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FISHji: New ImageJ macros for the quantification of fluorescence in epifluorescence images

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A B S T R A C T

Fluorescence in situ hybridization (FISH) is based on the use of fluorescent staining dyes, however, the signal intensity of the images obtained by microscopy is seldom quantified with accuracy by the researcher. The development of innovative digital image processing programs and tools has been trying to overcome this problem, however, the determination of fluorescent intensity in microscopy images still has issues due to the lack of precision in the results and the complexity of existing software.

This work presents FISHji, a set of new ImageJ methods for automated quantification of fluorescence in images obtained by epifluorescence microscopy. To validate the methods, results obtained by FISHji were compared with results obtained by flow cytometry. The mean correlation between FISHji and flow cytometry was high and significant, showing that the imaging methods are able to accurately assess the signal intensity of fluorescence images. FISHji are available for non-commercial use at http://pagina.fe.up.pt/~azevedo.

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1. Introduction

In science, digital images are a crucial piece of information. The ability to process and analyse the large volume of images produced by the plethora of microscopy techniques available raises the need for specialized software tools [1]. Images should be processed in a systematic and standardized way, such that results are comprehensible and reproducible. There are a number of commercial and open source image processing tools available, and the range of implemented functionalities is significant [2–10]. However, the general purpose of most of these tools hampers domain or application-specific use by laymen, i.e. it is not straightforward for a researcher to use such general software to analyse biological images, without understanding the specifics of the image processing and analysis algorithms. As such, the development of end-user biological image analysis tools is considered useful, especially for analyzing fluorescence results obtained by diagnostic methodologies for microorganisms. These methodologies or techniques use staining with fluorescent binding dyes which are then visualized using fluorescence microscopy. In order to have a quantitative analysis of the results, the fluorescence intensity needs to be accurately measured.

Fluorescence in situ hybridization (FISH) is one of the methodologies in which a quantitative validation of results is required. FISH has been used as a molecular tool for the analysis and detection of microorganisms [11]. Since the first application in 1989 by Delong et al. [12], this method has undergone several improvements in

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**Abbreviations:** FISH, fluorescence in situ hybridization; LNA, locked nucleic acid; LoG, Laplacian of Gaussian; MFI, mean fluorescence intensity; PI, propidium iodide; ROIs, regions of interest; TSA, trypticase soy agar; 2 OMe, 2′-O-methyl RNA.

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order to overcome limitations, such as low fluorescence intensity [13,14]. This evolution of the method has led to the use of new types of nucleic acids such as locked nucleic acid (LNA) and 2′-O-methyl RNA (2′OMe) [14,15]. Other chemical modifications within nucleobases, ribose or deoxyribose, and within phosphate moieties have also been introduced to improve the resistance to nuclease present in the cells or body fluids [16–18]. In FISH experiments, the evaluation of the results is based on microscopy, which the quantitative assessment is performed by the researcher, which is dependent of the observer interpretation [19], or on flow cytometry, that provides an automated quantitative assessment but requires the use of expensive equipment and is only applicable for cells in suspension. Therefore, computerized image processing tools offer the most promising and versatile approach to minimize costs and overcome the variability of human microscope analysis. Daim et al. developed a digital image analysis program (DAIME) especially used for microbial ecology [20]. Although, DAIME has different features such as analyzing 2D and 3D microscopy datasets of microorganisms stained by FISH, this program is complex, is generally used for environmental bacteria [21–23] and no validation studies against flow cytometry have been made so far. ImageJ is another well-known and publicly available image processing tool (http://rsbweb.nih.gov/ij/). It has released many plugins and macros useful to biomedical image processing [24], however, also in this case there are no comparative studies against flow cytometry to confirm the use of this software in fluorescence quantification of microscopic images.

This article describes five different methods for the quantification of fluorescent intensity in microscopic images (named FISHji). As a case study, LNA probes were applied in different hybridization conditions, and evaluated by microscopy and cytometry approaches. The biological goal was the detection of 16S rRNA in the clinically relevant bacterium Helicobacter pylori. The developed methods are freely available and integrated in the ImageJ software package.

2. Experimental procedures

2.1. Workflow of FISHji methods and validation

In this work, three semi-automatic and two automatic FISHji methods were created (Fig. 1). To validate these methods, the FISH procedure was performed both in bacterial suspensions and in attached bacteria. Eighteen LNA probes specifically designed for H. pylori were used at different temperatures, in order to obtain a large range of fluorescence intensities. All images obtained by microscopy (attached bacteria) were analysed by the five FISHji methods and these results were compared with cytometry data (bacterial suspensions) hybridized under the same conditions (temperature and buffer). Afterwards, all results were statistically analysed and the correlation between each FISHji method and flow cytometry was performed. Finally, an extra validation was then performed using another fluorescence staining procedure (propidium iodide) for the FISHji methods.

2.2. Fluorescence analysis and FISHji architecture

In terms of operational mode, there are three semi-automatic (FISHji1, FISHji2 and FISHji3) and two automatic approaches (FISHji4, and FISHji5) (Fig. 2). For all methods, the channels of the original RGB (red, green, and blue light) images were separated in order to analyse the channel where fluorescence is emitted.

The methods consist of three sequential steps: optimization, segmentation and analysis, and measurement steps. In the optimization module, pixel-based treatments are performed in order to highlight the regions of interest (ROIs) and allow the removal of artefacts. In the segmentation module a default black and white threshold supports cell segmentation. The ROIs are outlined and artefacts are discarded by the command “Analyze Particles”. The mean fluorescence intensity (MFI) is then calculated as the average of each ROI (Fig. 2).

For FISHji1, optimization treatments are not applied. In the FISHji2 and FISHji3, a background subtraction based on the “rolling ball” algorithm (radius = 80 pixels) is performed to the green channel in order to reduce background heterogeneity and the presence of artefacts [25]. In the FISHji2 method, the obtained image is segmented and further analysed to determine the MFI of the image. In FISHji3, the brightness and contrast features of the corrected image are manually adjusted using the “B/C Adjustment” (Brightness and Contrast) tool to further enhance the foreground. Subsequently, the adjusted image is thresholded and a binary mask is created. The ROIs present in the mask are redirected to the corrected image and MFI is quantified by the ROI manager interface.

In the automatic FISHji4 and FISHji5 methods, the corrected image is convoluted using a Laplacian of Gaussian (LoG) filter (9 × 9 kernel) [26]. In the FISHji5 method, an automatic B/C adjustment of the corrected image is performed before the convolution. First, a B/C adjustment based on the image’s histogram is automatically applied and then the convolution step was performed. In addition, the binary operation “Fill Holes” is applied to fill intra-aggregate spaces and maintain ROI integrity, providing an optimal binary mask. These optimization steps (except “Rolling Ball”) change the real pixel values of the image, including the ROIs. So, a duplicated image is taken from the corrected image in order to allow enhancement of foreground pixels without compromising MFI quantification. After segmentation, the ROIs defined by the binary mask are redirected to the duplicated image, and only then the MFI is calculated. Thus, the MFI is determined from the real pixel values of the Corrected Image, outlined by the binary mask overlay.

All FISHji methods were implemented as macros in ImageJ (version 1.49o or higher). The analysis parameters were set as 100–600 pixels for the minimum and maximum valid object size (i.e. pixel2) and 0.1–0.7 for the range of circularity i.e. 4 × π × (area/perimeter2). These values were established by experimental trials in H. pylori, and taking into account the real size and shape of the bacterium. Images were not calibrated since all had the same resolution (1392 × 1040 cm).

FISHji4 and FISHji5 macro can be found on the webpage http://paginas.fe.up.pt/~nazevedo/.

2.3. Overlap index

Overlap index Ω is a measure of activating regions’ spatial overlap and symmetry. This value was determined using two different edges (image processing technique for finding the boundaries of cells within images detecting discontinuities in brightness): the defined by the ImageJ method and the manual one as the ground truth [27]. After obtaining the binary images of each ROI using ImageJ, a Matlab® script (data not shown) was developed to overlap the images and subsequently to calculate the overlap index Ω according to the following equation [28]:

\[
Ω = \frac{TP}{TP + FP + FN}
\]  

where TP is the true positive area, FP is the false positive area (FP) of the ROI and FN is the false negative area or background of the ROI.

2.4. Oligonucleotide probe design and synthesis

Fluorescein-labeled locked nucleic acids oligonucleotides DNA/LNA and LNA/2′-OMe RNA chimeras with phosphates (PO)
or phosphorothioates (PS) backbones (Table 1) were designed based on previous reports [29,30]. Oligonucleotides were prepared as described previously [31]. LNA and 2′OMe monomers were purchased from Exiqon (Copenhagen, Denmark) and Ribotask (Langeskov, Denmark), respectively.

2.5. Bacterial strains and culture conditions

*H. pylori* strain 26695 obtained from the American Type Culture Collection (ATCC 700392, VA USA) was maintained on trypticase soy agar (TSA) supplemented with 5% (v/v) sheep blood (Becton Dickinson GmbH, Germany). Single colonies were streaked onto fresh media every 48 h and the plates were incubated at 37 °C under microaerobic conditions. Bacterial density was determined by the dilution of initial culture in water or saline and the absorbance was measured at 600 nm. All experiments were performed using bacteria in the same growth phase to avoid differences associated to cell permeabilization [32].

2.6. Hybridization conditions on slides and in suspension by FISH

To validate the FISHji methods, hybridization was assessed by fluorescent microscopy on slides in attached bacteria and by flow cytometry in bacterial suspensions. All probes from Table 1 were tested at temperatures between 37 °C and 65 °C and the fluorescence signal was quantified by both cytometry and FISHji methods. For both methodologies triplicate samples were used. In FISHji measurements we have also analysed nine images for each sample in order to study the entire sample. FISH in attached and suspended bacteria was performed as described in a previous study [31]. The cells were left at 4 °C until being analysed by cytometry. Probes were tested simultaneously at the same temperature in both samples types (attached and suspension bacteria), to minimize experimental variations between both methods (cytometry and FISHji).

2.7. Propidium iodide (PI)

Increasing concentrations of bacteria suspension (500 µL) were mixed with propidium iodide (PI) (Invitrogen, Eugene, USA) at a concentration of 50 µg/mL. For each concentration four samples were stained. Bacteria were incubated with PI for 10 min at room temperature. Afterwards, the samples were centrifuged during 5 min at 14,000 rpm and the supernatant was discarded. Then, two of the pellets were resuspended in water and analysed by microscopy and the other two were resuspended in saline and analysed by flow cytometry.

2.8. Microscope evaluation and quantitative analysis of fluorescence intensity

The slides with attached bacteria were evaluated on a Leica DM LB2 epifluorescent microscope. For image acquisition, a Leica

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**Table 1**

Probe sequences used in this study. LNA nucleotide monomers are represented with L superscript. 2′-OMe-RNA monomers in boldface letters, DNA nucleotides in capital letters, and phosphorothioate linkages by the symbol *.

<table>
<thead>
<tr>
<th>Probes analysed</th>
<th>Sequence (5′-FAM-3′)</th>
</tr>
</thead>
<tbody>
<tr>
<td>HP,18LNA,PO</td>
<td>GA*GAGTAAAGCCCTTGCTTT</td>
</tr>
<tr>
<td>HP,18LNA,PS</td>
<td>GA*GAGTAAAGCCCTTGCTTT</td>
</tr>
<tr>
<td>HP,18LNA,20Me,PO</td>
<td>GA*GAGTAAAGCCCTTGCTTT</td>
</tr>
<tr>
<td>HP,18LNA,20Me,PS</td>
<td>GA*GAGTAAAGCCCTTGCTTT</td>
</tr>
<tr>
<td>HP,15LNA,PO</td>
<td>GA*GAGTAAAGCCCTTGCTTT</td>
</tr>
<tr>
<td>HP,15LNA,PS</td>
<td>GA*GAGTAAAGCCCTTGCTTT</td>
</tr>
<tr>
<td>HP,15LNA,2,2</td>
<td>GA*GAGTAAAGCCCTTGCTTT</td>
</tr>
<tr>
<td>HP,15LNA,2,2</td>
<td>GA*GAGTAAAGCCCTTGCTTT</td>
</tr>
<tr>
<td>HP,15LNA,PS</td>
<td>GA*GAGTAAAGCCCTTGCTTT</td>
</tr>
</tbody>
</table>

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Fig. 1. A general view of this study. Firstly 18 LNA probes were designed and synthesised. FISH was then performed using these probes in different hybridization temperatures, in attached and suspended bacteria. The images obtained by microscopy from attached bacteria were analysed and quantified by four FISHji methods. Subsequently, these results were statistically compared with the data obtained by flow cytometry from bacteria in suspension. The FISHji methods that obtained high correlation values were then validated using a standard staining procedure (propidium iodide).
Fig. 2. The schematic workflow of FISHji methods. After microscopic acquisition of the fluorescent images, the green channel (the wavelength at which the probe fluoresces) was obtained and further analyzed in order to obtain the MFI. The FISHji methods consist of three steps: optimization, segmentation and analysis and measurements. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

Optimization step: optimization treatments were applied to eliminate possible background defects and/or to adjust the brightness and contrast features of the corrected image. Segmentation step: for FISHji1,2,3, the ROIs were segmented from the background by manual default black and white threshold, whereas for FISHji4,5 an automatic threshold was applied by the “Make Binary” option. Analysis and measurement step: after segmentation, the ROIs were outlined by the command “Analyze Particles”. Finally, the fluorescence intensity of each ROI was provided by the option “Measure” in the ROI interface and then the average value was determined by the “Summarize” command as well as the STD. It should be noted that the overlay of the ROIs with the original image represented by the “Outlined ROIs” images is only to demonstrate the efficiency of the ROI definition. In real action, this overlay is not necessary. B/C—Brightness and Contrast; LoG—Laplacian of Gaussian; ROI—region of interest; MFI—mean fluorescence intensity; STD—standard deviation.
DFC300 FX camera (Leica Microsystems GmbH, Germany) was used equipped with a filter capable of detect the FAM- labeling LNA probes (BP 450–490, FT 510, LP 515). The exposure time, gain and saturation values were fixed for all preparations. For image capture, Leica IM50 Image Manager, Image Processing and Archiving software was used. All experiments were performed in triplicate.

2.9. Flow cytometry and data analysis

The analysis of fluorescence intensities obtained by FISH in suspension was performed by flow cytometry. Flow cytometry analysis was performed using a Beckman Coulter Epics XL (Brea, USA) using the EXPOL32ADC software (Beckman Coulter, Brea, USA) equipped with a low-power air-cooled 15 mW blue (488 nm) argon laser. For each sample, 20,000 events were collected. All the experiments were performed in triplicate and negative controls (samples without probes) were included for each temperature in every analysis. Flow cytometric analyses of samples were performed based on forward scatter, side scatter and FL-1. Fluorescence was detected on the FL1 channel (BP530/30). Amplification was carried out using logarithmic scales.

2.10. Statistical and correlation analysis

Results are compared using one-way analysis of variance (ANOVA) by applying Tukey multiple-comparisons test, using GraphPad Prism version 5 software (GraphPad Software, San Diego, USA). All tests were performed with a confidence level of 95%.

3. Results and discussion

3.1. Comparison between FISHji measurements and quantification by cytometry

In this paper we developed and validated five semi-automatic or automatic fluorescent intensity quantification methods, called FISHji, capable of quantifying the MFI. The settings and characteristics of each FISHji method developed are described in Table 2. As ground truth, the MFI of the hybridization experiments was also quantified using flow cytometry (gold standard method in fluorescence quantification) at the same temperature as the hybridizations performed in attached bacteria.

In order to assess which FISHji method provide MFI values closer to the ones obtained by cytometry, the correlation between results from each FISHji (semi-automatic and automatic methods) and cytometry was calculated (Table 3). The three semi-automatic methods (FISHji 1, FISHji 2 and FISHji 3) present different results when compared to the cytometry. FISHji 1 showed the lowest correlation with flow cytometry (Table 3). FISHji 2 showed a better correlation than FISHji 1, which suggests that, the performance of the optimization steps improved MFI quantification efficiency. The FISHji 3 revealed to be the best semi-automatic method demonstrating that the application of a B/C adjustment after subtraction of the background contributed to a more effective MFI quantification.

Nevertheless, semi-automatic methods showed to be very time-consuming when a large number of samples have to be processed (Table 1), and since a routine protocol to perform fluorescence quantification should be fast and easy to work with, the need for an automatic method becomes clear. The automatic threshold definition is much more difficult to perform since one image is never equal to the other. Hence, has emerged the need for a more suitable and automatic segmentation technique. Previous studies have used LoG convolution to automatically enhance ROIs from the background with good outcomes [33]. Herein, FISHji4 was designed in which this approach was used to do both smoothing and edge detection. A high correlation between our plugin results and flow cytometry analysis for the automatic FISHji4 was observed (Table 3). However, the correlation factor is lower comparatively to the semi-automatic FISHji3. This might have been influenced by cells from aggregates that were selected and subsequently counted due to the elevated sensibility of the convolution (Fig. 2). Further, artefacts are not possible to discard in the ROI manager.

In an attempt to increase the automatic performance, a fifth method was designed (FISHji5) specifically to blurred regions after subtracted background, thereby connecting nearby cells (i.e. belonging to aggregates) and subsequently impairing the aforementioned problem. A high correlation between FISHji 5 and flow cytometry was also observed (Table 3). Contrary to expected, no significant differences emerged between both automatic methods. Visual image analysis allowed to conclude that the influence of aggregates in the present study was overestimated and auto B/C adjustment in ROIs with high fluorescence intensity increased the area of analysis, selecting false positive regions.

The results obtained for the correlation coefficient for FISHji 3, 4 and 5 are shown in detail in Fig. 3, where each data point consists of two parameters: the X-axis values corresponds to the mean of fluorescence intensity obtained by cytometry with a specific probe and temperature and the Y-axis values represent the mean value of fluorescence intensity obtained by FISHji in the same conditions. The average Pearson product moment correlation coefficient from all our biological repeats show a high level of correlation between the FISHji results and cytometry analysis, r = 0.819 (0.733–0.880) in FISHji3.

One of the main factors that is influencing the results is the implemention of optimization steps, that reflects on the efficiency of the definition of the edges of the ROI. In order to verify this influence in the MFI quantification, we calculated the overlap index Ω, as a measurement of the overlap performance between the edges defined by the implemented methods and the ground truth. This analysis is a commonly used method in several processing image studies, however it can suffer different adaptations depending on the purpose of the analysis [34,35]. FISHji1 showed the lowest mean Ω, indicating that the application of optimization steps before image segmentation improved the definition of the cell. In general, the overlapping results were congruent with the correlation factors for each method, indicating that the cell definition contributes largely to the MFI quantification.

To evaluate whether the choice of the optimization steps and the consequent image analysed had influence in the MFI quantification, FISH images were reanalysed by the FISHji4 and FISHji5 methods, however now redirecting the binary mask to the original image instead of the corrected one. The low correlation coefficients obtained (r = 0.317 and r = 0.231, for FISHji4 and FISHji5) respectively showed that the use of corrected images (after optimization steps) is essential in these analysis. Additionally, the original images can contain many different artefacts and autofluorescence signals that may blur or omit the real fluorescence value of the cell. The application of the “rolling ball” algorithm to the original green channel can correct background heterogeneity and eliminate some artefacts, contributing to a more realistic MFI quantification when compared to the cytometry. Therefore, the choice of the analysed image showed to be a major point in the MFI quantification, whereas the application of optimization steps before thresholding takes a relevant part in the efficiency of the definition of the cell, contributing in a smaller scale to the MFI quantification performance.

We also evaluated the influence of hybridization temperature in FISHji methods. Although there were differences between different hybridization temperatures, overall, the correlation coefficients were higher than those in the global analysis. As an example, we showed in Fig. 4 the results for hybridization performed at 45 °C and 55 °C for FISHji3. In both cases we observed a high correla-
Table 2
Comparisons between the five FISHji methods. The different methods were compared relatively to these general features.

<table>
<thead>
<tr>
<th>General features</th>
<th>FISHji1</th>
<th>FISHji2</th>
<th>FISHji3</th>
<th>FISHji4</th>
<th>FISHji5</th>
</tr>
</thead>
<tbody>
<tr>
<td>Operational mode</td>
<td>Semi-automatic</td>
<td>Semi-automatic</td>
<td>Semi-automatic</td>
<td>Automatic</td>
<td>Automatic</td>
</tr>
<tr>
<td>Image for analysis</td>
<td>Single image</td>
<td>Single image</td>
<td>Single image</td>
<td>Multiple images</td>
<td>Multiple images</td>
</tr>
<tr>
<td>Analysis speed per image</td>
<td>-50 s</td>
<td>-50 s</td>
<td>~1 min 20 s</td>
<td>4 s</td>
<td>4 s</td>
</tr>
<tr>
<td>Cell segmentation</td>
<td>Manual threshold</td>
<td>Manual threshold</td>
<td>Manual threshold</td>
<td>Automatic threshold</td>
<td>-</td>
</tr>
<tr>
<td>Rolling Ball</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>Manual B/C</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>+</td>
<td>-</td>
</tr>
<tr>
<td>LoG</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>Fill Holes</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>+</td>
<td>-</td>
</tr>
<tr>
<td>Analysed image</td>
<td>Original</td>
<td>Corrected</td>
<td>Corrected</td>
<td>Corrected</td>
<td>Corrected</td>
</tr>
</tbody>
</table>

Table 3
Comparison between each FISHji method values and flow cytometry data.

<table>
<thead>
<tr>
<th>Measurements</th>
<th>FISHji 1</th>
<th>FISHji 2</th>
<th>FISHji 3</th>
<th>FISHji 4</th>
<th>FISHji 5</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ω</td>
<td>0.46</td>
<td>0.51</td>
<td>0.54</td>
<td>0.63</td>
<td>0.62</td>
</tr>
<tr>
<td>Correlation coefficient (p-value)</td>
<td>0.456 (&lt;0.0001)</td>
<td>0.751 (&lt;0.0001)</td>
<td>0.819 (&lt;0.0001)</td>
<td>0.723 (&lt;0.0001)</td>
<td>0.717 (&lt;0.0001)</td>
</tr>
</tbody>
</table>

* Flow cytometry is used as a ground truth for calculating the correlation coefficient and the p-value.

Fig. 3. Correlation analysis between fluorescence intensity (A.U.F) of LNA FISH obtained by ImageJ method and flow cytometry. (A) FISHji3; (B) FISHji4; (C) FISHji5. Regression lines are drawn and a high correlation between both methods was observed for semi and automatic methods in this study. Each data point consists of two parameters: the X-axis values corresponds to the mean of fluorescence intensity obtained by cytometry with a specific probe and temperature and the Y-axis values represent the mean value of fluorescence intensity obtained by FISHji in the same conditions.

Fig. 4. Analysis of correction between fluorescence intensity (A.U.F.) obtained by FISHji3 method against fluorescence intensity (A.U.F.) obtained by flow cytometry, using hybridization temperatures. (A) 45 ºC; (B) 55 ºC. Regression lines are drawn and the r is displayed for each data comparison.
tion $r = 0.946 (0.724–0.990)$ for $45\, ^\circ \text{C}$ and $r = 0.907 (0.647–0.978)$ for $55\, ^\circ \text{C}$. This suggests that the lower correlation values obtained for the FISHji3 using the entire set of data might also be due to some experimental variation.

Because FISHji3 is a semi-automatic method, it requires manual definition of a threshold by the operator. In order to evaluate the influence of different operators in MFI quantification, two different operators have quantified the MFI of the same images. Our results (Fig. 5) did not show statistically significant differences between the MFI values determined by each operator ($p > 0.05$). The same results were obtained for FISHji1 and FISHji2 (see Supplementary material Fig. S1 in the online version at DOI: 10.1016/j.bej.2016.04.001).

While FISHji3 shows higher correlation with cytometer and is operator-independent for threshold, automatic methods FISHji4 or FISHji5 could be the key to development of a routine analysis for multiple FISH images. In this sense, automatic quantification without the need of expensive equipment (such as cytometer) would be recommended. Comparisons between FISHji3, FISHji4 and FISHji5 showed high levels of correlation in both cases (Fig. 6), suggesting that both FISHji4 and FISHji5 offer the ability to automatically analyse considerable amounts of image data.

In the present study, the main cornerstone of fluorescence intensity quantification was cell segmentation (i.e. definition of the ROIs). Indeed, segmentation can be very difficult due to the complexity of original images, represented by combination of different features like (1) autofluorescence; (2) background heterogeneity; (3) presence of non-cell objects and aggregates (herein considered as artefacts). Therefore, a very important issue to take in consideration when using this tool is the high dependence on image quality [3]. There are several factors which may interfere with the quality of the images, such as the numerical aperture (NA) of the objective, filed illumination of the lens, the refractive index of the embedding medium, use of coverslip, stability of the light source, the ratio between emission and excitation wavelength, amongst others [36]. Therefore, we always used the same settings in the microscope for every experiment to minimize these factors. Although all parameters associated with the image collection were kept constant, such as exposures times, illumination and focus, it was impossible to guarantee that the fluorescent lamp would produce the same light intensity during the whole of the experiments.

3.2. Correlation between FISHji measurements and quantification by cytometry using PI staining

In order to address the use of FISHji methods in measuring fluorescence obtained by means other than FISH, we performed the staining of H. pylori samples with PI. Therefore, we analysed the same sample by microscopy (and consequently the fluorescence intensity by FISHji 3, 4 and 5 methods) and cytometry. All
4. Conclusion

In biological sciences, fluorescence-based methods are crucial to study, analyse or identify microorganisms. Here, we present FISHi methods to quantify fluorescence in images from biological samples obtained by microscopy. We show that the semi-automatic FISHi3 and automatic FISHi4 and FISHi5 have high accuracy in different settings such as FISH and PI staining. Because FISHi4 presents better results comparatively to FISHi5, we consider that this method should be preferred for fluorescence quantifications. Nonetheless, it is important to bear in mind that variations in the types of samples (e.g. cell size or aggregation), might imply that for specific situations, FISHi5 might be more adequate.

Since automated methods are able to simultaneously analyse multiple images and are consequently faster, we provide an easy and freely available access not only to FISHi4 but also to FISHi5 for non-commercial use at http://pagina.fe.up.pt/~nazevedo/.

Author contributions

Conceived and designed the experiments: SF, DC, NFA. Performed the experiments: SF, DC. Analyzed the data: SF, DC, AL, NG, NFA. Tool development: SF, DC, AL; Contributed reagents/materials/analysis tools: SF, DC, PM, CF, NFA. Wrote the paper: SF, DC, AL, NG, PM, CF, NFA.

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