

SARS-CoV-2: High-Resolution Microscopy Study in Human Nasopharyngeal Samples.

Brian Mondeja (✉ brian.mr@cea.cu)

Center for Advanced Studies of Cuba <https://orcid.org/0000-0001-6196-3570>

Odalys Valdes

Institute of Tropical Medicine Pedro Kouri

Sonia Resik

Institute of Tropical Medicine Pedro Kouri

Ananayla Vizcaino

Center for Advanced Studies

Emilio Acosta

Center for Advance Studies

Adelmo Montalván

Center for Advance Studies

Amira Paez

Center of Advance Studies

Mayra Mune

Institute of Tropical Medicine Pedro Kouri

Roberto Rodríguez

Institute of Cybernetic, Mathematic and Physic

Juan Valdés

Center for Advanced Studies

Guelsys Gonzalez

Institute of Tropical Medicine Pedro Kouri

Daisy Sanchez

Center for Advanced Studies

Viviana Falcón

Center of Genetic Engineer and Biothecnology

Yorexis González

Center for Advanced Studies

Vivian Kourí

Institute of Tropical Medicine Pedro Kouri

The IPK Virology Research Group

Instutute of Tropical Medicine Pedro Kourí

Angelina Díaz

Center for Advanced Studies

Maria Guzmán

Institute of Tropical Medicine Pedro Kouri

Research

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Abstract

Background

The novel coronavirus SARS-CoV-2 is the etiological agent of COVID-19. This virus has become one of the most dangerous in recent times with a very high rate of transmission. At present, several publications show the typical crown-shape of the novel coronavirus grown in cell cultures. However, an integral ultramicroscopy study done directly from clinical specimens has not been published.

Methods

Nasopharyngeal swabs were collected from two Cuban individuals, one asymptomatic and RT-PCR negative (negative control) and the other from a COVID-19 symptomatic and RT-PCR positive for SARS CoV-2. Samples were treated with an aldehyde solution and processed by Scanning Electron Microscopy, Confocal Microscopy and, Atomic Force Microscopy. Improvement and segmentation of coronavirus images were performed by mathematic algorithms.

Results

The images of the negative control sample showed the characteristic healthy microvilli morphology at the apical region of the nasal epithelial cells. As expected, they do not display virus-like structures. The images of the positive sample showed characteristic coronavirus-like particles and evident destruction of microvilli. In some regions, virions gemmating through the cell membrane were observed. Microvilli destruction could explain the anosmia reported by some patients. Virus-particles emerging from the cell-surface with a variable size ranging from 80 to 400 nm were observed by scanning electron microscopy (SEM). A confocal study showed viral antigen recognition in the apical cells zone.

Conclusions

The integral microscopy study showed that SARS-CoV-2 has a similar image to SARS-CoV. The application of several high-resolution microscopy techniques to clinical samples can help to answer important questions its replicative cycle and immunopathogenic mechanism of this novel coronavirus, relevant for the development of new treatments against this disease.

Background

Coronaviruses family is a wide-ranging of viruses that infect many animal species including human beings(1). In December 2019, a new coronavirus disease emerged in China. Rapidly, the disease caused by the SARS CoV-2 extended to the whole world with a record in the number of cases and fatalities(2–5). SARS-CoV-2 is 96.2% identical to a bat coronavirus at the whole-genome level and it belongs to the species of SARS-CoV(6). Due to the genetic similarity to SARS-CoV, several characteristics have been assumed, but not clarified. Therefore, more deep investigations with a precise specific morphological description of this novel virus are necessary(5,7).

SARS-CoVs are enveloped viruses with a diameter ranging from 60 to 400 nm. The envelope is constituted by the envelope and the spikes (S-protein) proteins. The ARN is covered by the capsid. The overall structure looks similar to other viruses of the *Coronaviridae* family. The S-protein forms a clover-shaped trimer, with three S1 heads and a trimeric S2 stalk. The envelope-anchored spike protein guides coronavirus entry into host cells. This state of the spike on the mature virions is called “prefusion” (1,8–10). After viral replication and before the viral release, the spikes budding from the Golgi apparatus to cell membrane and the ribonucleic core start to evolved out-side with this S-rich membrane. Not microscopical evidence of this final stage was proved for SARS-CoV-2, but following its similarity with SARS-CoV, this characteristic has been assumed(8).

The structure identification of the SARS-CoV-2 virus is essential towards 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)(11,12). High-resolution microscopic studies are essential in identifying the etiological agent of several outbreaks. This technology is a useful tool in guiding subsequent laboratory and epidemiologic investigations mostly in new epidemic emergencies(13). Particularly, in the case of SARS-CoV-2, the Scanning Electron Microscopy (SEM) images provide fundamental data of the structural aspects of the virus and must be a guiding point in therapeutic development(11).

The application of the Atomic Force Microscopy (AFM) to viruses-associated to human pathologies might have a significant impact on the diagnosis and treatment. Nowadays, scanning probe microscopy is a well-established technique for the rapid visualization of pathogens, including viruses at high resolution(14).

Methods

This study aimed to describe the morphologic characteristics of the SARS-CoV-2 present in human nasopharyngeal specimens using high-resolution microscopy.

Clinical specimens. Two nasopharyngeal swabs tested by Real Time-PCR to SARS CoV-2 were studied. One, with a negative Real-Time -PCR result, was collected from a contact of a confirmed COVID-19 patient. The other belongs to a confirmed COVID-19 patient and resulted in a positive Real -Time-PCR. These samples were received and tested at the National Reference Laboratory of Viral Respiratory Infections of the Institute of Tropical Medicine for virologic diagnostic. Real Time-PCR was performed as previously described(15).

Inactivation of clinical specimens. 200 µL of the clinical specimens were inactivated for 12 hours in a solution of 25% formaldehyde and 5% glutaraldehyde before microscopy study. Inactivated samples were processed at the Center for Advanced Studies of Cuba by Scanning Electron Microscopy, Confocal Microscopy, and Atomic Force Microscopy.

Scanning Electron Microscopy. Ten microliters of the inactivated clinical specimen were placed in a glass-coverslip and dry-in air oven overnight. Then, the coverslips were fixed with 5% glutaraldehyde and dehydrated through a series of increasing concentrations (25–100%) of ethanol. Coverslips were further subjected to critical point drying for 1.5 h and left in a 37 °C oven overnight. Subsequently, the coverslips were sputter-coated with gold (thickness of 10 nm) and viewed under the MIRA3-TESCAN Scanning Electron Microscope (TESCAN, Czech Republic) at 10 kV.

Atomic Force Microscopy. Inactivated clinical samples were processed similarly. Normally, samples for the Atomic Force Microscopy should be subjected to minimal processing to maintain its original condition. However, because of the biohazard of SARS-CoV-2, fixed, and gold-coated samples were used for this technique. The di-Innova Scanning Probe Microscope (Veeco Instruments, Santa Barbara, California) was used in tapping mode. Golden silicon probes NSG30-A, supplied by NT-MDT (Zelenograd, Russia), with a curvature radius of 10 nm and a resonant frequency of 240–440 kHz were used.

Confocal Microscopy. For the immunofluorescence staining, inactivated clinical samples were hydrated for 10 min in PBS and incubated with PBS-Tween (PBS-T) for 20 min. To block non-specific antibody reaction, the best results were obtained by incubating the sections with 0.2% bovine serum albumin (free of IgG) (Sigma Chemical Co. St. Louis, Mo. USA), for 20 min. After two washes in PBS-T, samples were incubated for 1 h at 4 °C with the primary antibody (hyper-immune serum of the COVID-19 -convalescent Cuban patient, dilutions 1:40 in PBS-T). Incubations were followed by washes with PBS-T. The second incubation was accomplished with FITC-conjugated Anti-human Polyvalent Immunoglobulins (IgA- IgG- IgM (dilutions 1:40 in PBS-T, Sigma Co. St. Louis, Mo.USA) for 1 h. After three washes with PBS-T, the sections from all samples were counterstained with propidium iodide (dilution 1:1000, Vector laboratories, Inc. Burlingame CA., USA), followed by extensive washing in PBS-T. Immunostained samples were coverslipped in Vectashield mounting medium (Vector Laboratories, Inc. Burlingame, CA., USA), Fluorescent images were observed on a Confocal laser scanning microscope OLYMPUS FV1000 IX81.

Improvement and segmentation of coronavirus images by mathematics algorithm. The Gauss filter was used to diminishing the noise in the original images. The best performance was obtained using $\sigma = 3$. The used window size was 5×5 . Smaller dimensional windows produced a lot of noise, mainly due to the change of intensity. A larger dimensional window caused a loss of information in the images. A mathematical approach for the morphological grayscale reconstruction was the following:

Let J and I be two grayscale images defined on the same domain D_I , taking their values in the discrete set $\{0, 1, \dots, L-1\}$ and such that $J \leq I$ (i.e., for each pixel $p \in D_I$, $J(p) \leq I(p)$). L is an arbitrary positive integer. In this way, it is useful to introduce the geodesic dilations according to the following definition(16):

The elementary geodesic dilation of grayscale image $J \leq I$ "under" I (J is called the *marker* image and I is the *mask*) is defined as,

where the symbol \wedge stands for the pointwise minimum and $J \oplus B$ is the dilation of J by flat structuring element B . The grayscale geodesic dilation of size $n \geq 0$ is obtained by,

The grayscale reconstruction of I from J is obtained by iterating grayscale dilations of J “under” I until stability is reached [1, 2, 3], that is,

Results

The SEM images showed a general view of SARS-CoV-2 infected human cells. The virus-induced morphological changes demonstrated by the destruction of epithelial cells microvilli in the positive sample (Fig. 1). Uninfected cells show a typical rough surface morphology under the scanning electron microscope and no pseudohyphae are visible either on the cell edge or surfaces (Fig. 1A).

The viral budding through the cell membrane was evident particularly at the edge of these cells. Different stages of the gemmation process were observed (Fig. 1A-B). Some areas show a high density of S-spike gemmating and pseudohyphae projections appear to be related to an active zone of viral gemmation. The active gemmation and release of virions are more evident in the apical membrane of the epithelial cells. The whole viruses are well visualized in the lateral and basal membranes. Figure 2 shows the different stages of the viral budding process and urchin-shape of virions after the improvement and segmentation of coronavirus images by mathematic algorithms.

The viral particle size varied from 86 to 400 nm, but typical virions, observed in the cell-free space, had a size of approximately 80 nm. At a higher magnification, crown-shape is evident in some viral particles, resembling the typical morphology of the coronaviruses (Fig. 1C-1D). S-trimmers showed a typical tips clover-shape in gemmating and mature virions.

Non-lymphocyte-like cells were observed in any of the 50 viewed-fields, suggesting a low immune local inflammatory cell response (data not shown).

The clover-like structures on the virus surface were further confirmed by AFM. The extrusion process of SARS-CoV-2 particles from infect cells is shown in Fig. 3. It can be seen as a group of viruses in gemmation and others about to come out (arrows), which indicates the presence of different populations. This confirms the obtained results through SEM.

Figure 4 shows the results of the immunofluorescence staining by confocal-microscopy. Image A represents the negative control of SARS-CoV-2. Image B, correspond to the COVID-19 confirmed patient. The cells of the patient express the SARS-CoV-2 virus antigens, the antibodies present in the hyperimmune serum recognized the antigen and stuck. The positive signal of green-clouds was observed due to the presence of the viral antigens (white arrows). Clusters of antigens over the cell surface and in the free-cells space detected by confocal microscopy, like to be as the virus aggrupation found by scanning electron and atomic force microscopy.

Discussion

High-resolution SEM and AFM allow a three-dimensional holistic view of the virus and the infected cell surfaces. High-resolution microscopy is gaining popularity in other areas of the life sciences(14,17–19). Most of these studies were on purified macromolecules. However, the AFM has also become a virologic standard in recent years(14).

Images reconstruction provides a very efficient method to extract regional maxima and minima from grayscale SEM-images. Furthermore, the reconstruction technique extends to the determination of maximal structures, which will be call *h-domes* and *h-basins*. The h-dome transformation extracts light structure without involving any size or shape criteria. The only parameter (*h*) is related to the height of these structures. The improved images showed a wild quantity of details of virions and hypothetical cycle life stages of SARS-CoV-2 that were not seen in raw images.

The microvilli destruction observed in the infected epithelial nasal cells could be associated with the anosmia described by some patients. The active gemmation of the virus particles is observed mostly in the apical zone of the cells. It could be associated with the polarized translocation of immature viral particles from the Golgi apparatus as was described for SARS-CoV (6,11,20). Profuse complete-virus release areas were observed in some cells. In the same ones and rear of these active zones, a pseudohypha- formations were recorded with the hugest quantity of S-spike inserted in the membrane. This pseudohypha formation was reported during the multiplication of SARS-CoV in Vero cells(14).

One of the advantages of this finding is the possibility to use SEM for the follow up of long-term PCR-positive asymptomatic individuals. If the virus is actively replicating in these individuals, the presence of cluster spike in the cell membrane, pseudohypha formations, and or viral particle budding, will be noted. The lack of those findings in the presence of serials PCR -positive samples could be explained by the amplification of viral RNA fragments released from infected cells. However, this could be proved by a large-cohort study comparing the SEM and cell-assisted culture in this important group of patients.

The non-homogeneous diameter of virus particles was observed by SEM or AFM, possibly due to the same unsynchronized replication of the virus. However, in the majority of the cells, virion clusters rather than isolated particles were observed. This active replication could explain the high contagious rate of the virus that associated with an efficient mechanism of viral infection and replication conducting to total cell destruction(4). Billions of virus particles are released for a relatively long time, without much effect in the cell structure(2).

The AFM technique has shown to be a good tool for virus study. Details of the viral structure and the confirmation of the SEM finding was archived. The results of this technique in the SARS-CoV-2 study were similar to the morphological analysis of SARS-CoV previously reported(11,14). One explanation of the difference between SEM and AFM results of the virus-spike observation could be the previous sputtering of the sample with gold before AFM performance. For obvious reasons and the high risk of the work with non-inactivated samples carried SARS-CoV-2, the aldehyde-inactivation and gold coat were used. Fortunately, the inactivation methods do not affect the viral and cell morphology for SEM

consequently this procedure can be an option to the microscopy study of clinical samples in unknown or high virulent infectious pathogens.

Confocal studies confirmed the presence of SARS-CoV-2 in clinical specimens and the distribution of the virus into the cell membrane. This is a very useful technique for virus identification using a polyclonal hyperimmune serum of convalescence patients. It could be implemented as a diagnostic assay.

It is important to remark that this high-resolution microscopy study was done directly in positive SARS-CoV-2 clinical samples rather than viruses grown in cell cultures as the previous investigation of the topology of virions made in the past for SARS-CoV(11,14). However, our observation is very close to describing the real cellular pathology and damage of SARS-CoV-2 to the respiratory tissue in the patients. On the other hand, the use of hyperimmune serum of a convalescent patient could be able to introduce some background in the virus detection by confocal microscopy. This could be possible to cross-reactivity with another respiratory virus that could be present in the sample. If they use a different cellular receptor to ACE2 (e.g. influenza A virus), the coinfection is possible, and a false positive signal will be recorded and misunderstood as SARS-CoV-2(8,21–23).

Previous microscopy studies of SARS-CoV-2 were focused on viral infected cell cultures or applied only one of these advanced electron microscopy techniques to the study of clinical specimens(14). Our investigation on the morphology of the novel coronavirus SARS-CoV-2 is based on viruses on clinical samples using high-resolution microscopy (SEM, AFM, Confocal microscopy).

Further investigations should be aimed at the nanostructure of SARS-CoV-2 by High-Resolution SEM and imaging processing to bring us a better understanding of the viral structure in clinical samples. This will enhance the development of new inhibitors and innovative materials to control or destroy in-vivo viruses that could be used in protective clothes and medical devices.

Conclusions

The integral microscopy study showed that SARS-CoV-2 has a similar image to SARS-CoV. The application of several high-resolution microscopy techniques to clinical samples can help to answer important questions its replicative cycle and immunopathogenic mechanism of this novel coronavirus, relevant for the development of new treatments against this disease.

Abbreviations

SEM
scanning electron microscopy; AFM:atomic force microscopy; PBS-T:PBS-Tween- Phosphate Buffer Saline

Declarations

Ethics approval and consent to participate

The study protocol conformed to the ethical guidelines of the 1975 Declaration of Helsinki and was approved by the Ethics Committee of the Institute of Tropical Medicine Pedro Kourí.

Consent to publication

Not applicable.

Availability of data and material

Not applicable.

Competing interests

The authors declare that they have no competing interests.

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Figures

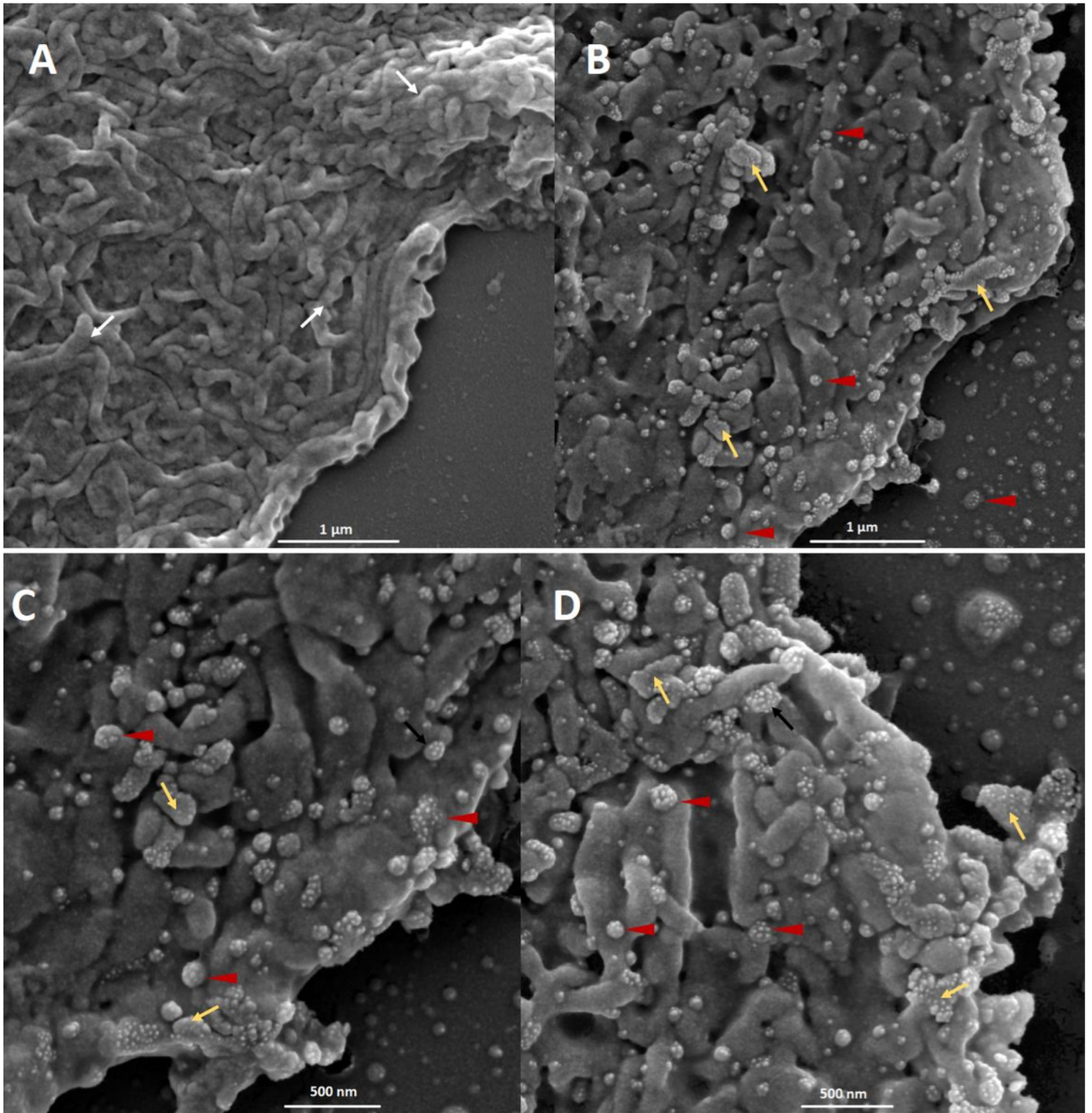


Figure 1

Microphotography performed with an Electronic Scanning Microscope of the apical surface of nasal mucosal epithelial cells. A- Sample from a negative SARS-CoV-2 individual; B, C, and D- sample from a SARS-CoV-2 positive patient. The white arrows point to the cell microvilli; red triangles point to the SARS-CoV-2 virus and yellow arrows point to the pseudohyphae formation. Bars 1μm- 500 nm.

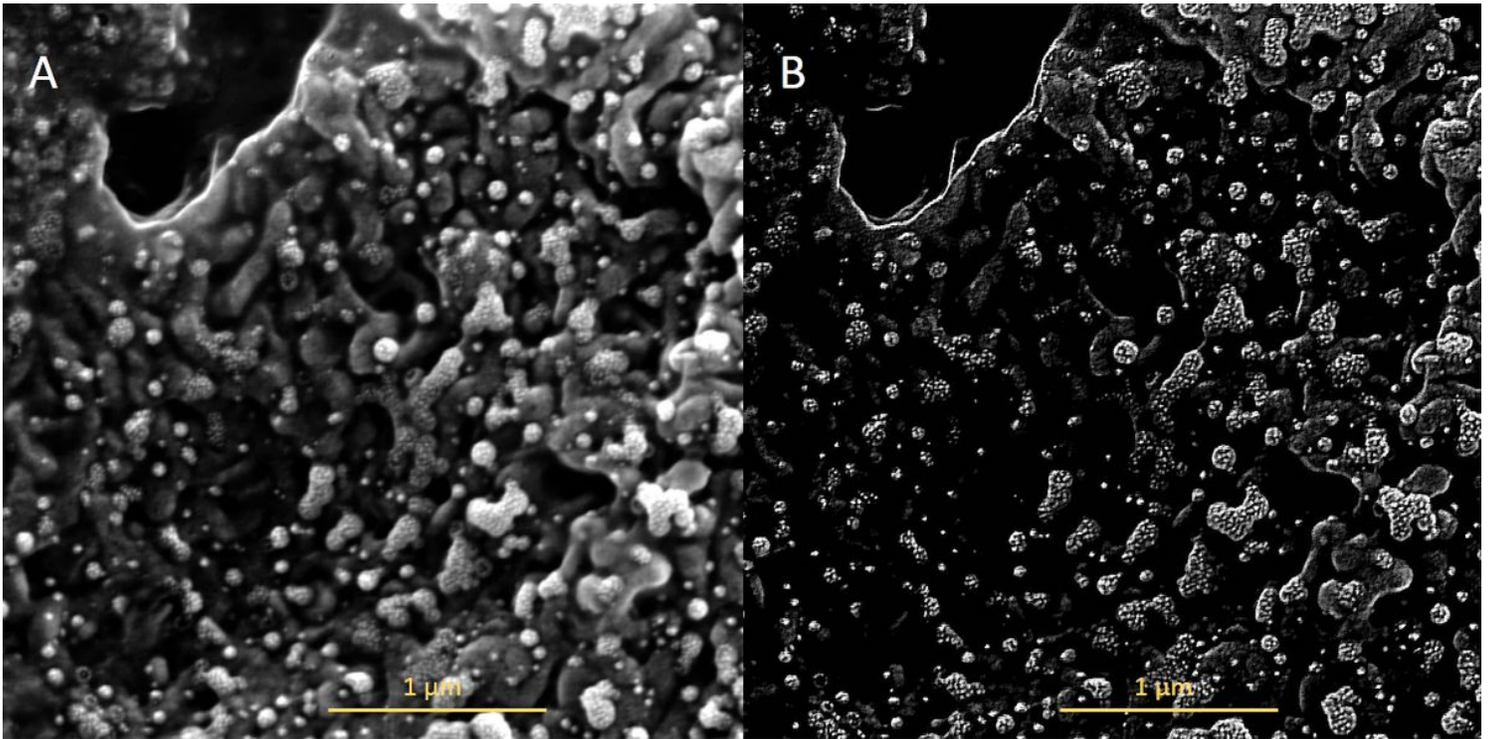


Figure 2

Improvement and segmentation of coronavirus images by mathematics algorithm. Microphotograph performed with an Electronic Scanning Microscope of the apical surface of an epithelial cell of the nasal mucosa. A- Original images; B- Improved images by mathematics algorithm. S-spikes are evident and virion particles shows the urchin-shape. Bar 1µm.

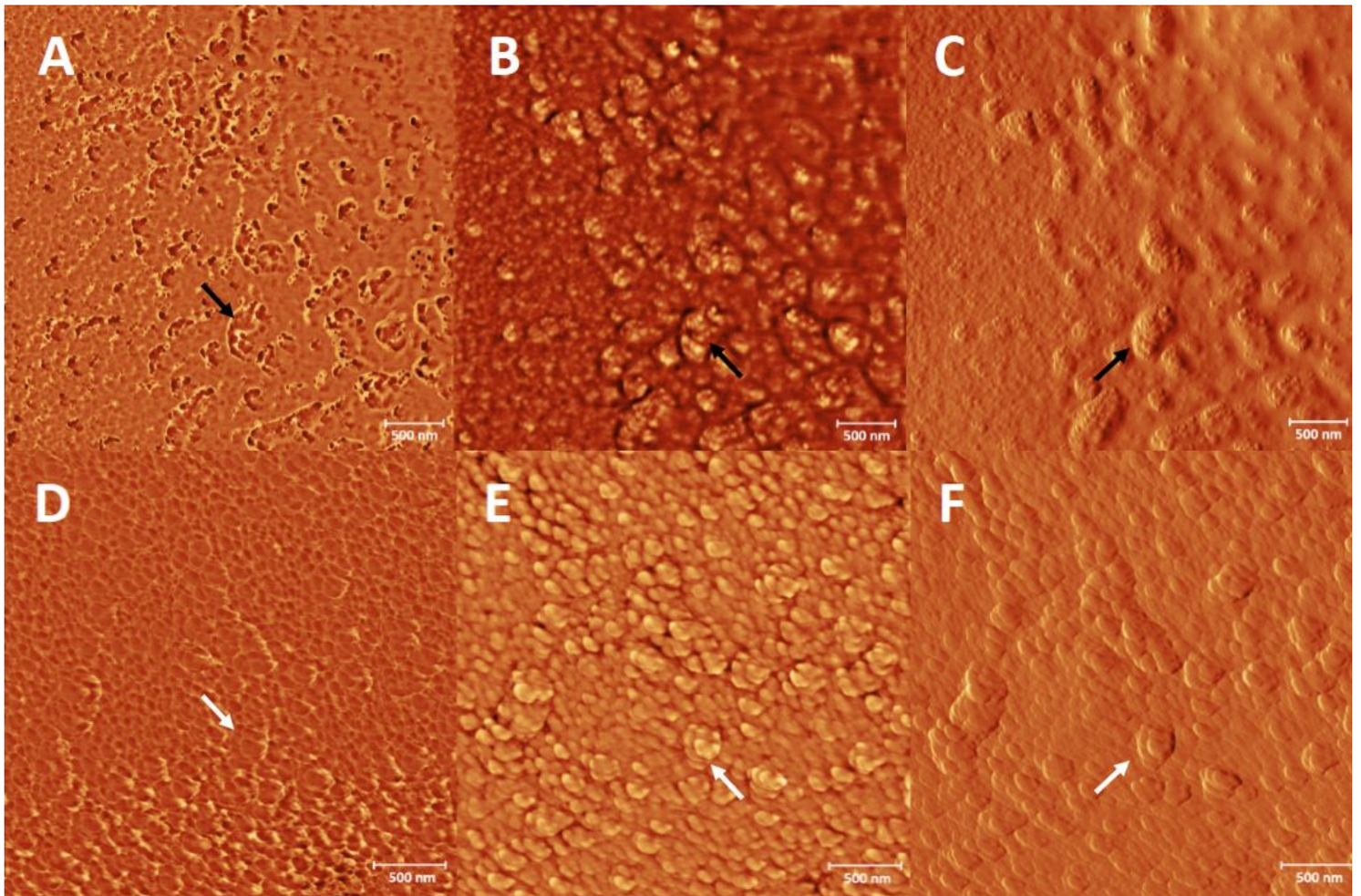


Figure 3

Atomic Force Microscopy images of an epithelial cell of the nasal mucosa of a SARS-CoV-2 infected patient. A and D- phase mode images; B and E- height mode images; C and F- amplitude mode images. The black arrows point to SARS-CoV-2 S-spike in the cell membrane and white arrows point an isolated viral particle show the typical morphology of coronavirus. Bars 500 nm.

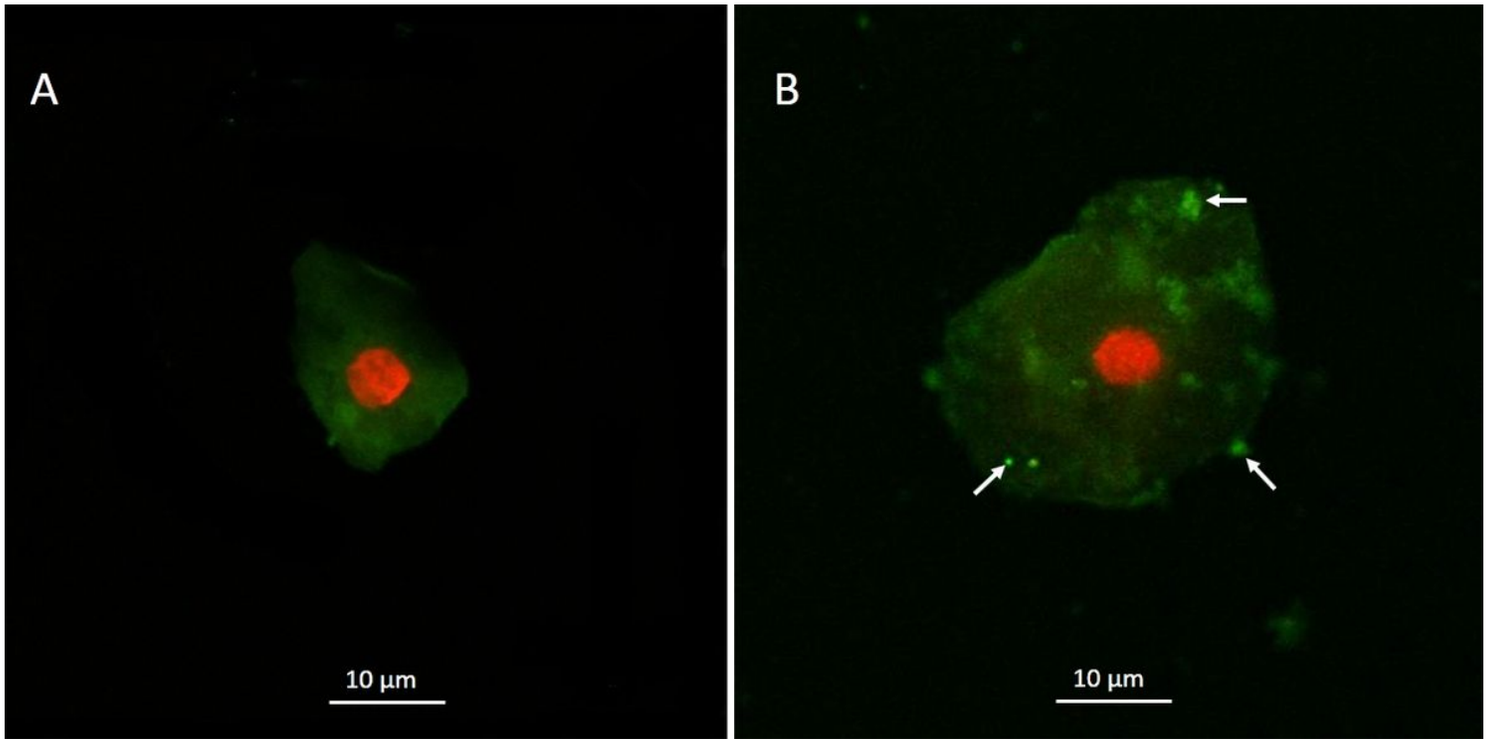


Figure 4

Confocal microphotograph of immunofluorescence staining of SARS-CoV-2 in epithelial cells of the nasal mucosa. A- Sample of PCR-negative individual; B - Sample of PCR-positive. White arrows point the virus antigen cluster coat the cell membrane. Barr 10 µm.

Supplementary Files

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