EFFECT OF ERUFOSINE ON MEMBRANE LIPID ORDER IN BREAST CANCER CELL MODELS

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ABSTRACT

Alkylphospholipids are a novel class of antineoplastic drugs showing remarkable therapeutic potential. Among them, Erufosine (EPC3) is a promising drug for the treatment of several types of tumors which has not been sufficiently characterized. While EPC3 is supposed to exert its function by interacting with lipid membranes, the exact molecular mechanisms involved are not known yet. In this work, we applied a combination of several fluorescence microscopy techniques (i.e., scanning fluorescence correlation spectroscopy, line-scan fluorescence correlation spectroscopy and generalized polarization imaging) to quantify the effect of EPC3 in biophysical models of the plasma membrane, as well as in cancer cell lines. Our results indicate that EPC3 affects lipid-lipid interactions in cellular membranes by decreasing lipid packing and increasing membrane disorder and fluidity. As a consequence of these alterations in the lateral organization of lipid bilayers, the diffusive dynamics of membrane proteins are also significantly increased. Taken together, these findings suggest that the mechanism of action of EPC3 might be linked to its effects on fundamental biophysical properties of lipid membranes in cancer cells.
INTRODUCTION

Erufosine (EPC3) is a novel derivative of erucylphosphocholine that belongs to a group of antineoplastic drugs based on alkyl ether lipids (1). EPC3 can effectively be applied intravenously, can cross the blood-brain barrier and shows antitumor activity in the μM range (1, 2). For these reasons, EPC3 is a promising drug for treatment of several types of tumors, including human urinary-bladder carcinoma, breast carcinoma, glioblastoma and multiple myeloma (2). Due to its hydrophobic nature, this molecule is supposed to interact with cellular membranes, but no detailed information regarding its molecular mechanism of action is available. On the other hand, other alkylphospholipids (APL) have been characterized more in detail. For example, it was shown that Edelfosine – one of the first characterized APLs – induces apoptosis in cancer cells via interactions with lipid rafts (1), i.e. lipid-protein domains of the plasma membrane (PM) which are enriched in sphingolipids and cholesterol (3-5) and are involved in several cellular functions (see e.g. (6, 7)). Such domains can also be characterized in protein-free model membrane systems (e.g. lipid vesicles) constituted of typical PM lipids (e.g. saturated sphingomyelin (SM), unsaturated phosphatidylcholine (PC) and cholesterol). It was shown in fact that these model membranes display a phase separation into a liquid-ordered (L_ο) and a liquid-disordered (L_d) phase (8). The L_ο phase is enriched in saturated lipids and cholesterol and, therefore, provides a simple physical model to study raft-like domains (9, 10). In this context, it was shown that APLs partition into lipid bilayers and directly interact with L_ο domains (11). The observed effects vary depending on lipid composition and the investigated APL (e.g. Edelfosine or Miltefosine) and include: disorganization of L_ο domains (11), moderate to significant increase of membrane fluidity (12, 13) and stabilization of SM/cholesterol domains (14). More specifically, studies on Erucylphosphocholine (which is more similar to EPC3, due to the shared unsaturated acyl chain) indicate that this APL increases the fluidity of both cellular and model membranes (12), while weakening SM-cholesterol interactions (15).

So far, studies regarding the molecular mechanisms connected to the antitumor activity of EPC3 have been limited. Recently, using spectroscopic ellipsometry as a novel technique to study solid-supported lipid model systems, we have shown that treatment of lipid films composed of PC, SM and cholesterol with EPC3 induces an increase in monolayer thickness (16).
In this work, we employed several methods based of fluorescence microscopy to investigate for the first time how EPC3 affect the physical properties of the PM.

First, we have applied line-scan fluorescence correlation spectroscopy (lsFCS) (17, 18) and scanning fluorescence correlation spectroscopy (sFCS) (19-21) to monitor the dynamics of membrane lipids and proteins, both in simple artificial membranes and directly in the PM of living cells. These methods belong to the family of fluorescence fluctuation techniques and were effectively used in the past to quantify membrane dynamics (and, indirectly, membrane order) (22). Furthermore, the use of supported lipid bilayers (SLBs) as membrane models allows the study of lipid-lipid interactions in a controlled environment and in specific relation to the precise lipid composition of the bilayer (23, 24).

Second, we quantified the influence of EPC3 on the membrane order for the PM of cancer cell models by using polarity-sensitive fluorescent probes (25). More in detail, we have investigated the spectral properties of two membrane probes (Laurdan and Di-4-ANEPPDHQ) which were shown to be influenced by lipid packing, membrane hydration and composition to different extents (25, 26).

Our results indicate that EPC3 affects lipid-lipid interactions both in lipid membrane models and in the PM of cancer cells. Furthermore, such alterations in membrane order appear to have a direct effect on membrane protein dynamics.
MATERIALS AND METHODS

Chemicals

Erufosine (EPC3) was synthesized in the Max Planck Institute for Biophysical Chemistry, Göttingen, Germany, and was most graciously provided by Prof. Martin R. Berger. It was dissolved in a PBS (phosphate buffer saline, pH 7.4) in 10 mM stock solution and kept at 4°C. Di-4-ANEPPDHQ and Laurdan (6-dodecanyl-2-dimethylaminonaphtalene) were from Molecular Probes (Eugene, OR). L-α-phosphatidylcholine chicken egg (EggPC), cholesterol from ovine wool (Chol), sphingomyelin from porcine brain (bSM), 23-(dipyrrometheneboron difluoride)-24-norcholesterol (TF-Chol) and 1,2-dioleoyl-sn-glycero-3-phosphoethanolamine-N-(lissamine rhodamine B sulfonyl) (Rhod–DOPE) were purchased from Avanti Polar Lipids, Inc. (Alabaster, AL).

Cell culture

MDA-MB-231 highly invasive breast cancer cells and MCF-7 epithelial cancer cells were acquired from the American Type Culture Collection (ATCC, USA). Both cell lines are derived from breast adenocarcinoma, with MCF-7 cells retaining some characteristics of the differentiated mammary epithelium. All cell lines were incubated in phenol red-free DMEM culture medium with 10% fetal bovine serum, 2 mM L-Glutamine, and 100 U/mL penicillin and 100 µg/mL streptomycin at 37 °C and 5% CO2. Cells were passaged every 3–5 days, no more than 15 times. All solutions, buffers and media used for cell culture were purchased from PAN-Biotech (Aidenbach, Germany).

Supported lipid bilayers

Supported lipid bilayers (SLBs) were prepared as previously described (27, 28). Briefly, lipids were mixed in organic solvent at the desired concentrations and dried on the walls of a glass vial. The lipid film was then rehydrated in PBS pH 7.4 and, after vigorous vortexing, sonicated to clarity in a bath sonicator (model). Typical concentrations during sonication were ~5-10 µM. The thus obtained small unilamellar vesicle suspension was then diluted ca. 10-fold and 100 µL were deposited on a small thin piece (~10 mm²) of freshly-cleaved mica glued to the surface of a glass coverslip (thickness #1). The mica and the vesicles suspension were confined using a 7 mm-plastic cylinder, also glued to the glass surface. Vesicle fusion
and bilayer formation were induced by addition of 3 mM CaCl$_2$. The volume was adjusted to 300 μL and the suspension was incubated for 10 min. Unfused vesicles were removed by addition and removal of 500 μL DPBS, performed 10 times. The treatment with EPC3 was performed by adding the drug at the desired concentration in PBS directly on top of the SLB and waiting ca. 30 min before performing fluorescence measurements.

**Plasmids**

The plasmid coding for human EGFR tagged with EGFP (hEGFR-EGFP) was a kind gift from Alexander Sorkin (Addgene plasmid #32751) (29). The hemagglutinin (HA) gene from influenza virus strain A/FPV/Rostock/34 (H7N1) tagged with mEGFP at the C-terminus (FPV-HA-mEGFP) was cloned based on the previously described FPV-HA-mEYFP (gift from Andreas Herrmann, Humboldt University Berlin) (30). Briefly, FPV-HA-mEYFP was digested using BglII and SacII (New England Biolabs GmbH, Ipswich, MA), and the obtained HA-insert was ligated into mEGFP-N1. This plasmid is now available on Addgene (#127810).

**Confocal microscopy imaging using di-4-ANEPPDQ and Laurdan**

The cells were seeded in 35-mm microscopy dishes (CellVis, Mountain View, CA) with an optical glass bottom (#1.5 glass, 0.16 - 0.19 mm) at a density of 5x10$^4$ cells/well for MDA-MB 231 and 1x10$^5$ cells/well for MCF-7. After a ~12 h incubation, MDA-MB-231 cells were treated with a 20 (or 30) μM EPC3 solution in culture medium, corresponding to the previously measured IC50 and IC75 values (31). MCF-7 cells were treated with a 40 (or 60) μM EPC3 solution (corresponding to IC50 and IC75 values, Tzoneva R., unpublished data) in culture medium. In both cases, the treatment lasted for 24, 48 or 72 h. Control cells were incubated just with culture medium, following the same protocols. After the incubation period, the cell medium was removed. The cells were then washed twice with PBS, pH 7.4. Afterwards, 2 ml of serum-free and phenol red-free DMEM and fluorescent dye (with final concentration of 1 μM for di-4-ANEPPHQ or 5 μM for Laurdan) were added to the cell dish. The cells were further incubated for 30 min at 37 °C.

A Zeiss LSM 780 system (Carl Zeiss, Oberkochen, Germany) was used to acquire the confocal images, with a pixel size of ca. 200 nm (512 x 512 pixels). Samples were imaged using a Plan-Apochromat 40x/1.2 Korr DIC M27 water immersion objective. The excitation sources were a 488 nm argon laser (for di-4-ANEPPDQ) or a 405 nm diode laser (for
Laurdan). Fluorescence was detected between 498-579 nm (Channel 1) and 620-750 nm (Channel 2) for di-4-ANEPPDQ, after passing through a 488 nm dichroic mirror, using a gallium arsenide phosphide (GaAsP) detector. Fluorescence of Laurdan was detected in the spectral ranges 410-463 nm (Channel 1) and 472-543 nm (Channel 2), after passing through a 405/565 nm dichroic mirror. Out-of-plane fluorescence was reduced by using a 42.4-μm pinhole in front of the detector.

**Image processing**

Confocal images were analyzed as previously described (25, 26). Briefly, for each experimental condition, ca. 5-10 confocal images of treated cells were acquired. In each image, several regions of interest (ROIs) were selected in correspondence of the plasma membrane of different cells. Experiments were performed as independent duplicates on different days so that, for each EPC3 concentration and each time point, at least 50 cells were analyzed in total. All the pixels from ROIs collected within equivalent samples (i.e. same EPC3 concentration and same time point) were pooled together. For each pixel, the GP value was calculated as defined in Refs. (26), using a custom-written Matlab (The MathWorks, Natick, MA) script and setting G=1. The obtained results are shown as normalized occurrence histograms of all selected pixels.

**Scanning Fluorescence Correlation Spectroscopy (sFCS)**

For one-color sFCS experiments, $8 \times 10^4$ MDA-MB-231 cells were seeded in 35-mm dishes (CellVis, Mountain View, CA) with optical glass bottoms (#1.5 thickness, 0.16 - 0.19 mm). After 24 h, cells were treated with medium containing 30 μM EPC3. After an additional incubation for 24 h, cells were transfected by using 200 ng (mp-mEGFP and mp-mEGFP(2x)) or 600 ng (hEGFR-EGFP or A/FPV-HA-mEGFP) plasmid DNA per dish with Lipofectamine™ 3000 according to the manufacturer’s instructions (Thermo Fisher Scientific). Plasmids were incubated for 15 min with 2 μl P3000 per μg plasmid and 2 μl Lipofectamine™ 3000 diluted in 50 μl serum-free medium, and then added dropwise to the cells.

After 24 h incubation, scanning fluorescence correlation spectroscopy (sFCS) measurements were carried out on a Zeiss LSM 780 system, equipped with a Plan-Achromat 40x/1.2 Korr DIC M27 water immersion objective. Samples were excited with a 488 nm argon laser and the fluorescence was detected between 499 and 597 nm, after passing through a 488 nm
dichroic mirror, using a GaAsP detector. To decrease out-of-focus light, a pinhole size of one airy unit (~ 39 µm) was used. To keep photobleaching below 20%, a laser power of 1.2 µW was chosen. For cell measurements, a line-scan of 256x1 pixel (pixel size 80 nm) was performed perpendicular to (i.e. across) the plasma membrane with a 403.20 µs scan time (0.67 µs pixel dwell time). For each measurement, 400000 lines were acquired in photon counting mode, and the total scan time was around 3 min per measurement. Scanning data were exported as TIFF files. At the beginning of each measurement day, the signal was optimized by adjusting the collar ring of the objective to the maximal count rate for an Alexa Fluor® 488 (AF488, Thermo Fischer, Waltham, MA) solution (50 µM dissolved in water) excited at the same laser power. For the focal volume calibration, a series of point FCS measurements was performed (ten measurements at different locations, each consisting of 15 repetitions of 10 s), and the data were fitted with a three-dimensional model including a triplet contribution. The structure parameter was typically around 6 to 9, and the diffusion time around 35 to 40 µs. The waist $\omega_0$ was calculated from the measured average diffusion time ($\tau_{d,AF488}$) and previously determined diffusion coefficient of the used dye at room temperature ($D_{AF488} = 435 \mu m^2 s^{-1}$) (32), according to the following equation:

$$\omega_0 = \sqrt{4\tau_{d,AF488}D_{AF488}}$$

Typical values were 200-250 nm. All measurements were performed at room temperature.

sFCS analysis follows the procedure described previously (19, 21, 33). Briefly, the TIFF files were imported and analyzed in Matlab using a custom-written code. All scanning lines were aligned as kymographs and divided in blocks of 1000 lines. In each block, lines were summed up column-wise and the x position with maximum fluorescence was determined, by fitting with a Gaussian function. This algorithm finds the position of the plasma membrane in each block and is used to align all lines to a common origin. The pixels corresponding to the membrane were defined as pixels which are within ±2.5σ of the peak. In each line, these pixels were integrated, providing the membrane fluorescence time series $F(t)$. A background correction was applied by subtracting the average pixel fluorescence value on the inner side of the membrane multiplied by 2.5σ (in pixel units) from the membrane fluorescence, in blocks of 1000 lines (34). In order to correct for depletion due to photobleaching, the fluorescence time series was fitted with a two-component exponential function and a mathematical correction was applied (17). Finally, the normalized ACF was calculated as:
\[ G(\tau) = \frac{\langle \delta F(t)\delta F(t - \tau) \rangle}{\langle F(t) \rangle^2} \]

where

\[ \delta F = F(t) - \langle F(t) \rangle \]

To avoid artefacts that can be caused by long-term instabilities of the system or single bright events, correlation functions were first calculated segment-wise (10 segments per time trace), and segments with distortions were manually removed before averaging the correlation functions. Eventually, a model for two-dimensional diffusion in the membrane and a Gaussian focal volume geometry (33) was fitted to the ACF:

\[ G(\tau) = \frac{1}{N} \left( 1 + \frac{\tau}{\tau_D} \right)^{-1/2} \left( 1 + \frac{\tau}{\tau_D S^2} \right)^{-1/2} \]

where \( \tau_D \) denotes the diffusion time, and \( N \) the number of particles. The structure parameter \( S \) was fixed to the value of the daily based calibration measurement. Diffusion coefficients \( D \) were calculated using the calibrated waist \( \omega_0 \) of the focal volume: \( D = \omega_0^2 / 4\tau_D \).

**Statistical analysis**

All data were analyzed using GraphPad Prism 5.0 (GraphPad Software Inc., San Diego, CA), and displayed as box plots indicating the median values and with Tukey whiskers. Quantities in the main text are expressed as mean ± SD. Sample sizes and \( p \)-values are indicated in figure captions. Statistically significant differences between control and test samples were determined using the one-way ANOVA analysis followed by the Bonferroni’s multiple comparisons test. A \( p \)-value <0.01 was considered indicative of statistical significance.

**Line-scan Fluorescence Correlation Spectroscopy**

Line-scan FCS (lsFCS) measurements on SLBs were performed as previously described (17). Briefly, data were acquired by scanning repeatedly the focal volume in a linear fashion in the plane of the membrane. Line scans of ca. 5-10 µm length were chosen so that both bilayer
phases (L\text{d} and L\text{o}) were scanned through. We typically acquired 250000 lines, each divided in 256 pixels, with a pixel time of 1.27 µs (line time: 763 ms). Intensity values were correlated along each line and between different lines, calculating the full spatio-temporal autocorrelation \( G(\zeta, \tau) \). To account for photobleaching, a mathematical correction was applied (17). Data analysis was performed with a custom-written script in Matlab, by fitting \( G(\zeta, \tau) \) using a weighted nonlinear least-squares fitting algorithm and a mathematical model taking into account the linear scanning and 2-d Brownian diffusion. To capture the statistical information at larger lag times, we also included in the evaluation (via global fitting) the analysis of the temporal autocorrelation curve \( G(0, \tau) \), calculated on a logarithmic scale with a multiple \( \tau \)-algorithm (17). We could thus obtain estimates for the waist \( \omega_0 \) and the diffusion coefficient \( D \) for a fluorescent lipid analogue (TF-Chol, 0.01 mol\%) added to the examined SLBs. Autocorrelation curves were calculated independently for the different lipid phases, which could be identified due to the distinct affinity of a fluorescent dye (i.e. Rhod-DOPE, 0.1 mol\%) to the L\text{o}- and the L\text{d}-phases. Line scans were performed using the same setup described above for sFCS, using a 488 nm argon laser (ca. 1.5 µW) for the excitation of TF-Chol. The signal originating from Rhod-DOPE (excitation: 561 nm, 561/488 dichroic mirror, emission collected between 571 and 650 nm) was collected only in order to distinguish the lipid phases, but not further used for lsFCS analysis. The \( D \) values for TF-Chol in the L\text{o} and L\text{d} phase of SLB, in the absence of EPC3, were calculated as average of three independent measurements performed in different days. These values were then used as normalization reference for \( D \) values measured in the presence of EPC3, so to emphasize the effect of the drug rather than day-to-day variations.
RESULTS AND DISCUSSION

EPC3 increases membrane fluidity in lipid bilayer models

In order to assess the influence of EPC3 on lipid-lipid interactions, we have first investigated its effects on controlled membrane models. More in detail, we have prepared SLBs mimicking the general composition of the outer leaflet of the PM. These bilayers were composed of a natural mixture of PC (i.e. EggPC), sphingomyelin (bSM) and cholesterol (4/4/2 molar ratio). Such ternary lipids mixtures are known to separate into a liquid disordered (L\(_d\)) and a liquid-ordered phase (L\(_o\)), thus providing a simple model for the study of phase separation occurring at the PM of living cells (22, 35, 36).

In this experiment, we first observed the effect of EPC3 on the general appearance of the L\(_o\)-L\(_d\) phase separation in SLBs, via confocal optical microscopy. For this purpose, SLBs were labelled with an unsaturated fluorescent lipid (Rhod-DOPE) which readily partitions in the L\(_d\) phase (37). As apparent in Fig. 1 A, SLBs showed bright patches enriched in Rhod-DOPE (i.e. L\(_d\) phase) and dark regions devoid of Rhod-DOPE (i.e. L\(_o\) phase). In the presence of increasing amounts of EPC3 in solution (Fig. 1 B-D), the surface occupied by the L\(_d\) phase increases. At the highest EPC3 concentration (Fig.1 D), only small patches of L\(_o\) phase are still visible. These results indicate that EPC3 can effectively insert into the bilayer and destabilize the L\(_o\) phase. A similar induction of lipid mixing was observed also for other alkylphospholipids in giant unilamellar vesicles (38).
Figure 1: Effect of EPC3 on phase-separating model membranes. Panels A-D show representative cLSM images of EggPC/bSM/Chol 4/4/2 molar ratio SLBs in the presence of 0, 1, 5 and 10 µM EPC3, respectively. The bilayers were labelled with 0.1 mol% Rhod-DOPE. Darker zones (devoid of Rhod-DOPE) indicate L<sub>o</sub> regions of the SLB. Images were acquired at RT. Scale bars are 10 µm. Panels E and F show the values of the diffusion coefficients D measured via lsFCS in the L<sub>d</sub> (E) and L<sub>o</sub> phase (F) of EggPC/bSM/Chol 4/4/2 molar ratio SLBs in the presence of EPC3. The reported values refer to the diffusion of a fluorescent cholesterol analogue (TF-Chol, 0.01 mol%). Independent experiments were repeated in triplicates and averaged after normalization (see Materials and Methods). Error bars represent standard deviations.

Next, we characterized the degree of order of the lipid bilayer, in each phase. To this aim, we quantified the diffusion coefficient (D) of a fluorescent membrane probe (TF-Chol) both in the L<sub>o</sub> and L<sub>d</sub> phases. Lipid dynamics are in general connected with membrane order, with lower D values being associated to tighter lipid packing and stronger lipid-lipid interactions (22). Using lsFCS, we measured D for TF-Chol in the L<sub>d</sub> (Fig. 1 E) and L<sub>o</sub> phase (Fig. 1 F), in the presence of increasing concentrations of EPC3. In the absence of the drug, D was ca. 0.1 µm<sup>2</sup>/s in the ordered phase and ca. 20-fold higher in the disordered phase, as expected from previous experiments (35). In the presence of EPC3, we observed a ca. 2-fold increase in lipid diffusion dynamics in the L<sub>d</sub> phase. For the L<sub>o</sub> phase, we observed a similar behavior, although the total change in D was minor compared to the data spread, in this case. These
results clearly suggest that EPC3 has a fluidizing effect on the lipid bilayer that leads to a significant increase of the diffusion of membrane components. On the other hand, it is not possible to determine from these data whether EPC3 interacts preferentially with (and inserts in the bilayer through) the $L_o$ or the $L_d$ phase. Of interest, saturated alkylphospholipids were suggested to partition at the boundary between $L_o$ and $L_d$ phases (38).

**EPC3 alters the fluidity of the plasma membrane of living cells**

In order to investigate the effects of EPC3 directly in the PM of living cells, we applied an approach based on the spectroscopic properties of two fluorescent molecules (i.e. Di-4-ANEPPDHQ and Laurdan) which are strongly sensitive to membrane order and lipid packing (25, 26). In the context of studies regarding the effect of APLs on lipid bilayers, Laurdan was used to detect and increase in the fluidity of model membranes induced by Miltefosine (39).

In this experiment, we characterized directly the PM of two breast cancer cell models: the high-invasive MDA-MB-231 cell line and the low-invasive MCF-7 cell line. In previous studies, we observed that EPC3 causes increased cytotoxicity, apoptosis and cytoskeleton reorganization especially in highly invasive MDA-MB-231 cells, compared to MCF-7 samples (40, 41). In addition, MDA-MB-231 cells showed cell cycle arrest after treatment with EPC3 (41).

The mechanism by which Laurdan and Di-4-ANEPPDHQ detect changes in the local membrane environment is similar for the two molecules: the less polar environment of ordered bilayer phase (e.g. $L_o$ phase or a raft-like domain in the PM) induces a blue shift in the emission maxima of both fluorescent probes. This shift can be quantified by calculating a ratiometric measurement of the fluorescence intensity observed in two spectral regions (or channels), known as a generalized polarization (GP) value. Higher GP values correspond to a relatively higher fluorescence emission in the shorter-wavelength spectral region and, therefore, a higher degree of membrane order and tighter lipid packing (26).

Fig. 2 shows normalized histograms for Di-4-ANEPPDHQ GP values measured in MDA-MB-231 cells, in the presence of 20 µM or 30 µM EPC3. Cells were observed via confocal fluorescence microscopy after 24 h, 48 h and 72 h (see Fig. S1). In all cases, the PM region of several cells was manually selected and the GP values for each pixel within these ROIs were calculated as described in the Methods section.
Figure 2: Di-4-ANEPPDQH GP values measured at the PM of MDA-MB-231 cells after EPC3 treatment.

Normalized histograms of GP values measured in pixels belonging to the PM of MDA-MB-231 cells labelled with Di-4-ANEPPDQH. Cells were treated with 20 µM (orange bars) or 30 µM (yellow bars) EPC3. GP values measured in cells not treated with EPC3 are shown as blue bars. Fluorescence intensity values were acquired 24 h (Panel A), 48 h (Panel B) and 72 h (Panel C) after the addition of EPC3. For each condition, GP values were pooled from ca. 50 ROIs selected at the PM of distinct cells, in two independent experiments. The total number of calculated GP values (and measured pixels) for each experimental condition was between ca. 10000 and 40000. Measurements were performed at RT. Representative images of treated cells and ROIs are shown in Fig. S1.

As shown in Figs. 2, S2 and Table S1, the fluorescence emission of di-4-ANEPPDQH indicates that EPC3 treatment of cancerous cells induces a time-dependent increase of PM disorder. Only minor differences were observed between IC50 and IC75 EPC3 concentrations (i.e. 20 and 30 µM for MDA-MB-231 cells, respectively). Performing the same experiment using MCF-7 cells (Figs. 3, S3 and Table S2), we observed that EPC3-induced membrane order alterations are stronger and occur faster in MDA-MB-231 cells.
Figure 3: Di-4-ANEPPDQH GP values measured at the PM of MCF-7 cells after EPC3 treatment. Normalized histograms of GP values measured in pixels belonging to the PM of MCF-7 cells labelled with Di-4-ANEPPDQH. Cells were treated with 40 µM (orange bars) or 60 µM (yellow bars) EPC3. GP values measured in cells not treated with EPC3 are shown as blue bars. Fluorescence intensity values were acquired 24 h (Panel A), 48 h (Panel B) and 72 h (Panel C) after the addition of EPC3. For each condition, GP values were pooled from ca. 50 ROIs selected at the PM of distinct cells, in two independent experiments. The total number of calculated GP values (and measured pixels) for each experimental condition was between ca. 10000 and 30000. Measurements were performed at RT. Representative images of treated cells and ROIs are shown in Fig. S1.

In the case of MCF-7 cells, in fact, no significant alteration in membrane order can be observed after 24 h of treatment, both at IC50 and IC75 EPC3 concentrations (i.e. 40 and 60 µM for MCF-7 cells, respectively). After 48h and 72 h however, the GP distribution appears significantly shifted to lower values, independently from EPC3 concentration (Figs. 3 B-C, S3 and Table S2).

Next, we have used a complementary assay quantifying the GP parameter, following analogous treatment of MDA-MB-231 and MCF-7 cells, using Laurdan as fluorescent probe instead of Di-4-ANEPPDHQ (Fig. S1).
Figure 4: Laurdan GP values measured at the PM of MDA-MB-231 cells after EPC3 treatment.

Normalized histograms of GP values measured in pixels belonging to the PM of MDA-MB-231 cells labelled with Laurdan. Cells were treated with 20 µM (orange bars) or 30 µM (yellow bars) EPC3. GP values measured in cells not treated with EPC3 are shown as blue bars. Fluorescence intensity values were acquired 24 h (Panel A), 48 h (Panel B) and 72 h (Panel C) after the addition of EPC3. For each condition, GP values were pooled from ca. 50 ROIs selected at the PM of distinct cells, in two independent experiments. The total number of calculated GP values (and measured pixels) for each experimental condition was between ca. 10000 and 40000. Measurements were performed at RT.

The results shown in Figs. 4, S4 and Table S3 indicate the presence of a concentration-dependent decrease in membrane order after 48h and 72h treatment with EPC3. We did not observe prominent alterations of the cell membrane order after 24h treatment. These observations are qualitatively similar to those performed using Di-4-ANEPPDQH as fluorescent probe. Also, in analogy to the results shown in Fig. 3 (i.e. labelling of MCF-7 cells with Di-4-ANEPPDQH), the observed changes in lipid packing were larger in MDA-MB-231 than in MCF-7 cells (Fig. S5, S6 and Table S4).

In summary, fluorescence microscopy measurements based on both Laurdan and Di-4-ANEPPDQH labelling indicate a decrease in PM lipid packing and order induced by EPC3, especially in the case of MDA-MB-231-cells. Interestingly, we noticed that the observed effects were much more noticeable when Di-4-ANEPPDHQ was used, instead of Laurdan. This difference might stem from two different factors. First, as previously described (42), Di-4-ANEPPDQH is more effective in specifically labelling the PM. Laurdan, on the other hand, penetrates more easily into the cytosol and labels also intracellular structures (see Fig. S1), thus making ROI selections more difficult and less precise, while generally increasing the spread in observed GP values. Second, Di-4-ANEPPDQH and Laurdan are sensitive to
different properties of the lipid bilayer (25). While Laurdan is sensitive to lipid packing, the spectroscopic properties of Di-4-ANEPPDQH are influenced by other factors, such as cholesterol content of the membrane or internal electric dipole potential of the bilayer. It is therefore tempting to assume that the changes in membrane order (observed by using Di-4-ANEPPDQH rather than Laurdan) might be connected to alterations in membrane compositions brought about by EPC3. This hypothesis is in agreement with the observation that certain APLs might influence lipid metabolism and lipid-mediated signaling cascades (1, 2, 43). Also EPC3 can reduce the synthesis of key raft-associated phospholipids, such as PC and SM ((44) and Tzoneva R., unpublished data). On the other hand, our experiments on model membranes clearly indicate that EPC3 can directly influence lipid-lipid interactions and membrane fluidity, even in the absence of any alteration in membrane composition (Fig.1). We conclude therefore that EPC3 modulates the physical properties of the PM, by e.g. directly influencing the interactions between cholesterol and other lipids and/or altering the internal electric dipole potential of the bilayer. Additional effects due to alterations in lipid metabolism and PM composition cannot, in general, be excluded.

**EPC3 increases diffusion dynamics of trans-membrane proteins in the PM**

We next investigated whether the alterations in lipid-lipid interactions caused by EPC3 and discussed at the end of the previous paragraph can have a specific influence on the behavior of membrane proteins. We therefore investigated the diffusion of two illustrative transmembrane proteins, namely the human EGF receptor (hEGFR) and the Hemagglutinin receptor from Influenza A FP virus (FPV-HA), in MDA-MB-231 cells treated with 30 µM EPC3. The hEGFR is a well-characterized membrane protein which partitions in lipid domains of the PM (45). The transmembrane glycoprotein HA of the influenza A virus is known to form trimers at the PM, where it localizes in raft domains (46, 47).

As shown in the previous paragraph, EPC3 (in this case, 30 µM in MDA-MB-231 cells) strongly influences the physical state of the PM, as reported by Di-4-ANEPPDQH. The two examined proteins were labelled with a fluorescent protein (GFP) and visualized via confocal fluorescence microscopy. As shown in Fig. 5 A, EPC3 does not alter the localization of either protein at the PM. We then applied sFCS to quantify the diffusion dynamics of the two fluorescently labelled proteins. Fig. 5 B shows representative autocorrelation curves for the
fluorescence signal recorded for hEGFR (upper panel) and FPV-HA (lower panel), also in the presence of EPC3.

Figure 5: Quantification of protein diffusion in EPC3-treated cells via sFCS. Panel A shows representative images of MDA-MB-231 cells expressing hEGFR-EGFP (upper row), and FPV-HA-EGFP (lower row), in the absence of EPC3 or after a 48 h incubation with EPC3. The yellow line in panel A represents an example of scanning path used for sFCS measurements. Scale bars are 10 µm. Panel B shows representative sFCS autocorrelation functions and fit curves obtained for cells expressing hEGFR-EGFP (upper graph) and FPV-HA-mEGFP (lower graph), in the absence of EPC3 or after a 48 h incubation with EPC3. Fit curves (solid line) were obtained by fitting a two-dimensional diffusion model to the data, as described in the Methods section. Panel C shows box plots of diffusion coefficients calculated from sFCS diffusion times, pooled from three independent experiments (each consisting of measurements on 20 cells). The value of diffusion coefficient D for hEGFR-EGFP significantly increased from $0.26 \pm 0.08 \mu m^2/s$ to $1.1 \pm 0.5 \mu m^2/s$ after EPC3 treatment. The value of diffusion coefficient D for FPV-HA-mEGFP significantly increased from $0.22 \pm 0.08 \mu m^2/s$ to $1.2 \pm 0.5 \mu m^2/s$ after EPC3 treatment. All measurements were performed at RT.

Both membrane proteins diffuse in the PM with a diffusion coefficient $D \sim 0.25 \mu m^2/s$, as expected from previous measurements (19-21). Furthermore, in the presence of EPC3, we observed an average $\sim$5-fold increase in diffusion dynamics, both for hEGFR and FPV-HA. Changes in protein diffusion were previously observed as a consequence of alterations in lipid-lipid and lipid-protein interactions. For example, it was shown that hEGFR and HA diffusion coefficients increase as a result of alterations in membrane composition or protein association to membrane domains (30, 45, 48). Our data are compatible with both possibilities. Nevertheless, as already mentioned in the previous paragraph, the results
obtained from our experiments in SLBs indicate that changes in membrane composition are not required for the EPC3-induced increase in membrane dynamics.

CONCLUSIONS

The increased amount of saturated lipids and ordered lipid domains in tumor cells (compared to healthy cells) (49) correlates with a reduction in membrane fluidity/dynamics and an increase in chemotherapy resistance (50, 51). In cancer cells, a wide range of signaling proteins and receptors regulating pro-oncogenic and apoptotic pathways are localized in lipid raft domains (6). It is now accepted that APLs exert their effect by interacting with lipid rafts, due to their participation in the regulation of cell survival and cell death pathways (52). In this context, we show for the first time that EPC3 causes a significant increase in PM disorder and its effect appear stronger in the high-invasive cancer cell line MDA-MB 231. The enhanced membrane fluidity has a direct effect on the dynamics of membrane components, as demonstrated by our measurements of membrane protein diffusion. Increased protein diffusion dynamics likely affects the likelihood of protein-protein interactions and, therefore, might have an effect on signaling pathways. Furthermore, it is possible that the decrease in lipid packing corresponds to a higher permeability of the PM and, thus, an increased intake of EPC3 through the cell membrane.

ACKNOWLEDGMENTS

This work was supported by the grant DN 11/1/2017 from the Bulgarian Science Fund and partially by the grants DO1-154/28/08/2018/ (Scientific Infrastructure on Cell Technologies in Biomedicine (SICTB) from the Bulgarian Ministry of Education and Science) and 254850309 (German Research Foundation (DFG)).
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