Angiogenesis Analyzer for ImageJ — A comparative morphometric analysis of “Endothelial Tube Formation Assay” and “Fibrin Bead Assay”

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Angiogenesis assays based on in vitro capillary-like growth of endothelial cells (EC) are widely used, either to evaluate the effect of anti- and pro-angiogenesis drugs of interest, or to test and compare the functional capacities of various types of EC and progenitor cells. Among the different methods applied to study angiogenesis, the most commonly used is the “Endothelial Tube Formation Assay” (ETFA). In suitable culture conditions, EC form two-dimensional (2D) branched structures that can lead to a meshed pseudo-capillary network. An alternative approach to ETFA is the “Fibrin Bead Assay” (FBA), based on the use of Cytodex 3 microspheres, which promote the growth of 3D capillary-like patterns from coated EC, suitable for high throughput in vitro angiogenesis studies. The analytical evaluation of these two widely used assays still remains challenging in terms of observation method and image analysis. We previously developed the “Angiogenesis Analyzer” for ImageJ (AA), a tool allowing analysis of ETFA-derived images, according to characteristics of the pseudo-capillary networks. In this work, we developed and implemented a new algorithm for AA able to recognize microspheres and to analyze the attached capillary-like structures from the FBA model. Such a method is presented for the first time in fully automated mode and using non-destructive image acquisition. We detailed these two algorithms and used the new AA version to compare both methods (i.e. ETFA and FBA) in their efficiency, accuracy and statistical relevance to model angiogenesis patterns of Human Umbilical Vein EC (HUVEC). Although the two methods do not assess the same biological step, our data suggest that they display specific and complementary information on the angiogenesis processes analysis.

Angiogenesis, the growth of new blood vessels from pre-existing ones, is a complex and critical process that takes place during vertebrate development, in specific physiological conditions in adult individuals and during different pathologies. Endothelial cells (EC) represent the main cell type engaged in this process. Under appropriate stimuli, these cells sprout from a root vessel, migrate, proliferate, then align and ultimately form tubes. Accurate and objective evaluation of these phenomena is necessary for the full comprehension of this process, for both fundamental approaches and pharmacological applications. For example, the use of anti-angiogenic molecules still represents a promising therapeutic strategy to impede the development of solid tumors. Other fields of interest, such as stem cell research, also require the evaluation of angiogenic capacities of EC progenitors as well as quantitative tools to monitor their proliferation and differentiation capacities. In this context, in vitro as well as in vivo experimental models have been developed to evaluate angiogenesis features, to screen a variety of new angiostatic molecules and to study their properties. Most in vitro angiogenesis models were designed based on the so-called “sprouting angiogenesis” differentiation process, whereby pseudo-capillary formation mimics...
The adaptation of the FBA model into 96-well plates allows high-throughput drug screening using small amounts of the tested products. This method has the advantage of being closely related, in terms of 3D sprouting freedom with a solid anchorage point, to an in vivo assay known as the "Rat Aortic Sprouting Assay". In addition, the FBA test avoids the use of animal experimentation and favors good repeatability.

Although the qualitative features of both ETFA and FBA have been well described, better quantification methods are needed. Indeed, these assays still present technical flaws, due to physical, optical and computer constraints. Among technical difficulties lays the focus default due to meniscus formed by culture gels, especially in small culture wells used for high-throughput microscopy in FBA experiments. Another difficulty is the choice of the most appropriate acquisition method (fluorescence or phase contrast microscopy) and associated software tool for image analysis. Finally, special consideration must be given to the morphologic parameters to be taken into account for optimal statistical analysis.

Here, it is proposed to compare ETFA and FBA models by using a customized version of the previously proposed "Angiogenesis Analyzer", a simple and precise tool built in the ImageJ environment and formerly used to quantify the FBA experiment images. The "Angiogenesis Analyzer" program was conceived to extract characteristic points and elements of endothelial cells network. This image analysis software was successfully used to characterize meshed and/or branched structures in more than 150 different studies (http://image.bio.methods.free.fr/ImageJ/?Angiogenesis-Analyzer-for-ImageJ&amp;artpage=6-6#outil_sommaire_6), which include endothelial in vitro ETFA cell differentiation in phase contrast, fluorescence microscopy, and in vivo studies using the mouse retina angiogenesis model. The "Angiogenesis Analyzer" has also been used to analyze branching in alternative meshed or branched biological scaffolds unrelated to angiogenesis, as reported for the characterization of diatom silicate structures in phytoplankton research, or for in vitro neuritic branching analysis in cultured neurons.

The "Angiogenesis Analyzer" software works as an integrated program, avoiding the requirement for image pretreatment using additional plugins, or online computation services on distant servers, such as required by other available solutions. The "Angiogenesis Analyzer" is free and open source, easy to customize and compatible with several OS platforms (Linux, MacOS and Windows) thanks to a Java execution environment. Moreover, it can be used to obtain basic hierarchical data in the form of series of vectorial objects by modeling the analyzed structure at different levels, while most other solutions afford basic measurements as total tree lengths and meshes areas under the form of binary measurements.

We previously separately validated "Angiogenesis Analyzer" for both ETFA and FBA models. Here, using the customized "Angiogenesis Analyzer", we compared the reliability, accuracy and statistical relevance of parameters obtained from two sets of FBA and ETFA experiments, using both activating and inhibiting conditions of angiogenesis on HUVEC cells. These cells spontaneously differentiated, aligned and branched in the FBA model, and formed an irregular meshed network in the ETFA experiments. Cellular organization images were acquired through phase contrast microscopy and analyzed with their corresponding algorithms that were described in details for both model. The FBA algorithm described in this work is the first solution available for FBA fully automated computer analysis using phase contrast microscopy, allowing thousands of microbead analysis per day on a personal computer, without user intervention. By comparative analysis of measurements we observed that sprouting initiation capacities in FBA seem related to meshing development in ETFA, while length of pseudo-vascular tree in FBA is rather associated to a later stage of development. Our data suggest that the two methods do not assess the same biological step and display specific and complementary information in the analysis of angiogenesis processes.

**Results**

**FBA image analysis.** Image analysis for FBA was performed using a program developed for the ImageJ software. This plugin is an extension of the "Angiogenesis Analyzer" for ImageJ written in the macro language of ImageJ. The analysis can be divided into three main steps: (1) sphere detection (Fig. 1), (2) tree detection (Fig. 2) and (3) tree structure analysis of Junctions and Extremities (Fig. 3) as well as Segments, Branches and Anchorage Junctions (Fig. 4):

Sphere detection (Fig. 1): Sphere analysis in ideal conditions does not present particular difficulties. However, sphere clustering, dirt, uneven lighting due to the meniscus of the culture medium and the walls of the culture wells introduce difficulties, especially defaults of object closure. A series of image manipulations were necessary to enable sphere detection using simple threshold methods as follows:

1. Light field correction (lfc) was performed using an artificial flat field obtained by applying a Gaussian convolution with a sigma value (σ) corresponding to the average sphere diameter. Uniform backgrounds were obtained by dividing the initial image (ima1) by the flat field with:
Strong edges enhancement (enh) was then performed by log transforming the image and applying a variance filter with a radius ($r$) corresponding to the apparent thickness of the sphere edges (Fig. 1B) with:

$$\text{Ima}1\,(\text{enh}) = \text{variance}[\log(\text{Ima}1\,(\text{fc}))]$$

Initial segmentation (spMask) was performed to get a coarse binary sphere mask by applying a threshold on ima1(enh) obtained using the « IsoData » method, and binary objects resulting from the analysis were then submitted to a cycle of closing-opening operations and filling holes (Fig. 1C) with:

$$\text{spMask} = \text{fill}\text{hole}[\text{Open}[\text{fill}\text{hole}[\text{Close}[\text{IsoData}(\text{ima}1\,(\text{enh}))]]]]$$

The interior of spheres was detected by a second segmentation (inMask) applied on the difference of images “spMask” and “ima1(enh)” (Fig. 1D). The result was then segmented by the “Minimum” threshold method and hole filling and opening binary operators were applied, with “r” the radius of the Maximum filter matrix (set to 2 pixels) (Fig. 1E) with:

$$\text{inMask} = \text{Open}[\text{fill}\text{hole}[\text{Minimum}\text{Threshold}[\text{Maximum},(\text{ima}1\,(\text{fc}) - \text{spMask})]]]$$

Sphere edges were detected using binary object analysis of “inMask”. Some circles were fitted to the edges and their diameters were enlarged to take into account the thickness of the sphere envelope. Figure 1F shows the sphere edges using a red circle vectorial overlay.

Tree detection (Fig. 2): Tree detection consists of segmentation, avoiding shadows due to phase contrast lighting and small acellular structures, followed by skeletonization, as follows.

Light field correction (lfc) was performed in the same manner as for sphere detection (§1), followed by noise removal using a band pass FFT (Fast Fourier Transform) filter (FFTbp).

$$\text{Ima}2(\text{fc}) = \text{FFTbp}_{(r1,r2)}[\text{ima}2/\text{Gauss}_a(\text{ima}2)]$$

Figure 1. Sphere detection in FBA. (A) initial image observed in phase contrast. (B) edge-enhanced image (§1-2). (C) first binary mask of spheres (§3). (D) difference between B and C, highlighting sphere interiors (§4). (E) binary mask delineating the interior of the spheres (§4). (F) final detection of the sphere edges (red) (§5). “§” labels refer to the different steps described in the Results section. Scale bar: 200 μm.
**Figure 2.** Tree detection in FBA. (A) initial sample image exhibiting the detected sphere as a red vector overlay ($§1-§5$). (B) enhancement of high gradients and background removal ($§6-§7$). (C) “Mean” threshold. (D) final binary segmentation ($§8$). (E) skeleton of the binary segmentation ($§9$). (F) final skeleton after clearing of the sphere interior ($§9$). “§” labels correspond to steps described in the Results section. Scale bar: 200 µm.

**Figure 3.** Detection of *Junctions* and *Extremities* in skeletonized trees obtained through different steps. (A to D) Represent four pixels (in red) each linked to three neighboring pixels (green). (E) The junction between the three binary segments is composed of the four nodes (red) materialized by circles of seven pixels in diameter. (F) The border of the overlaying of these circles (red spot) was defined as the *Junction*, materialized by its smoothed vectorial representation (blue). (G) An *Extremity* corresponds to a pixel linked to only one neighbor pixel. H: representation of one *Extremity*. See step “§10” in the Results section.
Figure 4. Vectorial objects detection in skeletonized trees. (A) Extremities, red dots surrounded in yellow (inset 1) as explained in the Methods section (§10). (B) detection of Branches (green) and Segments (magenta) (§11). Inset 2 shows an artifactual branch (cyan) (because too small) which will be removed by the program. Inset 3 shows a fusion between two nearby Junctions into a single one (blue edge). (C) representation of the final analysis, including the Anchorage Junctions (violet, inset 4), which intersect with the sphere limit (red). "§" labels correspond to steps described in the Results section.

where “ima2” is the original image (Fig. 2A), “v1” is the minimum size of structures to keep and v2 is a value corresponding to the size of the shadows to be removed. “v2” has the same value than “σ”. These values depending of the image resolution are typically v1 = 1.5 and v2 = 400.

§7 Edge enhancement (enh) was performed to increase the signal of the small cellular extensions. A variance (r) filter was used to suppress the alternatively positive-negative aspect of objects inherent in phase contrast imaging as well as to increase high gradient areas, with “r”, the radius corresponding to the minimum size of structures to detect, set to 2 pixels). The histogram modal value was then subtracted to reduce the background (Fig. 2B), with:

\[ima2(enh) = \text{variance}_c(ima2(fc))-\text{modal}[\text{variance}_c(ima2(fc))]]

§8 The image was then segmented using the “Mean” threshold method33, creating a binary mask (Fig. 2C). The mask was smoothed and made continuous using closing and dilation binary operators (Fig. 2D).

\[ima2(mask) = \text{dilate}[\text{close}[\text{MeanThreshold}(ima2(enh))]]

§9 The area within the circles was filled and the resulting binary mask skeletonized (Fig. 2E). Structures not in contact with the circles corresponding to the previously detected spheres were removed to avoid isolated pseudo-capillary objects. Circles in the skeleton were cleared to get the final tree (Fig. 2F).

Tree structure analysis: Junctions and Extremities detection (Fig. 3):

§10. Tree structure analysis started with Junctions and Extremities detection. To perform this, we defined three sub-elements: Nodes, which consisted in the minimum structure allowing a bifurcation in a skeletonized network, Junctions and Extremities. A pixel corresponded to a Node when it had at least 3 neighbors (Fig. 3 A-D). To avoid eventual further skeletonization artifacts, each pixel forming a Node was then replaced by a circular dot from 7 pixels of diameter (Fig. 3E). Junctions were formed by the group of dots associated to a bifurcation (Fig. 3F). A pixel was qualified as an Extremity when it had only one neighbor (Fig. 3G-H). Figure 3 summaries different configurations of Nodes leading to a Junction and an Extremity.

Tree structure analysis: Segments, Branches and Anchorage Junctions detection (Fig. 4):

§11. Result of step §10 (Fig. 4A) was submitted to the analysis of branch and segment content, according to the following definitions. Branches were lines, which are linked to one Junction and one Extremity (green elements, Fig. 4B) and Segments were lines connected to the main tree by two Junctions (magenta elements Fig. 4B). Touching Junctions, or too close Junctions (spaced less than 20 pixels apart) were fused into single Junctions to remove segments composed of less than two aligned cells (zoom inset 3, Fig. 4B). Artifactual small Branches were removed by an iterative pruning (zoom inset 2, Fig. 4B). Segments corresponding to Circle pieces intercepting circles selections were removed by a previously described method35. Using the same principle, the Junctions that intercepted Circles corresponding to sphere limits were defined as Anchorage Junctions (zoom inset 4 Fig. 4C). At this step, the automatic analysis resulted in a model of the sphere and its associated tree, consisting in a group of vectorial objects (Circles, Extremities, Junctions, Branches, Segments and Anchorage Junctions) (Table 1) which were counted and measured for statistical calculations.

ETFA image analysis. HUVEC cultured in Matrigel were treated to reduce noise and lighting inhomogeneity similarly than FBA images. The tree structure detection was done as in steps §6 to §9, except for the removal of the sphere area (Fig. 5A-C). Analysis of the binary mask was performed as explained in steps §10 and §11 except for the detection of Anchorage Junctions.

§12. Isolated Elements detection
angiogenic parameters derived from AA on EFTA are summarized in Fig. 7, panel A to E. A highly developed meshing, depending on the treatment conditions (Fig. 6). Quantitative analysis of different aspects after one day of culture, as observed by phase contrast microscopy, from short and isolated segments to figure and to the analytical step in the Methods section that describe each of the vectorial elements.

Table 1. Vectorial objects (elements) characterized by the software analysis. The “Text” column refers to the figure and to the analytical step in the Methods section that describe each of the vectorial elements.

| Vectorial element | Abbreviation | Short definition | Text |
|-------------------|--------------|------------------|------|
| Node              | Nd           | Pixel having at least three neighbors | Fig. 3, §10 |
| Junction          | Jnc          | Groups of nodes forming a bifurcation | Fig. 3, §10 |
| Extremity          | Ext          | Pixel having only one neighbor | Fig. 3, §10 |
| Segment           | Seg          | Binary line linked with two junctions | Fig. 4, §11 |
| Branch            | Br           | Binary line linked with one junction and one extremity | Fig. 4, §11 |
| Anchorage junction | AchJ         | Junction linking a branch or a segment to a sphere limit | Fig. 4, §11 |
| Isolated Element  | IsE          | Binary line connected to two extremities | Fig. 5, §12 |
| Mesh              | Msh          | Area enclosed by segments | Fig. 5, §13 |

Elements limited by two extremities were classified as isolated elements. They were removed when the size was lower than a user-defined value (10 pixels in our analysis) which corresponds to the minimum size of an isolated cell. §13. Meshes detection:

HUVEC in 3D gel grow according to a meshed network. To implement a parameter allowing to quantify this level of cellular organization, a supplementary vectorial object was defined in analysis - “Meshes” - corresponding to closed areas delimited by Segments and associated Junctions (Fig. 5D) that were detected and measured (Fig. 5E). The final representation (Fig. 5F) summarizes all the detected structures from the initial image. Vectorial objects and measurement definitions are summarized in Tables 1 and 2, respectively.

We further decided to test and compare this two algorithms on the reliability, accuracy and statistical relevance of some measurements obtained on two sets of FBA and EFTA experiments. These assays were performed in parallel on HUVEC cells, using both activating (VEGF-A) and inhibiting (sunitinib) conditions of angiogenesis. Images from cellular organization were acquired from EFTA and FBA through phase contrast microscopy (Fig. 6) and analyzed with the corresponding algorithms to detect the different vectorial objects described above (Table 1). We finally made a comparative analysis of the parameters derived from the object measurements (Table 2), (Fig. 7).

The endothelial tube formation assay response. HUVEC in culture wells present a wide range of aspects after one day of culture, as observed by phase contrast microscopy, from short and isolated segments to a highly developed meshing, depending on the treatment conditions (Fig. 6). Quantitative analysis of different angiogenic parameters derived from AA on EFTA are summarized in Fig. 7, panel A to E.

Compared to control condition, the mean size of HUVEC meshes (MMS) slightly increased when VEGF-A concentrations went from 5 to 10 ng/ml, and then decreased with higher concentrations (25 and 50 ng/ml) (Fig. 7A). Although this effect is not statistically significant (VEGF-A 10/Ctrl = 1.18, NS), the tendency is still clearly visible. The same biphasic effect of VEGF-A increasing dose was significantly observed when considering the Total Mesh Area (TMA) (VEGF-A 10/Ctrl = 1.36 (p < 0.05)) (Fig. 7C). While this tendency was still visible with Total Segment Length (TSL) (VEGF-A 10/Ctrl = 1.13 NS) (Fig. 7D), it became erratic in terms of Junction Number (JN) and Total Length (TL) values (Fig. 7B,E). This can be explained by considering the kinetic of the meshing network structuration during HUVEC culture in Matrigel. As previously observed in time-lapse recording of EC in similar culture conditions (video at this link: http://image.bio.methods.free.fr/ImageJ/?Human-Endothelial-Progenitor-Cells-in-vitro-tube-forming-analysis-using-Lens), the network establishment started (3–6h of culture) by the formation of a multitude of small and unstable meshes. The size of the meshes then progressively increased by fusion of proximal meshes, which occurred by segments (tube) regression or disruption. Indeed, disruption of a segment can lead to the transformation of two adjacent meshes in a single one, as regression of a segment can lead to regression of a mesh until its disappearance. Examples of this phenomenon are shown in Fig. 8. Red double-head arrows show segment disruption leading to meshes fusion by opening a breach in segment continuities. Residual branches maintained the constant Junction Number (JN) at this early stage of the regression and the decrease of the Total Length (TL) was negligible. Although the disruption leads to a higher decrease in Total Segment Length (TSL) by a length value that corresponds to the double head blue arrow (Fig. 8B), such phenomena is not sufficient to be significant in the TSL measurements in the VEGF-A response (Fig. 7D). MMS increased as TMA, segment disruptions and mesh fusion leading to an increase in residual meshes (Fig. 7A,C). This tendency reversed when mesh size became too high to be entirely visible in the image field or when meshes became broken. At this step (VEGF-A 25 and 50, Fig. 7A,C), contribution of big meshes decreased, as the mean values of MMS and TMA.

In the presence of 5 nM sunitinib, the capillary-like network was not significantly increased (MMS Sunit 5/Ctrl = 1.18 (NS)). Comparatively, the network was highly altered with 25 nM sunitinib (MMS Sunit 25/Ctrl = 0.06 (p < 0.0001)) (Fig. 6,8A). This apparent contradiction - there was a non significant increase of MMS with 5 nm sunitinib (Fig. 7), while a decrease would have been expected - can be explained by the process of mesh growing. As indicated above, the increase in mesh size occurs by segment disruption, thus leading to mesh fusion. A slight inhibitory effect affecting the tube organization as a function of time has the same effect - segment disruption and mesh fusion (Fig. 8A). This makes the use of MMS value inappropriate in the presence of a low concentration of inhibitor. Interestingly, the TMA and TSL reflected a slight decrease with 5 nM sunitinib (TMA Ctrl/Sunitinib = 1.20 NS) (Fig. 7C,D). This is consistent since mesh fusion did not increase the TMA whereas it decreased the TSL.
Figure 5. HUVEC network analysis in ETFA. (A) Initial image enlargement of the HUVEC network in culture. (B) Binary image after segmentation ($6$–$8$). (C) Skeleton of the binary segmentation ($9$). (D) Detection of Extremities (dark dots), Branches (green), Segments (magenta) and Junctions (blue). Twigs and Isolated Elements that were too small to be kept in the data are in cyan. (E) Map of the Meshes consisting in closed areas surrounded by Segments. (F) Summary of the detected and vectorized objects, superimposed to the initial image. "$\S$" labels correspond to steps described in the Results section. Scale bar: 75 µm.

Table 2. Summary of the measurement definitions performed from the vectorial objects detection.

| Parameter                  | Abbreviation | Short definition                                      | Text |
|----------------------------|--------------|-------------------------------------------------------|------|
| Mean Mesh Size             | MMS          | Mean value of Mesh size on each image (unit: pixel)   | Fig. 5 §12 |
| Total Mesh Area            | TMA          | Sum of Mesh area on each image (unit: pixel)           | Fig. 5 §12 |
| Total Segment Length       | TSL, TSL/S   | Sum of Segment length encountered in each image (ETFA) or per Sphere (FBA) (unit: pixel) | Fig. 4, §11 |
| Number of Junctions        | JN, JN/S     | Number of Junction per image (ETFA) or per Sphere (FBA) | Fig. 3, §10 |
| Total Length               | TL, TL/S     | Sum of Branches and Segments length per image (ETFA) or per Sphere (FBA) (unit: pixel) | Fig. 4, §11 |
| Number of Anchorage Junction | AJN/S        | Number of Anchorage Junction per Sphere                 | Fig. 4, §11 |

Figure 8). When segment disruptions occurred in large scale, a strong and significant inhibition was observed on TMA values (Ctrl/sunitinib 25 = 24 (p < 0.0001)) (Fig. 7C). The high level of inhibition (sunitinib 25 nM) led to a dramatic alteration of the network, as reflected by significant differences in other parameters: JN Ctrl/sunitinib 25 = 1.7 (p < 0.01) and TSL Ctrl/sunitinib 25 = 5.4 (p < 0.0001).

The fibrin bead assay response. In the FBA model, each microbead consists in a little local experiment in terms of EC loading and spatial HUVEC tube growing, and can be considered as an assay by itself. Pseudo-capillary growth on micro beads presents variability, smoothed by the number of analyzed elements (about 30 beads analyzable per well, 180 for each dose in this study). Depending on treatment conditions, images revealed diverse structural aspects going from a few number of small branches anchored to a sphere (Fig. 6 Sunit 25) up to developed trees composed of several segments and branches (Fig. 6, VEGF-A 25). All parameters exhibited a biphasic response to VEGF-A and a dose-dependent inhibition with sunitinib (Fig. 7). Within the best responses in terms of treated/control ratio, Junction Number/Sphere (JN/S), Total Length/Sphere (TL/S) and
Total Segment Length/Sphere (TSL/S) VEGF-A/Ctrl ratios were respectively 2.9 ($p < 0.001$), 1.8 ($p < 0.0001$) and 2.9 ($p < 0.001$) for the maximal activation at 25 ng/ml (Fig. 7G,I,J). Among these three measurements, TSL/S and JN/S gave the best results in terms of response amplitude to the treatment, under strong or low activator concentrations. This shows the superiority of using segment and junction detection instead of the basic total length of capillaries. The Number of Anchorage Junction/Sphere (AJN/S), which are specific measurements of FBA and can be associated to the capacity to initiate sprouting, gave the smallest SEM for the maximum activation at the dose VEGF-A 10 compared to control conditions (AJN/S VEGF-A 10/Ctrl = 1.35 ($p < 0.001$)). Another measurement associating a precocious phenomenon (initiating of sprouting) with a later one revealing the tree complexity, consists in the sum of AJN/S and JN/S (Fig. 7H). Values from these measurements gave a nearly similar inhibition/activation profile than AJN/S only, with a higher control ratio (1.6 ($p < 0.001$)), although with higher SEM. It is interesting to note that the maximum response to VEGF-A occurred at the dose of 10 ng/ml considering AJN/S (as for TMA in the ETFA assay) instead of 25 ng/ml for other measurements. The FBA assay gave consistent results with the sunitinib inhibitor, with significant inhibitions ($p < 0.0001$) for the higher dose of 25 nM using the three following measures: AJN/S, TL/S and AJN/S + JN/S (Fig. 7FGH), as expected. The dose of 5 nM showed also significant differences compared to the control condition with the TL/S and the AJN/S measurements with a control ratio of 2.1 ($p < 0.001$) and 1.7 ($p < 0.0001$), respectively.

Comparison of ETFA and FBA responses. The ETFA and FBA tests are not directly statistically comparable, since respective size of culture wells involved a different number of replicates: 3 wells for ETFA (190 mm², total of 570 mm² cultured), and 8 for FBA (32 mm², total of 256 mm² cultured). Even with a higher cultured area for ETFA compared to FBA (about 2-fold more), a significant difference compared to the control was only found with one VEGF-A treatment (VEGF-A 10 ng/ml, $p < 0.05$), and with TMA parameter only using ETFA. In the
In the case of FBA, several VEGF-A doses showed significant differences ($p < 0.01$ or lower), considering all parameters (Fig. 7). As mentioned above, each sphere can be assimilated to a single test in itself, and thus, the high number of analyzed spheres (about 30 per well) resulted in high reproducibility, which is an advantage for this method.

Another difference between the ETFA and FBA methods was the amplitude of the response amongst the various treatment doses in each method. The strongest response was found with a dose of 10 ng/ml VEGF-A in ETFA experiments, with a ratio VEGF-A 10/Ctrl$=1.36$ ($p < 0.05$) when measuring TMA (Fig. 7C). The same condition using the FBA method showed ratios going from 1.35 (for AJN/S) to 2.46 (for JN/S), and 2.24, 1.57 and 1.57 for TSL/S, TL/S and AJN$+\ JN$/S, respectively. Except for the Anchorage Number (AJN/S), other parameters exhibited a higher treatment/control ratio. This can be explained as the early meshing structure of HUVEC constrained differences in amplitude in ETFA. In the case of FBA, pseudo-capillary growth from the sphere surface is not spatially limited and therefore differences in measurements were bigger than with ETFA.

It is interesting to note that VEGF-A 10 induced nearly the same maximum response regarding AJN/S (ratio over control 1.35) in the FBA assay than for TMA (ratio over control 1.36) in the ETFA assay. While AJN/S can be assimilated to a measurement of sprouting of EC, a precocious phenomenon, other FBA parameters (TL/S, TSL/S and IN/S) reflect later development processes, like segment elongation and/or pseudo-vascular bifurcations. They exhibit a maximum response for the VEGF-A dose of 25 ng/ml.

When measuring inhibitory activities, FBA proved to be more powerful and sensitive than ETFA to measure small changes or weak inhibitors.
Discussion

In this comparative analysis between ETFA and FBA methods, we extended the “Angiogenesis Analyzer” to enable FBA analysis, including sphere detection and pseudo-vessel organization. The extended “Angiogenesis Analyzer” is written in the macro language of ImageJ, free image analysis software from the National Institutes of Health. While other software offers some solutions for analysis of images acquired by fluorescence complex confocal imaging, here we focused on the analysis of images issued from phase contrast high-throughput microscopy. Although three-dimensionally distributed, the extensions of micro-capillaries preferentially developed in the horizontal plane. Thanks to a short, but sufficient depth of field, microscopy observation in phase contrast was found convenient enough to get a good sampling of field deepness containing pseudo-capillary developments (not shown). Furthermore, although more difficult to analyze, phase contrast observation method does not require any previous treatment of cells, has no phototoxicity and can therefore be used for kinetic experiments and time-lapse recording.

Although widely used for ETFA analysis (more than 150 citations), the “Angiogenesis Analyzer” has never been described in details. Here, we report the algorithm and describe the meaning of the detected and modelled structures. This may facilitate the analysis of ETFA when following published protocols, which advise the use of “Angiogenesis Analyzer”. This method has the advantage of being easy to implement. In this work, we bring out some limitation in terms of sensitivity, especially for angiogenesis inhibitors with low activities that cannot be identified.

Similarly, we demonstrate here that the “FBA algorithm” is adapted for performing precise analyses of pseudo-capillary growing on microbeads. We overrode the constraint of using phase contrast and also implemented the method in 96 well plates with automatic image acquisition. By using this new approach, we dramatically increased the repeatability and precision of angiogenesis measurements.

We observed a biphasic profile of VEGF-A treatment with the two methods, ETFA and FBA. This biphasic effect of VEGF-A has already been reported in similar in vitro angiogenesis assays, and can be explained by a dose-dependent effect of the growth factor on the expression of its cognate receptors, or by the effect of accessory molecules interfering with VEGF-A signaling. While this similar profile can be observed with both methods, the amplitude of difference between control conditions and VEGF-A treatment is far more precise with the FBA method, and moreover by using a smaller total cell culture area. In ETFA, the measured parameters are mainly followed by the regression or growing of the meshing, which is spatially limited by the network itself. This limits the amplitude of the measured difference between a control condition and an effector. In the case of FBA, the growth in the first 3 days of culture is not spatially limited. This and the high number of analyzed spheres, each of them being considered as an assay in itself, yields accuracy and repeatability superiority to that obtained with ETFA. Remarkably the maximum of biphasic VEGF curves was the same in both cases (10 ng/mL ETFA-TMA and FBA-AIN/S), although the two sets of experiments were performed in two different laboratories, with two different HUVEC and VEGF sources.

An advantage of the FBA method is that it is easily adapted to small culture wells. Indeed, the meniscus of the gel medium imposes a relatively large area of culture to get a reasonable representative EC network sample into single image acquisition on ETFA. On FBA, the small size of beads and their associated endothelial cellular extensions, allow the encompassment of this limitation. Another advantage of the FBA, is to allow the analysis
of several aspects of angiogenesis: sprouting (Anchorage Junction detection), as well as elongation and bifurcations, which are measured through parameters qualifying and quantifying segments, junctions and branches organization of pseudo-capillaries. Moreover, the use of tightly packed cells on carrier microbeads represents characteristic features of an in vivo environment in this assay. Indeed, ETFA appears more limited to an early mix of sprouting and growing capacities occurring at the precocious meshing establishment. However, both assays can be run in a relatively high throughput manner, allowing vascular studies at a genetic, molecular, or pharmacological level.

In conclusion, we demonstrated that phase contrast images of ETFA and FBA experiments could be successfully analyzed by the customized ImageJ’s “Angiogenesis Analyzer”, from small image samples, to large batches of images. We also shed light on the advantages and disadvantages of both methods; ETFA in 24 wells plates being easy to implement, although less precise than FBA in 96 well plates, the latter approach requiring higher technicality. ETFA on the other side is faster to perform, with only 24 h of cell culture instead of 4 days for FBA. The present comparison of the ETFA and FBA methods demonstrates their robustness. More generally, our approach in terms of image analysis represents a valuable and robust tool to the study of basic branching systems, related to the angiogenesis process. It can also be used as alternative applications in the analysis of neural network or of any other biological or physical systems in which the accurate measurement of branching or meshing morphology complexity is of interest. As we observed that segments, junctions and branches provide a good representation of low developed networks and trees, we also recognize that more complex microvascular architectures in highly developed biological structures may require more elaborate descriptors. Finally, using similar approaches, it may be possible to upgrade our present version of ‘Angiogenesis Analyzer’ in order to analyze in vivo microvascular structures to better characterize tumor angiogenesis.

**Methods**

**Reagents.** FBA: HUVEC and NHDF, Endothelial basal medium-2 (EBM-2), complete fibroblast growth medium (complete FGM) culture media, Supplement Mix, human fibroblast growth factor-2 (FGF2), long R insulin-like growth factor-1 (R3-IGF-1), human epidermal growth factor (EGF) were from PromoCell (Heidelberg, Germany), fetal bovine serum (FBS) from Gibco. Ascorbic acid, hyaluronate and heparin were from Sigma Aldrich, penicillin and streptomycin from Invitrogen, Cytodex 3 microcarriers beads from GE Healthcare Europe GmbH (Freiburg, Germany). Sunitinib and human vascular endothelial cell growth factor-A (VEGF-A) were from R&D Systems (Minneapolis, USA). All reagents were of culture grade.

ETFA: HUVEC cells were purchased from ABCell-Bio (Evry, France), culture medium endothelial growth medium 2-microvascular (EGM2-MV) was from Lonza (Basel, Switzerland), Matrigel from BD (Le Pont de Claix, France) and VEGF-A was from Miltenyi (Paris, France). All reagents were of culture grade.

**Cell culture.** FBA: HUVEC were cultured in EBM-2 with 1% penicillin/streptomycin supplemented with 2% Supplement Mix, thus constituting the EGM-2 as previously described. In all experiments, cells between passages 2 and 7 were used. NHDF from juvenile foreskin were cultured in complete FGM supplemented with 5µg/ml insulin and 1ng/ml hFGF2 (human FGF2). NHDF were used between passages 2 and 9. All cells were cultured in 5% CO2 at 37 °C and media were replaced every 2 days.

ETFA: HUVEC were seeded at 5 000 cells/cm² and cultured in EGM-2 MV medium. Media was changed every 2 days. Assays were performed between passages 5 and 10.

**Endothelial tube formation assay.** Network formation in the ETFA was carried out by seeding HUVEC (10⁵ cells/well) on Matrigel (250μl/well) into a 24-well plate for 24 h at 37 °C with 5% CO2. Cells were suspended in EGM2-MV medium without VEGF-A (control condition), or complemented with 5, 10, 25 or 50 ng/ml VEGF-A. Sunitinib, a multitargeted tyrosine kinase inhibitor was added (5 nM or 25 nM) to culture medium as an angiogenesis inhibitor. Five pictures per well (center of the well and four cardinal points) were taken at time 24 h using a camera Nikon D5300 associated to an inverted microscope Nikon Eclipse TS100 using a 4× objective (NA 0.13) in phase contrast mode without fixation. For statistical analyses, three wells were seeded per conditions.

**Fibrin bead assay.** FBA was carried out using HUVEC on dextran-coated Cytodex 3 microcarriers beads embedded in fibrin gel as previously described. Dry Cytodex 3 microcarrier beads were hydrated in PBS for at least 3 h at RT and autoclaved for 15 min at 115 °C. The culture medium used in the assay was EGM-2. EGM-2 corresponded to EBM-2 medium supplemented with 2% [v/v] FBS, 10 ng/ml FGF2, 5 ng/ml EGF, 0.5 ng/ml VEGF-A, 20 ng/ml R3-IGF-1, 1 μg/ml ascorbic acid, 0.2 μg/ml hydrocortisone and 22.5 μg/ml heparin. HUVEC were mixed with Cytodex 3 microcarrier beads at a cell density of 400 cells per bead in a solution of 2500 beads per ml of EGM-2 medium. Beads and HUVEC were then co-incubated in a humidified incubator at 37 °C and 5% CO2 and gently manually shaken every 20 min for 4 h to allow cell adherence to the bead surface. Beads with adherent cells were transferred to a 75 cm² tissue culture flask and were further incubated for 24 h. Beads coated on beads were then washed three times with 1 ml of EGM-2 to remove non-coated cells and were resuspended at a density of 1000 beads/ml in a solution of fibrinogen type I (2.5 mg/ml) with 0.15 U/ml of aprotinin at a pH of 7.4. In order to polymerize the fibrin in the wells, four μl of 10 units/ml of thrombin were deposited into each well of a 96-well-optical plate (BD Falcon) suitable for high throughput imagers. Then, 80 μl of the fibrinogen type I-aprotinin-HUVEC coated bead solution were delivered over the thrombin drops, gently mixed by slowly pipetting up and down and allowed to clot for 2 min at room temperature and then at 37 °C and 5% CO2 for 30 min to promote gel formation. Eighty μl of EGM-2 medium were added to each well and allowed to equilibrate with the bead-containing gels for 30 min at 37 °C and 5% CO2. Afterwards, 3700 NHDF were added and allowed to adhere to the top of the gel. After 1 h, EGM-2 was replaced with fresh medium with or without VEGF-A or sunitinib.
Medium was changed one day after and then every two days. Sprouting was apparent between days 2 and 3 and cultures were imaged at day 4. In order to quantify the sprouting pseudo-microvessel network, samples were automatically scanned with a high-throughput imager (IXM, Molecular Device) at a 4x magnification with definition of 4 quadrants (images) per well to cover the entire surface of each well. Eight wells were analyzed per condition. The image analysis described further was performed on the largest squared area contained in the lightening area of quadrant images.

**Image analysis.** Image analysis for the FBA and ETFA were performed using a program developed for the ImageJ software. This plugin is an extension of the “Angiogenesis Analyzer” for ImageJ written in the macro language of ImageJ, which is described in the Results section.

**Statistical analysis.** Data are expressed as means ± SEM as previously described. One-way ANOVA was used for multiple comparisons in experiments with one independent variable. A Dunnet’s test was used for post hoc analysis of the significant ANOVA. A difference in mean values between groups was considered to be significant when p ≤ 0.05. Error bars correspond to the number of analysed images (n = 15 for ETFA, and n = 32 for FBA) ± SEM providing from one experiment performed with each method used.

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**Author contributions**

G.C. conceived and programmed “Angiogenesis Analyzer” and its customized version. Conceived the project, performed image analysis and wrote the paper. S.B. conceived the adaptation of the FBA method to high-throughput microscopy, participated to the paper redaction, conceived and performed FBA experiments and data analysis. S.F. performed ETFA experiments and data analysis. W.R. is the author of ImageJ. He adapted overlays (vectorial objects) functions and optimized skeletonize operation for the “Angiogenesis Analyzer”. M.C. supervised the FBA experiments and contributed to writing the paper. G.U. supervised the ETFA experiments. P.A. was at the origin of the ETFA experiments at the Gly-CRRET lab, she supervised the ETFA experiments and contributed to writing the paper.

**Competing interests**

The authors declare no competing interests.

**Additional information**

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