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ABSTRACT

The physical and mechanical properties of cells modulate their behavior such proliferation rate, migration and extracellular matrix remodeling. In order to study cell behavior in a tissue-like environment in vitro, it is of utmost importance to develop biologically and physically relevant 3D cell models. Here, we characterized the physical properties of a single cell type growing in configurations of increasing complexity. From one human skin biopsy, primary dermal fibroblasts were isolated and seeded to give monolayer (2D model), spheroid (3D model poor in extracellular matrix) and tissue-engineered cell sheet (3D model rich in endogenous extracellular matrix). Living native human dermis tissue was used as a gold standard. Nanomechanical and viscoelastic properties at the cell scale were measured by Atomic Force Microscopy (AFM) while biphoton microscopy allowed collagen detection by second harmonic generation and scanning electron microscopy helped in model morphological characterization. In all models, fibroblasts presented a similar typical elongated cell shape, with a cytoskeleton well-arranged along the long axis of the cell. However, elastic moduli of the tissue-engineered cell sheet and native dermis tissue were similar and statistically lower than monolayer and spheroid models. We successfully carried out AFM force measurements on 3D models such as spheroids and tissue-engineered cell sheets, as well as on living native human tissue. We demonstrated that a tissue-engineered dermal model recapitulates the mechanical properties of human native dermal tissue unlike the classically used monolayer and spheroid models. Furthermore, we give statistical evidence to indicate a correlation between cell mechanical properties and the presence of collagens in the models studied.

Key words: 2D, 3D, AFM, cell sheet, collagen, fibroblast, tissue engineering, self-assembly, spheroid
INTRODUCTION

Over the past century, cells cultured in two dimensions (2D) on glass or in Petri dishes served as an invaluable tool in cell biology [1]. Although still widely used and useful, growing cells in flat 2D monolayers on plastic surfaces does not accurately model the in vivo state. Nowadays, it is well accepted that 3D cell culture models are more relevant to in vivo situations than 2D cell cultures [2–6]. Indeed, 3D cell models exploit more representative spatial, biological, biochemical and biophysical parameters to bridge the gap between in vitro and in vivo experiments [7–9]. In addition, the presence of extracellular matrix in 3D models is more relevant, to mimic biochemical and biomechanical properties of cells within tissues [10,11].

Tissue engineering approaches have been widely used in vitro to produce 3D tissue equivalents, especially for skin substitutes [12,13]. Most of these methods rely on the use of biomaterials to provide a 3D structure, or scaffold, in which living cells are added. For example, fibroblasts, which are the main cell type within the dermis, can be embedded in different types of gel [14]. Although easy to carry out, this method presents some limitations. Fibroblasts seeded within collagen gels remodel them, leading to the phenomenon of collagen matrix contraction [15,16], making long-term culture unfeasible. By contrast, scaffold-free tissue-engineering approaches are based on the ability of cells to develop cell-cell interactions and to produce their own endogenous extracellular matrix. Spheroids were developed in the 1980s by Sutherland et al. and are based on cell aggregation properties when cells are cultured in suspension or in a non-adhesive environment [17]. Nowadays, spheroids are widely used in vitro to investigate many aspects of cell biology, especially in cancer studies [18,19]. However, some studies showed only a weak presence of extracellular matrix within spheroids [20,21]. The self-assembly approach is an attractive tissue-engineered strategy to produce in vitro, complex 3D tissue models rich in endogenous extracellular matrix [22]. The
reconstructed tissue produced by the self-assembly technology is called a cell sheet. Thus, it becomes possible to produce reconstructed human dermal tissue, devoid of any exogenous material [23–25]. We previously demonstrated that the primary human dermal fibroblasts reproduce a physiological environment by secreting and organizing a dense, native 3D extracellular matrix rich in organized collagens [26].

In this study, we analyzed the biophysical properties of human dermal fibroblasts grown under configurations of increasing complexity, going from a 2D classical monolayer to living native tissue. For this, we used a cutting edge multiparametric atomic force microscopy (AFM) in liquid which is particularly relevant to perform, morphological and mechanical analyses on live cells and useful to study cell interface properties in various areas, such as in microbiology, pharmacology and oncology [27–29]. However, while performing AFM experiments on cell monolayers is widespread, to the best of our knowledge we were the first to succeed in measuring mechanical forces by AFM in physiologic conditions, on a spheroid model, cell sheet model and fresh native tissue. Hence, our AFM experiments represented a technical challenge. Scanning Electron Microscopy (SEM) was used to characterize structural organization and architecture of the fibroblasts within the different models. Then, second harmonic generation (SHG) was used to visualize by two-photon microscopy, the organization of fibrillar collagens within extracellular matrix [30].

Our results provide a better understanding of the biophysical organization of each model presented in the next sections.
RESULTS

Cell morphological characterization.

Living human primary dermal fibroblasts seeded in distinct growth configurations were observed by optical microscopy during AFM measurements (Fig. 1). Fibroblasts seeded in monolayers appeared elongated and well spread out (Fig. 1a). Spheroids displayed a round shape with an average diameter of 250µm (Fig. 1b). Interestingly, no spheroids were taken off during AFM measurements by the passage of the tip. No artifacts with AFM tip were observed (Supplementary Fig.S1). Fibroblasts in cell sheets were narrower than those in monolayers (Fig. 1c). They grew in an orientated manner, parallel to each other. No picture of fresh native dermis could be taken as light was unable to pass through the thick tissue.

Scanning electron microscopy gave a better definition of each model at cell scale (Fig. 2). In all models, fibroblasts were bipolar and elongated. At higher resolution (images of 10 µm x 10 µm), cytoskeleton was detectable under the cell surface in all cell models. The cell surface was smooth and no collagen fibers being observed either in monolayers (Fig. 2a) or in the spheroid model (Fig. 2b), whatever the magnification. In the cell sheet model (Fig. 2c) and native dermis tissue (Fig. 2d), cells were embedded in a complex extracellular matrix and were partially covered by large collagen fibers.

Topographic properties of fibroblasts within models of increasing complexity.

Quantitative Imaging measurements were performed by AFM to study the morphology in living conditions of various models of human dermal fibroblast. Height AFM images exhibited fibroblast cells from different growth configurations (Fig. 3). For monolayer (Fig. 3a), spheroid (Fig. 3b) and cell-sheet models (Fig. 3c), cells displayed distinct edges, a smooth surface and had a bipolar, elongated shape. Unfortunately, due to its heterogeneous
surface, no 50 μm x 50 μm pictures was acquired for the dermis tissue, so no conclusion can be reach on global fibroblast shape within this model. Interestingly, in higher resolution images (images of 10 μm x 10 μm), fibers were observed in the whole region of the cell well-arranged along the long axis of the cell, as previously shown for dermal fibroblasts [31]. Similar fibers were also observed in native dermis (Fig. 3d). These observations confirmed the typical fibroblast morphology for all the in vitro cell models studied.

**Cytoskeleton motifs revealed by elasticity measurements.**

Comparative topographic and elasticity studies were carried out by AFM to investigate the cytoskeleton of cells in a monolayer configuration, spheroid, tissue-engineered cell sheet and native dermis (Fig. 4). Quantitative data were obtained by scanning the area shown in the height AFM images (Fig. 4 a). 139478 curves were recorded, translated into elasticity maps given in Young’s Modulus values at 200 nm depth (Fig. 4b), down to 1000 nm (data not shown). In all models, repetitive parallel motifs of cytoskeleton were noted in the cross section of Young’s Modulus versus distance (Fig. 4c). Analysis by fast Fourier transform and filtering highlights the motifs (Fig. 4d). It confirmed the parallel stiff motifs with a periodicity between 0.7 and 1.2 μm. This observation suggests that the motifs visualized correspond to cytoskeleton, as previously described by AFM in dermal fibroblasts grown in a monolayer [31] and observed by fluorescent microscopy in Supplementary Fig. S2.

**Cell membrane viscoelastic properties are correlated with collagen within the model.**

For each model, the Young’s Modulus values were calculated at 500 nm and 1000 nm of indentation from all the force curves (e.g. 36 780 curves from 9 maps in 3 independent monolayers, 53 246 curves from 13 maps in 3 independent spheroids, 36 466 curves from 9 maps in 3 independent cell-sheets and 12 986 curves from 6 maps in native dermis) and
expressed as histograms (Fig. 5ab). Statistical analyses (Table 1) revealed 2 groups of Young’s Modulus, a group of low elasticity (around 50 Pa) corresponding to dermis and cell-sheet configurations and a group of higher elasticity (between 100 and 175 Pa) corresponding to spheroid and monolayer configurations. Interestingly, old monolayers grown for 4 weeks without ascorbic acid were also statistically significantly different to the cell-sheet (Supplementary Fig. S3). Multiphoton imaging was used. This non-invasive technology allows at high-resolution imaging of organized fibrillar collagen structures in living models. Monolayer and spheroid models showed a lack of SHG signal (Fig. 5cd), which confirms results observed by scanning electron microscopy (Fig. 2ab). Cell sheets and native dermis were highly positive for SHG (Fig. 5ef), meaning that organized fibrillar collagens were abundant in these models, as previously observed by SEM (Fig. 2 cd) and by histological analyses of tissue cross sections (Supplementary Fig. S4). Collagen content in each cell model was quantified with Imaris software (Supplementary Fig. S5). Thus, collagens represent 0 ± 0.0% of monolayer’s volume, 0 ± 0.0% of spheroid’s volume, 7 ± 1.4 of cell sheet’s volume and 6 ± 2% of native dermis’s volume.
DISCUSSION

Extracellular matrix mechanical properties synergize with chemical cues to drive cell behavior in culture and acts to establish and maintain tissue homeostasis in vivo [32]. Indeed, extracellular matrix was previously thought to serve only a structural and mechanical role by maintaining tissue architecture and providing a support for cell migration. However, nowadays it is recognized that it also provides biomechanical cues influencing a wide range of cell activities (adhesion, differentiation, phenotypic modulation, survival) [10]. Understanding tissue homeostasis requires now an appreciation of cell and tissue level mechanics and is an absolute necessity when designing therapies to treat cancers, chronic diseases or delayed wound healing. Thus, correlating confocal microscopy and atomic force indentation reveals that metastatic cancer cells stiffen during invasion into collagen [33]. AFM applied on sectioned mice-skin enables the connection of the tissue function to the corresponded ultrastructure and revealed tissue structures with a nanoscale resolution, especially in a wound model [34].

It is of utmost importance to develop relevant in vitro models providing biomechanical cues. By using cutting-edge advanced microscopies, we examined morphological and mechanical properties of human dermal fibroblasts grown in various cell models, compared to native human dermis tissue.

Our experiments SEM experiments revealed that the cell sheet is the closest model to human dermis, especially in terms of extracellular matrix content. We assumed that the extracellular matrix was mainly composed by fibrillar type I collagen, which is biosynthesized and remodeled by dermal fibroblasts [35] Highly organized collagen fibrils and fibers are known to provide skin its structural framework and mechanical properties [36] Interestingly, a correlation between extracellular matrix, gene expression and cell elasticity was recently described by Kim et al in human dermal fibroblast monolayers [31] They showed that a
decrease in pro-collagen I mRNA with cell aging is correlated with increased stiffness of dermal fibroblasts.

In conclusion, we investigated and proved the feasibility of carrying out AFM force measurements on 3D models such as spheroids and tissue-engineered cell sheets, as well as on living native human tissue. We demonstrated that cell mechanical properties depend on the cell growth configuration and overall on the presence of extracellular matrix. We underlined the major role of collagens on cell mechanical properties and proved that the tissue-engineered cell sheet model rich in collagens is the closest model to human native dermis. Within context, further experiments should be done to understand the impact of extracellular matrix on various cellular fates, such as proliferation, migration and differentiation but also to investigate its impact both on genes and proteins expression in skin fibroblasts. Although there are an increasing number of studies published on the relevance of 3D models compared to classical 2D monolayers, their incomplete physical, mechanical and biochemical characterization limits their full potential exploitation.

Our observations offer perspectives in wound healing and aging studies with the relevant cell sheet model. Furthermore, developing relevant 3D cell culture models in vitro both from a biological and physical point of view complies with the ethical principles of animal research (3R’s philosophy) by proposing alternative methods to the use of laboratory animals.
MATERIAL AND METHODS

Dermis native tissue and human cell isolation. Primary human dermal fibroblasts were isolated from a 3-year-old child’s foreskin purchased from Icelltis (Toulouse, France) as previously described [23]. Briefly, after overnight incubation in Dispase II enzyme (10 mg/ml, Gibco-Invitrogen, Carlsbad, USA), epidermis was carefully peeled-off, and part of the fresh native dermis was used for direct analyses by AFM while another part was used for enzymatic isolation of fibroblasts with collagenase type 1 (1 mg/ml, Abnova, Taipei, Taiwan). 5 days after primary fibroblast isolation and expansion, cells at precocious passage 1, from the same batch, were used to seed and produce at least three independent monolayers, spheroids and cell sheet models. Cells were grown in Dulbecco’s Modified Eagle’s Medium (Invitrogen), supplemented with 10% of heat-inactivated fetal calf serum, 100U/ml penicillin and 100 µg/ml streptomycin. Cells were maintained at 37°C in a humidified atmosphere containing 5% CO₂.

Monolayer. For monolayers, cells were directly seeded (3 000 cells/cm²) onto removable, gelatin-coated 1-well Lab-tek glass slide (Fisher Scientific, Illkirch, France). Gelatin-coating allows better cell attachment and spreading. Three distinct cell monolayers were analyzed 5 days after seeding. Alternatively, cells grown in monolayer were cultivated for 4 weeks in order to compare their properties with cell sheet model.

Spheroid. We took advantage of our expertise in spheroid culture to include it in this comparative cell model study. Spheroids were generated by the non-adherent technique as previously described [37,38]. Briefly, 10 000 cells/well were seeded on ultra-low attachment
96-well plates (Fisher Scientific). At least three distinct spheroids were analyzed 5 days after seeding.

**Cell sheet.** Primary fibroblasts at passage 1 were seeded (15 000 cells/cm$^2$) onto 24-well plates as for monolayer, but grown for 4 weeks in complete cell culture medium supplemented with 50 µg/mL ascorbic acid (Sigma). When stimulated with vitamin C, cells are able to produce and assemble more collagen in the extracellular space. This trick is widely used in tissue engineering to produce tissue substitute by the self-assembly approach [22]. Indeed ascorbic acid, which is a co-factor of prolyl-hydroxylase and lysyl-hydroxylase enzymes, is well known to improve collagen production, secretion and maturation [39,40]. Three distinct cell sheets were analyzed 4 weeks after seeding.

**Scanning electron microscopy.** Cells were fixed with 2% glutaraldehyde in 0.1M Sorensen phosphate buffer pH7.2. The samples were prepared by the CMEAB platform (Toulouse, France). Images were visualized with an electron microscope QuantaTM 250 FEG (FEI, USA) at an accelerating voltage of 5 kV.

**Atomic Force Microscopy.** All AFM experiments were carried out in Phosphate Buffered Saline (Sigma-Aldrich) which is compatible with cell survival for at least 30min, as previously described by Chopinet *et al* [41]. AFM acquisitions lasted less than 15 min for each sample. For monolayers, cells were directly observed on gelatin-coated Lab-tek glass slide. Spheroids were immobilized for 15 min on a glass slide coated with polyethylenimine (PEI) (Sigma-Aldrich). Cell sheet and native dermis were directly deposited on the glass slide. AFM measurements were performed with a Nanowizard III (JPK Instruments, Germany) in
Quantitative Imaging mode (QI) [42]. The cantilever (CP-PNPL NanoAndMore) spring constant was measured by the thermal tune method [43] before each experiment and found to lie in the range of 0.05N/m to 0.2N/m. A force of 10 nN was applied. The loading frequency was 100 Hz and the indentation speed 400 µm/s, as used in previous studies [44]. Cell elasticity was determined from the Young’s Modulus values which were calculated with the JPK Data processing software (JPK) from square QI images of 10 µm using the Hertz Model [45,46] and according to the equation 1. Typical force distance curved used for Young’s Modulus calculations were shown in Fig. 6.

\[
F = \frac{4}{3} \cdot \frac{E}{1-v^2} \cdot \sqrt{r} \cdot \delta^2
\]

Where F is the applied force in N, E the Young’s Modulus in Pa, \( v \) Poisson’s ratio, \( r \) the tip radius of the probe in m and \( \delta \) the indentation of the sample in m. Here, we considered AFM tip as a sphere of 10 µm of radius and a Poisson ratio of 0.5 was applied.

Elasticity maps were obtained at 200 nm depth, as previously described in the literature [47,48]. Fast Fourier Transform (FFT) calculations were carried out with the free software Gwyddion 2.41.

**Statistical analysis** The elasticity maps were obtained by calculation of the Young’s Modulus values which were deduced from each approach curve (Fig. 6). The statistical analyses between each cell model were performed from the means of elasticity maps and by using the non-parametric Mann-Whitney test with the Prism5.04 software (GraphPad).
Collagens detection by second harmonic generation (SHG). As previously described [26], 3D stacks of 425x425x60 μm were acquired using a 7MP multiphoton laser scanning microscope (Carl Zeiss, Jena, Germany), equipped with a 20x objective and coupled to a Ti-sapphire femtosecond laser, Chameleon Ultra 2 (Coherent Inc) tuned to 800 nm. Organized fibrillar collagens were detected by second harmonic generation at a wavelength of 400 nm. The acquired 3D images and collagens quantification were analyzed off-line with Imaris software (Bitplane AG). Briefly the volume occupied by the collagen was calculated by measuring the volume occupied by the isosurface of the SHG in a cube of defined volume (80,1 x 80,1 x 80 μm). Three independent samples were used for each condition.

Histological staining

Fresh cell sheets and native dermis tissues were embedded in OCT and conserved at -80°C until use. 5µm sections were realized on a cryostat. After removal of OCT in deionized water, tissue sections were stained with Masson’s trichrome according to the manufacturer’s protocol (Sigma-Aldrich). The Masson’s trichrome method stains cells red and collagens within the dermal tissue in blue [49].
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FIGURES

Figure 1. Optical pictures of human dermal fibroblast in various growth configurations.

Pictures of fibroblast monolayer (a), spheroid (b) and cell sheet (c) taken during AFM experiments.
Figure 2. Scanning Electron Microscopy (SEM) pictures of cell models of increasing complexity from cell monolayer to native dermal tissue. SEM shows the cell architecture and organization in monolayer (a), spheroid (b), cell sheet (c) and native dermis (d). Inset exhibits the presence of naked cells on monolayer and spheroid and cell partially covered by collagens fibers on cell sheet and native dermis (white arrows). Black arrows indicate the cytoskeleton fibers.
Figure 3. Atomic Force Microscopy (AFM) height images of cell models of increasing complexity. Examples of topographic images obtained by AFM on monolayer (a), spheroid (b), cell sheet (c) and native dermis (d). The dashed lines revealed the positions of individual cells.
Figure 4. Visualization of the cytoskeleton motifs by AFM measurements. (a) Height images showed the surface morphology of cells. (b) Elasticity measurements on the same area revealed corresponding motifs for a depth of 200 nm. (c) Height (full lines) and elasticity (dotted line) cross sections were taken across the dashed lines in a and b. (d) Fourier transforms were obtained from elasticity maps. The filtering of Fourier Transform (red circles) exhibited the cytoskeleton motifs. Data were obtained from representative fibroblast cells in monolayer, spheroid, tissue-engineered cell sheet and native dermis. Scale bars: 2 µm.
Figure 5. Relationship between global elasticity and collagens. Histograms of elasticity measurements at 500 nm (a) and 1000 nm (b) of deep. Examples of Second Harmonic Generation (SHG) indicating organized collagen fibers. Collagens are absent from monolayer (c) and spheroids (d). By contrast collagen was clearly detected on cell-sheet (e) and native dermis (f).
Figure 6. Examples of Force distance curved used for Young’s Modulus calculations.

The Young’s Modulus were calculated with indentation curves (blue lines) on 1000 µm from monolayers (a), spheroid (b), cell sheet (c) and dermis (d). The Hertz fitted curves (red lines) were calculated from the Hertz model (equation 1).
Table 1. Statistical analyses of Young’s Modulus revealed 2 groups of elasticity. For each model, the mean values of Young’s Modulus were calculated at 500 nm and 1000 nm from the elasticity maps and their related curves. Non-parametric statistical analyses were performed with the Mann-Whitney test. 4!/4 pairs of test were performed to compare all the conditions with each other. The level of significance is given by the P value (* for <0.05; ** for <0.01; *** for <0.001; **** for <0.0001)). Non-significant differences of Young’s Modulus were obtained between the dermis and the cell-sheet and between the spheroids and the monolayer.

<table>
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