

# Could Your Smartphone Analyze the Coronavirus on Surfaces?

Project Chipscope brings nano-level microscopy into the field

**Koen Mortelmans**

European researchers are developing a new generation of optical microscopes for mobile phones where LEDs and tiny detectors in the camera can detect very small structures. The ChipScope project is funded by the European Union's Research Programme Horizon 2020.



SMILE platform for structured micro-illumination to enhance resolution of optical microscopes (Source: QubeDot)

Optical microscopy is entering a new world. Future microscopes will be chip-sized and have superresolution capabilities. This means they will be available in everyday life, not just in specialist laboratories. And it might become possible to detect SARS-CoV-2 and other viruses using a smartphone.

“This use of chip-integrated nanoLEDs will revolutionise the way science and applications related to optical microscopy will be done in the future,” says Angel Dieguez, professor at the University of Barcelona, Spain. “It can be used to investigate extremely small structures such as viruses, DNA or living cells, almost anywhere and in real time.”

The resolution of conventional light microscopes – still standard equipment in laboratories – is limited by diffraction: they cannot resolve structural features smaller than about two hundred nanometers. Until now, all technologies for going beyond this limit rely on complex setups, with bulky components and advanced laboratory infrastructure. They are not suitable for mobile research devices used in field or in remote areas.

The University of Barcelona is a partner in ChipScope, a project supported by the EU-programme FET – Future and Emerging Technologies; now part of EIC Pathfinder – which is developing an array of nano light emitting diodes (nanoLEDs) with individual pixel operation [1]. LEDs smaller than the diffraction limit will be used



Source: private

Prof Dr Angel Dieguez  
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as light sources for a microscope, which will be integrated on a chip. “This will lead to extreme miniaturization and simplicity for superresolution microscopy,” Dieguez remarks.

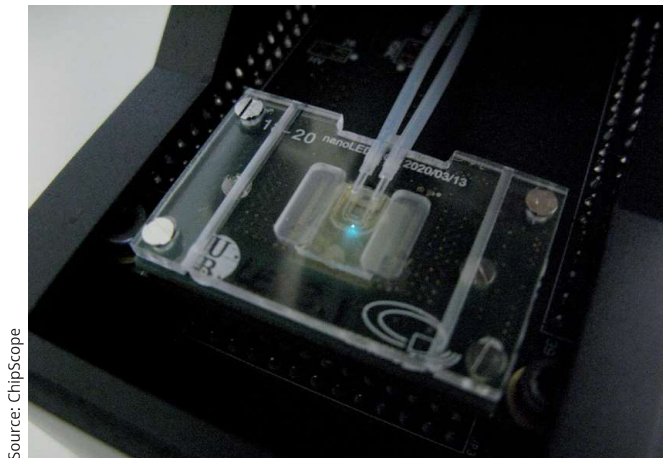
The project's members are universities and research centres as well as small and medium sized enterprises.

They intend to integrate the nanoLEDs into a real-time imaging device. Several European industry leaders in imaging and microscopy are represented on an advisory board.

Dieguez explains: “In conventional optical microscopy, the area that needs to be analyzed is illuminated simultaneously, collecting the light which is scattered from each point of the sample with the human eye or the sensor of a camera as a sensitive detector. Our idea, however, is based on a structured light source with tiny, individually addressable elements. The sample is put on top of this light source. Whenever single emitters are activated, the light propagation depends on the spatial structure of the sample.”

“This is very similar to shadow imaging in the macroscopic world. To obtain an image, the overall amount of light which is transmitted through the sample region is sensed by a detector, activating one light element at a time and thereby scanning across the sample space.”

The nanoLEDs, which are the first with individually addressable pixels in the submicrometer range to become available on a commercial scale, are



Source: ChipScope

Detail of the microfluidics over the nanoLED array with one LED switched on



Source: ChipScope

LED driver chips to control the nanoLED array

being developed at the Braunschweig University of Technology, Germany. “They are based on gallium nitride, a semiconductor commonly used for blue and white LEDs.”

Dieguez’ own team in Barcelona constructed the light detectors: “We now have a prototype device to conduct further experiments. Other projects will be tasked with making a marketable microscope. I don’t have a time scale in mind, but I’m convinced that this will revolutionize the microscope market and the ways to do field research. With chips instead of lenses, the devices will be very mobile extremely cheap. They also can be integrated into existing tabletop systems.”

### Taking the approach even further

Joan Daniel Prades, also a professor at the University of Barcelona, coordinates the project ‘Scalable Structured Micro-Illumination Light Engines’, SMILE, which is using the outcomes of ChipScope. The researchers are developing microLED devices that can bring innovation in microscopy and other research fields such as DNA chip fabrication and optogenetics.

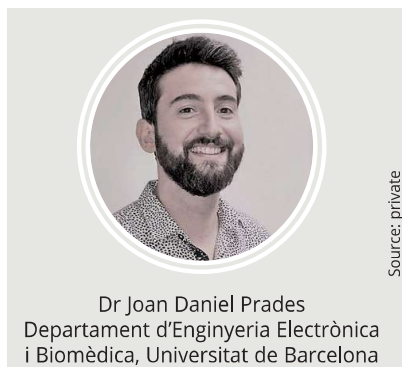
Particularly for optical microscopy, the researchers see possibilities to study the properties of large quantities of substances. “We try to combine holography and chip-based technology with microscopy to allow transparent cells to be visualized with classic cell culture plates. Today, we have a prototype that allows a resolution of 256 pixels per inch, which is available through our start-up partner QubeDot in Braunschweig, Germany. The project’s goal is to obtain 2,000 pixels.”

Another application is the fabrication of DNA biochips. Prades explains:

“DNA chips or DNA microarrays are an important tool for gene expression analysis. These chips consist of a solid surface with a collection of single-stranded DNA fragments [which is the known sequence, ed. note]. The DNA samples that need to be analyzed [the unknown sequence, ed. note] are fluorescently labelled and fixed onto the chip.” The complementary binding between the known and the unknown sequences is then analyzed.

Thanks to the microarray technology, the researchers can get information on thousands of genes simultaneously with just a single DNA chip. “A similar application already exists with lasers. However, these produce DNA samples point by point, whereas DNA chips will perform this task by blocks, which means they work much faster,” Prades continues.

The project also aims at bringing innovation into optogenetics, the bio-



Source: private

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logical technique that uses light to activate or deactivate individual neurons or neuronal networks in the brain of living animals by shining light on them. Up to now, complex fiber-optic systems are used to do this. “The time to generate a pixel and switch to the next one will be less than ten nanoseconds, at a full fram-

erate of one megahertz. So, the planned systems will encompass relevant existing restrictions regarding the maximal frequency of activation, minimum spot size and maximal power output. In addition, conventional microscopy systems are very costly and not readily available, leading to restrictions in the number of experiments to be performed per time, which limits scientific progress.”

Photolithography is another field of application: “Manipulation technologies enabled by large computing capacity allow high replication throughputs as demanded for in industrial micro-patterning,” Prades remarks.

The project is one of the EIC Transition to Innovation Activities, which aim at bringing innovative FET technologies to a level of development for business creation. “An end-user board, with experts from various branches – multinationals, SMEs and interdisciplinary research institutes – will perform tests and validations for quickly increasing markets with the prototypes we are going to make available in five pilot applications,” Prades concludes.

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Text: Koen Mortelmans, freelance journalist, Belgium, on behalf of FETFx ([www.fetfx.eu](http://www.fetfx.eu)).

### Project ChipScope

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[1] “A New Approach to Optical Microscopy”, *PhotonicsViews* 17 (2020) 3, 17, June / July