Scanning electron microscopy preparation and analysis of the cell cortex ultrastructure

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Abstract

The cellular cortex is a 200-nm-thick actin network that lies beneath the cell membrane. It is responsible for the mechanical properties of the cell and is involved in many cellular processes, such as cell migration and interactions with the environment. To develop a clear view of the structure of this meshwork, high resolution imaging is essential, such as electron microscopy. This technique requires complex sample preparation that can lead to artifacts like shrinkage or hole formation. We present a preparation method that reduces artifacts significantly. Here, the final drying step that is typically performed by critical point drying is replaced by hexamethyldisilazane drying. We quantitatively investigated sample integrity after both preparation methods, and show that there are significant advantages of hexamethyldisilazane drying compared to critical point drying. Furthermore, automated analysis of a network is classically performed by thresholding-based software programs, which are sensitive to noise and uneven brightness of images. The here presented analysis that we have developed is based on a vectorial node algorithm. It reproduces all kinds of networks sufficiently to allow derivation of quantitative network-specific parameters, such as mesh hole size. We use this analysis to compare the network structure of cells prepared by these two drying methods, and show that hexamethyldisilazane drying leads to fewer artificial mesh holes compared to critical point drying. We thus present here a significantly improved method to quantitatively investigate the actin cortex of cells, and show that hexamethyldisilazane drying leads to more accurate imaging compared to critical point drying.

Insight Box

The highest resolution for imaging the cellular actin cortex is provided by electron microscopy. Scanning electron microscopy samples require a drying process, usually

achieved by critical point drying, which is critical for the sample integrity. We compare the structural defects in the actin cortex of hTert RPE1 cells after critical point drying and a chemical based method, namely hexamethyldisilazane drying. In order to characterize the actin network, we also developed a new vectorial based tracing software. We bring here new tool, both experimental and analytical, which will help to streamline studies of the actin

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cortex.

Introduction

The actin cell cortex assembles under the cell membrane with a typical thickness of 200 nm.

It is the main determining factor for the cellular mechanical properties. Understanding the

structure of this thin dynamic meshwork has attracted great interest in recent years [1, 2],

especially under conditions where its adaptation is crucial, such as for mitosis [3], cell

migration [4, 5] and differentiation [6].

Given the mesh size of the cortex (<100 nm) [1, 2, 7-9], its structural properties are difficult

to address with conventional light microscopy. While some recent studies have produced

more accurate representations of this network using super-resolution microscopy [10, 11] and

rapid atomic force microscopy (AFM) [8, 12], the main technique applied in such studies

remains scanning electron microscopy (SEM). It is possible to extract quantitative data on the

superstructure of the cellular cortex from electron micrographs; e.g., the mesh size of the

network, and the higher organizational structures, such as bundles of filaments and assembled

patterns [13].

Most SEM preparation protocols involve the crucial step of sample drying. The preferred

method for this step is critical point drying (CPD), which was introduced by Anderson in

1951 [14]. CPD became an important part in the preparation of biological samples for

electron microscopy. The principle of this technique relies on the transition from liquid to

vapor phase of CO₂ without any phase transition beyond the critical point [15, 16]. The

absence of a brutal phase transition induces a better preservation of thin super-structure within

the sample. However, it is a highly invasive method that can influence the shape and structure

of biological samples through temperature, pressure, and osmolarity changes, coupled with

relatively high cost due to the special equipment needed [17, 18]. As the structure of the cell

cortex is finely regulated by actin nucleators (e.g., the formins and Arp 2/3 [19, 20], actin

capping proteins [1], and motors, such as myosin), it is important to be able to reduce drying

artifacts and discriminate them from real induced changes in the cortex structures.

An alternative drying method is the use of hexamethyldisilazane (HMDS). This drying

method has a negligible influence on the sample temperature and is performed under normal

pressure conditions, and it is coupled with moderate costs due to the use of the standard

chemical HMDS [18]. In contrast to CPD, the physical background of the mode of action of

HMDS drying is poorly understood [21-24]. Empirical studies have shown that HMDS drying

leads to similar results to those of CPD under specific biological conditions [18, 21, 25].

Typically, these are studies of relatively large structures and their integrity, or the localization

of metallic particles. Consequently, HMDS drying is a widely used method for large

structures, while small structures like the actin cortex of cells are commonly investigated

using CPD [1, 2, 26]. At the cellular level, both CPD and HMDS drying are known to lead to

cell shrinkage [17, 25, 27-30].

The visualized structures need to be analyzed for quantitative measurements. The

reproducibility of structural analyses of networks has been addressed using different

approaches. The classical thresholding and segmentation approaches are widely available. For

example, plug-ins such as Diameter J are implemented in Image J, and allow correct tracing if

the threshold used reproduces the network sufficiently well, which is only the case if the

network differs strongly from the background [31, 32]. For SEM images of the cellular

cortex, the three-dimensional structures of the actin fibers lead to significant variations in the

gray value of single fibers. Therefore, algorithms based on thresholding typically

overestimate or/and underestimate the actual network.

To increase the accuracy of the analysis of our imaged networks, we developed a method for

network recognition using a vectoral tracing algorithm in combination with (starting) nodes

[33, 34]. Such methods are, for example, based on second order Gaussian derivatives of each

pixel, to follow high or low pixel intensities, in order to correctly recognize fibers correctly.

This powerful technique has already been used successfully by Sato and co-workers [35].

Here, we show that HMDS drying better preserves the structural integrity of the cell cortex

than CPD and produces fewer artifacts. Consequently, HMDS drying leads to better

reproducibility and reduces time and costs for the preparation of small-scale structures, such

as the dense actin network. Additionally, our analysis shows that HMDS drying is more

robust than the classical thresholding method, and its flexibility allows its use in many

different contexts.

Materials and methods

Cell culture

Immortalized retinal pigmented epithelium (hTERT-RPE1)cells were grown in DMEM/F12

medium supplemented with 10% fetal bovine serum (Thermo Fisher, MA, USA), 1%

Glutamax, and 1% PenStrep under 5% CO₂ at 37 °C, in cell culture flasks (Cellstar, Greiner

Bio-One, Austria). For fluorescent imaging, RPE1 cells that stably expressed mCherry

LifeAct (kind gift from Matthieu Piel, Paris) were used under the same conditions as the

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RPE1 wild-type cells.

Electron microscopy preparation prior to final drying

The electron microscopy samples were generally prepared following the protocols of Svitkina, Chugh, and others [1, 26, 36, 37]. In more detail, the cells were detached from the culture flasks using trypsin, re-suspended in growth medium, and left to adhere onto glass slides (Thermo Fisher Scientific, MA, USA) for at least 24 h. The cells were then rinsed three times with serum-free Leibovitz medium (Thermo Fisher Scientific, MA, USA). To dissolve the cell membranes and to pre-fix the cellular cortex, two extractions were performed. The first membrane extraction and pre-fixation solution was composed of 0.5% Triton X-100, 0.25% glutaraldehyde, and 10 µM phalloidin in buffer M (50 mM imidazole, pH 6.8, 50 mM KCl, 0.5 mM MgCl₂, 0.1 mM EDTA, 1 mM EGTA) and was added to the cells for 5 min. The second extraction was composed of 2% Triton X-100 in 1% CHAPS in milliQ water, and was added to the cells for 5 min, with no rinsing of the cells between the first and second extractions. This step was followed by rinsing the cells three times with buffer M before adding the fixation solution, which contained 2% glutaraldehyde (EM grade; Science Service GmbH, Germany) and 2% paraformaldehyde (EM grade: Science Service GmbH, Germany) in 100 mM sodium cacodylate (pH 7.3) overnight. Then 0.1% aqueous tannic acid was added for 20 min, without prior rinsing. After the removal of the tannic acid, the cells were washed with distilled water three times. Then 0.2% aqueous uranyl acetate (Science Service GmbH, Germany) in distilled water was added. After 20 min, the cells were rinsed three times with distilled water. In order to affect the actin cortex structure, 100 nM Latrunculin A was used for 30 min at 37 °C prior to cell pre-fixation step. Unless otherwise indicated, all chemicals were purchased from Sigma Aldrich.

Afterwards, the samples were ethanol dried using two syringe pumps (with 60-mL syringes).

One syringe added ethanol, the second removed the ethanol solutions in order to increase the

ethanol concentration continuously to avoid discrete steps. In this manner, 60 mL 50%

aqueous ethanol was added over the time course of 1 h, followed by the same procedure with

100% ethanol. In the end, two manual changes (without using the syringe pumps) with 100%

ethanol dried over a molecular sieve were used, with an incubation time of 20 min after each

exchange.

After the HMDS drying and CPD, the samples were sputtered with a layer of 5-6 nm platinum

(coater: Model 681; Gatan, USA).

Hxamethyldisilazane and critical point drying protocols

After dehydration of the samples through the ethanol series, the cells were either dried using

HMDS (>98%; Roth, Austria; for HMDS drying to 98%; >99.9%; Sigma Aldrich, Germany;

for drying at 100%), or CPD (Quorum K850; UK) (Table 1). In normal conditions, the HMDS

drying was performed at room temperature (~23 °C). In order to reproduce the temperature

ramps the samples are exposed to during CPD, HMDS drying was also tested with a

temperature shift from 4 °C to 37°C by using a fridge and an incubator. Different HMDS

protocols, involving different rates of exchanges were tested and are summarized in Table 1.

The HMDS10 protocol turned out to be the most efficient one.

Preliminary experiments have been carried out to fine control the CO₂ exchange rates.

However, this fine control did not show any significant differences in the results. With the

critical point dryer used for the experiments presented here (K850-CPD; Quorum

Technologies, UK), ethanol was exchanged for CO₂ following 10 exchange cycles (minimum

50% of the filling level of the chamber). The chamber was then flushed twice with excess

CO₂ for 20 min per flush, followed by the final temperature increase to 37 °C. This was

accompanied by a pressure increase up to 90 bar. With the temperature kept constant, the

pressure was slowly decreased to 1 bar over 40 min.

Imaging

All electron microscopy images were obtained with an environmental surface scanning

microscope (Quanta 400; FEI, USA), using high vacuum mode. For nuclei and cells surface

measurements, cells were imaged with a fluorescent microscope (Ti Eclipse; Nikon, Japan),

equipped with an incubation chamber (37 °C, 5% CO2 in air) for live cell imaging. Nuclei

were stained with 50 ng/mL Hoechst, followed by three rinses with phosphate-buffered saline.

The nuclei and cell areas, both in the case of electron and fluorescence microscopy were

analyzed by thresholding the images in combination with the particle analysis function of

ImageJ.

Atomic force microscopy

Cell height was determined by AFM (Nanowizard 3; JPK Instruments, Bruker, USA), used

with MLCT type C cantilevers (Bruker, USA). A setpoint force of 2 nN and an approach

velocity of 5 μm/s were used. Comparing the contact point on the cell with the contact point

of the substrate next to the cell of interest resulted in cell height measurements.

Image analysis by classical thresholding and used parameters

Images were obtained with a scanning electron microscope (Quanta 400; FEI, USA). The

images were analyzed using Image J [32] according to the following procedure: to compare

the threshold quality directly with the original image a threshold was set manually by

overlaying the original image with the threshold image. The threshold was chosen in such a way that small mesh holes were not discernable (diameter, <30 nm), as they were not required for the next step of the analysis. At the same time, the big mesh holes were outlined precisely (diameter, >50 nm). In the next step, the *analyze particle* tool was set to only analyze the particles with diameters ≥110 nm (calculated via the areas). According to previous studies [2], the diameters of the mesh hole size of different cell lines was assumed to be <100 nm. Consequently, a threshold of 110 nm includes the meshes of the expected diameters plus a certain standard deviation. Therefore, every mesh hole size >110 nm that was then collected by setting this diameter threshold was assumed to be an artefact that had resulted from the electron microscopy preparation procedure. Additionally, almost all mesh holes >110 nm showed broken filaments pointing to the inside of the holes. This encourages the definition of artificial mesh holes as sample preparation artifacts.

Image analysis using home-made tracing software

The aim of our algorithm is to trace fibers on a grayscale image to increase the precision compared to classical thresholding. This was realized by populating the image with a defined number of starting nodes on the fibers. A circle was taken around each node. An algorithm defined where the fibers were localized on the circle using the second order derivative of the Hessian matrix (see Supplementary Information). At this location, it set a second node that was connected to the first one. This procedure was performed for every node until the whole network was traced. In this way, each node can be connected to an arbitrary number of other nodes, which allows the branching of the traced network and the formation of closed loops.

This procedure led to several potential output parameters, such as the total length of the traced network (respectively, fibers), the branching frequency, and the mesh hole sizes of the

network. The mesh hole size is obtained by realizing closed loops without nodes insight of the

loops. In dense networks, almost all of the fibers are parts of closed loops. The traced fibers

that are not parts of the closed loops are often at the edges of such images, as the fibers

closing the loop lie outside the image.

Details of the tracing algorithm (SI)

Before the image convolution by our algorithm, the image brightness is rescaled to be in-

between 0 and 255 (greyscale). To compute the smoothed second-order derivatives of the

image brightness, the tracing algorithm works on the basis of the Hessian of the Gaussian

transform. The convolution with the three second-order derivatives of the Gaussian transform

yields to the Hessian matrix for each pixel. Using the Hessian matrix, the directional second-

order derivative for any pixel and any direction can be computed easily. Generally, negative

second-order derivatives with large absolute values indicate the presence of a fiber. This fiber

travels perpendicular to the direction of this directional second-order derivative. An automatic

function sets several starting nodes. From these nodes, the algorithm traces the whole network

without multiple tracing of the fibers. The algorithm itself is available here:

https://github.com/SRaent/Actin.

Results

HMDS drying versus CPD

To compare the final drying methods during the electron microscopy preparation, cells were

prepared as described above. Since the preparation methods prior to drying are similar, the

first steps were done such as no sample was attributed a drying method a priori. During CPD

the cells undergo a temperature change from room temperature down to <10 °C, and then up to 37 °C or higher. To find out if this temperature shift influenced the preparation artefacts, the HMDS drying was performed both at room temperature (~23 °C) and with temperature changes analogous to the CPD, as 4 °C to 37 °C. Representative images are shown in Figure 1 at different magnifications (low, medium and high magnification being respresented on the top, middle and bottom line respectively). In the medium magnification images (Fig. 1, middle panels), it is already possible to see that there are fewer 'cracks' (holes) around the nucleus of the standard HMDS dried sample (i.e., at room temperature), compared to HMDS drying at 4/37 °C and CPD. The highest magnification images (Fig. 1, bottom panels; edge length, 3.14 µm) show the periphery area (i.e., between the nucleus and the edge of the cell). Eye inspection of such representative images, as well as thorough quantification as depicted in Fig 2(b) and (c), indicated that CPD resulted in more and larger artificial mesh holes (AMHs; defined below) than HMDS drying, both with and without the temperature shift. Nonetheless, for all of these preparation protocols, there were cells that were in good conditions (i.e., very few holes, if at all) and in bad condition (i.e., with many holes). Therefore, we carried out quantitative analysis of randomly chosen cells.

Artificial mesh holes were defined as holes >110 nm in diameter, as proposed by Chugh and co-workers [2]. The diameter of the mesh holes was derived from the measured perimeter of each hole assuming circular mesh holes for the diameter calculation. To quantify these AMHs, classical thresholding was used, as well as our algorithm. As both methods provided similar data for AMHs >110 nm in diameter, only data from our algorithm are shown in Fig 2. In this method, any mesh holes <110 nm in diameter were not taken into account. A representative image of AMHs of larger diameter is given in Figure 2a for the HMDS drying. The analysis of the diameters and AMH area fractions for all three conditions (HMDS,

HMDS 4/37 °C, CPD) are shown in Figure 2b, c. There are no significant differences between

the AMH diameters for HMDS drying at room temperature and at 4/37 °C, which indicates

that temperature has little influence at this preparation step. The AMH diameter for HMDS

ranged from 300 nm to 500 nm. In contrast, the AMH diameter distribution for CPD lied

between 400 nm and 800 nm and was statistically different to the one obtained for HMDS

drying method (p <0.001). At the same time, the area fraction of the AMHs, in terms of the

proportion of AMH area compared to the total image area, was not significantly different for

HMDS and HMDS 4/37 °C, averaging between 2% to 25% for both conditions. In contrast,

the area fraction of the AMHs for CPD was significantly higher (p <0.001), representing a

fraction between 10% and 50% of the image area (Fig. 2c). Of note, the temperature decrease

as a result of the HMDS evaporation at room temperature was measured to be less than 2 °C

(data not shown). Since we did not show an influence of the temperature cycles during HMDS

drying at 4/37 °C, these data lead to the conclusion that the AMHs cannot be solely attributed

to the temperature cycles during CPD.

Moreover, the exchange concentration and the rates from 100% ethanol to 100% HMDS

significantly influenced the AMH diameter measurements. There were no differences in the

AMH diameters between the HMDS1 and HMDS10 protocols (see Table 1), as shown in

Figure 2d. In contrast, the HMDS25 and HMDS50 protocols (see Table 1) lead to

significantly larger AMHs than for the HMDS1 and HMDS10 protocols. Due to

practicability, we decided to use the HMDS10 protocol for all of the following presented

experiments.

Correlation between artificial mesh holes around the nuclei and at the cell periphery

Artificial mesh holes close to the nuclei can be very large compared to AMHs at the cell periphery. The AMH sizes within a 2 µm large ring around the nuclei were compared to AMHs at the periphery (defined as the area between the outer part of the 2µm ring and the cell edge, see Figure 3b). Figure 3a shows the size distribution of AMHs >110 nm around the nucleus for both HMDS drying and CPD. AMHs at the periphery and AMHs close to the nuclei are clearly correlated (Fig. 3c, d). For both HMDS drying and CPD, the total and the mean size of the AMHs at the periphery increased with the AMH size around the nuclei. This correlation between AMHs around the nuclei and at the periphery is eventually helpful to rapidly define the quality of SEM-samples. Rapid imaging of the perinuclear region (2µm ring) is enough to determine whether a user should spend time on one particular sample to focus on and to optimize the parameters for the imaging at the periphery.

Origin of artificial mesh holes around nuclei

In most cases AMHs were seen around the nuclei, as it was already shown in other studies [1, 2, 26, 36, 37]. However, the origin of these AMHs has, to the best of our knowledge, not been investigated to date. As these AMHs are around the nuclei, our hypothesis was that these AMHs appear during strong shrinkage of the nuclei compared to the rest of the cell. Therefore, we measured the nuclei and cell sizes for living and fixed cells by fluorescent microscopy, as well as after the final drying with either HMDS drying or CPD by electron microscopy. Additionally, to determine whether the cell shrinkage also affects the height of the cells, cell height above the nucleus and at the cell periphery was probed by AFM. The sole fixation of cells with 2% paraformaldehyde and 2% glutaraldehyde in 0.1M cacodylate buffer had no influence on the cell area, as shown in Figure 4a. In contrast, the nuclear area was significantly reduced after HMDS drying and CPD compared to living and fixed cells. We also observed that the nuclear area was significantly larger after HMDS drying, compared to

cells which had been dried by CPD (Fig. 4b). Consequently, we concluded that the AMHs

around the nuclei are caused by significantly greater shrinkage of the nuclei compared to the

periphery of the cell. The smaller shrinkage of the nuclei induced by the HMDS procedure

explains why samples dried with this protocol have, on average, fewer and smaller AMHs. As

such, HMDS drying is less invasive and alters the cellular morphology less than CPD.

However, both methods resulted in significant shrinkage in cell height during the fixation

step. Additionally, decrease in cell height was also measured during the final drying step for

both of the drying methods (Fig. 4c, d). While the nuclear height was reduced more with

HMDS drying, the periphery height showed no significant reductions in height comparing

HMDS drying and CPD (Fig. 4c, d). This leads to the conclusion that the significant

shrinkage of the nuclei but no shrinkage of the cell parallel to the surface is the main driver

for AMHs at the cell periphery.

The fixation and drying influence on the cells is illustrated in Fig. 4e.

New algorithm for fiber detection and mesh hole size determination of the cellular actin

We developed an algorithm to trace the cortex structures in these SEM images. This tracing

identifies cortical structures more accurately than the classical thresholding method, which

leads always to overestimation or underestimation of the actual network. An exemplary

tracing for wild-type RPE1-cells is shown in Figure 5 (top row images) and compared to

Latrunculin A treated cells, a condition in which the actin polymerization is partially

inhibited, leading to bigger mesh holes (bottom row images). By analyzing all mesh holes

present in the cortex, we found a mesh hole size of 64 +/- 14 nm (mean +/- SD) for RPE1-wt

cells and 68.4 +/- 28 nm (mean +/- SD) for Latrunculin A treated cells (Fig. 5d). In addition,

the number of branches/nodes decreases when cells were treated with Latrunculin A (Fig. 5h).

To quantify the quality of our tracing algorithm, we measured the maximum density of the

fibers that are traceable within an acceptable error (10% SD). To do so, we digitally created

an orthogonal grid of white fibers on a black background. For this network we increased the

number of fibers, and thereby the density of fibers, from 26% coverage with white pixels

(74% black pixels) up to 92% with white pixels (8% black pixels), as illustrated in Fig. 6a.

The relative root square deviations were taken for the comparisons of the known mesh-hole

sizes versus the measured mesh-hole sizes. As shown in Figure 6b, the expected mesh hole

size is in good agreement with the measured mesh hole size analyzed by our algorithm. We

conclude that our algorithm allows a relatively accurate description of porous and dense

networks, with up to 92% coverage of the fibers.

Contrary to our self-generated grid, electron micrographs always display a certain level of

noise. Consequently, we investigated the maximum signal to noise ratio that our program can

handle while giving a good measurement of mesh hole sizes. Therefore, digital noise was

added to the grid created with 73% black pixels. An example of the tracing results for these

images is given in Fig. 7a. Analysis of the relative root square deviations indicated that a

signal-to-noise ratio of 0,83 dB is still traceable, within an error of below 10% (Fig. 7b). As

comparison the signal-to-noise ratio of the electron microscopy images of the cortex are in the

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range between 5 and 15 dB.

Variety of applications of the tracing algorithm

While we developed our algorithm in order to quantify actin cortex meshes from electron micrographs, it still can be adapted to the recognition of different fiber-like structure imaged by different techniques. To demonstrate this, several images of random objects that included fiber-like networks were taken with a camera at different magnification and analyzed with our algorithm: spaghetti, a three-dimensional tunnel painted on a wall, the structure of leather, a line drawing of a person, human leg hairs, and handwriting (Fig. 8). These examples demonstrate that this software can be used for analysis of a wide variety of network images, which proofes it as a powerful tool for all network quantification across many applications.

Discussion

Little is known about the precise origin of AMHs. There are several explanations possible that can define this process for the artefacts observed. One explanation indicates that this might arise from water or ethanol remaining within the sample before the final drying process. During CPD neither water nor ethanol traces would reach their critical point (374 °C at 221 bar for water and 241 °C, at 60 bar for CO2), thus the transition from liquid to gas of either compound would be accompanied with a sudden change of density leading to potential fractures of the thin structures of the sample. In the case of HMDS drying, the HMDS itself combines both the low surface tension of ethanol (18mN/m for HMDS and 22 mN/m for ethanol) and the low vapor pressure of water (23 mmHg for HMDS; 17,5 mmHg for water). These properties translate into a lower rate of drying and the liquid-gas transition induces less mechanical stress on the microstructures. The potential crosslinking properties of HMDS towards biological sample have also been suggested as a reason of observing less shrinkage in biological tissues during drying [24]. Nonetheless, HMDS drying does not involve critical

point, contrary to CPD, and thus the liquid to gas transition will always be accompanied by

changes in densities.

In addition, local shrinkage of samples was shown during HMDS drying and CPD up to 30%

for large scales, such as in tissues [14, 15, 18]. This shrinkage will always lead to local

tensions in the sample, which can potentially affect microstructures such as the actin cortex.

A further effect might arise from the exchanges of solutions during the preparation. The

osmolality of these solutions can vary from 10 to 50 mOsmol/L (for MilliQ water) to over 300

mOsmol/L (for medium, Leibovitz, first extraction solution), and up to 700 mOsmol/L (for

the fixation solution). We aimed to minimize these effects by the successive and slow

addition of each solution in order to reduce the osmotic shock. While this effect was reduced,

it can still not be completely excluded.

Finally, CPD has two additional aspects that are not part of HMDS drying. First, the pressure

during CPD varies between 1 bar to 80 bar, or even higher. In principle, this should not have a

significant influence on the sample, as long as there are no air-filled cavities. While we

haven't noticed such cavities, one cannot fully discard the potential presence of nanometer

size air-pocket. The compression of ethanol during the pressure increase up to 50 to 60 bar at

the beginning of CPD is in the range of a 0.5% volume compression, and as such, the impact

on the sample should be neglectable. However, the degassing of CO₂ to reduce the pressure to

atmospheric pressure is a particularly delicate step. If this happens too quickly, the gas can

expand and potentially rupture the ultrastructure, which will lead to AMHs. The second

aspect is the wide temperature change during CPD, from ~5 °C to 40 °C, while HMDS drying

is carried out at room temperature, and only shows a temperature decrease of 2 °C below

room temperature at maximum. However, simulation of the temperature difference of CPD

for HMDS drying showed that the temperature had no significant effects on AMH sizes.

All in all, while it would be mostly impossible to avoid AMH in HMDS drying due to local

surface tension, the kinetic of the reaction makes those AMH smaller and fewer in number

and sample preparation is less sensitive to traces of water or ethanol as well as the rate of

sample degassing.

Conclusions

To examine the intact cellular cortex using SEM, the preparation follows four essential steps:

membrane removal; fixation; dehydration; and drying. This process often leads to artifacts,

and these appear to arise during the final drying process, which is commonly performed as

CPD [1, 2, 26, 36, 37]. HMDS drying represents an improved alternative to CPD. We have

shown here that both of these drying methods lead to artifacts, but that for HMDS drying, the

AMHs are significantly smaller than for CPD.

Moreover, we have shown that the sizes of the AMHs around the nuclei and at the periphery

of cells are correlated. Our explanation for this correlation relates to the shrinkage of the

nuclei in three dimensions, coupled with no change in the size of the outer edge of the cell

during the preparation. We postulate that this collapse of the cellular cortex around the nuclei

will lead to increased tension towards the periphery, and therefore AMHs appear to reduce

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this local tension.

To accurately analyze such network structures as the cellular cortex, we used a vectoral

tracing algorithm that accurately traces a wide diversity of networks. These networks can be

accurately traced for up to 92% fiber coverage and a signal-to-noise ratio of 0,83, assuming a

maximum error of 10%. The algorithm itself is limited to white fibers of approximately

uniform thickness on a dark background.

Finally, we were able to measure the average mesh hole diameter of the actin cortex of RPE1

wild-type cells and found this size to approximately be 64 nm. After actin polymerization

disturbance by using LatrunculinA, the mesh hole size of the cortex increased significantly by

almost 7% confirming that the structural properties of the cell cortex is directly linked ot the

actin dynamic. Altogether we propose here a combination of technical and analytical tools

and their limitations. These tools will be of great help to improve the rate of success as well as

the standardization of the measurement of actin cell cortex in the future.

Conflicts of Interest

The authors declare that they have no conflicts of interest regarding the publication of this

work.

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Table 1. Hexamethyldisilazane drying protocols, carried out at room temperature (~23 °C), 4 °C, and 37 °C.

Protocol	Primar	y HMDS proc	edure	Secondary HMDS procedure				
code	Step size	Incubation	Total	Wash	Wash time	Completion		
	(%)	time (min)	steps					
HMDS1	1	0.5	100	100%	20 min, ×2	Evaporation		
HMDS10	10	5	10	100%	20 min, ×2	Evaporation		
HMDS25	25	5	4	100%	20 min, ×2	Evaporation		
HMDS50	50	5	2	98%	10 min, ×1	Evaporation		

Figure legends

Figure 1: Scanning electron microscopy images of hmds drying and CPD. Representative

scanning electron microscopy images for three different conditions for the final preparation

step: HMDS dried (left column), HMDS dried at 4/37 °C (middle column), and critical point

dried (right column). Top: Whole cell images, indicating (blue boxes) the sensitive area

around the nucleus magnified in the middle images. Bottom: magnification of the periphery

(area between nucleus and cell edge) Scale bars: 20 µm (top row); 5 µm (middle row); 1 µm

(bottom row).

Figure 2: AMH analysis upon different preparation procedures. (a) Analysis of mesh

holes overlaid on an original image. The sample was HMDS dried at room temperature (see

Fig. 1, bottom left). Scale bar: 1 µm. (b) Mean diameter of artificial mesh holes (AMH)

calculated via the perimeters for around 30 images (i.e., 30 cells). (c) Total area fraction of the

analyzed areas based on whole image size. (d) AMH diameters for different HMDS samples

(see Table 1). * p <0.05, ** p <0.01, *** p <0.001, **** p <0.0001, n.s. not significant (t-

tests).

Figure 3: Comparison of AMH close to the nucleus and in periphery. (a) Representative

image of a cell nucleus (outlined in green) of a cell prepared by CPD. The surrounding

artificial mesh holes are indicated in red, to a 2 µm distance from the edge of the nucleus.

Scale bar: 10 µm. (b) Representative image of a cell prepared by hmds drying. Green outline

refers to the nucleus, red outline refers to the 2 µm distance from the edge of the nucleus. The

area defined as periphery lays between the red and black outline. Scale bar: 10 µm (c, d)

Comparisons of total AMH areas around the nuclei (e.g., red outlines in (a)) with the total and

averaged areas of the cell periphery (area between red and black outline in (b)). Lines, linear

approximations. Squares, CPD; circles, hmds10.

Figure 4: Influence of fixation and final drying. Geometry changes of the cell during

preparation for scanning electron microscopy. (a, b) Measurements from scanning electron

microscopy of cell areas (a) and nucleus areas (b) by fluorescent microscopy of these living

and fixed cells. (c, e) Measurements from atomic force microscopy of the nucleus heights (c)

and cell periphery heights (d). (e) Illustration of the four conditions investigated (i.e., living

and fixed cells, and after HMDS drying and CPD). * p <0.05, *** p <0.001, n.s. not

significant (t-tests).

Figure 5: Mesh hole size analysis. Representative images of the cellular cortex of adhered

wild-type RPE1-cells (a) and adhered RPE1-cells treated with 100 nM Latrunculin A for 30

min (e) with networks traced using our tracing algorithm (b, f) and the magnification of the

blue rectangle in (c, g). The analysis of the mesh hole areas below 110 nm in diameter (d) and

the analysis of the number of junctions/nodes per fiber length (h) are compared for the wild-

type RPE1-cells (control) and Latrunculin A treated cells. Red tracing lines, closed loops;

green tracing lines, open loops. Scale bars: 1 µm for (a, b, d, e); 100 nm for (c, f).

Figure 6: Accuracy of the tracing algorithm. (a, b) Created quadratic black mesh holes

with increasing mesh hole size (as indicated) as designed (a) and traced using our software in

(b). (c) Known mesh hole size versus the mesh hole size calculated via the tracing algorithm

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with a linear approximation.

Figure 7: Influence of noise on tracing accuracy. (a) Artificial mesh holes (73% black

pixels) as designed with noise addition (left column) and as traced using our algorithm (right

column), with signal-to-noise ratios as indicated. Red tracing lines, closed loops; blue tracing

lines, open loops. (b) The measured mesh holes shown in (a) were compared without and with

noise additions, assuming no tracing as 100% standard deviation, to provide relationship

shown between normalized standard deviation and signal-to-noise ratio. At 0,83 dB and

higher, the normalized standard deviation is <10%, so a signal-to-noise ratio ≥0,83 dB is

recommended for using our algorithm.

Figure 8: General examples of network tracing with our algorithm. Images of examples

of tracing of different kinds of networks using our algorithm. All images were color modified

for bright 'fibers' and dark backgrounds. Left, original image; right, traced image. The images

shown and traced are a dense network of spaghetti (a), a porous network of spaghetti (b), a

three-dimensional tunnel painted on a wall (c), the structure of leather (d), a line drawing of a

person (e), human leg hairs (f), and handwriting (g). Red tracing lines, closed loops, blue

tracing lines, open loops.

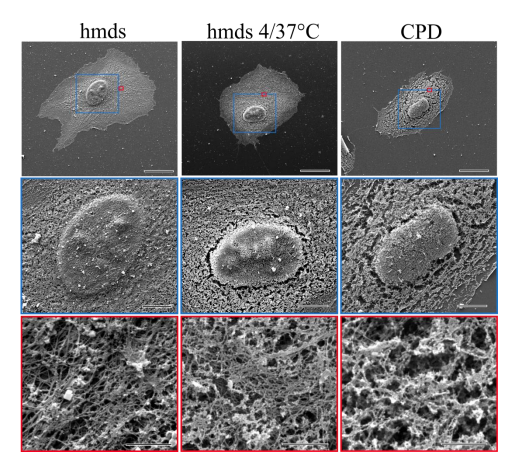


Figure 1

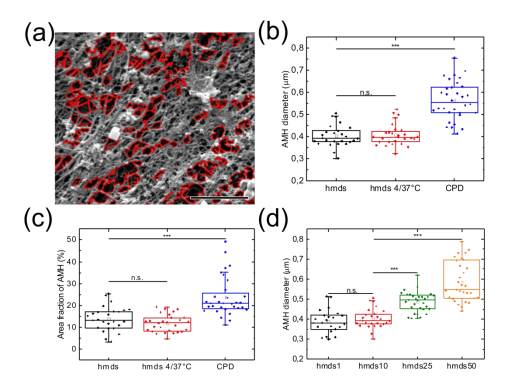


Figure 2

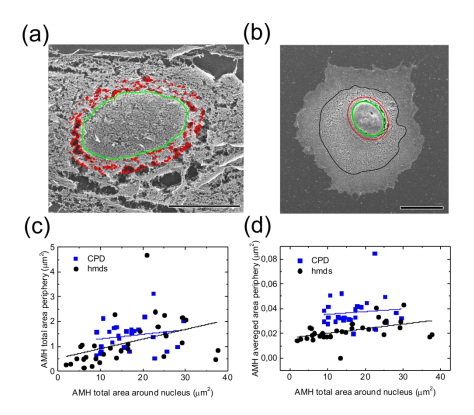


Figure 3

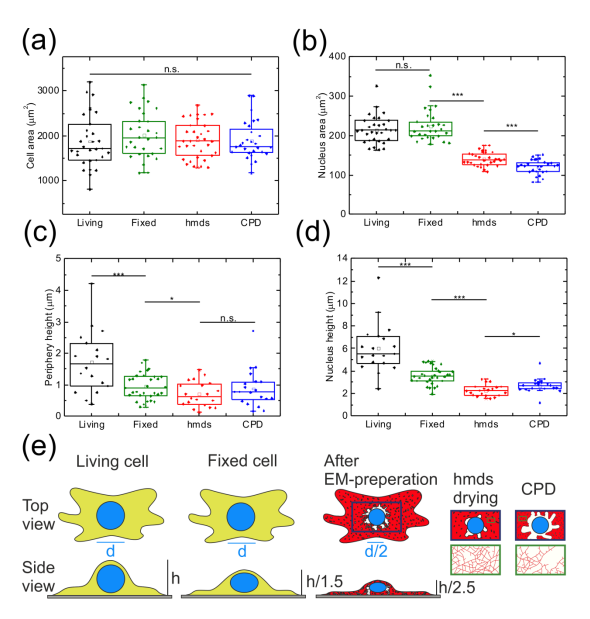


Figure 4

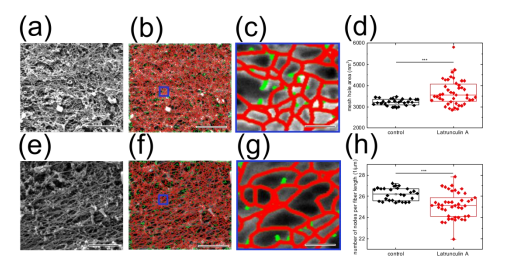


Figure 5

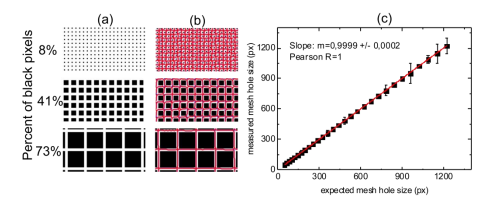


Figure 6

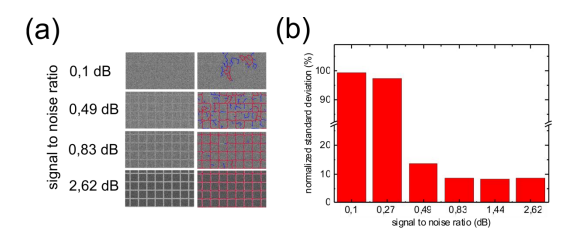


Figure 7

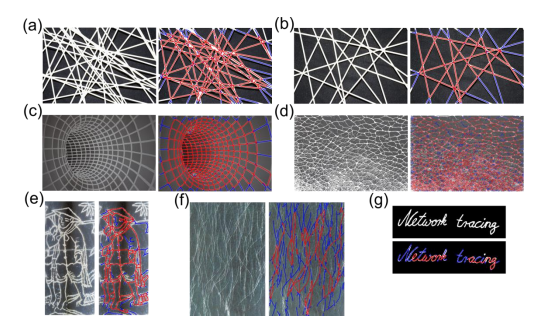


Figure 8