The waterborne pathogenic bacteria, especially the enteric bacteria of human fecal origin, have become currently a global public health issue. The detection and quantification of drinking water microorganisms have been an essential part of any quality control or water safety management plan interconnected to enteric bacterial pathogens such as Salmonella spp., Shigella spp., Vibrio cholerae or to non-faecal bacterial pathogens such as Pseudomonas aeruginosa. The standard methods of detecting waterborne pathogenic bacteria are time-consuming due to the bacterial growing step in a specific culture media, followed by isolation, microbiological and/or serological identification and in some cases followed by subspecific characterization. This study aimed to develop a faster, powerful, more sensitive and reproducible diagnostic tool to monitor a specific pathogen contamination in drinking water (Pseudomonas aeruginosa) by specific antibody-antigen interactions followed by a computerized technology to identify bacteria as a digital image from water samples. Digital Image processing was carried out using National Instruments Vision Assistant Software. Pseudomonas aeruginosa was specifically detected by immunofluorescence technique with fluorophore tagged antibodies, and then the image formation in fluorescence microscopy was processed by computer vision software functions organized in pipeline-like data analysis processing. Overall, these techniques proved to be a reliable one, time-effective and sensitive for diagnosis and prevention of drinking water quality and waterborne bacterial disease.

**Keywords:** antibody-antigen interaction, bacteria, immunofluorescence, high-tech technology
In the past few years, nucleic-acid-based technology achieved widespread use in the field of pathogen detection, with a particular focus in polymerase chain reaction (PCR) assays developed mainly to detect every relevant bacterial pathogen [19] based on recognition of specific nucleotide (DNA or RNA) sequences of microorganisms. Currently, quantitative PCR (qPCR) is widely available and provides as good as or better sensitivity for the detection of molecular targets than all other currently available detection strategies, including newly developed assays such as: microarrays and prosequencing [20-23]. Furthermore, modern advances in micro- and nanofabrication technology have led to the development of a wide range of nucleic-acid based biosensors that capitalize on the new capabilities of microfluidic technologies and micro total analysis systems in order to reduce reagent and power consumption, enhance analytical performance, and enable portability. Many of these technologies have been extensively studied [17, 18, 24-26], successfully commercialized, and are currently widely used in clinical and research laboratories. In today’s world, the genetically engineered whole cell-based bacteria have usually utilized to generate and amplify the luminescence signal. Bacteria incorporate reporter genes that code for signaling elements that emit bioluminescent, fluorescent, or colorimetric formation in fluorescence microscopy could be processed, digitalized and wireless sent to organizations and/or environmental fields due to their high specificity, reliability and accuracy. There have been many ways to detect the interaction between antibody and bacteria, but among them, the immunofluorescence detection based on the fluorochrome tagged antibody have been one of the most straight one [25]. The bacterial detection image formation in fluorescence microscopy could be processed and be switch to a digital signal which could be wireless sent to health and/or environmental monitoring specialized organization, in order to rapidly prevent any potential bacterial outbreaks. The image processing has been carried out using NI Vision Assistant Software [33] which is based on pipeline type. The fluorescence digital image is composed of three channels in RGB color space (alpha channel not used) and the digital image could be obtained through a pipeline type.

In this paper, we described an immunofluorescent fast, specific and reliable alternative bacterial detection method to the classical and laborious bacterial identification method based on repeatedly bacterial growth on selective medium. A gram-negative bacterium *P. aeruginosa* with a pathogenic potential was specifically immunofluorescent detected and the immunofluorescent image was processed, digitalized and wireless sent to organizations involved in health and environmental monitoring.

**Experimental part**

**Bacterial growth.** The bacterial strains *P. aeruginosa* (ATCC 29212) and *Enterococcus faecalis* (*E. faecalis*) were purchased from ATCC (International Center for the Authentication, Storage and Production of Microorganisms and Cell Lines). The strains were grown on tryptone soy agar (Oxoid, UK) and incubated overnight (O/N) at 37°C. One colony was then transferred in 5 ml acetamide nutrient broth (*P. aeruginosa*) or sodium azide broth (*E. faecalis*) in a shaking incubator (New Brunswick Scientific, Innova 44) at 37°C for 24h. Bacterial growth was monitored by measuring the absorbance at 600 nm using UV-Vis spectrometer (VWR International, USA).

**Bacterial immunofluorescence detection.** The 0.4 OD<sub>600nm</sub> bacterial suspension in 250 µL PBS was incubated O/N at 4°C in presence of monoclonal antibody anti-*P. aeruginosa* tagged with Alexa Fluor 647 (Novus Biologicals, CO, USA) (dilution of 1:100). After the incubation, the sample was centrifuged for 5 minutes at 5000 rpm and 4°C, and the bacterial pellet was washed trice in PBS. The bacterial sample was resuspended in 20 µL mounting medium for further microscopic analysis. The microscopic detection of fluorescent bacteria was performed using LEIKA DMi8 inverted microscope at a 63X magnification. The sample was observed in both fluorescence and transmitted light and the negative control consisted on *P. aeruginosa* incubated in absence of monoclonal antibody anti-*P. aeruginosa* tagged with Alexa Fluor 647 or on *E. faecalis* incubated in presence of antibody anti-*P. aeruginosa*.

**Computerized analysis and measurement of pathogenic bacteria.** Computerized analysis was carried on immunofluorescence digital images obtained by standard microscopic investigations. The computerized analysis consisted of a pipeline processing image based on two stages: i) creating the binary mask (fig. 1) and ii) bacteria identification the in bright field microscopic image.

The image processing flow of the binary mask creation was described below:

Step #0: The fluorescence image was loaded from disk into the NI Vision Assistant Software application.

Step #1: Calibration by associating a numerical value of microns to a numerical value of a distance expressed in pixels [34].

Step #2: The red color plane was extracted from the fluorescence image.

Step #3: The image histogram was displayed onscreen.

Step #4: A threshold was applied on grey level in order to obtain a binary image. That was the most important computation step. Two approaches were used: global threshold and local threshold. Global threshold enforced a unique threshold value over the entire image. Local threshold took into consideration a user defined neighborhood around each pixel in image. That rectangular windows is moved over the entire image.

Local Mean value and variance of a image contained in a neighborhood of MxN size were:

\[
\mu = \frac{1}{M \times N} \sum_{i,j} A(i,j)
\]

\[
\sigma^2 = \frac{1}{M \times N} \sum_{i,j} [A(i,j) - \mu]^2
\]

In addition, Niblack algorithm [35, 36] was used on the threshold calculus, where:

**Fig.1. Pipeline processing for obtaining image mask and blobs’ data analysis**
Threshold \((i,j) = m(i,j) + k \cdot \sigma(i,j)\) where \(k \in [0..1]\)

If \(I(i,j) > \text{Threshold} (i,j)\)
then \((i,j)\) belongs to an object (particle)
else \((i,j)\) belongs to the background.

Step #5: Remove small objects. This morphological operation was used to eliminate some particles using one iteration of 3 x 3 erosions. The remaining particles had exactly the same shape as in the immunofluorescent source image.

Step #6: Particles removing was continued by imposing a minimum area value.

Step #7: The binary image was dilated, then eroded. The effect was compacting the scattered pixels and maintaining the overall shape of particles.

Step #8: Pixels belonging to inner bacterial areas were assigned white color.

At this step, the binary image contained white blobs (bacterial areas), firmly demarcated from background.

Step #9: Every blobs (bacterial areas) got an unique ID. The numerical values of the selected bacterial areas were displayed in a form table (table 2).

The antibody targeted bacteria was also identified in bright field by superimposing the digital masked fluorescent image on the bright image.

**Results and discussions**

**Specific bacterial detection with fluorescent tagged antibody.** One colony from each bacterial strain isolated from the solid growth medium was incubated in specific broth and both bacterial strains (\(E. faecalis\) and \(P. aeruginosa\)) showed a robust growth curve after about 3 hours on incubation at 37°C reaching a high optical density up to 1.5OD measured at 600 nm (table 1).

A bacterial density of 0.4 OD\(_{600}\) was incubated in presence or absence of a monoclonal antibody anti-\(P. aeruginosa\) tagged with Alexa Fluor revealed the specificity of those antibody to the \(P. aeruginosa\) strains (fig. 2). No detection (red spots) were detected in the \(E. faecalis\) bacterial sample, but an abundant red spots presence on the \(P. aeruginosa\) strain. This showed a very robust and specific interaction between the antibody and the targeted bacteria. The red spots, observed in the fluorescent image, seemed to overlap to the bacterial shapes identified in the transmitted light (bright field) which suggested an even interaction all over the surface between antibody and targeted bacterial molecule (antigenic properties) from the bacterial surface.

**Processing of the bacteria immunodetection forming image into a digital signal**

Both brightfield and fluorescence images were processed by a pipeline processing protocol (fig. 1) in order to achieve a fast and specific digital signal linked to the bacterial identification.

Initially, the fluorescent image of the \(P. aeruginosa\) strain was processed to obtain a grey level image which contains just one plane by extracting the R (red) channel. A binary digital image was obtained by applying a threshold value to the grey level image (pixels’ values spread in 0.255 domain), having background (black) and foreground bacteria regardless if it was the bacterial strain used as a negative control (\(E. faecalis\)) or the bacterial strain of interest (\(P. aeruginosa\)) (fig. 2). The shaped identified in the bright field were consistent with the shapes of each bacterial strain used in this test.

The fluorescent analysis of those bacterial strains incubated in the presence of the monoclonal antibody anti-\(P. aeruginosa\) tagged with Alexa Fluor revealed the specificity of those antibody to the \(P. aeruginosa\) strains (fig. 3). No detection (red spots) were detected in the \(E. faecalis\) bacterial sample, but an abundant red spots presence on the \(P. aeruginosa\) strain. This showed a very robust and specific interaction between the antibody and the targeted bacteria. The red spots, observed in the fluorescent image, seemed to overlap to the bacterial shapes identified in the transmitted light (bright field) which suggested an even interaction all over the membrane surface between antibody and targeted bacterial molecule (antigenic properties) from the bacterial surface.

### Table 1

| Bacterial strain | Growth medium | Absorbance at 600 nm |
|------------------|---------------|---------------------|
| \(E. faecalis\)  | Broth medium - simple concentrated sodium azide | 1.250 |
| \(P. aeruginosa\)| Acetamide nutrient broth | 1.500 |

**Fig. 2.** Microscopic analysis in transmitted light (bright field) of bacterial strains (63x magnification): A-\(E. faecalis\), negative control; B-\(P. aeruginosa\)

**Fig. 3.** Microscopic analysis in fluorescence of bacterial strains: A-\(E. faecalis\), negative control (red filter, 635 nm); B-\(P. aeruginosa\) (63x magnification)

**Fig. 4.** Threshold modification of a grey level image after extracting the red of the in the original fluorescence image.
A good criteria for threshold value is to choose a local minima in the histogram. The binary digital image was further enhanced by several morphological filters and it was carried out for foreground area, in order to eliminate the noise. For small window width and height, Niblack criteria worked well around compact group of pixels, but produces artefacts between those groups, as expected. In order to obtain good results when applied to the whole image, a sliding window having dimension a fourth of the image size was chosen and \(k=1\). Background correction (BCG) worked well on sliding windows having also lesser dimensions (fig. 5). An immediate conclusion is the following: the existence of blobs in the image involves the presence of analyzed bacteria in the bright image.

At the end, a proper binary image was obtained, containing compact blobs (bacterial areas) where fluorescence markers have been detected. The binary digital picture was used for measuring the areas of the detected blobs (bacterial areas) (table 2).

This table containing area of each bacterial areas (expressed both in pixels and square microns) and the percent of each blob area/image area was automatically generated. The sum of blob areas relative to the whole image area gives a percentage indicator of the presence of bacteria in the whole bright image.

The overlapping of the masking bright image with the binary mask image only showed the bacteria recognized by the antibodies eliminating the contaminants and other bacteria which are not specifically recognized by immunofluorescence (fig. 6).

When masked bright image is saved to file, masked area are saved unchanged while blue area are turn to black. This new digital image contains information related only to specific studied bacteria. Figures 5 and 6, as well as the results of the blobs in fig 4 were information on which the final report on the microbiological quality of the analyzed water sample are based on.

Conclusions

A specific and rapid identification of \(P.\) aeruginosa strains from water sample was performed by immuno-fluorescence in presence of monoclonal antibody anti-

\(P.\) aeruginosa tagged with Alexa Fluor 647. The microscopic image (63 fold magnification) displayed a clear identification of bacterial presence both in fluorescence and transmitted light.

A high-tech detection technology proposed an identification and quantification of pathogenic bacteria from drinking water sample based on the fluorescent image that was digitalized, so the results could be sent wireless to all organizations involved in health and environment monitoring.

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