SARS-CoV-2: enhancement and segmentation of high-resolution microscopy images—Part I

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Abstract
Possibly, and due to poor eating habits and unhealthy lifestyle, many viruses are transmitted to human people. Such is the case, of the novel coronavirus SARS-Cov-2, which has expanded of exponential way, practically, to whole world population. For this reason, the enhancement of real microscopic images of this coronavirus is of great importance. Of this way, one can highlight the S-spikes and visualizing those areas that show a high density, which are related to active zones of viral germination and major spread of the virus. The SARS-Cov-2 images were captured from nasopharyngeal samples of Cuban symptomatic individuals (RT-PCR positives for SARS-CoV-2) and processed via scanning electron microscopy. However, many times these microscopic images present some blurring problems, and the S-spikes do not look well defined. Therefore, the aim of this work is to propose new computational methods to carry out enhancement and segmentation of SARS-Cov-2 high-resolution microscopic images. The proposed strategy obtained very satisfactory results, and we validated its performance, together with specialist physicians, on a set of 1005 images. Due to the importance of the obtained results, this first work will be addressed to the application of the proposed algorithm. A second paper will deeply analyze the theory related to these algorithms.

Keywords Image enhancement · Segmentation · Algorithms · Coronavirus · SARS-CoV-2 · Microscopy

1 Introduction

Probably, due to poor eating habits (for instance; ingestion of semi-processed or not very well-cooked foods) and unhealthy lifestyle (few hand hygiene, unnecessary crowding of people, and no use of masks, among others), many types of viruses are transmitted to humans. Thus, this novel virus, named coronavirus due to visual appearance—under electron microscopy—similar to a crown [1], is the cause of an infectious disease by severe acute respiratory syndrome (SARS-Cov-2) [2], and named by the World Health Organization (WHO): COVID-19. This disease has expanded of exponential way, practically, to the whole world population, and already today, it is considered a pandemic.

The fight against COVID-19 has motivated many scientific institutions and researchers from all the specials in the world to seek effective methods and techniques to help end this pandemic. In this direction, the computer vision community has not been left behind, and many articles have been published, mainly for the improvement of computed tomography (CT) images using deep learning [3–7].

In this sense, it draws attention the small number of published papers for the improvement of microscopic images of the novel coronavirus SARS-Cov-2, when is known the blurring problems inherent in these types of images, which often makes it difficult to analyze correctly them. Therefore, the aim of this work is to propose new computational methods to carry out enhancement and segmentation of SARS-Cov-2 high-resolution microscopic images, which will allow the S-spikes to be highlighted and visualizing those areas that show a high density, which are related to active zones of viral germination and major spread of the virus.
It is known that the microscopy images are widely used for the diagnosis of various diseases, such as infections and tumors. These facilitate basic biomedical research that has been continuously generating new knowledges into the causes of human diseases, and for that reason, they are of great importance in improving of quality of health. However, the obtaining high-quality microscopic images are not always a trivial process, even in the best microscopes, due to physical phenomena that originate between the glass coverslip, light, and microscope optics. Some of these phenomena are scattering, out-of-focus, imperceptible vibrations, voltage disturbances, among others, which causes certain blurring in the microscopic images. Examples of some of them can be seen in [8–10].

With the rapid development of computer vision, where its application has covered practically all fields of modern life, and basically, digital image processing which has been widely utilized in the medical field, including structure segmentation and image enhancement, has provided great support for diagnosis and better medical prognosis [11, 12].

The importance of this study is that the new developed algorithms will be applied to the enhancement of microscopic images of the novel coronavirus SARS-Cov-2, which contributes, due to very little information appears in the literature on this issue, to the novelty of this work.

This first research will be addressed to application of the proposed algorithm, which it obtained very satisfactory results, and we validated its performance, together with specialist physicians, on a set of 1015 images. A second paper will deeply analyze the theory related to these algorithms, and we will carry out a quantitative evaluation of them. Here, by space problems are not possible.

The rest of the paper is organized as follows: In Sect. 2, the materials and methods are given. Section 3 slightly outlines some theoretical and algorithmic aspects. A second paper will deepen into these issues. Section 4 contains the obtained results and discussion. We describe our conclusions in Sect. 5.

2 Materials and methods

2.1 Medical methodology

We captured the samples from nasopharyngeal swabs collected from Cuban individuals with COVID-19 symptomatic and RT-PCR positive for SARS-CoV-2, and we treated the same (samples) with an aldehyde solution and processed by scanning electron microscopy [13]. Finally, we completed a database of more than 1015 microscopic images.

The structure identification of the SARS-CoV-2 virus is essential toward understanding its historical resemblance to other coronaviruses (mode of infection, mechanism of entry at tissue site of infection, and the replication process in the infected cells) [14]. Particularly, high-resolution microscopic studies are essential in order to identify the etiological agent of several outbreaks. In the case of SARS-CoV-2, the Scanning Electron Microscopy (SEM) images provide fundamental data of the structural aspects of the virus and can be a guiding point in therapeutic development [13].

The specialists inactivated clinical specimens for 12 h in a solution of 25% formaldehyde and 5% glutaraldehyde before microscopy study. Finally, we processed the inactivated samples by SEM. The coverslips were sputter-coated with gold (thickness of 10 nm) and viewed under the MIRA3-TCAN scanning electron microscope at 10 kV [13].

2.2 Characteristics of studied images

The dataset has more than 1015 images, but by space problems, we only give eight examples. Figure 1 shows typical coronavirus images. Figure 2 shows other examples.

In Figs. 1 and 2, one can observe several notable characteristics of these microscopic images, which are common to typical microscopic images we encounter in the study of the novel coronavirus. For example, in Fig. 3, one can see a horizontal profile through images of Figs. 1b and 2d. This is a plot of the pixel intensities along a single row, where one can note:

1. The intensities are slightly clearer within the virus than in the local surrounding background (Observe in Fig. 3, the “y” axis).
2. One can observe high local variation of intensities both, within the coronavirus and within the background (see Fig. 3).
3. It is possible to see (Figs. 1 and 2) high variability of coronavirus in both, size and shape.
4. The contrast of intensities between the coronavirus and the background is low. This is evident from the intensities of histogram (see Fig. 4). This histogram is unimodal, reflecting the low separation in average intensities of the virus and background.

Due to the acquisition protocol and to the already mentioned physical phenomena, a lot of noise corrupts the microscopic images. One can see the ripples in the histograms and profiles (see Figs. 3 and 4), and the blurring that exists in all these images. Across the entire image, changes in intensity levels cannot reliably distinguish S-spikes, due to possible non-uniformity of the average background intensity and the low contrast between virus and background. However, within a region of microscopic images, changes in intensity levels can effectively distinguish S-spikes, since locally a virus is clearer than its surrounding background.
In other words, one can notice that coronavirus areas (the S-spikes) are hyper-dense regions.

The previous paragraph points out the need of working in the region of interest, that is, locally. This strategy ensures that one taken into account the correlation among pixels of the S-spikes, which is of great importance for getting good results, and in addition, it helps notably in the design of the algorithms [15].

In Fig. 5, we represent a 3D projection of a region of interest (ROI) from Figs. 1b and 2d. One can see that the crests are smooth and peaks are not well distinguished, which it indicates the blurring existing in these microscopic images.

The characterization of the study images is of great importance, and a basic principle to have in mind. Many times, this determines the strategy to follow [16].

3 Some theoretical and algorithmic aspects

In this section, due to the importance of the obtained results, we only outline lightly some theoretical and algorithmic aspects. A second paper will deeply analyze the theory related to the proposed algorithm.

3.1 Some definitions

**Definition 1** (Regional maximum) A regional maximum at altitude $h$ of grayscale image $I$ is a connected component $C$ of $T_h(I)$ such that $C \cap T_{h+1}(f) = \emptyset$.

We consider the successive thresholds $T_h(I)$ of $I$, for $h = 0$ to $L-1$ [16],

$$T_h(I) = \{ p \in D_I / I(p) \leq h \}$$

They constitute the threshold decomposition of $I$, where these sets satisfy the following inclusion relationship:

$$T_h(I) \subseteq T_{h-1}(I) \quad \forall \quad h \in [0, L-1],$$

where $L$ is the number of gray levels.

In Fig. 6, one can observe a representation of regional maxima in a grayscale image. In this case, we considered the regional maximum as the S-spikes, where one can see that are justly the hyper-dense areas (see Figs. 1 and 2).

**Definition 2** (Grayscale reconstruction) The grayscale reconstruction $\rho_f(J)$ of $I$ from $J$ obtained by iterating...
grayscale dilations of $J$ “under” $I$ until stability is reached, that is,

$$\rho_t(J) = \bigvee_{n \geq 1} \delta^{(n)}_t(J)$$

(3)

**Definition 3** The h-dome image $D_h(I)$ of the h-domes of a grayscale image $I$ given by

$$D_h(I) = I - \rho_t(I - h)$$

(4)

The $h$-dome transformation extracts light structures without involving any size or shape criterion. The only parameter ($h$) related to the height of these structures. In the case of segmentation of the S-spikes of novel coronavirus, this parameter played a vital role.
All researches carried out with the proposed algorithm worked locally. This was the principal characteristic of designed algorithms for segmenting and enhancing the microphotographs of the novel coronavirus.

In effect, when locally working the spatial information is taken in consideration (and implicitly the correlation among the pixels), and one obtains a major stability in performance of strategies. One of the problems that have the classical thresholding operators, it is to assume that the pixels are statistically independent [15].

A deep explanation of the theory of the Algorithm No.1 will appear in a second paper. Here, we will give a brief review of it.

The Gaussian filter was used to diminishing the noise in the original microscopic images. However, the MSHi algorithm has a performance more precisely [16], but it consumes more computational time. In the case of the Gaussian filter, the best performance was obtained using \( \sigma = 1.5 \). The used window size was \( 3 \times 3 \). A larger dimensional window caused a loss of information in the microphotographs.

The value of the \( h \) parameter determines the processing characteristics. For a small value of \( h \), the \textit{S-spikes} were improved (more contrasted) in relation to the original image. While for a large value of \( h \), the \textit{S-spikes} were isolated (segmented), where the \textit{S-spikes} appeared clear on a black background. The importance of improving or segmenting the \textit{S-spikes} is because the areas that showed an \textit{S-spike} high density are related to an active zone of viral germination [13].

### 4 Experimental results. Discussion

In computer vision, image enhancement and particularly segmentation are recognized as one of the most important steps in any image analysis system.

In Fig. 7, we show a first example of the obtained result of applying the Algorithm No.1 for different values of \( h \) parameter on the image of Fig. 1a.

One can see (in Fig. 7) that all the processed images were enhanced; that is, contrast increased, and the \textit{S-spikes} are evident and virion particles show the urchin-shape. Note that for small values of \( h \), we obtained a better-contrasted image. However, as we increased the \( h \) value, the \textit{S-spikes} begin to segment, where one can see white contrasted areas on a black background. It is possible to observe the edges more defined. Of this way, we can to carry out an automatic analysis of the density of \textit{S-spikes} (number of \textit{S-spikes} / area unit); that is, automatic count of white points on a black background area.
When analyzing the profiles of the processed images there are remarkable differences with those of Fig. 3. In effect, one can observe in Fig. 8 that the peaks of the profiles, for different $h$ values, begin to flattening. In other words, peaks with parallel lines to the "x" axis are indicative of equal intensity levels (segmented zone). In addition, distanced peaks show the isolation of the S-spikes in the different areas where the coronavirus is. The intensity of background reached zero value.

Moreover, on Fig. 7, one can have the following question: Which image to choose as the best-processed image? The answer to this question is not direct, because it depends on the aim of observer or from the application itself; that is, segmentation is heavily dependent on the application [18]. A theoretical analysis of the influence of $h$ parameter will appear in an upcoming publication. Figure 9 shows the obtained result of processing the image of Fig. 1b for three values of $h$ parameter.

Note that the algorithm has very similar behavior to the result of Fig. 7, which shows the stability of this strategy.
In Fig. 10a, we can better appreciate the blurring in the original image, where the S-spikes seem to be out-of-focus. Note in the 3D representation that the crests are flattened (see image 10 (b)). However, in Fig. 10c, one observes the S-spikes and the crown shape is evident in some viral particles, resembling the typical morphology of the coronaviruses [13]. The image has better contrast and edges more defined. The 3D representation of this processed image shows acute, separated crests and high peaks (see Fig. 10d).

Other processing examples carried out with the Algorithm No. 1, for two $h$ parameter values, appear in Fig. 11. One can note the stability of this strategy for different images. See that the enhanced and segmented images the S-spikes are easy to distinguish. However, in the original microscopic images, one cannot say the same. Here, the S-spikes appear very blurred. We carried out more than 750 experiments on different images and we obtained the same results.

### 4.1 Comparisons with other methods

In this section, we will compare our strategy with two recognized classical techniques. Namely, a histogram equalization (contrast improvement method) and the Otsu’s method (segmentation). Such comparison is necessarily incomplete, given the information volume existing in the literature. Instead, we selected two recognized classical techniques, which are suitable for the kind of used images in our study. In Fig. 12, we show the obtained results with our algorithm and the histogram equalization (HE) and Otsu methods.

In a visual comparison with specialist physicians (more than 10), one can notice that the histogram equalization was not able to discriminate the S-spikes and produced an over-saturation. Very similar happened with the Otsu’s method. However, our algorithm accentuated and isolated the S-spikes. Here, we will show a qualitative comparison carried out by ten experts (see Table 1). We utilized as qualitative evaluation criteria: good (g), regular (R), and poor (P).

Table 1 reflects that our Algorithm had better performance than the classical methods in the enhancement and sharpens of the S-spikes. One can see that from a sample of 750 images, the experts rated of “Good” to 738 images enhanced with our algorithm.

Other examples of comparisons with other images appear in Figs. 13 and 14.
We presented a strategy for enhancement and segmentation of microscopic images of the novel coronavirus. The basic idea was to take advantage of characteristics of microscopic images of the novel coronavirus (hyper-dense zones) for the creation, with a minimum of parameters, of a new algorithm. The proposed strategy enhanced and segmented the S-spikes. In this sense, we proposed a variant of the h-dome transformation that consisted in subtracting an $h$ parameter to separate hyper-dense areas and locally working with high spatial frequencies. The proposed algorithm showed accuracy and had good performance and stability in the enhancement and isolation of S-spikes of the SARS-CoV-2 coronavirus. Finally, we compared our algorithm with two well-established classical methods and proved that in all cases, it was superior in the processing of microscopic images. In that sense, we carried out a qualitative comparison in a database of 750 images, where ten experts evaluated 738 results obtained by our algorithms as “Good.” A next publication will carry out a theoretical analysis of these results.

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