs of the S-components of all level sets, paired to the parent relation allows us to test the inclusion of any D-components belonging to the stack \mathcal{S} .

Definition 6: The *D-component hierarchy* of \mathcal{S} is the graph whose nodes are the S-components of \mathcal{S} and such that there is an arc from a S-component (j, Y) of \mathcal{S} to a S-component (i, X) of \mathcal{S} if

- (j, Y) is the parent of (i, X) ; or
- $j = i$ and (Y, X) is an arc of the DAG $\mathfrak{D}(\mathfrak{G}_i)$ of S-components of \mathfrak{G}_i .

For instance, the D-components of the stack in Fig. 5 are depicted in the second row of Fig. 5. The associated D-component hierarchy is depicted in Fig. 6(c).

As a corollary to Th. 5, there is an isomorphism between the order induced by the D-component hierarchy of the stack \mathcal{S} and the partial order on the D-components of \mathcal{S} such that $(i, X) \sqsubseteq (j, Y)$ if $i \leq j$ and $X \subseteq Y$. In particular, the S-component (j, Y) is the parent of (i, X) if and only if $B_{\mathfrak{G}_j}(Y)$ is the minimal element (for the inclusion relation) among all the D-components of \mathfrak{G}_j that include the D-component $B_{\mathfrak{G}_i}(X)$. A direct consequence of this isomorphism is that the D-component hierarchy of \mathcal{S} is a DAG. In particular, two S-components at the same level set cannot be linked by a cycle since the DAG of S-components of a graph is acyclic. It can also be seen that two S-components of two distinct level sets cannot be linked by a cycle either since a S-component of a given level set cannot be both an ancestor and a descendant of a S-component of another level set.

3.4 Generalization of tree structures

The framework presented in this section for handling the components of a stack of graphs generalizes the handling of

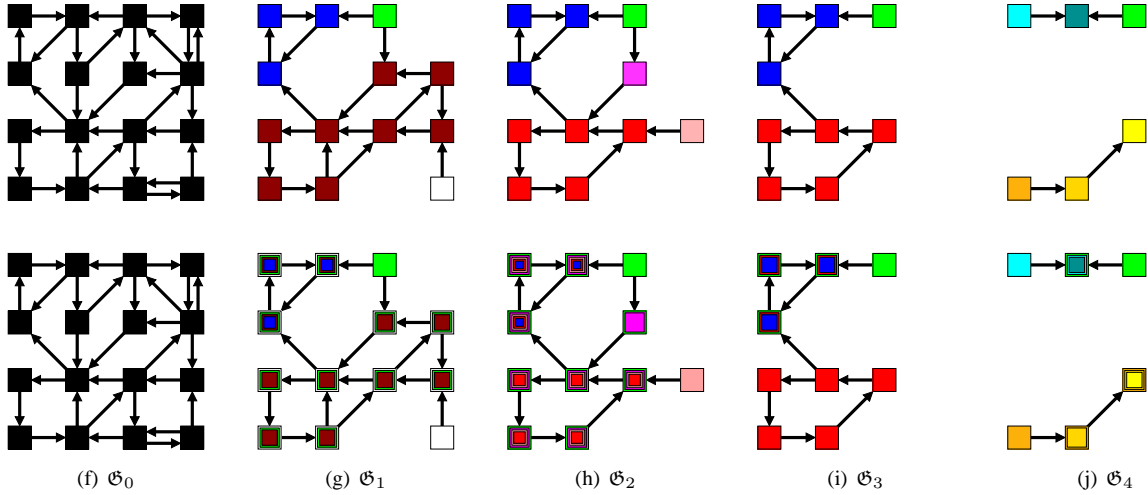


Fig. 5. A stack $\mathcal{S} = (\mathcal{G}_0, \mathcal{G}_1, \mathcal{G}_2, \mathcal{G}_3, \mathcal{G}_4)$. First row: Each color represents a S-component. Second row: Each color represents a D-component (vertices with more than one color belong to all the associated D-components).

connected components via component trees, in both edge- and vertex-weighted undirected graphs.

Indeed, it can be seen that if a graph is symmetric, then a set of vertices is a D-component if and only if it is a connected component in the associated undirected graph. Furthermore, such a set is a D-component if and only if it is a S-component. Hence, in the case of a stack whose level sets are all symmetric graphs, the D-component hierarchy and the S-component tree are indeed the same. Moreover, if a stack is upper (resp. lower) induced by an image, then its D-component hierarchy is also the max- (resp. min-) tree of that image. If a stack is upper (resp. lower) induced by an arc similarity measure, then its D-component hierarchy is the max- (resp. min-) tree of the associated undirected edge-weighted graph. In this last case, if the stack is furthermore completed, then the D-component tree is exactly the partition tree [18] (also known as the quasi flat zones hierarchy [38], [52], [53] or α -tree [54]) of the image. As shown in [18], completed stacks also allow us to retrieve the binary partition trees [13] and hierarchical minimum spanning forests or watersheds [55] [11, Ch. 9] [56].

4 BUILDING D-COMPONENT HIERARCHIES

In this section, we describe how to build the D-component hierarchy of a stack of graphs $\mathcal{S} = \{\mathcal{G}_0, \dots, \mathcal{G}_\ell\}$ (Sec. 4.1), and we discuss the computational cost of this process (Sec. 4.2).

4.1 Algorithm

For the sake of concision, we assume here that the stack \mathcal{S} is constructed from a vertex-weighted graph $\mathcal{G} = (V, \mathcal{A})$ (see Sec. 3.1). We also assume that graphs are represented by adjacency lists: for each vertex x of V , we store the list of vertices y of V adjacent to x (i.e., such that (x, y) is in \mathcal{A}). This representation allows us to access to the list of vertices adjacent to a given vertex in constant time.

The overall construction procedure is described in Alg. 1. Its results consist of: a labeling of each level of the stack \mathcal{S} into S-components ($Label_i$), the adjacency lists of the DAGs

of S-components at each level of the stack (Suc_i), and the parent relation between the S-components of successive levels of the stack (PAR_i).

For each level i of the stack, the algorithm consists of three steps: (1) label the vertices of \mathcal{G}_i into S-components; (2) construct the DAG of S-components of \mathcal{G}_i , i.e., the adjacency lists representing the DAG; and (3) define the parent relation between these S-components and those at altitude $i + 1$.

Step (1) is carried out by either the Tarjan [57] or Kosaraju-Sharir [58] algorithms, which both produce a labeling in S-components of the vertices of a directed graph in linear $\mathcal{O}(|V| + |\mathcal{A}|)$ time. We assume that the labels are integers and that the labels at the different levels are all distinct (i.e., $Label_i \cap Label_j = \emptyset$ for $i \neq j$); so they can be used as array indices. For the sake of readability, we consider that the result $Label_i$ is at the same time the set of labels of $\mathcal{D}(\mathcal{G}_i)$, denoted by $Label_i$, and the map that associates a label, denoted by $Label_i[x]$, to each vertex x of $V(\mathcal{G}_i)$.

Step (2) is performed by Alg. 2. It produces the adjacency list of each vertex of the DAG of the S-components of the level set of \mathcal{G} at altitude i . To this end, it successively scans each S-component of \mathcal{G}_i . For any scanned S-component $SCC[v]$ of label v , the adjacent vertices of all the vertices in $SCC[v]$ are considered. If one of these adjacent vertices belongs to another S-component $SCC[v']$ of label v' (i.e., if $v' \neq v$, line 11), and if $SCC[v']$ has not yet been reached from $SCC[v]$ (i.e., if $Flag[v'] \neq v$, line 11), then the label v' is added to the adjacency list of v (line 12), and v' is flagged as having been reached from v (line 13). The two outer loops visit each vertex once, and for each vertex, its adjacency list is scanned. The algorithm can thus be run in linear $\mathcal{O}(|V| + |\mathcal{A}|)$ time.

Step (3) is performed by Alg. 3 which produces an array such that, for every label v of the S-component labeling at level $i + 1$, the element of index v in the array is the label of the S-component in the level i that includes the S-component v . The algorithm loops through each vertex x in $V(\mathcal{G}_{i+1})$ of the previous level $i + 1$, and finds the labels $Label_{i+1}[x]$ and $Label_i[x]$ of its associated S-components at levels $i + 1$

Algorithm 1: D-component hierarchy construction.

Input: $\mathcal{S} = \{\mathcal{G}_0, \dots, \mathcal{G}_\ell\}$, a stack of graphs.
Output: $Label_i$, S-component labeling for each i in $\{0, \dots, \ell\}$.
Output: Suc_i , array of adjacency lists for each i in $\{0, \dots, \ell\}$.
Output: PAR_i , parent relation for each i in $\{0, \dots, \ell\}$.

```

1 for  $i$  from  $\ell$  to 0 do
2    $Label_i \leftarrow \text{S-component labeling}(\mathcal{G}_i)$ 
3    $Suc_i \leftarrow \text{adjacency lists}(\mathcal{G}_i, Label_i)$ 
4   if  $i \neq \ell$  then
5      $PAR_{i+1} \leftarrow \text{parent relation}(\mathcal{G}_{i+1}, Label_{i+1}, Label_i)$ 

```

Algorithm 2: Adjacency lists construction.

Input: \mathcal{G}_i , a graph.
Input: $Label_i$, S-component labeling of \mathcal{G}_i .
Output: Suc_i , array of adjacency lists of \mathcal{G}_i .

```

1 foreach  $v \in Label_i$  do
2    $Suc_i[v] \leftarrow \emptyset$ 
3    $SCC[v] \leftarrow \emptyset$ 
4    $Flag[v] \leftarrow \text{undefined}$ 
5 foreach  $x \in V(\mathcal{G}_i)$  do
6    $SCC[Label_i[x]] \leftarrow SCC[Label_i[x]] \cup \{x\}$ 
7 foreach  $v \in Label_i$  do
8   foreach  $x \in SCC[v]$  do
9     foreach  $(x, y) \in \mathcal{A}(\mathcal{G}_i)$  do
10       $v' \leftarrow Label_i[y]$ 
11      if  $v' \neq v$  and  $Flag[v'] \neq v$  then
12         $Suc_i[v] \leftarrow Suc_i[v] \cup \{Label_i[y]\}$ 
13         $Flag[y] \leftarrow v$ 

```

Algorithm 3: Parent relation definition.

Input: \mathcal{G}_{i+1} , a directed graph.
Input: $Label_{i+1}$, a labeling of the S-components of \mathcal{G}_{i+1} .
Input: $Label_i$, a labeling of the S-components of \mathcal{G}_i .
Output: PAR_{i+1} , parent relation on $Label_{i+1}$ and $Label_i$.

```

1 foreach  $x \in V(\mathcal{G}_{i+1})$  do
2    $PAR[Label_{i+1}[x]] \leftarrow Label_i[x]$ 

```

and i . It then defines the label $Label_i[x]$ as the parent of the label $Label_{i+1}[x]$. In order to achieve a linear $\mathcal{O}(|V|)$ complexity, the labels have to be determined in constant time which is achieved by storing them for each vertex during the S-component labeling.

4.2 Complexity analysis

The time complexity of Alg. 1 is $\mathcal{O}(\ell \cdot (|V| + |\mathcal{A}|))$. In particular the algorithm is efficient if ℓ is small, for instance in the case of 8-bit images. As there are at most $|V|$ levels (each vertex can have a different weight), this complexity is bounded by $\mathcal{O}(|V|(|V| + |\mathcal{A}|))$. However, if we do not restrict ourselves to vertex-weighted graphs, there are at most $|V| + |\mathcal{A}|$ levels (each vertex and arc is added one after the other), so in the worst case, this leads to a complexity of $\mathcal{O}((|V| + |\mathcal{A}|)^2)$. However, one can generally assume that $\ell \ll |V| + |\mathcal{A}|$.

The algorithm can be improved by observing that a S-component at level i is strongly connected at level $i - 1$. Thus, we can use a more complex algorithm to build the subgraph

at level i . Instead of considering the graph induced by the vertices of weights higher than i , we can add to the DAG of S-components at level $i - 1$, the vertices and the arcs that appear at level i . This requires the use of a union-find structure to dynamically manage the S-components. This generates an additional cost leading to a $\mathcal{O}(\alpha(|V|) \cdot (|V| + |\mathcal{A}|)^2)$ worst case complexity, where α is the extremely slowly growing inverse of the Ackermann function. Practically, we have verified that the improved algorithm is about six times faster than the basic one.

5 SELECTING NODES

Similarly to connected operators, D-connected operators consist of processing a hierarchical data structure, namely the D-component hierarchy. This processing requires selecting or discarding nodes according to criteria that are specifically defined according to the considered application (Sec. 5.1). The selected nodes can then be used, *e.g.*, to obtain a segmentation or to filter an image. Some applications will be described in Sec. 6. Since D-component hierarchies are not tree structures, D-connected operators are more difficult to develop than classically connected ones (Sec. 5.2). In particular, they require specific regularization strategies (Sec. 5.3).

5.1 Node selection criteria

A node selection criterion σ is a predicate associating a Boolean value to each node/component of a hierarchical data structure. Given a component C , we say that the criterion σ *holds true* (resp. *false*) for C , or that C *satisfies* (resp. *violates*) σ if $\sigma(C)$ equals true (resp. false).

A classical criterion σ_A that discards small nodes, often associated to noise, is defined by

$$\sigma_A(C) = \begin{cases} \text{true} & \text{if } \text{Area}(C) > t \\ \text{false} & \text{otherwise} \end{cases} \quad (7)$$

where Area is a measure of the area of the component, for example the number of vertices in C , while t is the area threshold value.

Many other criteria have been proposed in the literature. Most are obtained by replacing Area by some other attribute in Eq. (7). Proposed attributes focus on different aspects: (1) the shape of the component, *e.g.*, the geometrical moments or the compactness; (2) the gray level content of the component, *e.g.*, the volume or the entropy; (3) the topology of the hierarchy, *e.g.*, the number of children of the component; or (4) combination of the previous types of attributes, *e.g.*, the dynamic or the Mumford-Shah energy. It is also possible to replace a constant threshold by a more complex process in the definition of the criterion, *e.g.*, criteria based on energy minimization (Viterbi in [12]) or on shape-space filtering [42].

5.2 The case of D-connected operators

Thinking in terms of D-connected operators, one may desire to mark each D-component as selected or discarded. However – in contrast to the case of connected operators – we may fall into situations such as the one depicted in Fig. 4(b), where two

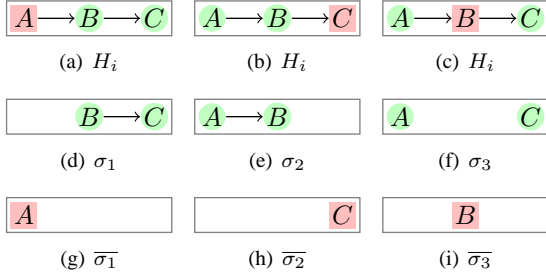


Fig. 7. First row: a graph (DAG) H_i , composed of nodes that are selected (in green) or discarded (in red) with respect to some criterion σ_j (see Sec. 5.2). Each node corresponds to a S-component (A , B and C , respectively), and models a D-component ($A \cup B \cup C$, $B \cup C$ and C , respectively). Second row: subgraphs induced by the nodes that satisfy σ_j (green circles). Third row: subgraphs induced by the nodes that violate σ_j (red squares).

D-components overlap. This creates an ambiguous situation if one of them is selected, while the other is discarded. It is not obvious how to proceed with the S-components that correspond to overlapping D-components. A first partial answer to this question consists of considering the criterion on the S-components instead of the D-components. This choice better suits the data structure constructed in Sec. 4. As well, due to the bijection between D-components and S-components (Th. 3), this strategy is information lossless.

In the following, we consider a stack $\mathcal{S} = \{\mathfrak{G}_0, \dots, \mathfrak{G}_\ell\}$ of graphs. We also consider, for any i in $\{0, \dots, \ell\}$, the DAG at altitude i , denoted by H_i , defined as the subgraph of the D-component hierarchy of \mathcal{S} induced by the S-components of \mathcal{S} at altitude i (i.e., induced by the set of components $\{(i, C) \in \text{SCC}_{\mathcal{S}}\}$). We observe that there is an arc in H_i from the component (i, C_1) to the component (i, C_2) whenever there is an arc from C_1 to C_2 in the DAG $\mathfrak{D}(\mathfrak{G}_i)$ of S-components of \mathfrak{G}_i . Thus, when no confusion may occur, if $C = (i, C')$ is a S-component of \mathcal{S} , we use the symbol C , instead of C' , for the associated strong component C' of \mathfrak{G}_i .

For simplicity – and without loss of generality – we consider the example of the graph H_i , depicted in the first row of Fig. 7, such that $V(H_i)$ is $\{A, B, C\}$ and $\mathcal{A}(H_i)$ is $\{(A, B), (B, C)\}$. Each column of Fig. 7 corresponds to a different criterion: σ_1 holds true for B , C , and holds false for A ; σ_2 holds true for A , B , and holds false for C ; and σ_3 holds true for A , C , and holds false for B . The second line shows the graphs associated to each criterion, i.e., the subgraphs induced by the nodes that satisfy the criterion. The third line shows the graphs induced by the negation of the criterion, i.e., the graphs induced by the nodes that violate the criterion. For the first criterion, it can be seen that the D-components of the graph induced by σ_1 are also D-components of H_i , but the D-component of the graph induced by $\bar{\sigma}_1$ is not a D-component of H_i . The converse situation is true for the second criterion: the D-component of the graph induced by $\bar{\sigma}_2$ is also a D-component of H_i , but the D-components of the graph induced by σ_2 are not. Finally, with the third criterion, neither the D-components of the graphs induced by σ_3 nor by $\bar{\sigma}_3$ are D-components of H_i .

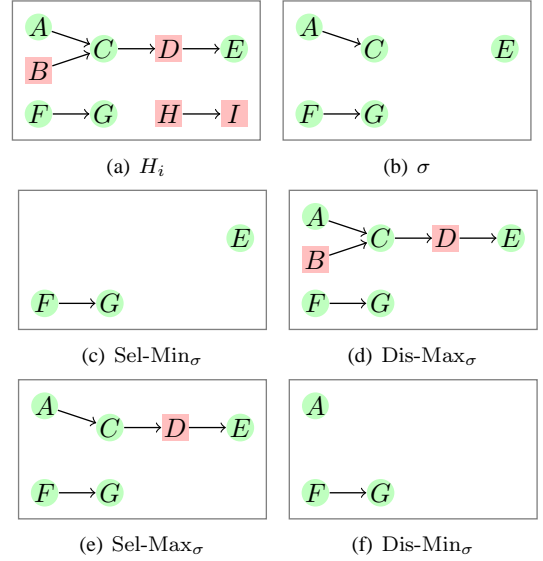


Fig. 8. Node selection (regularization) strategies. (a) The DAG H_i of the S-components $\{A, \dots, I\}$. The components represented as green circles (resp. red squares) satisfy (resp. violate) σ . (b–f) The graphs induced by σ : Sel-Min $_{\sigma}$, Dis-Max $_{\sigma}$, Sel-Max $_{\sigma}$, and Dis-Min $_{\sigma}$.

Thus, we identify two desirable, yet generally exclusive, properties. Given a criterion σ we say that:

- σ is *selective* if the D-components of the graph induced by σ on H_i are also D-components of H_i ;
- σ is *discarding* if the D-components of the graph induced by $\bar{\sigma}$ on H_i are also D-components of H_i .

Furthermore, we say that a D-component C of H_i is *selected* (resp. *discarded*) by σ if C is also a D-component of the graph induced by σ (resp. $\bar{\sigma}$). Thus, C is selected (resp. discarded) if the criterion σ holds true (resp. false) for every S-component contained in C . Nevertheless, we have seen in Fig. 7 that in general, a criterion σ is neither selective nor discarding. Consequently, we propose several regularization strategies that transform any criterion into a selective or a discarding criterion.

5.3 Regularization strategies

Given a criterion σ , we propose four different regularized criteria of σ . Two of them are selective, namely Sel-Min $_{\sigma}$ and Sel-Max $_{\sigma}$. The other two are discarding, namely Dis-Min $_{\sigma}$ and Dis-Max $_{\sigma}$. Fig. 8(a) shows a graph H_i , while Fig. 8(b) shows the result of a non-selective and non-discarding criterion σ on H_i . Then, we have the following regularized criteria:

- Sel-Min $_{\sigma}$ selects the D-components such that every contained S-component satisfies the criterion σ (Fig. 8(c)).
- Dis-Max $_{\sigma}$ discards the D-components such that every contained S-component violates the criterion σ (Fig. 8(d)).
- Sel-Max $_{\sigma}$ selects the D-components whose root S-component satisfies the criterion σ (Fig. 8(e)).
- Dis-Min $_{\sigma}$ discards the D-components whose root S-component violates the criterion σ (Fig. 8(f)).

Thus, for any level i of the hierarchy and for any component C of the DAG H_i at altitude i , we have

$$\text{Sel-Min}_\sigma(C) = \bigwedge_{C' \in \text{DCC}_{H_i}(C)} \sigma(C') \quad (8)$$

$$\text{Dis-Max}_\sigma(C) = \bigvee_{C' \in \text{DCC}_{H_i}(C)} \sigma(C') \quad (9)$$

$$\text{Sel-Max}_\sigma(C) = \bigvee_{C' \in \text{DCC}_{H_i}(C')} \sigma(C') \quad (10)$$

$$\text{Dis-Min}_\sigma(C) = \bigwedge_{C' \in \text{DCC}_{H_i}(C')} \sigma(C') \quad (11)$$

where \bigwedge and \bigvee are the Boolean “and” and “or” operators.

Property 7: Let σ be a criterion on the D-component hierarchy of the stack $\mathcal{S} = \{\mathfrak{G}_0, \dots, \mathfrak{G}_\ell\}$. The regularized criterion Sel-Max_σ (resp. Dis-Min_σ) with respect to \mathcal{S} is the same as the regularized criterion Dis-Max_σ (resp. Sel-Min_σ) with respect to the transpose stack $-\mathcal{S} = \{-\mathfrak{G}_0, \dots, -\mathfrak{G}_\ell\}$, where $-\mathfrak{G}_i$ is the transpose of the graph \mathfrak{G}_i . More precisely, for any DAG H_i at altitude i of \mathcal{S} and for any component C of H_i , we have:

$$\bigvee_{C' \in \text{DCC}_{H_i}(C')} \sigma(C') = \bigvee_{C' \in \text{DCC}_{-H_i}(C)} \sigma(C') \quad (12)$$

$$\bigwedge_{C' \in \text{DCC}_{H_i}(C')} \sigma(C') = \bigwedge_{C' \in \text{DCC}_{-H_i}(C)} \sigma(C') \quad (13)$$

Remark 8: The simplest criteria – that include in particular the criterion σ_A – are those that are *increasing*. We say that a criterion σ is increasing if, for any two S-components C and C' such that the D-component rooted in C is included in the D-component rooted in C' , $\sigma(C) = \text{true}$ implies that $\sigma(C') = \text{true}$. So, given an increasing criterion σ and a S-component C , if σ holds true for C , we immediately know that all the predecessors of C also satisfy σ . Conversely, if σ holds false for C , we immediately know that all the successors of C violate σ , or, in other words, all the S-components contained in the D-component of root C violate the criterion. Thus, any increasing criterion σ is discarding. In this case, σ is equal to Dis-Min_σ and to Dis-Max_σ .

The previous discussions focused on node selection for a single level of the hierarchy. Nevertheless, a similar challenge exists in order to ensure result consistency between the different levels of the hierarchy, *i.e.*, in order to avoid “holes” between two or more levels.

The previously defined regularization rules can also be used on the S-component tree, by considering the ancestors (resp. descendants) instead of the predecessors (resp. successors). These new – but similar – hierarchical criteria are denoted Sel-Max-H , Sel-Min-H , Dis-Max-H , and Dis-Min-H :

- Sel-Max-H_σ holds true for all the descendants of a node that satisfies σ .
- Sel-Min-H_σ holds false for all the ancestors of a node that violates σ .
- Dis-Min-H_σ holds false for the descendants of a node that violates σ .
- Dis-Max-H_σ holds true for all the ancestors of a node that satisfies σ .

In effect, the Dis-Min-H_σ (resp. Dis-Max-H_σ) strategy is the analogue of the usual min (resp. max) filtering rules of the classical component trees [12].

Remark 9: Increasing criteria are consistent with the parent-child relation. Given a criterion σ and two S-components C and C' such that C' is a descendant of C (*i.e.*, $C' \subseteq C$), the D-component rooted in C' is included in the D-component rooted in C , and thus σ is increasing if $\sigma(C') = \text{true}$ implies $\sigma(C) = \text{true}$ (which is the usual definition of an increasing criterion). Thus, for an increasing criterion σ , all the proposed regularization strategies of σ yield the same result as σ due to the tree structure of the S-components at the different levels (by opposition to the DAG of the S-components at a single level).

6 EXPERIMENTS AND DISCUSSION

In this section we illustrate the relevance of the proposed D-component hierarchy framework. This relevance derives from its compliance with efficient image processing paradigms already proposed in the literature. This is notably the case of the non-local approaches, as discussed in Sec. 6.1, and shown by an experimental validation in the context of retinal image analysis. The versatility of the D-component hierarchy framework is exemplified by proposing two applications in Sec. 6.2: one in neurite image filtering and the other in cardiac image segmentation.

6.1 D-component hierarchy and non-locality

Local approaches for image processing rely on the assumption that neighboring pixels are often strongly correlated. In the graph-based formalism, this is generally interpreted by considering standard – symmetric – adjacency relations on \mathbb{Z}^2 or \mathbb{Z}^3 , that model such spatial neighborhoods [59].

In contrast, non-local approaches consider correspondence between pixels that are closely related from a statistical – instead of spatial – point of view. In this framework, all pixels are adjacent to one another. Each adjacency link is weighted by a distance that models the similarity between pixels, or more typically between regions around them (a patch). This approach was popularized in [60] and [44] for image segmentation and filtering and extended to segmentation, reconstruction [61] and classification [62].

A common issue with this approach is its algorithmic complexity. In the graph-based formalism, non-locality implies mapping a complete graph onto the processed image, leading to prohibitive computational costs. Practically, the non-local approach is approximated in a fashion similar to k -nearest neighbors, by *a priori* limiting potential pairing between pixels, for instance based on spatial distance. In so doing, the graph modeling the image becomes sparse, and it also becomes directed. Indeed, a pixel b can be within the k -nearest neighbors of a pixel a while a is not within those of b , for instance due to window size restrictions [44]. In particular, involving non-local approaches in hierarchical frameworks naturally leads to handling D-component hierarchies.

In previous works, these asymmetric, directed adjacencies are not fully exploited, for instance in [60] where the graph

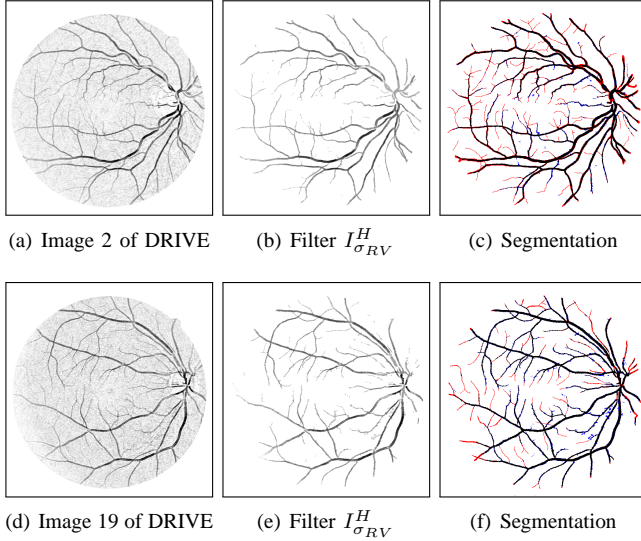


Fig. 9. Segmentation results on the DRIVE database. On each row, from left to right: pre-processed image, filtering result, and evaluation of the segmentation.

is symmetrized. We propose to use this directed information in the D-component framework to achieve improved results.

Retinal image (or eye fundus) analysis (see Figs. 9 (a) and (d)) necessitates to perform blood vessels segmentation, for measuring features like length, width, tortuosity or branching complexity, that can help physicians diagnose and follow-up several pathologies. The difficulty of retinal images lies in the separation of the faint and thin vessels from the background noise. These vessels appear as disconnected groups of pixels that can only be distinguished from the background by their spatial coherency. In order to solve this issue, we propose to construct a non-local directed adjacency relation that allows us to reconnect those groups of pixels, retaining the possibility to reject spurious groups of pixels whose spatial arrangement do not resemble a vessel.

We consider the D-component hierarchy H of the stack $S = \{\mathfrak{G}_0, \dots, \mathfrak{G}_\ell\}$ lower-induced by the image on a non-local graph $\mathfrak{G} = \mathfrak{G}_0$ (Sec. 3.1). The vertex set of \mathfrak{G}_0 is thus the domain of the image (*i.e.*, a rectangular subset of \mathbb{Z}^2). The arc set of \mathfrak{G}_0 corresponds to the traditional 4-neighborhood augmented with non-local arcs based on the k brightest pixels around each pixel.

Formally, given a pixel $x = (x_1, x_2)$, let $\mathcal{N}_4(x)$ be the set of 4-adjacent neighbors of x : $\mathcal{N}_4(x) = \{y = (y_1, y_2), |x_1 - y_1| + |x_2 - y_2| = 1\}$ and let $\mathcal{B}_{k,N}(x)$ be the k brightest pixels in a $N \times N$ window centered on x (x excluded). Then the arc set of \mathfrak{G}_0 is given by: $\mathcal{A}(\mathfrak{G}_0) = \bigcup_{x \in V(\mathfrak{G}_0)} (\{(x, y), y \in \mathcal{N}_4(x)\} \cup \{(y, x), y \in \mathcal{B}_{k,N}(x)\})$. In all the following experiments we set $k = 3$ and $N = 9$. The idea behind this construction is that the brightest pixels are more likely to belong to a vessel. A *jump* is then a possible extension of this vessel.

Fig. 10(a) is a close up on a distal vessel of a retinal image and Fig. 10(b) shows a *critical* threshold (selected manually) with the associated adjacency relation. As expected, there are many non-symmetric links (red arrows). Moreover,

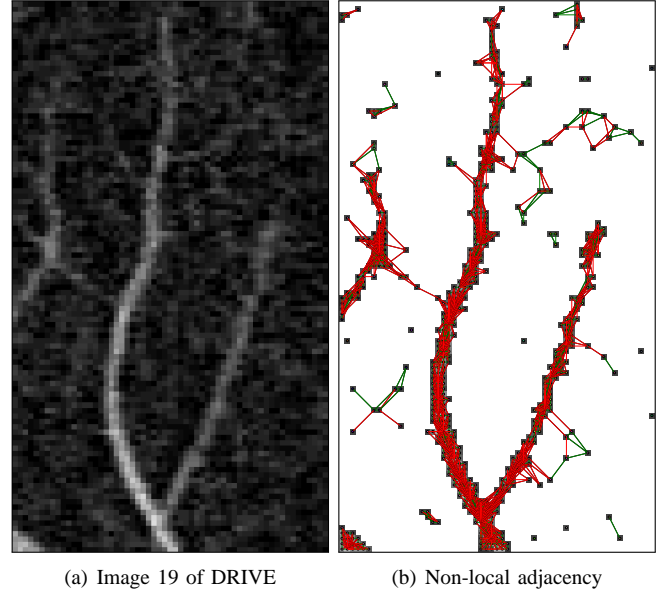


Fig. 10. Non-local adjacency relation. (a) Distal vessels of the 19th retinal image of the DRIVE database. (b) Adjacency relation shown on a critical threshold of (a). Green links represent symmetric edges while red arrows are asymmetric relations. Each S-component is associated to a color printed in a small circle inside each pixel.

we see that the ends of the vessels form thin elongated D-components while noise pixels appear as less spatially structured D-components. The problem can thus be formulated as the deletion of non-vessel-like D-components.

This leads to a selection criterion aiming at removing small scale structures that are not elongated (using a moment based elongation measure) and large scale structures (using the first moment of Hu [63]). Given a set of pixels X , we denote by $\mu_{pq}(X)$ the central moment of X of order pq : $\mu_{pq}(X)$ is equal to $\sum_{(x,y) \in X} (x - \bar{x}(X))^p (y - \bar{y}(X))^q$ with $\bar{x}(X)$ and $\bar{y}(X)$ the coordinates of the centroid of X . Then, we have $\text{Elongation}(X) = \sqrt{\lambda_1(X)/\lambda_2(X)}$ with $\lambda_k(X) = \mu_{20}(X) + \mu_{02}(X) + (-1)^k \sqrt{4\mu_{11}^2(X) + (\mu_{20}(X) - \mu_{02}(X))^2}$. Finally, the first Hu moment of X is defined by $\text{Hu}_1(X) = (\mu_{20}(X) + \mu_{02}(X))/\text{Card}^2(X)$.

Then, given a node C of the hierarchy (*i.e.*, a S-component of \mathfrak{G}_i for some $i \in \llbracket 0, \ell \rrbracket$) and the D-component DCC_C rooted in C (*i.e.*, $\text{DCC}_C = B_{\mathfrak{G}_i}(C)$), the criterion σ_V is defined by

$$\sigma_V(C) = \begin{cases} \text{false} & \text{if } \text{Area}(\text{DCC}_C) \leq t_1 \\ \text{false} & \text{if } t_1 < \text{Area}(\text{DCC}_C) \leq 1000 \\ & \text{and } \text{Elongation}(\text{DCC}_C) > t_2 \\ \text{false} & \text{if } 1000 < \text{Area}(\text{DCC}_C) \\ & \text{and } \text{Hu}_1(\text{DCC}_C) > t_3 \\ \text{true} & \text{otherwise} \end{cases} \quad (14)$$

with t_1 , t_2 and t_3 three thresholds that will be adjusted independently in each experiment. The first case in Eq. (14) discards very small components for which geometric measures are not reliable. We also consider the regularized criterion $\sigma_{RV} = \text{Dis-Min}_{\sigma_V}$ which discards a whole D-component

whenever its root S-component does not satisfy σ_V .

The D-component hierarchy can be used to perform image filtering, *i.e.*, to obtain a new image based on the node selection procedure. This requires to define the *reconstruction* I_σ^H of the hierarchy H with respect to the criterion σ as a function that maps each vertex v of \mathfrak{G}_0 to the altitude of the smallest node of H that satisfies σ and that contains v

$$I_\sigma^H(v) = \max\{i \in \llbracket 0, \ell \rrbracket \mid (i, C) \in V(H), v \in C, \sigma((i, C)) = \text{true}\} \quad (15)$$

Figs. 9 (b) and (e) show the results of retinal image filtering using the criterion σ_{RV} . The resulting filtered image is quite clean. A final segmentation is obtained by selecting every non-zero pixel from this filtered image.

The following results rely on the DRIVE (Digital Retinal Images for Vessel Extraction) database [64], which is composed of 20 test images. Each image of size 565×584 pixels is encoded in 24 bits RGB and comes with 2 expert segmentations and a mask of the eye fundus. The segmentation of an image is evaluated with 3 measures (pixels outside the eye fundus mask do not count): the true positive rate (TPR) (resp. true negative rate (TNR)) is the number of true positive (resp. negative) pixels divided by the total number of positive (resp. negative) pixels, and the accuracy is the sum of true positive and negative pixels divided by the total number of pixels. The score on the base is the mean of the image scores plus the standard deviation of the accuracy (that measures the stability of the algorithm). In our experiments, we used only the green channel, which is pre-processed using a black top-hat (difference between a closing and the original image) with a disk structuring element of diameter 5 pixels. This operation flattens the background of the image and inverts its contrast.

Method	TPR	TNR	Accuracy (σ)
2nd expert	0.7761	0.9725	0.9473 (0.0048)
RNL D-components (15,0.2,1)	0.7079	0.9790	0.9442 (0.0063)
NL D-components (15,0.2,1)	0.7046	0.9790	0.9439 (0.0064)
NL S-components (30,0.15,1.3)	0.7024	0.9789	0.9434 (0.0070)
NL symmetric (Max) (15,0.2,1.3)	0.6528	0.9828	0.9404 (0.0083)
NL symmetric (Min) (15,0.15,1.35)	0.6980	0.9786	0.9425 (0.0067)
Xu [65] (local symmetric)	0.6924	0.9779	0.9413 (0.0078)
Mendonça [66]	0.7344	0.9764	0.9452 (0.0062)
Staal [64]	0.7193	0.9773	0.9442 (0.0065)

TABLE 1

Result comparison on the DRIVE retinal image database.

The methods of the first group are all connected filters.

The last two methods are the state of the art.

We have conducted a set of experiments (see Tab. 1) in order to clearly identify the benefit of each aspect of the method. The evaluated methods are (the best found parameter values (t_1, t_2, t_3) for the criteria σ_V and σ_{RV} are given in Tab. 1 after the method name):

- 1) *RNL D-components*: regularized criterion σ_{RV} on the non-local asymmetric adjacency;
- 2) *NL D-components*: criterion σ_V on the non-local asymmetric adjacency;
- 3) *NL S-components*: same as above but where the criterion σ_V computes the features (area, elongation, Hu_1) on the S-components instead of the D-components;

- 4) *NL symmetric (Max)*: criterion σ_V on a non-local symmetric adjacency where the edge (p, q) belongs to the adjacency if the edges (q, p) or (p, q) are in the asymmetric adjacency;
- 5) *NL symmetric (Min)*: criterion σ_V on a non-local symmetric adjacency where the edge (p, q) belongs to the adjacency if (p, q) and (q, p) are in the asymmetric adjacency;
- 6) *Xu*: connected attribute filter proposed in [65] on a local and symmetric graph with a complex morphological filtering strategy;

Mendonça [66] and Staal [64] provide the state of the art methods. It can be noted that, with Mendonça, the parameters of the method are adjusted independently for each image, their methodology differs thus from the classical one.

The results show that the proposed method 1) achieves the same performances as Staal. In 2) when one removes the regularization strategy, the accuracy goes a bit down showing that the notion of D-component is indeed a good model for vessels in this application. In 3) all the information about D-components is dropped: thus, the method relies only on the S-component tree. The accuracy is lower than in 1) and 2) which shows the interest of exploiting the adjacency relation among the S-components (*i.e.*, the DAG of S-components). In 4) and 5) we measure the gain due to the asymmetric approach. For this, we construct two symmetric non-local adjacency following two classical strategies: min (only symmetric edges are kept) and max (every edge is symmetrized). In both cases the score is lower than in every other experiment. While the max strategy leads to connecting a lot of noise to the vessels (with no mean to disconnect it), the min strategy is indeed quite close to a local approach as most non-local edges of the asymmetric adjacency are asymmetric. 6) is a connected attribute filters (local symmetric adjacency) with a complex node selection strategy recently proposed in [65] which provides a reference score for classical connected filters.

All these experiments show the importance of each element of the method – non-locality (1, 2, 3, 4, and 5), asymmetry and S-components (1, 2, and 3), D-component (1 and 2), regularization (1) – with a gradual improvement of the performances when they are combined.

6.2 Other application examples

6.2.1 Neurite filtering

The proposed framework was also used to filter a sample image of a neuron with associated neurites (*i.e.*, its axon and dendrites), grown in vitro (Fig. 1(a)). We rely on a vesselness-like local object characterization [43], which enables us to classify pixels into tubes, blobs and background. We constructed an undirected pixel adjacency graph where a pixel classified as a tube is linked to its 4-adjacent neighbors classified as blobs but not the other way around. However, two 4-adjacent pixels of the same class are linked in both directions, while any background pixel is linked to all its 4-adjacent pixels. The vertices of this directed graph are then weighted by the gray values of the corresponding pixels and the associated D-component hierarchy is built. Relevant nodes

of the hierarchy are selected by a criterion using two heuristics: (1) a tube must have a (directed) connection with at least two other structures; and (2) a tube is connected at its extremities: the length of the interval between a tube and the structures it is connected to should be small. Similar criteria cannot be designed in the framework of component tree on undirected graphs, since in that framework a component cannot be connected to another (otherwise the two connected components would not be maximal connected sets). Then, from the selected components, a filtered image can be reconstructed (Fig. 1(b)). More details on this illustration are provided in an appendix section, which can be found on-line as supplemental material.

6.2.2 Marker based segmentation

In this section, a D-component hierarchy is used in a marker based image segmentation procedure. To this end, we build a directed arc-weighted graph from the image which is to be segmented. Intuitively, this weighted graph corresponds to a directed gradient. Then, we construct the D-component hierarchy associated to this graph and select D-components in order to obtain a segmentation. The D-components are selected using user-provided markers of the object of interest and of the background. More precisely, a D-component is selected whenever it is rooted in a pixel marked as object and it does not contain any pixel marked as background. The resulting segmentation is the union of the selected D-components.

Fig. 11 illustrates this procedure for the segmentation of the myocardium in a MRI of the heart. A weighted directed 4-adjacency graph is obtained from a rough pre-classification of the image pixels that produces a lot of false negative background pixels but tends to minimize the false positive (see Fig. 11(c)). This classification is obtained by excluding the extremal intensity values, which corresponds to blood and fat for the brightest pixels and to lungs for the darkest. Then, the weight of an arc (x, y) between two 4-adjacent pixels x and y is obtained as the absolute difference of intensity between x and y if y is not “pre-classified as background” or it is set to K times the absolute difference of intensity between x and y if y is pre-classified as background. Figs. 11(d) and (f) show the results obtained when $K = 1$ and $K = 1.5$. Note that when $K = 1$ the weights of an arc (x, y) and of its symmetric (y, x) are always the same and correspond to the magnitude of a simple intensity gradient. In this case, our method is the same as [13], [67] for undirected graphs with symmetric edge-weights. However, when K is greater than 1, the arc weights are not symmetric and the proposed weighting strategy tends to facilitate the connection of the pixels pre-classified as background to the background marker. Fig. 11 clearly shows this behavior and therefore illustrate the benefits of the directed non-symmetric method over its undirected symmetric variant. More details on this method, including a link with the notion of connection value [68], [69] and the oriented IFT segmentation framework [5] can be found on-line as supplemental material.

7 CONCLUSION

We have introduced and investigated a notion of directed connectedness. This has led us to the proposal of new (directed)

connected operators, no longer based on partition hierarchies organized as trees, but on partition covers organized as DAGs.

From a theoretical viewpoint, we have provided a relevant way to generalize various tree-based connected operators previously proposed in the literature. This may lead to a better understanding of the common properties between these operators, and also helps to clarify some subtle differences between those that lie in the framework of directed connected operators, and those that do not, such as hyperconnections. In this context, it is relevant to develop an axiomatization of directed connectedness such as was done in [70], in order to compare it to the axiomatizations already proposed for connections [45, Ch. 2] and hyperconnections [71].

From both the theoretical and algorithmic viewpoints, it may also be useful to compare the links that exist between the DAGs induced by directed connectedness, with other non-tree structures that have been recently introduced to extend the framework of connected operators, for instance in the case of hypertrees [37] or component graphs [72], [73], that constitute an extension of component trees to multivalued images.

From a methodological viewpoint, we have shown that the cover hierarchies obtained when considering directed connectedness can be efficiently handled by taking advantage of the intrinsic links that exist between directed connected and strongly connected components, the latter being organized in trees. Based on these properties, the complexity of the initial algorithm proposed in this article for building cover hierarchies, can be improved by using the recent incremental algorithm proposed in [74] for building the DAG of strongly connected components in $\mathcal{O}(N^{3/2})$ time complexity. Moreover, beyond the standard attribute-based anti-extensive filters developed in this article, other approaches initially devoted to tree structures can be adapted to the case of directed connected operators, and in particular the optimal tree-cut segmentation paradigms initially proposed in [40], and further formalized in the framework of connected operators [75].

From the applicative viewpoints, we have shown that the directed connectedness framework is suitable for efficiently handling non-local image processing paradigms, in their – standard – k nearest neighbor version. The directed connectedness framework is also quite versatile and then useful for various filtering and segmentation tasks. Applications will be more extensively proposed in further works, in particular by comparisons with similar approaches proposed in the literature, for instance in [5].

Source code corresponding to this article is available at the following url: <http://www.esiee.fr/~perretb/dc-hierarchy.html>.

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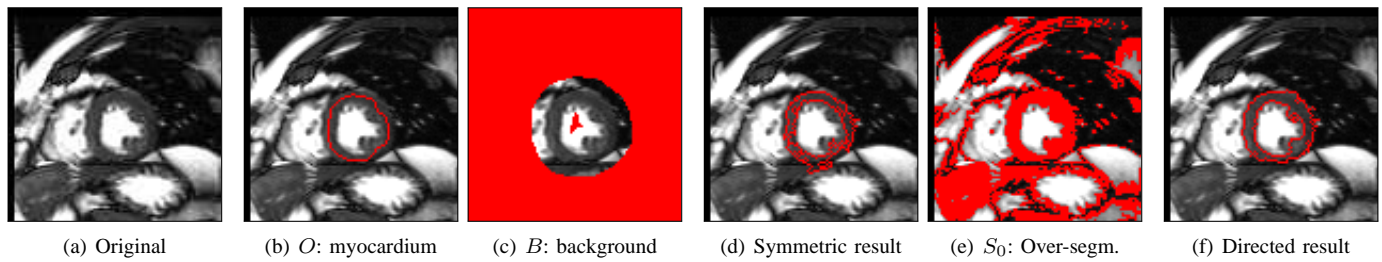


Fig. 11. Segmentation based on the D-component hierarchy. (b,c,e) The considered sets are superimposed in red to the original image. (d,f) The internal border of the segmentation results are superimposed in red to the original image.

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Directed connected operators: Asymmetric hierarchies for image filtering and segmentation

Supplemental Material

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A NEURITE FILTERING

In this example, we consider a sample image of a neuron grown in vitro (Fig. 1(a)), with associated neurites (i.e., its axon and dendrites). The objective is to derive measures of neurite tree complexity, which are useful in various toxicology assays, called neurite outgrowth assays [1]. Whereas neurites do form a tree, apparent overlap in vitro complexify the layout. The challenges of such images are low contrast of neurite vs. background elements unrelated to neurite structures as well as noise, making it complex to segment based only on an intensity criterion.

In our application, we rely on a vesselness-like local object characterization [2], which enables to classify regions into tubes, blobs and background (Fig. 1(c)). This allows us to construct an asymmetric adjacency relation where tubes can be linked to blobs but not the other way around. However, each of the three classes is linked to its own class, while background is linked to all classes. Then, we take into account both intensity and geometrical classification in order to filter the image. By imposing asymmetric blob-to-vessel connection, we exploit the fact that the tube classification is under-segmented and we seek to complete the missing information by searching the connections from the blob and the background classes with this more robust tube class.

A.1 Filtering criteria

We propose an original criterion σ_1 which relies exclusively on the specific structure of the D-component hierarchy, compared to the classical component tree. The criterion aims to hold true for neurite tubes using the following two heuristics: (1) a tube must be connected at least to two other structures; and (2) a tube is connected at its extremities: the length of the interval between a tube and its adjacent structures should be small.

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Formally, for any level i of the hierarchy and for any component C of the DAG H_i at altitude i , the criterion σ_1 relies on two attributes: the number of adjacent nodes $\text{NAdj}(C)$ of C and the number of arcs $\text{NOut}(C)$ going out of C into an adjacent node. For example, in Fig. 3 in [3], the S-component $Y_1\{a, b, c\}$ has one adjacent S-component $\{d, e\}$, but it has two arcs going out of it, (c, d) and (c, e) .

$$\text{NAdj}(C) = |\{C' \in V(H_i), (C, C') \in \mathcal{A}(H_i)\}| \quad (1)$$

$$\text{NOut}(C) = |\{(c, c') \in \mathcal{A}(\mathfrak{G}_i), c \in C, c' \notin C\}| \quad (2)$$

Then, the criterion σ_1 is defined by

$$\sigma_1(C) = \begin{cases} \text{true} & \text{if } \text{NAdj}(C) \geq 2 \text{ and } \text{NOut}(C) \leq 20 \\ \text{false} & \text{otherwise} \end{cases} \quad (3)$$

One can note that the second condition $\text{NOut}(C) \leq 20$ was adjusted empirically. Nevertheless, it roughly corresponds to a frontier of 6 pixels with a 8-neighborhood (3 arcs going out of each pixel on the frontier). More complex measures could be proposed in order to obtain a scale invariance property.

A.2 Result

The filtered image $I_{\sigma_1}^H$ of Fig. 1(a) according to σ_1 is represented in Fig. 1(d). It can be seen that the reconstructed structures are mostly thin and elongated, as expected. Nevertheless, we still miss a few neurite tubes: this problem can be solved by regularizing the criterion σ_1 (see Sec. 5.3 in [3]). Indeed, the chosen adjacency allows us to say that whenever the criterion holds true for a node, then its successors (which can only be classified as blobs or tubes by the vesselness) are also part of the neurite. Thus, we want to select the D-components whose root S-component is selected by σ_1 , which corresponds to the regularized criterion σ_2 given by $\text{Sel-Max}_{\sigma_1}$. Furthermore, in order to fully reconstruct the brightness of the selected S-components, we can perform a hierarchical regularization, setting the final criterion σ_3 to $\text{Sel-Max-H}_{\sigma_2}$. The filtered image $I_{\sigma_3}^H$ of the image Fig. 1(a) according to σ_3 is represented in Fig. 1(b). Moreover, Fig. 1(e) shows the non null pixels of $I_{\sigma_1}^H$ (Fig. 1(d)) that were classified as background with the

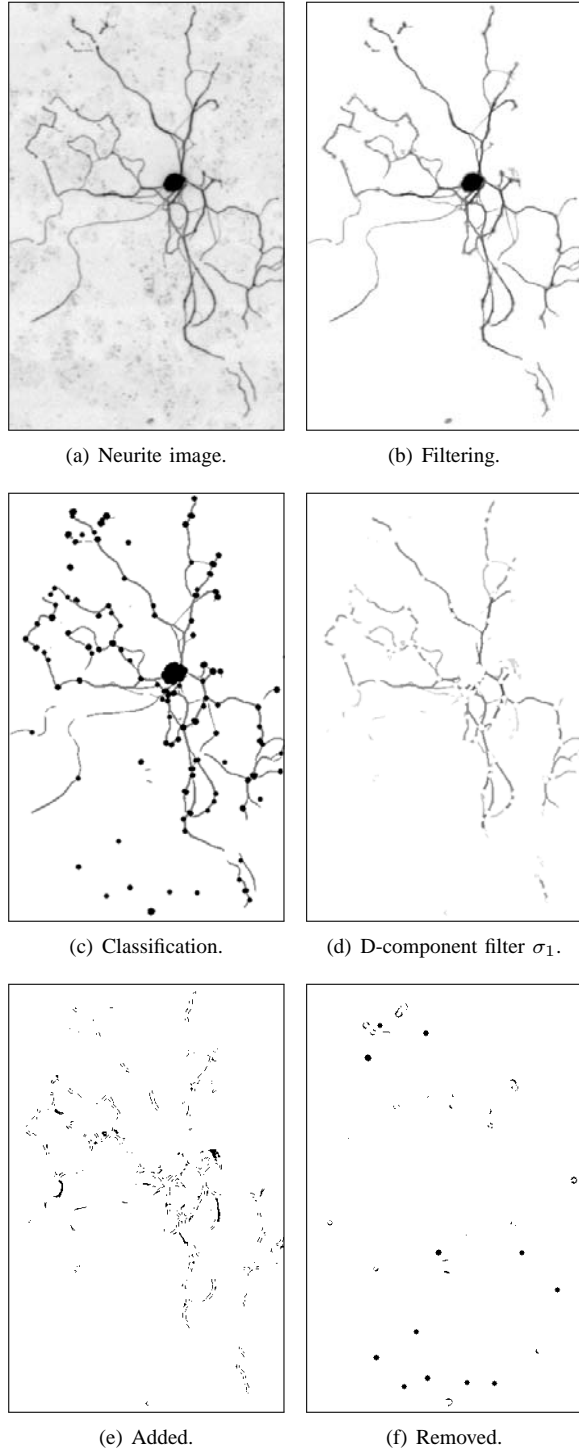


Fig. 1. D-component filter on a neurite image. (a) Neurite image. (b) Directed connected filtering. (c) A classification of (a) into neurite “tubes” (gray), “blobs” (black), and “background” (white) used to generate the directed adjacency. (d) Results of the filtering criterion σ_1 . (e) What has been added in (b) compared to (a). (f) What has been removed from (a) in (b).

vesselness (Fig. 1(c)), i.e., the false negatives. We can observe that the selection process was able to recover several neurite tubes that were classified as background. Conversely, Fig. 1(f) shows the null pixels of $I_{\sigma_1}^H$ (Fig. 1(d)) that were not classified as background with the vesselness (Fig. 1(c)), i.e., the false positives. We can see that isolated structures were correctly identified as non-neurite parts.

B MARKER BASED SEGMENTATION

In this section, we illustrate the use of the proposed framework in a marker-based image segmentation procedure. To this end, we consider the magnetic resonance image of the heart shown in Fig. 2(a). From this image, we aim at segmenting the left ventricular myocardium (see the segmented result in Fig. 2(f)) from the two markers of the myocardium and of the background that are shown in red in Figs. 2(b,c). For this illustration, the markers were manually overlaid on the image. However, in a clinical context, these markers may be obtained via an automated procedure (see, e.g., [4]).

As is classically done with graph based segmentation methods, we consider an arc weighted graph (\mathcal{G}, w) such that the arc weights function w is a dissimilarity measure based on the gradient magnitude of the image. The vertex set of \mathcal{G} is the domain of the image to be processed (i.e., a rectangular subset of \mathbb{Z}^2) and the arc set of \mathcal{G} is given by the 4-adjacency relation: the pair (x, y) is an arc of \mathcal{G} if $|x_1 - y_1| + |x_2 - y_2| = 1$, where $x = (x_1, x_2)$ and $y = (y_1, y_2)$. The graph considered in our illustration is symmetric but the weight of an arc (x, y) from a pixel x to a pixel y is not necessarily equal to the weight of the symmetric arc (y, x) from y to x . The setting of the weight map w will be given after the description of the segmentation procedure and a property of its result that guided us for defining the map w .

Given the arc-weighted graph (\mathcal{G}, w) and two sets of pixels, denoted by O and B , which correspond to the marker of the object (myocardium, see Fig. 2(b)) and of the background (see Fig. 2(c)), respectively, our method consists of two steps:

- 1) build the D-component hierarchy of the weighted graph (\mathcal{G}, w) (or more precisely of the completed stack lower induced by the similarity measure w , see Sec. 3.1 in [3]);
- 2) select from this hierarchy all D-components rooted in a pixel marked with the label object (i.e., a vertex that belongs to O) and that does not contain any pixel marked as background (i.e., any vertex that belongs to B).

The resulting segmentation, denoted by $S_{O \rightarrow B}$, is the union of the selected D-components. More formally, the segmentation is made of the components selected by the regularization $\text{Sel-Max}_{\sigma_4}$ of a simple criterion σ_4 defined for any level i of the hierarchy and for any component C of the DAG H_i at altitude i by

$$\sigma_4(C) = \text{true} \iff C \cap O \neq \emptyset \text{ and } \text{DCC}_{H_i}(C) \cap B = \emptyset \quad (4)$$

Step 1 is performed by the algorithm of Sec. 4 in [3], whereas Step 2 is performed by recursive traversals of the hierarchy.

The segmentation result can be characterized owing to a directed version of a measure widely used for segmentation purposes in undirected weighted graphs, namely the

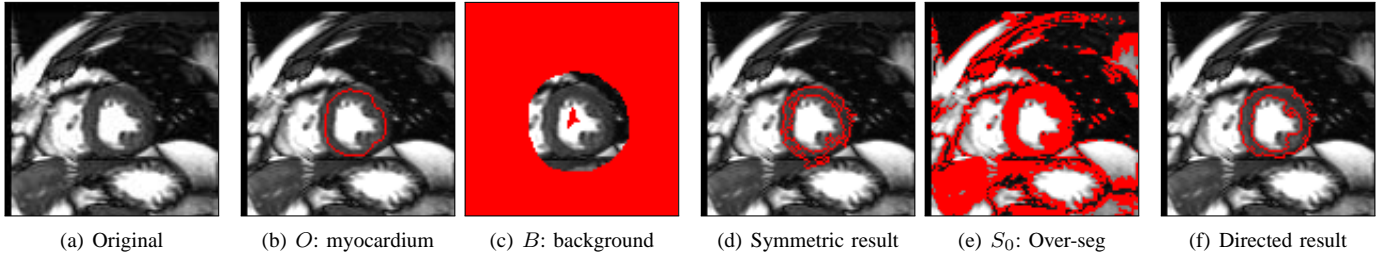


Fig. 2. Segmentation based on the D-component hierarchy. (b,c,e) The considered sets are superimposed in red to the original image. (d,f) The internal border of the segmentation results are superimposed in red to the original image.

connection value [5], [6] (also called degree of connectivity [7] or fuzzy connectedness [8] up to an inversion of w). If $\pi = (x_0, \dots, x_\ell)$ is a path in \mathfrak{G} , the *connection value* $\Upsilon_w(\pi)$ of π is the maximum weight of the arcs of π

$$\Upsilon_w(\pi) = \begin{cases} \max\{w(x_i, x_{i-1}), i \in \llbracket 1, \ell \rrbracket\} & \text{if } \pi \text{ non-trivial} \\ 0 & \text{otherwise} \end{cases} \quad (5)$$

The (*directed*) connection value $\Upsilon_w(x, y)$ from a vertex x of \mathfrak{G} to a vertex y is then defined as the minimum of the connection values of the paths from x to y

$$\Upsilon_w(x, y) = \min\{\Upsilon_w(\pi), \pi \text{ is a path from } x \text{ to } y\} \quad (6)$$

Hence, by the very definition of a D-component hierarchy, it can be shown that the segmentation result $S_{O \rightarrow B}$ is the set of all vertices whose directed connection value from a vertex marked as object is less than the one to a vertex marked as background, i.e., for any $x \in V(\mathfrak{G})$, we have

$$x \in S_{O \rightarrow B} \iff \min_{o \in O} \{\Upsilon_w(o, x)\} < \min_{b \in B} \{\Upsilon_w(x, b)\} \quad (7)$$

From the previous characterizations of the proposed segmentation method, we can say intuitively that the weights of the arcs should be higher in the border of the object to be segmented than inside and outside this object. In order to produce such a map and, following the work of [9], the weight $w(x, y)$ of the arc (x, y) is the product of a symmetric term $\nabla(x, y)$ and a non necessarily symmetric term $\delta(x, y)$

$$w(x, y) = \nabla(x, y) \cdot \delta(x, y) \quad (8)$$

The first term $\nabla(x, y)$ plays the role of a gradient magnitude. For our illustration, it is defined as the absolute difference of intensity between the pixels x and y . If I denotes the grayscale image to be segmented we have

$$\nabla(x, y) = |I(x) - I(y)| \quad (9)$$

when no further knowledge on the objects to be segmented exists, symmetric weights can be used. In this case the second term $\delta(x, y)$ is set to a constant value. For instance, the image of Fig. 2(d) is obtained by the proposed method from the markers O and B shown in Figs. 2(b,c) when a constant value is used for the map δ .

However, when domain knowledge is available, the second term $\delta(x, y)$ can be used to ease the connection of a pixel either to the background or to the foreground marker. From Eq. (7), we remark that if the weight of every arc (x, y)

ending at a vertex y is set to an infinite value (i.e., a maximal weight), then the vertex y does not belong to the segmented object $S_{O \rightarrow B}$. Indeed, in this case, the connection value from any point o of the marker O to the vertex y is maximal and therefore, by Eq. (7), the vertex y does not belong to the segmented object $S_{O \rightarrow B}$. On the other hand, when the weight of every arc starting at x is maximal (i.e., in the cases where the connection value from the object marker is not also maximal), the point x generally belongs to the segmented object $S_{O \rightarrow B}$. These observations allow us to design a strategy to take into account some prior information in the arcs weights in order to ease the connection of a given point either to the background marker or to object marker. Indeed, if we have good confidence that a pixel x belongs to the object (but not high enough to put it in the object marker), we can increase the weights of the arcs going out from x in order to harden its connection to the background. Conversely, if we are confident that a pixel x belongs to the background, we have to increase the weights of the arcs ending in x .

Following this strategy, we set the term $\delta(x, y)$ in Eq. (8) based on a first classification of the image pixels that produces many false positives but tends to minimize the false negatives. In other words, this classification tends to produce a superset, denoted by S_0 , of the expected segmentation result. In our illustration, S_0 is obtained by excluding the extremal intensity values, which correspond to blood and fat for the brightest pixels and to lungs for the darkest ones (see Fig. 2(e)). Hence, we are confident that the points that do not belong to S_0 do not belong to the object that we seek to segment. Then, the terms $\{\delta(x, y)\}$ are set up in order to ease the connection of the pixels that do not belong to S_0 , to the background marker B . This is done by multiplying by a constant value the weight of any arc (x, y) ending at a pixel classified as background

$$\delta(x, y) = \begin{cases} 1 & \text{if } y \notin S_0 \\ K & \text{otherwise} \end{cases} \quad (10)$$

where $K > 1$.

The result of our method (with $K = 1.5$) for the myocardium segmentation is presented in Fig. 2(f).

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