Descriptive Image Gradient from Edge-Weighted Image Graph and Random Forests

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Abstract—Creating an image gradient is a transformation process that aims to enhance desirable properties of an image, whilst leaving aside noise and non-descriptive characteristics. Many algorithms in image processing rely on a good image gradient to perform properly on tasks such as edge detection and segmentation. In this work, we propose a novel method to create a very descriptive image gradient using edge-weighted graphs as a structured input for the random forest algorithm. On the one side, the spatial connectivity of the image pixels gives us a structured representation of a grid graph, creating a particular transformed space close to the spatial domain of the images, but strengthened with relational aspects. On the other side, random forest is a fast, simple and scalable machine learning method, suited to work with high-dimensional and small samples of data. The local variation representation of the edge-weighted graph, aggregated with the random forest implicit regularization process, serves as a gradient operator delimited by the graph adjacency relation in which noises are mitigated and desirable characteristics reinforced. In this work, we discuss the graph structure, machine learning on graphs and the random forest operating on graphs for image processing. We tested the created gradients on the hierarchical watershed algorithm, a segmentation method that is dependent on the input gradient. The segmentation results obtained from the proposed method demonstrated to be superior compared to other popular gradients methods.

I. INTRODUCTION

Image segmentation may be considered as a semantic task and it is an active topic of research [1]. This task consists in partitioning perceptually similar pixels into sets of regions representing areas of interest. Usually this task is done in two stages: (i) the extraction of image characteristics that facilitates interpretation and further analysis; and (ii) the mapping of these characteristics into coherent regions. A coherent region is a subjective concept, but according to [2], it must present characteristics such as: (i) uniformity; (ii) continuity; (iii) contrast between adjacent regions; and (iv) well-defined boundaries. Independently on how well-designed a mapping method is, most of them are limited by the characteristics extracted on the first stage. For instance, taking the grey-level contrast on the first stage produces a great variation between regions, but have absolute values very distinct, making it harder to determine which value actually represents a region change.

Image gradients are known to facilitate the analysis by enhancing desirable properties, extracting structural elements of an image usually based on pixel intensities. Fast and

![Input image](a) Ground-truth
![SED](c) Sobel
![Laplacian](e) Proposed

Fig. 1. Examples of gradient computation, for the input image (a). In (c), the SED gradient presents reinforced fuzzy borders of the main objects and small details are in large ignored. Sobel (d) presents very thin edges for both large and small objects, while large uniform regions, such as the asphalt and vegetation are discretely represented. For Laplacian in (e), it is perceived a large amount of noise for objects, edges and patterns. The proposed in (f) computed enhanced borders for both large and small objects and image textures are firmly represented with different simplified patterns.
inexpensive to compute, they are commonly used as a pre-
processing step in multiple applications, such as medical
analysis [3], [4], text extraction [5], video processing [6],
[7] and segmentation [8], [9]. Even with the advent of deep
networks, gradient use continues to be relevant due to its
performance. Also, the gradients are used as support for some
networks, providing enhanced features or reducing computation
complexity [10]–[12]. Traditional gradient methods, such as
Laplacian and Sobel, are kernel filters for local variation,
highlighting the borders of objects and are usually very
sensitive to abrupt changes on the original image. In 2014, [13]
proposed a method for structured edge detection (SED) that is
fast and precise to predict object edges, and became a common
approach as image gradient creator for the segmentation task.
SED extends the Random Forest (RF) [14] formalism to
a general structured output space, using local patches to
map similar structured labels to the same discrete label. The
gradients produced by SED have enlarged fuzzy borders of
the main objects present on the image, while small details
and other regions are in large omitted. In Fig. 1, we illustrate
the gradient computed by SED, Sobel and Laplacian (Fig. 1
(c), (d) and (e) respectively) from the input image in Fig. 1(a).

The kernel methods and SED are widely used [3]–[12],
achieving their goal of enhancing the borders and partially
the contrast between regions. In this work, we argue that
although the borders constitute an important characteristic
of the objects depicted in images, other properties that reflect the
uniformity, homogeneity and continuity are also important for
the interpretation of coherent regions, particularly on the task
of image segmentation.

To reflect these characteristics, we propose a novel method
to create gradients that firmly depicts edges of large and
small objects as well as uniform regions and patterns on
the image, making it a very descriptive image gradient. It
is proposed to use edge-weighted graphs aggregated with the
Random Forest (RF) [14]. This approach is motivated because
the spatial connectivity of the pixels of an image gives us a
structured representation of a grid graph, creating a particular
transformed space close to the spatial domain of the images,
but strengthened with relational aspects. The edge-weighted
graph as an image gradient operator acts as a transformation
filter on the image, but as in the case of many spatial filters
based on local differences, it is subjected to respond strongly
to noise. We expect that the attribute selection and implicit
regularization process [15] on the RF trees can mitigate this
aspect while reinforcing desirable characteristics.

As in many machine learning algorithms, RF requires a
systematic input and a strategy must be placed to deal with the
dynamic structure of the graphs. The goal is to maintain the
main characteristics of the graph—e.g., topology, relationships,
esential features—without losing too much information. A
popular approach is graph embedding [16]–[21] that creates
good vectorial representations, but is extremely expensive for
image graphs in terms of computational resources and time.

Alternatively to graph embedding, authors working with
graphs as input for the RF also proposed to create a regular
representation by the use of: (i) graph adjacency matrix [22];
(ii) graph topology measures (e.g., centrality, community) [23],
[24]; (iii) selection of attributes [25], [26]; (iv) pairwise com-
parison of graph components [27]; and (v) feature inference
methods [28]. We propose to make a selection of attributes
that preserves the main features of the original image and
incorporates the relational aspects of the graph.

The main contributions of this work are three-fold:

1) The use of graph representations of images as input of
Random Forests, in which: (i) the regions of analysis are
delineated in the neighborhoods inside the graph; (ii) the feature space is defined in terms of graph attributes; and (iii) the discrete label attribution is centered on the components, therefore the entirety of a neighborhood is assigned to a single label. It aims to keep the main features of the original image and the information about relational aspects of the graph. Also, the standard discrete label is considerably faster to train than the complex structured output space mapping in SED—to be precise, experiments training each tree of the proposed method and SED took approximately 2 seconds for the former and 4 hours for the latter, using the same CPU.

2) A straightforward strategy to create a systematic input to implement machine learning on grid graphs, a topic of great interest recently due to: (i) its autonomy—the source data becomes virtually irrelevant once the learning system operates on graphs; (ii) the multiple possibilities of applications; and (iii) the graph capacity to represent multivariate information.

3) The description of how RF could be used for image processing by mapping the RF predictions back to the image space in the form of very descriptive image gradients. The quality of the obtained gradients is assessed qualitatively and quantitatively on a segmentation task. Some segmentation samples are presented in Fig. 2, obtained from the proposed method, SED and Sobel.

From now on, we will refer to the proposed method as \textbf{graph-based image gradient (GIG)}, illustrated in Fig. 1(e).

This work is organized as follows. In Section II, we give a brief description of graphs, their components and terminologies. An overview on RF and the proposed strategy to apply the graph structured input with attribute selection for learning are given in Section III. In Section IV, we provide some experimental results to demonstrate how these proposed gradients could be successfully used to improve segmentation results of one well-known segmentation algorithm, called hierarchies of watershed [29], in contrast with other popular methods for the gradient, illustrated in Fig. 1. And finally, in Section V, we draw some conclusions.

II. \textbf{Edge-Weighted Image Graphs}

The main concern in graph theory is the interconnection of objects, depicting many data. In this section, we provide a brief description of its components and terminologies, mostly following the notations in [30].

\textbf{Definition 1.} A (undirected) graph \(G = (V, E)\) consists of a finite nonempty set of vertices, denoted by \(V\), and a finite set of edges \(\{(u,v) \mid u, v \in V\}\), denoted by \(E \subseteq V \times V\).

The notion of vertices relates to the representation of the basic components of the data and the edges to the connections and dynamic between them. Furthermore, multiple functions could be associated with each vertex and/or edge, in order to enhance the relational aspects, interpretation of different adjacency relations and insertion of metric properties.

\textbf{Definition 2.} An \textbf{edge-weighted} graph could be denoted by \((G, F)\) in which \(F : V \times V \to \mathbb{R}\) is a function that weights the edges of \(G\). The set of all functions that could be used to weight the edges of a graph is denoted by \(\mathcal{F}(E)\).

The preserved characteristics on the graph depend on the nature of \(F\) and the problem of selecting a function to weight an edge could be considered as a problem of measuring the similarity between two finite sets of points.

\textbf{Definition 3.} The set \(E\) induces a unique \textbf{adjacency relation} \(\Gamma\) on \(V\), which associates \(u \in V\) with \(\Gamma(u) = \{v \in V \mid (u,v) \in E\}\). \(\Gamma\) is reflexive \((u,u) \in \Gamma(u)\) and symmetric \((u,v) \in \Gamma(u) \iff (v,u) \in \Gamma(v)\).

The discussions about graph creation and manipulation can be made generic enough to model any data, but, for instance, we are interested on image graphs, in which the spatial connectivity of the pixels gives us a structured representation of a grid graph, close to the spatial domain and strengthened with relational aspects. For the image graph \(G\) defined on the image domain, the adjacency relation \(\Gamma\) between the pixels is typically obtained by a structured adjacency relation, such as 4- or 8-adjacency in a grid form, and the set of vertices \(V = \{v_1, v_2, \ldots, v_N\}\) represents the \(N\) pixels of the image.

The set of functions associated with each vertex is denoted by \(f : V \subset \mathbb{Z}^2 \to \mathbb{R}\). Common functions in \(f\) include low-level descriptors, variations in the color space or in the gray-scale magnitudes. The latter being notably important as the most common source to calculate the weighting function.

For the set of weighting functions \(\mathcal{F}(E)\) of \(G(V, F)\), the best candidates are the ones that could characterize similarities, and for such, the Euclidean distance is the most common, defined in \(E\) as \(\mathcal{F}_{\text{euclidean}}(u,v) = \sqrt{(f(u) − f(v))^2}\). The edge weights may represent the local variation around a vertex, and serve as an image gradient operator bounded by the adjacency relation. The interaction between the image data and the preserved characteristics on the edge-weighted graph is conditioned by the topology choices, such as the adjacency relation and the properties of the weighting function.

III. \textbf{Image Gradients from Random Forest Predictions}

Weighting edges as an image gradient operator, like many gradient operators, acts as a transformation filter on the image creating a transformed space by changing the contrast of the original image and spreading the intensity levels.

\textbf{Definition 4.} The graph-based gradient operator for edge-weighted graph \((G, F)\) at vertex \(u\) could be defined as:

\[
\nabla_{f} f(u) = (\partial_{v_1} f(u), \ldots, \partial_{v_i} f(u)), \forall v_i \in \Gamma(u) \quad (1)
\]

where \(\partial_{v_i} f(u)\) is the edge derivative of \(f\) at a vertex \(u \in V\) along the edge \(e = (u, v) \in E:\)

\[
\partial_e f(u) = \left. \frac{\partial f}{\partial e} \right|_u = \mathcal{F}(u, v) \quad (2)
\]
As in the case of many spatial filters based on local differences, the graph-based gradient operator defined for $F_{euc}$ is subject to respond strongly to noise. We expect that the attribute selection on the RF trees can mitigate this aspect and also any eventual poor topology choice while reinforcing desirable characteristics.

A. Random Forest as regularizers

A RF is a non parametric machine learning method that can be used both for classification and regression. The RF predictor consists of $M$ randomized trees. The core of RF algorithm, as proposed by [14], is the randomization of sampled data distributed to supervise the training of independent decision trees, and the aggregation of the results for the final prediction.

In each internal node $k$ of a tree in the forest, there is a split function $h(x, \theta_k)$ for a query point $x$ with parameters $\theta_k$. During training, the parameters $\theta_k$ are learned, usually by maximizing the information gain $I_k$ to split the data samples covered by $k$ into two subsets with the maximum proportion of instances belonging to the same label. On the test phase, an unseen set of data is applied to $h$ at each split node and the result of the test determines the path the data will perform until it reaches a terminal node with the label prediction.

RF are empirically successful in suppressing noise, although the statistical and mathematical properties of the procedure are still obscure [31]. Some consensus is that the randomness in RF performs as an implicit regularization process, behaving as interpolating classifiers that encourage large consistent regions and reduce the effect of noise [15].

B. Applying Random Forest to edge-weighted graphs

To use the RF implicit regularization process with the local variation representation of the edge-weighted graph, we propose to use the information on the graph edges and vertices to represent the graph on the framework. We represent the regular input of the RF as:

\[
D_n = ((X_1, Y_1), \ldots, (X_n, Y_n)) \text{ with } n \leq |V| \text{ samples of vertices of the edge-weighted graph } (G, F), \text{ each represented as a vector } X \in \mathbb{R}^p \text{ and label } Y.
\]

In our application, edge-weighted graphs are created from images, each vertex thus corresponds to a pixel. $X$ is a vector with dimension $p = |G_{att}|$ for $G_{att}$ representing a set of selected attributes of the vertices of $(G, F)$. In this work, the selected attributes belong to two categories:

- **vertex attributes** ($X_V$), belonging to the set of vertices functions $f$. Each $v \in V$ is mapped into a set of low-level color descriptors proposed in [32]: from RGB colors of an image pixel, 3 color channels in CIE-LUV color space, 2 normalized gradient magnitude channels and 8 gradient orientation channels are calculated;
- **edge weights** ($X_F$), for a given vertex $v$ and all its adjacent vertices, it is represented by the set of edge weights between them. Therefore $X_F = \{F_{euc}(u, v) \mid \forall u \in \Gamma(v)\}$. In this work, we go further the immediate neighbours of $v$ and include also the neighbours in the adjacency of the immediate neighbours. Therefore,

\[
X_F = \{F_{euc}(u, v), F_{euc}(w, u)\}
\]

for all $u \in \Gamma(v)$ and $\forall w \in \Gamma(u)$.

We thus end with $X = G_{att} = \{X_V, X_F\}$, by concatenating the two sets of selected attributes.

C. Using Random forest to compute gradients

In order to obtain gradients, RF is trained on an edge detection task. Because each vertex of the graph is created from a pixel of the image with a unique label on the ground-truth, all the $n$ entries $X$ have a unique discrete label $Y \in \{0, 1\}$ on the task of edge detection.

To obtain the image gradient on this framework, all vertices of a test graph are subjected to the estimations of the RF trained on the input data sample $D_n$. At inference, instead of
taking the label prediction, we use the RF as a regression estimator, producing an estimate $m_{M,n}$ from the $M$ trees composing the RF.

Following the notations presented in [33], for a query point $X$, the $j$-th tree in $M$ gives the estimate $m : X \rightarrow \mathbb{R}$:

$$m(X; \Theta_j, D) = \sum_{i \in D^*(\Theta_j)} \frac{1_{X_i \in A(X; \Theta_j, D)} Y_i}{N(X; \Theta_j, D)},$$  \hspace{1cm} (3)

where:
- $\Theta_j$ is the random variable for re-sampling and selecting the split directions;
- $D^*(\Theta_j)$ is the set of sampled data points for the tree construction;
- $A(X; \Theta_j, D)$ is the tree terminal node containing $X$;
- $N(X; \Theta_j, D)$ is the number of data points falling in this terminal node.

The final estimated value, $m_{M,n}(X)$, obtained by averaging the $M$ estimates $m_i(X; \Theta_j, D)$ is thus taken as a confidence value that a certain vertex $X$ indeed represents an edge. These estimated values for vertices are mapped back to the image coordinates as an intensity value to create the image gradient.

An illustration of the proposed framework is provided in Fig. 3 with simplified examples.

IV. EXPERIMENTS

We evaluate the proposed GIG both qualitatively and quantitatively, and compared it to the widely used gradients from SED, Sobel and Laplacian methods. We perform the evaluation through a hierarchical image segmentation task performed on the Berkeley Segmentation Dataset and Benchmark (BSDS500) [34], consisting of 500 RGB images (200 train, 100 validation and 200 test), each with human labeled segmentations and boundaries. For the qualitative analysis, we provide some resulting gradients and segmentation images and discuss the impact of the different gradients in hierarchical image segmentation. For the quantitative analysis, we use two image partition interpretation metrics on the BSDS dataset.

A. Experimental setup

From the input images, the graphs are created as undirected graphs with structured 8-adjacency relation. For the vectorial representation of the vertex attributes, we explore the low-level descriptors discussed in Sec. III, therefore $|X_V| = 13$. For the weighted edges attributes, we use $F(E) = \{F_{euc}\}$ to weight the 8 direct adjacent vertices of a certain vertex and each of their subsequent neighbors, therefore $|X_E| = 64$. Not all values are unique in this representation as the vertices on the path share some neighbours. For the vertices that do not have all neighbors considered on the path, such as the ones created from the pixels on the border of the image, we added a padding value to the missing neighbors.

The training set $D_n$ is composed of $n = 7,720,050$ vertices, corresponding to a balanced sub-sample of 25% of all vertices from the 200 training images, each labeled with the pixel ground-truth label for the boundaries. We explored training parameters on the validation set. The RF is trained with $M = 150$. All the 77 input features of $X$ are considered at each split during training (no random feature selection). The quantitative results are presented on the test set, the gradient images being created by the estimated values predicted by the trained RF.

In this work, we do not propose a segmentation approach, we present instead a strategy to extract image characteristics that facilitate interpretation and further analysis. Therefore, in order to evaluate the quality of the gradients, we propose to apply the compared methods on the watershed hierarchies [29].
This intuitive algorithm maps image gradients to segmentation and its performance depends on the gradient input, making it the ideal candidate to evaluate our approach. For the segmentation step, we have used a hierarchical segmentation: the watershed by area. It is worth to mention that, thanks to this hierarchical structure, it is very easy to compute segmentation with an exact number of regions, for instance, from 2 to 5000 regions. This allows us to analyze a small number of regions closer to the ground-truth, as well as a medium number of regions for region consistency, and very large number of regions (1000 and 5000), in which results are similar to a super-pixel segmentation method.

B. Qualitative analysis

We present in Fig. 4 the gradient images obtained from the compared methods for the input images on first row. As SED is a method for edge detection, it generally produces gradient images with soft edges close to the ground-truth boundaries, which guaranties its success on the edge detection task. Nonetheless, other aspects present on the input image, such as textures and small details, are wildly ignored. Sobel present more details, without big distinction (in terms of magnitude of values) for components other than the main object. Laplace in turn is permeated by noise on the object and background (see Fig. 4, 1rst row, for computed gradients). It is important to consider that Sobel and Laplacian depend on parameters definition, such as kernel size. For Sobel we represent the gradient magnitude with the \( L_2 \) norm and kernel of size of 3 calculated from the gray-scale image. For the Laplacian we represent the zero-crossing with threshold at 0.04 of maximum value. For the proposed GIG, we have a balance between highlighted strong edges, and different textures and uniform regions presented with homogeneous values distinguishing them.

In Fig. 5, we present more examples of segmented images from GIG, SED and Sobel gradients, illustrating some variation on the number of regions, including the super-pixel effect with large number of regions (3rd column). On the 1st column, we present a successful instance of the proposed method in which the presence of strong borders and large uniform regions on the input image, captured by the GIG gradient, created a better segmentation. Using SED, the fuzzy edges limit the delineation of the main object, while the noise in Sobel prevents its detection. An observed limitation of the proposed method is presented on the 2nd column: when the input image presents objects with patterns of high-contrast, such as zebras and tigers, the detail of the GIG gradient works against the distinction of the object. This is also partially
observed on Sobel, but not with the SED gradient, in which the pattern details are softly represented inside the object. On the super-segmented images in the 3rd column, once again the soft edges on SED works does not produce a good segmentation, while large regions with Sobel are indistinct, producing a lot of very small regions on the main object with little to none on large parts of the duck, the water and the shadow.

In general, the observed results of the proposed framework are very descriptive image gradients in which: (i) object boundaries are highlighted (including the one from very small components); (ii) image textures are firmly represented with different simplified patterns; and (iii) large regions are uniform with distinction of shadow regions. Limitations are perceived for images with objects with patterns of high-contrast.

C. Quantitative analysis

In terms of training time, in the standard discrete label and the graph attribute selection on GIG, we trained all the 150 trees on the RF in less than three minutes, while for SED, in the same CPU, each tree (of eight trees for the presented results) takes approximately four hours. The inference for all the compared methods are similar: a fraction of a second for each image.

For the quantitative metrics, we use two types of image partition interpretation measures, as categorized and defined in [35]: (i) Precision-recall for regions, using a pixel-wise comparison for an overall performance in terms of F-measure; and (ii) Probabilistic Rand Index (PRI), a pixel-wise measure that takes into account the multiple ground-truths presented for each image on the BSDS500 dataset.

Results for both metrics are presented in Table I. The F-measure results for regions are presented in terms of the optimal dataset scale (ODS), optimal image scale (OIS) and average precision (AP) through all scales. Perfect score=1.

![Fig. 6](image)

**V. CONCLUSIONS**

In this work, we explored the outcomes of a novel framework operating on an edge-weighted graph coupled with Random Forest estimates to create very descriptive image gradients. We also outlined the challenges of machine learning on graphs and proposed a strategy to create a systematic input for the random forest framework from the key attributes in an edge-weighted image graph. A qualitative analysis of the produced gradients showed that the proposed method produces gradients where boundaries are highlighted, including very small components, image textures, large uniform regions and distinction of shadow regions. The proposed method and other popular gradient methods were used as input for the watershed hierarchies segmentation method that relies on a good image gradient as input. A quantitative analysis of the produced segmentations confirmed the visual results and demonstrated that the proposed gradient is a better candidate to create image gradients for segmentation. Finally, the proposed approach on the structured input proved to be not only descriptive, but also considerably faster to train than the structured output. For further works, we will study the behaviour of GIG with different hierarchical methods, moreover, we will apply our gradient to region adjacency graphs.

**TABLE I**

<table>
<thead>
<tr>
<th>Gradient</th>
<th>F-measure for regions</th>
<th>PRI</th>
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