Marine Streptomyces sp. Isolated From the Brazilian Endemic Tunicate Euherdmania sp. Produces Dihydroeponemycin and Analogs With Potent Antiglioma Activity

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Marine natural products have emerged as an important source for drug development, notably in the field of anticancer therapy. Still, the limited effectiveness of current therapies for central nervous system tumors indicates the need to identify new therapeutic targets and also novel pharmacological agents. In this context, proteasome inhibitors are appearing as a promising new treatment for these diseases. Herein, cytotoxic extracts produced by four marine bacteria recovered from the Brazilian endemic ascidian Euherdmania sp. were screened to evaluate their potential as proteasome inhibitors. The extract from marine Streptomyces sp. BRA-346 was selected for further investigation due to the potent proteasome inhibitory activity it displayed. Bioassay-guided fractionation led to an enriched fraction (proteasome inhibition IC₅₀ = 45 ng/mL), in which the presence of dihydroeponemycin (DHE), known for its proteasome inhibitory effect, and related compounds were annotated by mass spectrometry and further confirmed by comparison with DHE standard. Both DHE and the epoxyketone-containing fraction were evaluated in glioma cell lines, displaying high cytotoxicity in HOG and T98G cells (GI₅₀ of 1.6 and 1.7 ng/mL for DHE, and 17.6 and 28.2 ng/mL for the BRA-346 fraction, respectively). Additional studies showed that the epoxyketone-containing fraction (at GI₅₀ levels) led to an accumulation of ubiquitinated proteins and up-regulation of genes related to ER-stress response, suggesting treated cells are under proteasome inhibition. DHE induced similar effects in treated cells but at concentrations 25 times its GI₅₀, suggesting that the other epoxyketone compounds in the bacteria extract derived fraction may contribute to...
INTRODUCTION

The marine environment encompasses more than 90% of the volume of the biosphere, housing an enormous and still majorly unknown number of species. The most optimistic estimate from the Census of Marine Life points out to barely 10% of properly described species in the oceans1 (Snelgrove, 2010; National Oceanic and Atmospheric Administration, 2020). However, if a broader concept of diversity is considered, one that includes genetic and chemical diversities, it becomes even more evident that this surface has hardly been scratched (Mora et al., 2011; Snelgrove, 2016). Indeed, from a chemical diversity perspective, the possibilities of exploring this multitude of molecules are virtually limitless, once a marine organism, such as an invertebrate, may be contemplated as a complex ecosystem in itself, with billions of associated microbial cells interacting and responding to biotic and abiotic factors through the production of their metabolites to warrant the holobiont’s homeostasis (Cheng et al., 2020; McCauley et al., 2020).

The uniqueness of the metabolites produced by marine species gathered interest from the scientific community for their potent biological activities, resulting in the development of thirteen drugs, of which most are being used to treat cancer2 (Jimenez et al., 2020; Midwestern University, 2020). In such realm, tunicates – filter-feeding animals distributed across the ocean – are among the most resourceful organisms considering the return of pharmacological capital (Palanisamy et al., 2017; Bauermeister et al., 2019), being recognized for the ability to produce nitrogen-containing compounds, like alkaloids, peptides and their derivatives (Watters, 2018). Didemnin B, a depsipeptide isolated from Trididemnum solidum, was the first marine natural product to enter clinical trials, back in 1986, having been later discharged due to marked toxicity and lack of efficacy (Rinehart et al., 1990; Markham, 2020). Both didemnin B and trabectedin are recognized as products of the metabolism of symbiotic bacteria, emphasizing the importance of the associated microbiota on the chemical versatility previously attributed to tunicates (Tsukimoto et al., 2011; Xu et al., 2012; Schofield et al., 2015; Bauermeister et al., 2019).

Although the majority of current anticancer drugs have been obtained from natural sources (Newman and Cragg, 2020), only a few marine compounds are included in this arsenal. However, marine natural products have already distinguished themselves from their outnumbered terrestrial counterparts in terms of their mode of action (Pereira et al., 2019). Plitidepsin, for example, is a first in the class drug that impairs de novo protein synthesis by blocking the function of eEF1A2 (eukaryotic elongation factor 1A2), a protagonist protein in delivering amino-acylated tRNA to ribosomes and thereby promoting polypeptide elongation (Mateyak and Kinzy, 2010; Losada et al., 2016). However, a secondary role to eEF1A2, which favors cancer cell apoptosis, involves the elimination of misfolded proteins through clustering them for lysosome degradation or tagging them for proteasome degradation. Therefore, inhibition of this elongation factor causes the accumulation of toxic misfolded proteins in the cytoplasm, which then leads to cell death (Hotokezaka et al., 2002; Losada et al., 2020). In fact, provoking cell buildup of misfolded proteins revealed to be a successful anticancer strategy, and one such approach involves hampering the catalytic function of the proteasome itself.

The proteasome is a multicatalytic enzyme complex responsible for most non-lysosomal cellular proteolysis. In fact, the proteasome can be considered the headquarter of protein turnover, influencing many cellular processes such as cell cycle, angiogenesis, immune response, apoptosis, and gene expression. The altered proteostasis of cell cycle regulators (cyclins, tumor suppressors, and oncogenes) and apoptosis-related proteins influence cancer development. For example, non-degradation of cyclin-dependent kinase inhibitors and Ik-Bα prevents cell cycle progression and activation of NF-κB, which switches on prosurvival genes (BCL-2) (Catalgol, 2012; Thibaudeau and Smith, 2019). In light of this, inhibition of the 20S proteasome core particle (20S-CP) is a validated strategy for cancer therapy, where impairment of the chymotrypsin-like activity (ChfL, beta 5) is sufficient to achieve cancer cell apoptosis with a favorable therapeutic window (Kisseley et al., 2012).

In this context, microorganisms have been proven as a promising source of proteasome inhibitors, including the α-′β′-epoxyketones epoxomicin, eponemycin and related compounds (Sugawara et al., 1990; Hanada et al., 1992), and the β-lactone-γ-lactam marizomib (Feling et al., 2003; Potts et al., 2011). Among them, epoxomicin led to the development of carfilzomib

Keywords: marine natural products, cytotoxicity, proteasome inhibitors, epoxyketones, ER-stress response

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1https://oceanservice.noaa.gov/facts/ocean-species.html
2www.midwestern.edu/departments/marinepharmacology.xml
3www.ebs.tga.gov.au
4www.fda.gov/drugs

enhance proteasome inhibition and further cellular effects in glioma cells. These findings revealed the molecular pathways modulated by this class of compounds in glioma cells and, moreover, reinforced the potential of this marine bacteria in producing a cocktail of structurally-related compounds that affect the viability of glioma cells.
(kyprolis®, Onyx Pharmaceuticals), approved by the United States Food and Drug Administration (2020) in 2012 for the treatment of patients with relapsed or refractory myeloma (Muchtar et al., 2016). Additionally, marizomib, produced by the marine free-living actinobacteria Salinispora tropica, is undergoing phase III clinical trials for the treatment of hematological and central nervous system tumors (Jimenez et al., 2020). The α′-β′-epoxyketone inhibitors bind to the β5-subunit of the proteasome, exhibiting a potent and highly specific inhibitory effect on the chymotrypsin-like activity (Kim et al., 1999). Marizomib, in turn, inhibits three catalytic subunits of the proteasome, with, generally, a greater affinity toward the β5 subunit (Potts et al., 2011).

Herein, we evaluated the presence of proteasome inhibitors in cytotoxic extracts of bacteria isolated from the Brazilian endemic tunicate Euherdmania sp. The genus Euherdmania comprises merely 13 species and none have been so far characterized for their secondary metabolism nor associated microbiota (Monniot, 1983; Lotufo and Silva, 2006). Still, a previous effort to access the biological activity of crude extracts obtained from tunicates collected on the coast of Ceará State (Jimenez et al., 2003) highlighted the extract from Euherdmania sp. for a prominent cytotoxic activity against leukemia, melanoma, and colorectal tumor cell lines. Using a HPLC-MS/MS analysis and comparison with standard compounds, the presence of a series of α′-β′-epoxyketones related to eponemycin, including dihydroeponomycin (DHE), were verified in Streptomyces sp. BRA-346 extract along with a potent inhibitory activity of the proteasome. Then, the antiglioma activities of DHE and BRA-346 fraction-containing a cocktail of α′-β′-epoxyketones were evaluated focusing on the modulation of pathways related to the unfolded protein response (UPR).

MATERIALS AND METHODS

Bacteria and Extracts

Four bacteria isolated from the ascidian Euherdmania sp., BRA-339, BRA-341, BRA-346, and BRA-386, collected at Talha beach, Brazilian northeast coast (3°34.125’S; 38°54.469’W), were used for this work and the procedure used was previously described in Velasco-Alzate et al. (2019). Briefly, ascidian samples were sorted for epibionts and debris, dipped in ethanol 70% for superficial decontamination and kept under sterile conditions. Samples were suspended in sterile seawater (1:1 v/v), macerated, and separated into two aliquots: the first was heated to 55°C over 10 min to minimize the growth of Gram-negative bacteria prior to inoculation; the second was directly inoculated onto 90 mm Petri dishes filled with starch-casein agar (SCA) prepared with reconstituted seawater. Cycloheximide (0.1 mg/mL) was added to isolation plates to reduce fungal contamination. Colonies owning an actinomycete-phenotype (color, presence of spores, texture, etc.) were picked out, transferred to fresh A1 (soluble starch, peptone, yeast extract, reconstituted seawater, and agar) medium dishes and subjected to subsequent replatings onto A1 dishes for purification.

Purified bacteria strains were cryopreserved in the microorganism strain bank of the Laboratory of Marine Bioprospecting and Biotechnology at the Federal University of Ceará. The selection of strains was based on the high cytotoxic activity observed against colon carcinoma cell lines (Supplementary Table 1). A license for the collection of tunicates was granted by Biodiversity Authorization and Information System (SISBIO, authorization number 48522-2). A license for genetic access was granted by the National System for the Management of Genetic Heritage and Associated Traditional Knowledge (SISGen, authorization number AC0781C).

The extracts were obtained from the growth of bacteria strains in Erlemeyer flasks (250 mL) containing 100 mL of A1 liquid medium (peptone, soluble starch, yeast extract, and reconstituted seawater) for 7 days at 140 rpm maintained at 26°C. To recover the extracts, 100 mL of ethyl acetate was added to the bacteria culture for 1 h under 60 rpm at 26°C. The organic phases were separated from culture bacteria media and collected. The solvent was then removed under reduced pressure and extracts were recovered.

MTT Assay

The colon carcinoma cell line [HCT-116 (ATCC CCL-247)] was used for the MTT assay. This assay measures cellular metabolic function based on reduction of tetrazolium salt, MTT 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide, to the formazan by mitochondrial dehydrogenases of viable cells. An increase in the formazan concentration indicates greater cell proliferation (Mosmann, 1983).

Cells were cultured in RPMI media supplemented with 10% SBF and 1% penicillin-streptomycin and seeded in 96-well plates (1 × 10^3 cells per well) 24 h before the test. The extracts and fractions were dissolved in DMSO and tested at increasing concentrations (0.0032–50 µg/mL) for 72 h of incubation at 37°C, 5% of CO2. DMSO 0.5% was used as a negative control, reflecting 100% cell viability. After the incubation period, the media was removed and an MTT solution (5 mg/mL) was added to plates for an additional incubation (3 h). Next, the supernatant was discarded and the precipitate (formazan) was dissolved in DMSO to measure absorbance at 570 nm. Three independent experiments were performed and absorbance values were normalized using blank as 0%, and negative control (DMSO, 0.5%) as 100% of cell proliferation. Inhibitory concentration mean (IC50) values and their 95% confidence intervals (CI95%) were calculated using non-linear regression by GraphPad Prism 7.0. (GraphPad Software, Inc., San Diego, CA, United States).

Proteasome Inhibition Assay

The EtOAc bacteria crude extracts, obtained as described above, were assayed against the purified 20S-CP from Saccharomyces cerevisiae cells. Yeast 20S proteasome complex was purified in-house from S. cerevisiae using affinity, ion exchange, and size exclusion chromatography, as previously described (Souza et al., 2018). The assay procedures were carried out according to Souza et al. (2018) with minor modifications. Briefly, bacteria crude extracts were diluted in DMSO and assayed by pipetting 1 µL in 384-well black v-shape microplates containing 19 µL of the
yeast proteasome at 1 μg/mL, in Tris 10 mM pH 7.5, SDS 0.01%. The yeast proteasome was exposed to the extracts for 2 h before addition of 5 μL of the fluorogenic proteasome substrate Suc-LIYV-Amc at 200 μM (final substrate concentration was 40 μM). The fluorescence was monitored in each well for 1 h, every 2 min, using the Clario (BMG Labtech) plate reader. Liquid handling was assisted by the Versette (Thermo Scientific) automated pipetting system. Each experiment was conducted in triplicates. Remaining proteasome activity (%) for test samples was calculated relative to the DMSO control group, carried out in the same assay plate, however, in the absence of the tested samples. The control group corresponds to 100% proteasome activity and was obtained by incubating the enzyme in the reaction buffer with 5% DMSO before adding the substrate. The intrinsic fluorescence of the samples was also accounted for in the measurement of the fluorescence in each well before the addition of the substrate. Initial experiments were carried out at single concentrations (50 μg/μL) of the crude extracts. Inhibition curves were performed at ten inhibitor concentrations, with final concentrations ranging from 0.003 to 50 μg/mL, in a dilution factor of threefold. The data were plotted as average ± SD of enzyme activity (%) as a function of logarthmic concentrations of the inhibitor. Curves were fitted using the normalized four-parameter concentration-response equation with variable slope implemented in Prism (GraphPad, San Diego, v. 7.0).

To assess the inhibitory potential of dihydroeponemycin (DHE) (ApexBio, A8172), epoxomicin (ABCAM, ab144598), and epoxyketone-containing fraction obtained from the fractionation of the BRA-346 extract (BRA-346 fraction) on human purified proteasome catalytic subunit, the 20S Proteasome Activity Assay (Merck Millipore, Cat. No. APT280) was used. DMSO (0.3%) was used as the negative control. The entire test was performed according to manufacturer protocol (Zhang et al., 2011). To calculate the half-maximum inhibitory concentration (IC50) of epoxyketone-containing fraction, the sample was evaluated in graded concentrations (0.001, 0.01, 0.1, 1, and 10 μg/mL). The emission fluorescence converted values (%) were compared to the negative control. Assays were analyzed using GraphPad Prism 7.0 software.

Molecular Identification of BRA-346
For molecular identification, genomic DNA of the isolated BRA-346 strain was obtained from pure culture as previously described by Pinto et al. (2020) using phenol:chloroform:isoamyl alcohol-based protocol. The gene 16S rRNA was amplified by polymerase chain reaction (PCR) using primers 27F (5′- AGAGTTTGATCCTGCTCAG-3′) e 1494R (5′-ACGGCTACCTTGTTACGACTT-3′) (Heuer et al., 1997). The PCR reaction was also performed as previously described by Pinto et al. (2020). The PCR product was stained using SYBR™ Safe DNA Gel (Life Technologies), analyzed by electrophoresis on agarose gel 1%, and purified with ExoSAP-IT (Affymetrix). Sequencing of amplicons was conducted at the Laboratory of Human Molecular Genetics (LGMH), Department of Genetics of Federal University of Pernambuco (UFPE) using Bigdyce TM terminator v3.1 cycle sequencing kit (Thermo Fisher). Sequences forward and reverse were analyzed and edited using Geneious Prime 2020.2 software. The obtained sequence was deposited in Genbank under the number MW342808 and compared with sequences from EzBiocloud database.

Fractionation of BRA-346 Extract
BRA-346 strain was grown in Erlenmeyer flasks (30 × 1 L) containing 250 mL of A1 medium at 26°C, 180 rpm. The crude extract (1.35 g) recovered with ethyl acetate was dissolved in 100 mL of MeOH/H2O (8:2, v/v) and partitioned with n-hexane and EtOAc (2 mL × 100 mL), followed by treatment with anhydrous Na2SO4. The organic solvents were dried by distillation under reduced pressure and, subsequently, by compressed airflow, providing their respective fractions: BRA-346 hexane (506.3 mg) and BRA-346 ethyl acetate (416.5 mg). The resultant MeOH/H2O mixture was dried by solvent distillation under reduced pressure and lyophilized, yielding BRA-346 MeOH/H2O (375.6 mg).

The BRA-346 EtOAc fraction (400.5 mg) was fractionated on a Sephadex LH-20 column (50 mm × 30 mm i.d.) using a mixture of MeOH/CH2Cl2 (8:2, v/v) to provide 20 fractions of approximately 10 mL each, which were monitored by thin layer chromatography (TLC) to yield five fractions, F1-4 (5.6 mg), F5-8 (25.0 mg), F9-11 (65.1 mg), F12-14 (217.5 mg), and F15-20 (86.5 mg), which were screened for cytotoxicity. F12-14 showed the most significant activity, therefore, an aliquot of this fraction (20.0 mg) was subjected to solid phase extraction (SPE) eluted with A (H2O), B (MeOH/H2O 1:1, v/v), C (MeCN/H2O 7:3, v/v), D (MeCN), and E (CHCl3/MeOH 9:1, v/v) to provide their respective fractions, A (0.7 mg), B (5.0 mg), C (9.0 mg), D (0.9 mg), and E (0.3 mg) as described by Cutignano et al. (2015). All obtained fractions were evaporated under reduced pressure to remove solvents and diluted in DMSO before screening for cytotoxicity. Fraction C was selected for sequential studies based on its proteasome inhibitory activity and HPLC-MS/MS analysis.

Supplementary Table 2 shows the yield and biological activity data of extract and fractions.

HPLC-MS/MS Data Acquisition and Molecular Networking
BRA-346 crude extract and fractions A, B, C, D, and E were analyzed by HPLC-MS/MS for metabolic profiling. Dihydroeponemycin (DHE) (ApexBio, A8172), epoxomicin (ABCAM, ab144598) were used as standards for comparison. The analyses were acquired on an HPLC System (Shimadzu) coupled to a micrOTOF QII mass spectrometer (Bruker Daltonics), equipped with an electrospray ionization source (operating in positive ionization mode), and a quadrupole and a time of flight (TOF) analyzers. Chromatographic separation was performed on a Supelco Ascentis Express C18 column (5 μm, 150 mm × 3.0 mm), equipped with a pre-column. The column temperature was maintained at 40°C. The elution was carried

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4https://www.geneious.com
5www.ncbi.nlm.nih.gov/genbank
6www.ezbiocloud.net
out with a linear gradient of the mobile phase, starting with 5–100% acetonitrile for 25 min with a flow rate of 0.7 mL/min. Both solvents (H₂O and ACN) contained 0.1% formic acid. The injection volume was 15 µL (1.0 mg/mL). The parameters used in the ionization source were as follows: voltage: 3.5 kV, cone: 4.5 kV, drying gas: 9.0 L/min, gas temperature: 220°C, m/z range: 50–1,500. A data-dependent acquisition (DDA) method was used, which selects to fragment, separately, all ions above 1.0E3 of intensity, in this method a collision energy ramp of 20–75 eV was used to fragment the ions.

The HPLC-MS/MS data were converted to the .mzXML format, a file type compatible with the GNPS platform (Wang et al., 2016), using the MSConvert software. The molecular network was constructed following the workflow on the GNPS platform. GNPS is an online platform, which deposits and shares a mass spectrometry data library (MS-MS), allowing the identification of natural products, mainly of marine origin, and which has been widely used in metabolomics works. Briefly, the data (.mzXML) was transferred to the GNPS server (ucsd.edu). The MS/MS spectra were grouped by spectral similarity using cosine score, and the parameters used are described in Supplementary Table 3. After the construction of the molecular network, the file was viewed and analyzed in the Cytoscape software (v. 3.2).

Mechanistic Studies in Glioma Cells
Sulforhodamine B (SRB) Assay

Due to the presence of proteasome inhibitors in the BRA-346 fraction, which can interfere on the turnover protein, and possible over and underestimation of cell viability by the MTT assay in glioblastoma cell lines (Stepanenko and Dmitrenko, 2015), SRB assay was used to evaluate initial cytotoxicity studies of the BRA-346 epoxide-containing fraction and dihydroeponemycin in glioma cell lines.

This is a colorimetric assay developed in 1990 which quantifies the total proteins and can be used for investigating cell proliferation and screening of the anticancer drugs. It entails the binding of sulforhodamine B dye to basic amino-acid residues of cell proteins, allowing measure increase or decrease of cellular mass (Skehan et al., 1990). SRB assay setup permits the identification of three main values: mean growth inhibitory concentration (GI₅₀) when the treated cells growth 50% in relation to “time-zero” plate cells; total growth inhibitory concentration (TGI), when treated cells do not growth in relation to cells seeded in “time-zero” plate; and mean lethal concentration (LC₅₀), when 50% cells died in relation observed cells concentration in “time-zero” plate. Cell proliferation was calculated as: % of control cell growth = 100 × (absorbance sample value − absorbance “time-zero” plate)/(absorbance negative control − absorbance “time-zero” plate) for 50% of cell proliferation; and absorbance sample value = absorbance “time-zero” plate for 0% of cell proliferation (Vichai and Kirtikara, 2006; NCI, 2021*), and each drug concentration was related. The percentage of proliferation was set range to −100% to +100% to facilitate finding GI₅₀ and LC₅₀ values. These values can be observed in the screening methodology applied by USA-NCI (National Cancer Institute, 2021).

Glioma cell lines [HOG (CVCL_D354) and T98G (ATCC CRL-1690)], certified by Rede PREMIUM FMUSP, São Paulo, Brazil (ATCC – STR Profile Report method), were used. Both cell lines were cultured in RPMI media supplemented with 10% SBF and 1% penicillin-streptomycin and seeded in 96-well plates (6 × 10⁴ cells per well). 24 h later, a plate designated by “time-zero,” containing only cells, was fixed with 100 µL/well of 10% trichloroacetic acid (TCA) and stored at 4°C for 1 h. Meanwhile, DHE and the epoxide-containing fraction, both dissolved in DMSO, were tested on the test plates at increasing concentrations (0.0032–50 µM or µg/mL) for 24, 48, and 72 h of incubation at 37°C and under a 5% CO₂ atmosphere. As a negative control, i.e., 100% cell growth, 0.5% DMSO was used. After each incubation period, the medium of test plates was removed, cells were fixed with 100 µL/well of 10% TCA and plates were stored for 1 h at 4°C. This solution was removed and all plates were washed three times with deionized water. After that, 100 µL of 0.4% SRB solution was added to each well and kept at 37°C and 5% CO₂ for 30 min. The SRB solution was removed by washing three times with 1% acetic acid and, to solubilize the bound SRB, fixed cells were dissolved in 200 µL/well of 10 mM Tris Base before the absorbance of each well was measured at 570 nm. Three independent experiments were performed. Mean growth inhibitory concentration (GI₅₀), total growth inhibitory concentration (TGI), and mean lethal concentration (LC₅₀) were calculated by interpolation of normalized data by GraphPad Prism 7.0. (GraphPad Software, Inc., San Diego, CA, United States).

Western Blotting

Accumulation of ubiquitinated proteins and expression of HSP70 were evaluated by Western blotting as molecular markers of proteasome inhibition. With this purpose, HOG and T98G cells were seeded in 100 mm dishes at 1 × 10⁶ cells/mL. After 24 h, the cells were treated with DHE or BRA-346 epoxide-containing fraction at concentrations corresponding to GI₅₀ values obtained after 48 h incubation (1.6 and 1.7 ng/mL for DHE, and 17.6 and 28.2 ng/mL for the BRA-346 fraction for HOG and T98G, respectively). Incubation time was set at 36 h to evaluate the onset of the effects. DHE was further tested in both cell lines at 40 ng/mL, approximately 25 times the GI₅₀ values obtained after 48 h incubation. Proteins of treated and untreated cells were extracted using a buffer containing 100 mM Tris (pH 7.6), 1% Triton X-100, 150 mM NaCl, 2 mM PMSF, 10 mM Na₃VO₄, 100 mM NaF, 10 mM Na₄P₂O₇, and 4 mM EDTA and their quantification was performed by the Bradford assay (Bradford, 1976) with a BSA curve (1 mg/mL) as standard. The proteins were submitted to SDS-PAGE then transferred to a nitrocellulose membrane for immunodetection. Antibodies against ubiquitin P4D1 (#3936) and α-tubulin (#2144) were obtained from Cell Signaling, while that against HSP70 (#648002) from BioLegend.
independent experiments were performed. Band intensities were quantified by the UN-SCAN-IT gel 6.1 program that captures and quantifies the pixels in each band. Values associated to test proteins were normalized by standard α-tubulin for the relative expression measure.

RNA Extraction and qRT-PCR

HOG and T98G cells were seeded in 100 mm dishes at $1 \times 10^6$ cells/mL and 24 h later treated with DHE or BRA-346 epoxyketone-containing fraction at concentrations corresponding to GI$_{50}$ values obtained after 48 h incubation (1.6 and 1.7 ng/mL for DHE, and 17.6 and 28.2 ng/mL for the BRA-346 fraction for HOG and T98G, respectively). Incubation time was set at 36 h to evaluate the onset of the effects. DHE was further tested in both cell lines at 40 ng/mL, approximately 25 times GI$_{50}$ values obtained after 48 h incubation. For the quantitative PCR analysis, total RNA was extracted from cells using Trizol (as recommended by the manufacturer). Sixteen genes related to UPR and to the ubiquitin-proteasome system (UPS) were analyzed in control and treated cells. The primer sequences are described in Supplementary Table 4. RNA samples were quantified and used for the production of cDNA that was performed according to the High-Capacity cDNA Reverse Transcription Kit protocol (Thermo Fisher Scientific) with the addition of ribonuclease inhibitor RNaseOUT™ (Thermo Fisher Scientific), according to the following steps: 10′ at 25°C; 2 cycles from 60′ to 37°C and one cycle from 5′ to 85°C. Sybr Green PCR Master Mix and solution of the forward and reverse primers were added to the cDNA. Hypoxanthine phosphoribosyltransferase 1 (HPRT1) and beta-actin (ACTB) were used as reference genes. Quantitative PCR (qPCR) was performed using an ABI 7500 Sequence Detector System and the qPCR reaction volume was 12 µL, which steps were 2′ at 50°C and 10′ at 95°C for the maintenance stage, 40 cycles of...
15' at 95°C and 1' at 60°C for the PCR stage; and 15' at 95°C, 1' at 60°C and 15' at 95°C for the melting curve stage. Three independent experiments were performed. For the analysis of relative expression, the values from the Cycle Threshold (CT) average of each gene were used and the values were normalized by the average reference genes values by the 2^{−ΔΔCT} method (Livak and Schmittgen, 2001). The software MeV (Multiple Experiment Viewer10) was used to construct the heatmap. Network construction was performed using modulated genes in both glioma cell lines (1.4-fold-change) and the GeneMANIA database11.

**Statistical Analysis**

Statistical analyses were performed using GraphPad InStat 7.0. (GraphPad Software, Inc., San Diego, CA, United States). For comparisons, ANOVA and Tukey multiple post-test were used. A p-value < 0.05 was considered statistically significant.

**RESULTS**

**Screening of Euherdmania sp. Associated Bacteria**

Since our prior study showed the cytotoxic potential of an extract obtained from the Brazilian endemic ascidian Euherdmania sp. (Figure 1A) (Jimenez et al., 2003), we proceeded with the isolation of four bacteria strains (Figure 1B and Supplementary Table 1) associated with the ascidian and evaluation of the cytotoxicity of their respective extracts in the colorectal carcinoma cells HCT-116. All tested extracts presented cytotoxicity with IC_{50} values varying from 0.03 (BRA-346) to 5.15 µg/mL (BRA-341) (Figure 1C and Supplementary Table 1). The extracts were further evaluated for their inhibitory potential against the proteasome chymotrypsin-like (ChTL) catalytic subunit (Figure 1C) for which only the extract from BRA-346 significantly inhibited this enzymatic activity, with IC_{50} of 0.42 µg/mL (Figure 1D).

Therefore, BRA-346 was selected for further studies, including molecular identification of the bacteria, bioassay-guided fractionation of the extract and mechanism of action (MoA). Through 16S rRNA sequencing and comparison with the EzBioCloud database, BRA-346 was identified as Streptomyces sp. (MW342808), with close similarity (above 99%) to Streptomyces amphotericinicus (KX777593) (National Center for Biotechnology Information, 2020).

The BRA-346 strain was grown in liquid media and extracted with ethyl acetate. The crude extract was then subjected to a fractionation process guided by cytotoxicity against HCT-116 cell line, assessed through the MTT assay (Supplementary Figure 1 and Supplementary Table 2). Ultimately, fraction C, obtained after a SPE step applied to a mid-polarity fraction eluted from the ethyl acetate partition subjected to a Sephadex LH-20 column, was shown to produce the highest cytotoxicity, with IC_{50} of 30 ng/mL, being 20 times more potent than the second most active one, fraction B (Supplementary Figure 1), and was selected to the MoA studies.

**Proteasome Inhibitors Detected in Streptomyces sp. BRA-346 Extract and Fractions**

Studies with BRA-346 fraction C were conducted to confirm the proteasome inhibitory activity and to access its chemical profile. Known proteasome inhibitors, including DHE and epoxomicin, were included in this analysis as positive controls. BRA-346 fraction inhibited 92.6% of the ChTL enzymatic activity at 1 µg/mL, while comparable inhibition of this enzyme by DHE and epoxomicin, by 94.6 and 93.9%, were reached at 12 and 16.6 µg/mL, respectively (Figure 2A), which is equivalent to nearly 30 µM of either compound. BRA-346 fraction showed an IC_{50} of 0.04 µg/mL (Figure 2B), a potency 10 times greater than that of the crude extract (IC_{50} = 0.42 µg/mL) when tested against the proteasome ChTL subunit.

**Chemical Characterization of Eponemycin Analogs**

HPLC-MS/MS analysis of the extract and fractions A, B, C, D, and E were conducted to investigate the chemical nature of the proteasome inhibitor present in the BRA-346 extract. Data was used to build a molecular network using the GNPS platform (Figure 2C). Two known proteasome inhibitors, DHE and epoxomicin, were included in the HPLC-MS/MS analysis for comparison. The fractionation was efficient and allowed the detection and identification of several compounds that were not observed in crude extract. The ion of m/z 401.26, compatible with DHE, was observed in DHE (Figure 2C) and exclusively in fraction C (Figure 2C and Supplementary Figure 2), while epoxomicin compatible signals were not detected in the extract or fractions. The molecular networking actually highlighted the presence of a large family of epoxyketone derivatives mainly in BRA-346 fraction C, but also in other fractions. Therefore, aiming at the identification of these compounds we decided to investigate their fragmentation pathway.

Present MS/MS data and that of Schorn et al. (2014) were used to propose the fragmentation pathways for DHE (Figure 3), and to guide the dereplication of compounds from the same family. As expected, the elimination of water makes up the picture of major ions, since there is an OH group in the central part of the molecule that can be easily eliminated due to its proximity to a more acidic alpha-carbonyl hydrogen. This route is represented in pathway C (Figure 3). The second water elimination could occur directly by a mechanism suggested in pathway E, however, a primary carbocation would have been formed, which does not occur due to its large instability. Thus, the alternative route goes through opening of the epoxide by an initial hydrogen rearrangement, affording an enol ion and presumably balancing the keto form (Li et al., 2007). Through this path, the elimination of a second water (m/z 365) can occur expending low energy. An additional loss of H_{2}CO confirms the actuality of this pathway. Furthermore, elimination of an amide from the same

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10http://mev.tm4.org
11https://genemania.org/
FIGURE 2 | Evaluation of proteasome inhibition by dihydroeponemycin, epoxomicin, and BRA-346 epoxyketone-containing fraction, and molecular networking profile of proteasome inhibitors, BRA-346 crude extract, and fractions. (A) Comparative graph of inhibition of the catalytic activity of the Chymotrypsin-like (ChTL) subunit of the proteasome by dihydroeponemycin and epoxomicin (at 12 and 16.6 µg/mL, respectively), and BRA-346 fraction (fraction C Supplementary Figure 1) (at 1 µg/mL) through the enzymatic assay. Data were analyzed using GraphPad Prism 7.0 software by ANOVA followed by Tukey multiple comparison test, ****p-value < 0.0001. (B) Dose-response curve of the proteasome inhibition test by BRA-346 fraction, with IC$_{50}$ value = 0.045 µg/mL, confidence interval (CI 95%) = 0.02 to 0.07. All values were obtained from three independent experiments. (C) Molecular network profile of dihydroeponemycin, BRA-346 crude extract and fractions A, B, D, and E. Compounds were analyzed by grouping from the fragmentation profile data of the compounds obtained by the ESI positive mode (ESI+). Only clusters related to dihydroeponemycin are shown. Nodes represent the ions present in the samples, which are indicated by related colors. Highlighted nodes refer to dihydroeponemycin (m/z 401.2) and analog ions present in the BRA-346 fractions.
Analyses were carried out on Bruker Daltonic machines and the isolated protonated molecule was excited at a CID chamber applying N₂ as the collision gas. The ion at m/z 109 can also be produced directly after loss of water from the epoxide moiety (fragmentation mechanism is present at Supplementary Material, Supplementary Figure 4 and Supplementary Scheme 2).

Fragmentation pathways A, B, D, and E also start as previously described (Schorn et al., 2014) and some important fragments...
with m/z 214 (A), m/z 258 (B), m/z 188 (D) m/z 275, and m/z 257 (E) can be useful to define possible modifications at the molecule moiety. Additional important information can be assessed by fragmentation of ion m/z 188. After elimination of the side chain through a retro-heteroene reaction (Crotti et al., 2005), the amine has freedom to undergo a long hand displacement reaction (McLafferty and Turecek, 1993), thus affording a water elimination product (ion at m/z 170, Figure 3). Now, two mechanisms for water elimination can occur from the same part of the molecule. The open-ring form of the ion at m/z 170 can lose H₂CO affording a final ion at m/z 140, confirming, once again, opening of the epoxide ring. The cyclic ion at m/z 170 can be opened by an anhmeric assistance (Lopes et al., 2002) and, at the same concerted process, the epoxide is also opened. Sequential loss of water and of CO affords the ion at m/z 96, that can be helpful to find new derivatives in combination with the previously described ions.

Once the main fragmentation mechanisms were defined, we started to analyze the LC-UV-MS/MS data for the structural determination of possible derivatives (Figure 4 and Supplementary Figures 3–13 in the Supplementary Material shows de MS/MS data of all identified compounds). The first step was to confirm the spectrum of DHE (compound 13) detected in BRA-346 fraction C at retention time (tR) 12.0 min. The perfect spectral match between the commercial DHE and the ion at m/z 401 present in BRA-346 fraction C undoubtedly confirmed its occurrence (see spectra in Supplementary Figure 2). A substance with m/z 401 with two losses of water and the ion at m/z 109 with significant intensity (see spectra in Supplementary Figure 3) were observed at tR 10.6 min. This initial analysis suggests that it could be a slightly more polar isomer. Additionally, the second H₂O elimination is the most intense ion in the spectrum (which does not occur in the DHE spectrum) and there is a lower intensity ion from a third H₂O elimination. These information suggest an expected reduction of the carbonyl group in the side chain and a double-bond in the same position as eponemycin 12. MS/MS analysis also shows an ion at m/z 151, which confirms the double-bond, as proposed for compound 9. As expected, eponemycin (m/z 399) was detected at a lower tR (11.4 min) than DHE. Only the initial pathway A shown in Figure 3 could not be observed in the MS/MS analysis for eponemycin (Scheme 2 in Supplementary Material). Also, the ion at m/z 109 was formed directly from the ion at m/z 363 (losses of two molecules of water). The majority of other ions have two mass units less than the observed for DHE, confirming the chemical structure.

Previous work from Schorn et al. (2014) showed several analogs of eponemycin with lower side chains. In the present work compounds with tR at 9.0 min (compound 2, m/z 371), 9.6 min (compound 4, m/z 329), 10.2 min (compound 7, m/z 315), 10.7 min (compound 10, m/z 387), and 10.8 min (compound 11, m/z 313) showed spectral feature consistent with this family of compounds. For all compounds, the major fragmentation pathway starts with the loss of water and ends with the diagnostic ion at m/z 109 (see Supplementary Figure 10). Pathway A (green) presented in Scheme 3 (Supplementary Material) was important to confirm the size of the side chain and a double-bond replacing the hydroxyl group present in DHE. These information suggest N-(1-(2-(hydroxymethyl)oxiran-2-yl)-4-methyl-1-oxopentan-2-yl)-2-propionamidoacrylamide as the structure for compound 11 (Figure 4). For compound 10, the fragment related to the side chain elimination was key for structure elucidation. Therein, demetallation may occur only at the isobutane section and the ion at m/z 109 also confirms that there are no modifications in the epoxide chain. Based on this, we propose compound 10 as N-(3-hydroxy-1-((1-(2-(hydroxymethyl)oxiran-2-yl)-4-methyl-1-oxopentan-2-yl)amino)-1-oxopropan-2-yl)-6-methylheptanamide (Figure 4). Compound 7 has two more mass units than compound 11, but in this case, the hydroxyl group remains in the structure and the side chain has only one alpha carbonyl carbon. Therefore, it was not possible to observe elimination from the side chain and all ions were afforded from the initial pathway C (Scheme 6 in the Supplementary Material). In this case, the ion at m/z 109 was a minor signal, but all ions collectively allowed confirmation of compound 7 as 2-acetamido-3-hydroxy-N-(1-(2-(hydroxymethyl)oxiran-2-yl)-4-methyl-1-oxopent-4-en-2-yl)propionamide (Figure 4). Like 10, compound 4 also owns a demetallation at the same site, probably by an oxidation of the metabolites followed by CO elimination, resulting in a loss of carbon for the molecule moiety. Once again, pathway C is not held as the most important, but pathway D (Figure 3 and Supplementary Scheme 8) contributed to the final elucidation. The initial ion from pathway D has less restrictions and an additional ion can be observed at m/z 130. The mechanism of all other ions formation are in agreement with the proposition displayed in Figure 3 and we confirm the compound as N-(3-hydroxy-1-((1-(2-(hydroxymethyl)oxiran-2-yl)-1-oxopent-4-en-2-yl)amino)-1-oxopropan-2-yl)butyramide. Compound 2 is another compound with an eponemycin core, owning a small side chain and a second carbonyl group. The presence of a second carbonyl group was already described in some analogs annotated by Schorn et al. (2014). The ions observed in the spectra and the major fragmentation pathways (Supplementary Figure 12 and Supplementary Scheme 10, respectively, in Supplementary Material) are fully in agreement with a protonated ion of N-(3-hydroxy-1-((1-(2-(hydroxymethyl)oxiran-2-yl)-4-methyl-1-oxopent-4-en-2-yl)amino)-1-oxopropan-2-yl)-3-oxopentanamide.

The last group of compounds had higher mass values than eponemycin. In all cases, we observed the increased number of oxygen atoms and no modifications in the carbon skeleton. We observed two ions with m/z 417 with identical spectra and their only difference is the tR. Comparison to a previously published spectrum (Schorn et al., 2014) confirms the presence of the same major ions. Detailed analysis in Supplementary Figure 7 and Supplementary Scheme 5 indicate no other possibility than the expected addition of water to the double-bond. In this case, we can assume that one compound is similar to that previously published and the other is an isomer of position. Therefore, we believe that compound 6 can be the first to be eluted, but keeping in mind that this order can be reversed with compound 8. Compound 3 (3-hydroxy-N-(3-hydroxy-1-((1-(2-(hydroxymethyl)oxiran-2-yl)-4-methyl-1-oxopent-4-en-2-yl)amino)-1-oxopropan-2-yl)-6-methylheptanamide) has been
FIGURE 4 | LC-MS/MS analysis of BRA-346 fraction C (BRA-346 epoxyketone-containing fraction) and proposed structures for eponemycin derivatives.
previously identified (Schorn et al., 2014) and the ions observed herein are in full agreement (Supplementary Figure 11 and Supplementary Scheme 9). Compound 5, a minor signal in the HPLC-MS/MS analysis, has an additional hydroxyl group and an expected reduction of the carbonyl group in the side chain. Therefore, we verified the occurrence of pathway E (Supplementary Figure 9 and Supplementary Scheme 7) and the ions at m/z 275 and 255 confirmed we had an unmodified residue of 3-hydroxy-N-(1-(2-(hydroxymethyl)oxiran-2-yl)-4-methyl-1-oxopentan-2-yl)propanamide. Moreover, the ion at m/z 109 reinforces this information. Hence, we propose compound 5 as 2-[(1,3-dihydroxy-6-methylheptyl)amino]-3-hydroxy-N-(1-[(2-hydroxymethyl)oxiran-2-yl]-4-methyl-1-oxopentan-2-yl)propanamide. Finally, compound 1 showed the highest oxidation level for all series. In this case, the core is still the same as eponemycin and a new hydroxyl group was observed in the terminal isobutyl group and in the side chain. While it is not possible to define the position of all groups, it is indeed possible to confirm the presence of a terminal hydroxyl group, which afforded the fragment ion from the heteroene reaction. So, a putative structure is presented in Figure 4 and details on spectrum and fragmentation pathways are offered in the Supplementary Figure 13 and Supplementary Scheme 11.

Cytotoxicity and Molecular Markers of Proteasome Inhibition of DHE and Streptomyces sp. BRA-346 Epoxyketone-Containing Fraction: Studies in Glioma Cells

The activities of DHE and Streptomyces sp. BRA-346 epoxyketone-containing fraction were further evaluated in glioma cell lines HOG and T98G, since proteasome inhibitors are emerging as promising compounds in the treatment of glioblastoma multiforme, which remains a poorly manageable disease with high mortality rates (World Health Organization, 2020). The analysis was based on the assessment of cytotoxicity and the evaluation of molecular markers related to proteasome inhibition. Table 1 and Supplementary Figure 14 show the results of the SRB analysis after 24, 48, and 72 h incubation. Generally, growth inhibitory activity was amplified 10-fold when incubation time increased from 24 to 48 h for both DHE and BRA-346 epoxyketone-containing fraction but remained constant when the incubation time was further increased to 72 h. The sensitivity of both cell lines was similar for DHE, with 48 h incubation GI_{50} values of 1.6 and 1.7 ng/mL for HOG and T98G cells, respectively, while T98G cells were slightly less sensitive to BRA-346 epoxyketone-containing fraction, with GI_{50} of 28.2 ng/mL when compared to 17.6 ng/mL in HOG cells.

To investigate the molecular changes incited by DHE and BRA-346 epoxyketone-containing fraction, glioma cells were exposed to the samples and incubated for 24, 36, or 48 h. Protein expression analysis demonstrated an increase in HSP70 and ubiquitinated proteins levels in both HOG and T98G cells treated with the BRA-346 epoxyketone-containing fraction, but no significant changes were observed when cells were treated with DHE at GI_{50} levels (Figures 5A,B and Supplementary Figure 15). Nonetheless, at concentration 25 times higher (40 ng/mL or 100 nM, Figure 5C) and 36 h of incubation, DHE induced comparable cellular effects with those of BRA-346 epoxyketone-containing fraction at 17.6 and 28.2 ng/mL in HOG and T98G cells, respectively (Figures 5A,C and Supplementary Figure 15).

We then turn to evaluate gene expression of treated cells, revealing a similar profile for both DHE at 25 times GI_{50} and BRA-346 epoxyketone-containing fraction (Figures 6A,C). DHE, at GI_{50} levels, had limited effects on the genes analyzed as also observed for protein alterations. Genes with a 1.4-fold-change were included in the network by GeneMANIA (see text footnote 11), which highlighted UPR as the main modulated mechanism for both DHE and BRA-346 fraction (Figures 6B,D). At 40 ng/mL, among the sixteen genes analyzed, DHE increased the expression of 9 and 13 genes in HOG and T98G cells, respectively (Figures 6B,D and Supplementary Table 5). BRA-346 epoxyketone-containing fraction, on the other hand, upregulated 12 and 5 genes at 17.6 and 28.2 ng/mL.

### Table 1 | Evaluation of cell growth inhibition of HOG and T98G cell lines exposed to dihydroeponemycin (DHE) and BRA-346 epoxyketone-containing fraction through SRB assay after 24, 48, and 72 h treatment.

| Substances          | HOG | T98G |
|---------------------|-----|------|
|                     | GI_{50} | TGI  | LC_{50} | GI_{50} | TGI  | LC_{50} |
| Dihydroeponemycin   |       |      |         |       |      |         |
| 24 h                | 15.2 (19.8) | 333.9 (377.7) | – | 16.1 (40.2) | 694.9 (1735.0) | – |
| 48 h                | 1.6 (4.0) | 18.9 (47.3) | 221.9 (554.1) | 1.7 (4.2) | 32.6 (81.5) | 630.0 (1573.1) |
| 72 h                | 1.5 (3.8) | 19.9 (49.7) | 262.6 (655.8) | 1.5 (3.7) | 15.8 (39.5) | 169.3 (422.6) |
| BRA-346 fraction    |       |      |         |       |      |         |
| 24 h                | 232.3 | 6135.4 | – | 197.2 | 12757.5 | – |
| 48 h                | 17.6 | 201.6 | 2307.0 | 28.2 | 404.3 | 5800.5 |
| 72 h                | 29.1 | 297.3 | 3033.4 | 7.8 | 105.2 | 1414.1 |

The mean values in ng/mL (nM) were obtained from 3 independent experiments and reflect concentrations calculated for half-maximum growth inhibition (GI_{50}), total growth inhibition (TGI) and half-maximum lethal concentration (LC_{50}). Some time-points did not allow the acquisition of values of LC_{50}. 
FIGURE 5 | Dihydroeponemycin (DHE) and BRA-346 epoxyketone-containing fraction led to the accumulation of ubiquitinated proteins and HSP70 in glioma cell lines HOG and T98G. (A) Glioma cells treated with BRA-346 fraction at GI50 levels (17.6 and 28.2 ng/mL for HOG and T98G, respectively) for 3 different times of exposition (24, 36, and 48 h). (B) Glioma cells treated with DHE at GI50 levels (1.6 and 1.7 ng/mL for HOG and T98G, respectively) for 3 different times of exposition (24, 36, and 48 h). (C) Glioma cells treated with DHE at 40 ng/mL (100 nM) for 36 h of exposure. Negative controls (DMSO 0.1%) were set at 0, 36, and 48 h. Experiments were repeated three times.

in HOG and T98G cell lines, respectively. Six genes were upregulated for both treatments in HOG – ATF6, CALR, DDIT3, HSPA4, HSPA5, and PPP1R15A – and five genes – CANX, DDIT3, HSPA5, MAPK8, and PPP1R15A – in T98G cells (Figure 7). The analysis of functions related to the modulated genes showed that both DHE and BRA-346-fraction are inducing a UPR phenotype, as expected by the inhibition of proteasome function, and ultimately leading treated cells to apoptosis (Figure 7).

DISCUSSION

Marine bacteria have been studied in the last years due to their large potential to produce molecules with interesting biological activities, including antibacterial, antifungal, antitumor, anti-inflammatory, antimalaria, and antiviral actions (Manivasagan et al., 2014). The chemical diversity of such bioactive marine natural products range from peptides, alkaloids, terpenes, lactones, and steroids (Hu et al., 2015) to complex secondary metabolites that have offered new mechanisms of action to be untangled by pharmacological sciences (Reen et al., 2015). Of the four extracts tested, three displayed a relevant cytotoxicity against HCT-116 cells and may be promising sources of anticancer compounds. However, the extract derived from BRA-346, besides its high cytotoxicity, also presented a potent inhibition of proteasome ChTL activity. Proteasome inhibitors have been shown to be important pharmacological resources for the treatment of central nervous system tumors. Therefore, this extract was selected herein for a further investigation of its chemical composition and antiglioma potential.

Herein, we present a Streptomyces bacterium recovered from an ascidian collected in the northeast coast of Brazil, Euherdmania sp., produces antiproteasome compounds. Evaluation of the most potent fraction obtained through a bioassay-guided fractionation protocol of this extract revealed potent proteasome inhibitory properties along with the presence of a family of eponemycin related compounds, including dihydroeponemycin (DHE). Both DHE and the bacteria epoxyketone-containing fraction showed potent cytotoxicity against human oligodendrogioma and glioblastoma cell lines (HOG and T98G, respectively), resulting in significant changes in expression levels of genes and proteins related to ER-stress, unfolded protein response (UPR) and the ubiquitin-proteasome system (UPS).

The proteasome is a multicatalytic enzyme complex dedicated to recycling intracellular proteins. The main function of such organelle is to degrade damaged or unnecessary proteins, thus maintaining cell proteostasis (Voges et al., 1999; Bard et al., 2018). In the context of cancer, a malady inflicted by intense protein synthesis at the cellular level, the proteasome has been established as a valuable target, as its downmodulation implies poor handling of unwanted and unfolded proteins, along with further cell perturbations leading to cell-apoptosis (Manasanch and Orlowski, 2017; Jang, 2018). Importantly, proteasome inhibitors display a favorable therapeutic window, selectively killing cancer cells in humans, and are within the most tolerated chemotherapeutic options for multiple myeloma, for example. Proteasome inhibitors that reached the clinic are
FIGURE 6 | Molecular changes promoted by dihydroeponemycin (DHE) and BRA-346 epoxyketone-containing fraction in glioma cells. (A,C) Heatmap of gene expression of HOG and T98G cells treated with DHE and BRA-346 epoxyketone-containing fraction for 36h. DHE was evaluated at GI$_{50}$ ($n = 1$) and 25xGI$_{50}$ (40 ng/mL) ($n = 3$), while the BRA-346 fraction was tested at GI$_{50}$ levels (17.6 and 28.2 ng/mL) ($n = 3$). Values for relative expression were established in comparison to control values and the heatmap was constructed by MeV 4.9.0 software (http://mev.tm4.org). All genes analyzed are related to ER-stress, except for BAX, which mostly relates to apoptosis. (B,D) The networks illustrate interactions of 1.4-fold-change in genes modulated by DHE at 40 ng/mL and BRA-346 epoxyketone-containing fraction (17.6 and 28.2 ng/mL) treatment in HOG and T98G. The up-regulated genes in the quantitative PCR analysis are illustrated as crosshatched circles and the interacting genes included by software modeling are indicated by non-crosshatched circles. The main networks and function of genes are indicated by colored lines and color in circles are described in the legend.
small peptides with an electrophilic warhead that can bind to the catalytic subunits of the proteasome and impair the proper function of the enzymatic complex (Micale et al., 2014; Park et al., 2018). Examples are bortezomib and carfilzomib, the latter belonging to the epoxyketone class, which was inspired by the actinobacteria-derived natural product epoxomicin. Many other bacterial natural products have been reported with these features, including carmaphycin, eponemycin, and other non-ribosomal peptides. These compounds form covalent bonds with threonine residues in the N-terminal region present in the catalytic active β subunits of the proteasome, leading to irreversible binding and, thus, inhibition of proteasome activity (Borissenko and Groll, 2007). Proteasome inhibitors use, including bortezomib and carfilzomib, is indicated for multiple myeloma patients, but many studies have highlighted its potential in the treatment of gliomas (Areeb et al., 2016; Wang et al., 2018; Tang et al., 2019).

Nonetheless, the proper functioning of the ubiquitin-proteasome system is also relevant for other nervous system diseases, including neurodegenerative ones, where the accumulation of unfolded or mutated proteins generating protein aggregates and inclusion bodies trigger the death of neurons (Cacabelos, 2017; Montaz et al., 2020). Thus, proteasome inhibitors may impact the therapy of many diseases, highlighting the importance of better understanding their mechanism of action and the pathways modulated by these compounds in treated cells.

The occurrence of the epoxyketone proteasome inhibitor, dihydroepoxymycin (DHE), in fractions obtained from the Streptomyces sp. BRA-346 extract was suggested by comparison of the HPLC-MS/MS spectra with the commercially available DHE. In fact, detailed dereplication studies showed the presence of several ions related to eponemycin in BRA-346 fraction. Eponemycin was first isolated from the extract of the bacterium Streptomyces hygroscopicus, recovered from a soil sample from the Philippines, showing cytotoxicity toward HCT116 cells with IC₅₀ value of 9.7 ng/mL (Sugawara et al., 1990). Herein, BRA-346 epoxyketone-containing fraction restrains proliferation of HCT116 cells, displaying IC₅₀ value of 30 ng/mL. The activity of this fraction against the proteasome ChTL catalytic subunit was also confirmed by enzymatic assay, revealing an IC₅₀ value of 45 ng/mL. DHE and tetrahydroepoxymycin was first described

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**FIGURE 7** Schematic representation of upregulated UPR-related genes in HOG and T98G cells by DHE or BRA-346 epoxyketone-containing fraction. Each gene selected for validation acts on the unfolded protein response (UPR) and endoplasmic reticulum stress (ER-stress) and shows different regulation in HOG and T98G cells for each treatment. Treatments were performed with DHE at 40 ng/mL (both cell lines) and BRA-346 epoxyketone-containing fraction at 17.6 and 28.2 ng/mL for HOG and T98G, respectively. Genes in green were found to be upregulated following exposure to both DHE and BRA-346 epoxyketone-containing fraction; genes in red were upregulated only by BRA-346 epoxyketone-containing fraction; genes in pink were upregulated only by DHE; and genes in black were not upregulated. Data of relative expression were obtained from average values for each gene normalized by the average values for reference genes by the \(2^{-\Delta\Delta CT}\) method from three experiments (Supplementary Table 5).
as the product of hydrogenation of eponemycin over palladium carbon (Sugawara et al., 1990), and while DHE showed IC<sub>50</sub> of 13 ng/mL in HCT116, tetrahydroeponemycin was not active at 1 µg/mL, reinforcing the importance of the epoxy group in the biological activity. DHE was also shown to inhibit the proteasome activity in micromolar concentrations (Meng et al., 1999). More recently, Schorn et al. (2014) described the biosynthetic gene cluster responsible for the production of eponemycin and, through heterologous expression in Streptomyces albus J1046, described several putative epoxyketone derivatives, including DHE. Herein, we compare the obtained MS/MS data of BRA-346 fraction C with that described by Schorn et al. (2014). As expected, a comparison of MS/MS data with previously published spectra shows there are slight differences in the initial fragmentation pathways (Schorn et al., 2014). Distinct ESI-MS/MS equipments can apply a variety of collision gases which results in a non-linear transfer of collisional energy in the CID chamber, therefore resulting in different numbers of initiating pathways since the protonated molecule (Demarque et al., 2016). All ions observed have their fragmentation pathways rationalized through the ESI-CID-MS/MS gas-phase chemistry concepts (Demarque et al., 2016). Thus, we started our analysis by proposing the expected fragmentation pathways from DHE and compared MS/MS data for the ions grouped in the same cluster, leading to the identification of thirteen eponemycin analogs in BRA-346 fraction C (designated, from now on, as BRA-346 epoxyketone-containing fraction), including DHE and eponemycin itself.

Meng et al. (1999) described that DHE antitumor activity, as also observed for structurally related compounds, is explained through inhibition of the proteasome function, but epoxyketone-containing antitumor natural products may differ in their antiproliferative activity, proteasome subunit binding specificity, and rates of proteasome inhibition. Considering the intriguing structure-activity relationship for this class of compounds (Kim et al., 1999; Groll et al., 2000) and the emerging knowledge on the antiangioma effects of the proteasome inhibitors (Di et al., 2016; Zhang et al., 2018; Veggiani et al., 2019), we further assayed DHE and BRA-346 epoxyketone-containing fraction in glioma cells HOG and T98G, revealing their potent antiproliferative effects in the ng/mL-range and the modulation of ER-stress related and pro-apoptotic related genes, along with unfolded protein response and the ubiquitin-proteasome system itself.

BRA-346 epoxyketone-containing fraction at GI<sub>50</sub> levels promoted an increased expression of HSP70 and ubiquitinated proteins, while DHE required higher concentrations, corresponding to 25 times the GI<sub>50</sub>, to achieve a comparable effect. Accumulation of ubiquitinated proteins is regarded as one hallmark feature of proteasome inhibition, since misfolded proteins are labeled by multiple ubiquitin molecules to be degraded by the proteasome (Myung et al., 2001). Therefore, proteasome inhibitors prevent the degradation of ubiquitinated proteins, promoting an accumulation of these in the cytoplasm, as demonstrated in HUVECs and neuronal cells (Canu et al., 2000; Mye ku and Figueiredo-Pereira, 2011). Furthermore, HSP70 is a family of chaperones related to proper functioning and maintenance of the cell as it supports the folding of new proteins, improving the transport of proteins out of the ER. The high expression levels of such chaperones in both cell lines promoted by proteasome inhibitors, as observed herein for both DHE and BRA-346 epoxyketone-containing fraction, can be interpreted as the ultimate attempt to restore homeostasis by handling excessive unwanted proteins, which, in turn, may be a consequence of proteasome inhibition (Bush et al., 1997; Mayer and Bukau, 2005). The difference in the potency of DHE and the BRA-346 fraction, considering the relation of GI<sub>50</sub> levels and the accumulation of ubiquitinated proteins, may be explained by the presence of several epoxyketone in the fraction, some owning, supposably, higher potencies that enhance the observed effects. The presence of other compounds not related to eponemycin cannot be excluded at this moment, since the studied fraction is indeed very complex.

That said, we then turned to evaluate the effects of DHE and BRA-346 epoxyketone-containing fraction on genes related to pathways that showed modulation of ER-stress response in gliomas (Markouli et al., 2020). While notable differences were observed between responses elicited by each cell line, a similar molecular profile, in which most of the assessed genes were found to be upregulated by both DHE and the BRA-346 epoxyketone-containing fraction was revealed, reflecting a compensatory mechanism undergoing in cells exposed to either treatment to overcome ER disturbances (Kadowaki and Nishitoh, 2013). Figure 7 illustrated the genes upregulated for both treatments (in green) or individually (DHE: pink; BRA-346 fraction: red) in both cell lines. Despite slight differences, the main functions impaired by either treatment are related to unfolded protein response (UPR) in glioma cell lines. Indeed, UPR is triggered by an overload of unfolded proteins in the ER lumen and accounts, in principle, for a transitory state aimed at re-establishing normal ER function, mainly characterized by synthesis, maturation, and export of proteins (Walter and Ron, 2011). Still, as many natural compounds have been shown to induce transitory or sustained UPR, we wish to highlight that to the best of our knowledge, this is the first report on the effects of DHE on glioma cells assessed to this extent and, moreover, in comparison to a complex natural fraction containing a myriad of structurally related epoxyketones.

HSPA5 refers to the gene that encodes GRP78 or BiP protein, a key chaperone for newly synthesized proteins, supporting co-translational modifications, which is an essential component of the UPR and an important target to cancer therapy (Wang et al., 2009; Lu et al., 2020). As a matter of fact, GRP78, a member of the HSP70 family of proteins, is the common initial response that detaches and activates the downstream ER-stress sensors, therefore a typical marker to UPR (Kopp et al., 2019). However, different cells may, indeed, launch then handle UPR in distinct manners, even if subjected to similar stressors, as the intrinsic capacity for protein synthesis varies among cell types (Bertolotti et al., 2000; Carrara et al., 2013). In thyroid cancer cells, proteasome inhibitors such as PSI, epoxomicin and MG132 induced upregulation of GRP78 and CHOP at mRNA and protein levels (Wang et al., 2007). HSPA4, in turn, encodes a large heat-shock chaperone from the HSP110 family, HSPA4 (also referred to as Apg-2), which works in...
concert with HSP70s to restore proper protein folding activity and counter ER-stress (Zuo et al., 2016). While HSPA4 has not been as broadly assessed as their closely related HSP70 assemblage, it has been associated to maintenance of neuronal functions under stress conditions (Okui et al., 2000) and a significant correlation was observed between the expression of this chaperone and increased levels of the antiapoptotic marker Bcl-2 in inflammatory cells, further suggesting an interplay of the protein in cell survival pathways (Adachi et al., 2015). Herein, both genes, HSPA4 and HSPA5, which respond to misfolded or unfolded proteins, as shown in the networks in **Figure 6**, were found upregulated in HOG cells exposed to either DHE or BRA-346 epoxyketone-rich fraction, indicating these cells are under proteotoxic stress due to proteasome malfunction and signaling an attempt to neutralize misfolded proteins. In T98G cells, however, expression of HSPA4 was not affected by the treatments. Thus, it may be suggested that T98G cells represent a more resistant cell line to ER-stress elicited by compounds of this chemical class. Indeed, the product of the XBPI gene, a typical downstream response of the dimerization and activation of the ER transmembrane protein IRE-1 UPR sensor further stimulating the UPR (Ron and Walter, 2007; Kadowaki and Nishitoh, 2013), also remained mostly unaltered in T98G cells.

PERK is a second transmembrane protein that acts as a sensor for ER-stress and, mainly, blocks general protein translation and cell cycle while also stimulating antioxidant systems to compensate for ER saturation. ATF4, a transcription factor encoded by the gene ATF4, is a key downstream marker for activation of this pathway which can contribute to survival or cell death (Okada et al., 2002; Iurlaro and Muñoz-Pinedo, 2016; Rozpedek et al., 2016). Studies highlighted the modulation of PERK-eIF2a-ATF4 pathway by proteasome inhibitors like bortezomib in human myeloma and squamous carcinoma cell lines, evidencing that this class of compounds may activate UPR-related molecular markers (Fribley et al., 2009; Iurlaro and Muñoz-Pinedo, 2016). Furthermore, the increase in CHOP by bortezomib, amplifies cytotoxicity in multiple myeloma and breast cancer cells when combined with inhibitors of early and intermediate autophagy processes, modulating apoptosis and the autophagic pathway (Kawaguchi et al., 2011; Komatsu et al., 2012). Actually, autophagy, the lysosome-mediated degradation of defective proteins, protein aggregate and even damaged organelles, is also included among cell survival strategies during ER stress, and is actively interconnected to UPR (Rashid et al., 2015). The changeover signal between pathways, however, is largely unknown. Still, besides CHOP, MAPK8 and closely related MAPK9, also known by JNK-1 and JNK-2, respectively, may be partly to blame in such a process and the duration of the signaling may be determinant cell fate (Wei et al., 2008). Interestingly, the treatments employed herein have distinguished between the expression of both these genes, as only DHE notably increased expression of MAPK9, but MAPK 8 increased under either treatment.

Nonetheless, unrepaired or aggravation of ER-stress promotes cell death for the benefit of the organism and, in such a situation, UPR is no longer portrayed as a survival mechanism (Tabas and Ron, 2011; Hetz, 2012; Urrea et al., 2013). Alike launching and handling, there is also variability considering the stress levels a particular cell type can endure before veering UPR to apoptosis (Luo and Lee, 2013; Peñaranda Fajardo et al., 2016). In this sense, overactivation of the UPR, either through proteasome inhibition or by other mechanisms, has been a recurring subject of investigation and also used therapeutically to treat cancers. Moreover, many natural products have been shown to induce ER-stress-mediated apoptosis (Kim and Kim, 2018), highlighting these compounds for their remarkable anticancer potential.
CONCLUSION

Herein, we described the recovery of the ascidian-associated bacterium Streptomyces BRA-346 that produces dihydroepromemycin and related compounds with proteasome inhibitory activity. Furthermore, we described the potent cytotoxicity and the pathways modulated by DHE and BRA-346 epoxyketone-containing fraction in glioma cell lines. While the present study advances the knowledge on the mode of action of DHE in gliomas cells, reinforcing that the antiproteasome effect is indeed in play thereof, then leading to ER-stress reflected by manifestation of UPR, it also reveals comparable signaling pathways are activated following exposure of cells to the epoxyketone-rich fraction. Nonetheless, at this point, we cannot exclude the participation of other minor compounds, including dihydroepromemycin derivatives, also present in the fraction in the displayed cellular effects.

DATA AVAILABILITY STATEMENT

The datasets generated for this study can be found in online repositories. The names of the repository/repositories and accession number(s) can be found below: https://www.ncbi.nlm.nih.gov/genbank/, MW342808.

AUTHOR CONTRIBUTIONS

All authors listed have made a substantial, direct and intellectual contribution to the work, and approved it for publication.

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