Tubastatin A, an inhibitor of HDAC6, enhances temozolomide-induced apoptosis and reverses the malignant phenotype of glioblastoma cells

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Introduction

Glioblastoma or grade IV astrocytoma is the most common form of glioma in adults, accounting for 60-70% of all gliomas, as well as the most aggressive brain tumor (1). Glioblastoma treatment currently involves extensive surgical resection followed by external-beam radiation and concomitant temozolomide chemotherapy (2). Temozolomide is an alkylating agent that forms O6-methylguanine in DNA, which mis-pairs with thymine during the next DNA replication cycle (3). These critical lesions progress to lethal DNA cross-links, which inhibit cell replication and result in cell death (4,5). Despite this therapeutic effort, numerous glioblastoma cases develop resistance to chemotherapy, which results in a poor prognosis for the patients, who only have a median survival time of 14.6 months after diagnosis (5). O6-methylguanine-DNA methyltransferase (MGMT), a DNA repair enzyme, is responsible for inducing temozolomide resistance (6,7). MGMT removes alkyl groups from the O6 position of guanine (8). MGMT promoter methylation has been observed in 40-57% of glioblastoma cases; additionally, when this occurs, MGMT is not transcribed and cannot repair DNA damage caused by temozolomide. Left unrepaired, these chemotherapy-induced lesions trigger cytotoxicity and apoptosis, thus resulting in more efficient treatment (9). For this reason, MGMT promoter methylation status can be considered as an important predictive factor for good therapeutic response and hence survival of patients with glioblastoma.

Histone deacetylases (HDACs) are primarily involved in the deacetylation of histones, but a number of HDACs, including HDAC6, can also affect the function of cytoplasmic non-histone proteins, becoming key regulators of cancer signaling pathways (10,11). A major substrate of HDAC6 is acetylated α-tubulin (10), the structural protein of the microtubules that form a whole variety of cellular structures, including the primary cilium (12). HDAC6 overexpression, which has been demonstrated to occur in numerous glioblastoma cases (13), rapidly deacetylates α-tubulin, resulting in depolymerization of microtubules and disruption of the primary cilium (12).

Abstract. Glioblastoma or grade IV astrocytoma is the most common and lethal form of glioma. Current glioblastoma treatment strategies use surgery followed by chemotherapy with temozolomide. Despite this, numerous glioblastoma cases develop resistance to temozolomide treatments, resulting in a poor prognosis for the patients. Novel approaches are being investigated, including the inhibition of histone deacetylase 6 (HDAC6), an enzyme that deacetylates α-tubulin, and whose overexpression in glioblastoma is associated with the loss of primary cilia. The aim of the present study was to treat glioblastoma cells with a selective HDAC6 inhibitor, tubastatin A, to determine if the malignant phenotype may be reverted. The results demonstrated a notable increase in acetylated α-tubulin levels in treated cells, which associated with downregulation of the sonic hedgehog pathway, and may hypothetically promote ciliogenesis in those cells. Treatment with tubastatin A also reduced glioblastoma clonogenicity and migration capacities, and accelerated temozolomide-induced apoptosis. Finally, HDAC6 inhibition decreased the expression of mesenchymal markers, contributing to reverse epithelial-mesenchymal transition in glioblastoma cells.

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Abbreviations: EMT, epithelial-mesenchymal transition; HDAC6, histone deacetylase 6; MGMT, O6-methylguanine-DNA methyltransferase

Key words: HDAC6, tubastatin A, temozolomide, glioblastoma, sonic hedgehog
Aberrant ciliogenesis is a common defect, as determined in five glioblastoma cell lines, which may contribute to the phenotype of these malignant cells (14,15). Cilia have been implicated in numerous signaling pathways important in embryonic development and disease, including the Hedgehog (Hh) (16,17), Wnt (18) and platelet-derived growth factor (17) pathways.

In vertebrate cells, the sonic Hh pathway requires primary cilia (19). A probable mechanistic function of the cilium is to regulate the Hh pathway by increasing the local concentration and bringing pathway components together for key protein-protein interactions required for regulation, including the presence of Smo in the primary cilium being required for the genesis of Gli activated forms, while the absence of Smo will generate Gli repressor forms (20-22). Among Gli targets, a number of proteins, including cyclins D1 and D2, insulin like growth factor binding protein-6, B-cell lymphoma-2, GLI1 and Myc-N, can be observed (20). Furthermore, GLI1 has been associated with increased expression of MGMT (23). Cyclopamine, an effective Smo antagonist, competitively binds the Smo receptor and subsequently inhibits the Hh pathway (23).

However, those cancer cells that lack cilia would not be responsive to this type of Smo inhibitors, and instead would be required to be treated with downstream inhibitors, such as the Gli antagonists (21). Based on this, only ciliated cells, or cells that had been previously exposed to cilia formation promoters, such as tubastatin, would respond to cyclopamine.

Epithelial-mesenchymal transition (EMT) is the biological process by which cells lose their epithelial characteristics and acquire a mesenchymal cell phenotype, which includes enhanced migratory capacity, invasiveness, elevated resistance to apoptosis and increased production of extracellular matrix components (22,24-26) (Table I). EMT is considered a promoter of metastasis, due to the transformations through which cells acquire motility (27). EMT is induced by growth factors, including transforming growth factor (TGF)-1, that decreases expression of E-cadherin (25,28,29), but at the same time increases expression of HDAC6 (30). Following this reasoning, inhibition of HDAC6 by a selective inhibitor, such as tubastatin A, would reduce TGF-induced downregulation of E-cadherin, and therefore EMT.

For the present study, two glioblastoma cell lines, LN405 and T98G, were treated with a selective HDAC6 inhibitor, tubastatin A, to determine whether this treatment modulates the sonic Hh pathway, reduces tumor cell clonogenicity and migration capacities, counteracts EMT and sensitizes glioblastoma cells to chemotherapy with temozolomide.

### Materials and methods

**Glioblastoma cell lines.** The present study was performed using T98G and LN405 glioblastoma cell lines in all experiments conducted. T98G cells were obtained from the European Collection of Cell Cultures (Salisbury, UK) and are derived from a glioblastoma multiform tumor from a 61-year-old Caucasian male. LN405 cells were purchased from The Leibniz-Institute DSMZ (German Collection of Microorganisms and Cell Cultures, Braunschweig, Germany) and correspond to glioblastoma cells established from an astrocytoma tumor (grade IV) of a 62-year-old female in 1986. The cell lines were cultured in Gibco RPMI-1640 GlutaMAX™ medium, supplemented with 10% fetal bovine serum (FBS) and 1% penicillin/streptomycin (all from Thermo Fisher Scientific, Inc., Waltham, MA, USA). These cells were grown as a monolayer in 75 cm² flasks and maintained in an incubator at 37°C in an atmosphere containing 5% CO₂.

**MTT (thiazolyl blue tetrazolium bromide) assay.** The MTT tetrazolium reduction assay is frequently used to estimate the number of viable eukaryotic cells and calculate the median lethal dose (LD50) for screening collections of compounds to determine if the test molecules exhibit direct cytotoxic effects that eventually result in cell death. Viable cells with active mitochondrial metabolism convert MTT substrate into a purple colored formazan product with an absorbance maximum ~570 nm. When cells die, they lose the ability to convert MTT into formazan, thus color formation serves as a useful and convenient marker (presumably directly proportional) of only the viable cells. This reaction generally involves NADH as a cofactor (31). This method requires the incubation of the MTT substrate at a final concentration of 0.5 mg/ml for 1.5 h at 37°C, with a population of viable cells that have previously been cultured in 96-well plates until reaching a confluence of 5,000 cells/well, and treated with each tested drug: For cyclopamine and tubastatin A the concentrations used were 0, 5, 10, 15, 20, 25, 30, 35, 40, 45 and 50 μM; while for temozolomide the concentrations used were 0, 50, 100, 150, 200, 250, 300, 350, 400, 450, 500, 550, 600, 650, 700, 750, 800, 850, 900, 950 and 1000 μM. After 1.5 h, MTT substrate was discarded to avoid cytotoxicity due to the reagent [as the conversion to formazan by cells in culture is time-dependent (31)] and dimethyl sulfoxide (DMSO) was added to each sample. The

**Table I. Common epithelial and mesenchymal markers for epithelial-mesenchymal transition evaluation.**

<table>
<thead>
<tr>
<th>Types of proteins or RNA</th>
<th>Epithelial markers</th>
<th>Mesenchymal markers</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cell-surface proteins</td>
<td>E-cadherin</td>
<td>N-cadherin</td>
</tr>
<tr>
<td></td>
<td>ZO-1</td>
<td>OB-cadherin</td>
</tr>
<tr>
<td></td>
<td></td>
<td>α5β1 integrin</td>
</tr>
<tr>
<td>Cytoskeletal markers</td>
<td>Cytokeratin</td>
<td>Vimentin</td>
</tr>
<tr>
<td>ECM proteins</td>
<td>α1 (IV) collagen</td>
<td>β-catenin</td>
</tr>
<tr>
<td></td>
<td>Laminin 1</td>
<td></td>
</tr>
<tr>
<td>Transcriptional factors</td>
<td>Snail</td>
<td></td>
</tr>
<tr>
<td>miRNAs</td>
<td>miR-200 family</td>
<td>ZEB1</td>
</tr>
<tr>
<td></td>
<td>miR-10b</td>
<td>Slug</td>
</tr>
<tr>
<td></td>
<td>miR-21</td>
<td></td>
</tr>
</tbody>
</table>

ZO-1, Zonula occludens-1; Slag., Slag family transcription repressor 2; ZEB1, zinc finger E-box binding homeobox 1; ECM, extracellular matrix; miRNAs, microRNAs.
resulting absorbance was monitored at 550 nm wavelength using the Multiskan EX reading spectrophotometer. Following MTT tests, cyclopamine, tubastatin A and temozolomide were used in LN405 cells at final concentrations of 27.5, 32.5 and 400 µM, respectively. In T98G cells, cyclopamine and tubastatin A were used at final concentrations of 20 and 30 µM, respectively. Treatments were conducted at 37˚C for 72 h. Temozolomide LD50 could not be achieved in T98G cells; therefore, a final concentration of 400 µM was selected (Table II).

**2D colony formation assay.** The aim of the 2D colony formation assay was to investigate the attachment-dependent growth of cells when exposed to different drugs. Cells were treated at 37˚C for 72 h as follows: LN405 with 27.5 µM cyclopamine, 32.5 µM tubastatin A and 400 µM temozolomide; and T98G with 20 µM cyclopamine, 30 µM tubastatin A and 400 µM temozolomide. Concentrations and durations for the combination of cyclopamine and tubastatin A, and the combination of temozolomide and tubastatin A, were as for the single treatments. Additionally, 1% DMSO was used as a vehicle control. Subsequently, 300 cells/well were cultured in six-well agarose plates, with 3 wells/condition at 37˚C in a 5% CO₂ incubator for 10 days. Subsequently, cells were fixed with 4% paraformaldehyde for 40 min at room temperature and stained with crystal violet (Sigma-Aldrich; Merck KGaA, Darmstadt, Germany) for 15 min at room temperature. Resulting colonies were counted using a Suntex 560 Colony Counter (Gemini, Apeldoorn, The Netherlands).

**3D colony formation assay in soft agar.** The 3D colony formation assay was conducted to investigate the attachment-independent growth capacity of the cell lines when exposed to different drugs. Cells were treated at 37˚C for 72 h as follows: LN405 with 27.5 µM cyclopamine, 32.5 µM tubastatin A and 400 µM temozolomide; and T98G with 20 µM cyclopamine, 30 µM tubastatin A and 400 µM temozolomide. Concentrations and durations for the combination of cyclopamine and tubastatin A, and the combination of temozolomide and tubastatin A, were as for the single treatments. Additionally, 1% DMSO was used as a vehicle control. Subsequently, 10,000 cells/well were cultured in six-well agarose plates, with 3 wells/condition at 37˚C in a 5% CO₂ incubator for 2 weeks. Previously, 2 ml agarose 0.5% (cat. no. 8016; Pronadisa, Laboratorios Conda, Torrejón de Ardoz, Madrid, Spain) with Dulbecco’s modified Eagle’s medium (DMEM; Sigma-Aldrich; Merck KGaA) were added to the wells. When this first layer had gelled, 5,000 cells/well, with 4 wells/condition. After 24 h, a scratch was produced in the middle of the well and medium was changed to one containing 2.5% FBS in order to avoid proliferation and apoptosis of the cells. Images were captured at 0, 8, 24, 32 and 48 h after scratching with a Nikon SMZ18 light microscope, at x10 magnification.

**Wound healing assay.** This assay was performed to investigate the migration capacity of the cells following treatment. Cells were treated at 37˚C for 72 h as follows: LN405 with 27.5 µM cyclopamine, 32.5 µM tubastatin A and 400 µM temozolomide; and T98G with 20 µM cyclopamine, 30 µM tubastatin A and 400 µM temozolomide. Concentrations and durations for the combination of cyclopamine and tubastatin A, and the combination of temozolomide and tubastatin A, were as for the single treatments. Additionally, 1% DMSO was used as a vehicle control. Subsequently, cells were cultured at 37˚C in a 5% CO₂ incubator in 24-well plates, at a concentration of 250,000 cells/well. After 24 h, a scratch was produced in the middle of the well and medium was changed to one containing 2.5% FBS in order to avoid proliferation and apoptosis of the cells. Images were captured at 0, 8, 24, 32 and 48 h after scratching with a Nikon SMZ18 light microscope, at x10 magnification.

**Cell death detection ELISAPLUS.** The Cell Death Detection ELISAPLUS (Roche Diagnostics GmbH, Mannheim, Germany; cat. no. 11544675001) was used in order to investigate the effect of different drugs on apoptosis. This assay is based on a quantitative sandwich-enzyme-immunoassay-principle using mouse monoclonal antibodies directed against DNA and histones. This allows the specific determination of mono- and oligonucleosomes in the cytoplasmic fraction of cell lysates. For this experiment, cells were cultured at 37˚C in a 5% CO₂ incubator in 96-well plates at a concentration of 5,000 cells/well, with 4 wells/condition. After 24 h, treatments were added to each corresponding well, and cells were treated at 37˚C for 72 h as follows: LN405 with 27.5 µM cyclopamine, 32.5 µM tubastatin A and 400 µM temozolomide; and T98G with 20 µM cyclopamine, 30 µM tubastatin A and 400 µM temozolomide. Concentrations and durations for the combination of cyclopamine and tubastatin A, and the combination of temozolomide and tubastatin A, were as for the single treatments. Additionally, 1% DMSO was used as a vehicle control. Apoptosis was measured at 24, 48 and 72 h following the manufacturer’s protocols.

### Table II. Mutational status of PTEN and p53 in the glioblastoma cell lines, and concentrations of cyclopamine, tubastatin A and temozolomide.

<table>
<thead>
<tr>
<th>Cell line</th>
<th>PTEN</th>
<th>p53</th>
<th>Cyclopamine</th>
<th>Tubastatin A</th>
<th>Temozolomide</th>
</tr>
</thead>
<tbody>
<tr>
<td>LN405</td>
<td>MUT</td>
<td>MUT</td>
<td>27.5 µM</td>
<td>32.5 µM</td>
<td>400 µM</td>
</tr>
<tr>
<td>T98G</td>
<td>WT</td>
<td>MUT</td>
<td>20 µM</td>
<td>30 µM</td>
<td>400 µM</td>
</tr>
</tbody>
</table>

MUT, mutated; WT, wild-type; PTEN, phosphatase and tensin homolog.
Table III. Sequences and melting temperatures of primers used for reverse transcription-quantitative polymerase chain reaction.

<table>
<thead>
<tr>
<th>Gene</th>
<th>Forward primer (5’-3’)</th>
<th>Reverse primer (5’-3’)</th>
<th>Temperature (˚C)</th>
</tr>
</thead>
<tbody>
<tr>
<td>18S</td>
<td>GTAACCCCGTTGAACCCCATTT</td>
<td>CCATCCAACTCGGTATAGGCCG</td>
<td>63</td>
</tr>
<tr>
<td>Gli1</td>
<td>AAGCGTGAGCCTGAATCTGT</td>
<td>CAGCATGTACTGGCTTTGA</td>
<td>61</td>
</tr>
<tr>
<td>PTCH1</td>
<td>AGTTGCGCACAGAATCCACT</td>
<td>GCATAGGGCCGACATGTAAG</td>
<td>63</td>
</tr>
<tr>
<td>Snail</td>
<td>GGGTCTTTCCGCTAATCGCT</td>
<td>TAGGGCTGCTGAAAGGTAAA</td>
<td>63</td>
</tr>
<tr>
<td>Slug</td>
<td>CATTTCACGCTCTCAAAAA</td>
<td>GGAATGGAGCAGCGGTAGT</td>
<td>63</td>
</tr>
</tbody>
</table>

Caspase-Glo 3/7 assay. The Caspase-Glo® 3/7 assay (Promega Corporation, Madison, WI, USA) is a homogeneous, luminescent assay that measures caspase-3 and caspase-7 activities. The kit provides a luminogenic caspase-3/7 substrate, which contains the tetrapeptide sequence DEVD. The addition of this reagent results in cell lysis, followed by caspase cleavage of the substrate and generation of a glow-type luminescent signal produced by luciferase. Luminescence is proportional to the amount of caspase activity present. Cells were cultured at 37˚C for 72 h in Gibco RPMI-1640 GlutaMAX™ medium, supplemented with 10% FBS and 1% penicillin/streptomycin, in 96-well plates at a concentration of 5,000 cells/well, with 4 wells/condition. After 24 h, cells were added to each corresponding well, that were treated at 37˚C for 72 h as follows: LN405 with 27.5 µM cyclopamine, 32.5 µM tubastatin A and 400 µM temozolomide; and T98G with 20 µM cyclopamine, 30 µM tubastatin A and 400 µM temozolomide. Concentrations and durations for the combination of cyclopamine and tubastatin A, and the combination of temozolomide and tubastatin A, were as for the single treatments. Additionally, 1% DMSO was used as a vehicle control. Caspase-3/7 activation was measured at 24, 48 and 72 h, following the manufacturer’s protocols.

RNA extraction. Total RNA extraction from 72 h treated cells was conducted following the TRIzol® reagent (Invitrogen; ThermoFisher Scientific, Inc.) protocol, which allows sequential precipitation of RNA, DNA and proteins from a single sample. Cells were treated at 37˚C for 72 h as follows: LN405 with 27.5 µM cyclopamine, 32.5 µM tubastatin A and 400 µM temozolomide; and T98G with 20 µM cyclopamine, 30 µM tubastatin A and 400 µM temozolomide. Concentrations and durations for the combination of cyclopamine and tubastatin A, and the combination of temozolomide and tubastatin A, were as for the single treatments. Additionally, 1% DMSO was used as a vehicle control. Following homogenization of the samples with TRIzol reagent, chloroform was added to separate (after 15 min centrifugation at 12,000 x g and 4˚C) into a clear upper aqueous layer containing RNA, an interphase and a red lower organic layer containing DNA and proteins. RNA precipitation was then achieved by the addition of isopropanol and centrifugation for 30 min at 12,000 x g and 4˚C. Subsequently, isopropanol was removed, and 1 ml 75% ethanol in diethyl pyrocarbonate (DEPC) water was added, vortexed and centrifuged for 5 min at 7,500 x g and 4˚C. The process was repeated once more. Subsequently, ethanol was removed, the RNA pellet was left to dry, and resuspended in 15 µl DEPC water, to finally be stored at -80˚C for use in downstream applications. Total RNA quantification in each sample was measured using NanoDrop™ Microvolume Spectrophotometer (Thermo Fisher Scientific, Inc.).

Reverse transcription-quantitative polymerase chain reaction (RT-qPCR). For reverse transcription, 2 µg RNA were mixed with 1 µl random primers (250 ng/µl) and 1 µl dNTPs mix (10 µM) in a final volume of 12 µl water. Random primers (cat. no. 48190011) were purchased from Thermo Fisher Scientific, Inc. This mixture was incubated for 5 min at 65˚C. Subsequently, 4 µl first strand buffer and 2 µl DTT were added, and this was incubated for 2 min at 25˚C. Following this, 1 µl SuperScript II Reverse Transcriptase (Invitrogen; Thermo Fisher Scientific, Inc.) was added for the synthesis of cDNA and this final mixture was incubated for 10 min at 25˚C, 50 min at 42˚C and finally 15 min at 72˚C. Furthermore, 80 µl water were added and the cDNA was stored at -20˚C.

RT-qPCR was used to analyze the expression of genes associated with the sonic Hh pathway and EMT in six different conditions (DMSO as control, tubastatin A, cyclopamine, temozolomide, tubastatin A plus temozolomide and tubastatin A plus cyclopamine). Cells were treated at 37˚C for 72 h as follows: LN405 with 27.5 µM cyclopamine, 32.5 µM tubastatin A and 400 µM temozolomide; and T98G with 20 µM cyclopamine, 30 µM tubastatin A and 400 µM temozolomide. Concentrations and durations for the combination of cyclopamine and tubastatin A, and the combination of temozolomide and tubastatin A, were as for the single treatments. Additionally, 1% DMSO was used as a vehicle control. Amplification reactions were conducted in an IQ5 multicolor real-time PCR detection system (Bio-Rad Laboratories, Inc., Hercules, CA, USA). Subsequently, 1 µl of each sample cDNA was used in a total volume of 20 µl/well, with a reaction mix containing 10 µl IQ™ SYBR® Green supermix (Bio-Rad Laboratories, Inc.). An initial denaturation step at 95˚C for 30 sec was followed by 40 cycles of amplification alternating between 95˚C for 10 sec, the corresponding annealing temperature for each gene for 30 sec and 72˚C for another 30 sec (Table III). Each sample was assayed in triplicate. Forward and reverse primers were designed using Primer3Plus (http://www.bioinformatics.nl/cgi-bin/ primer3plus/primer3plus.cgi). Sequences and melting temperatures of each primer pair are depicted in Table III. Ribosomal 18S gene was used as a housekeeping reference gene for the relative quantification of cDNA amount.

PTCH1, Patched 1; Slug, Snail family transcription repressor 2.
using the comparative Cq method (33), also known as the $2^{ΔΔC_q}$ method.

**Protein extraction.** Cells were treated at 37°C for 72 h as follows: LN405 with 27.5 µM cyclopamine, 32.5 µM tubastatin A and 400 µM temozolomide; and T98G with 20 µM cyclopamine, 30 µM tubastatin A and 400 µM temozolomide. Concentrations and durations for the combination of cyclopamine and tubastatin A, and the combination of temozolomide and tubastatin A, were as for the single treatments. Additionally, 1% DMSO was used as a vehicle control. Total protein extraction was then conducted using radio immunoprecipitation assay lysis buffer (50 mM Tris-hidroximetil-aminometano-HCl, pH 8.0, 150 mM NaCl, 0.5% Triton® X-100 and 0.5% sodium deoxycholate).

**Western blot analysis.** A total of 20 µg of each bicinchoninic acid-quantified protein sample were separated in 12% SDS-PAGE and then transferred to a nitrocellulose membrane. Following blocking with TBS-Tween 0.1% and 5% non-fat milk for 1 h at room temperature, membranes were incubated overnight with the primary antibody at 4°C. After three washes with TBS-Tween 0.1%, membranes were incubated with the corresponding secondary antibodies at room temperature for 1 h. To visualize the presence and quantity of protein, Lumi-LightPLUS Western blotting substrate (Merck KGaA) was used. The primary antibodies used in the present study were: Acetylated α-tubulin (cat. no. T6793; 1:10,000; Merck KGaA) and β-actin (cat. no. A5441; 1:10,000; Merck KGaA). The secondary antibody used was horseradish peroxidase-conjugated anti-mouse (cat. no. SC-516102; 1:10,000; Santa Cruz Biotechnology, Inc., Dallas, TX, USA).

**Statistical analysis.** Values are expressed as the mean ± standard deviation or as the mean ± standard error of the mean. GraphPad 7.0 Software (GraphPad Software, Inc., La Jolla, CA, USA) was used to analyze the statistics of the results obtained from the experiments. The statistical tests used were the one-way analysis of variance and Tukey's multiple comparison test. *P*<0.05 was considered to indicate a statistically significant difference.

**Results**

**Tubastatin A increases acetylated α-tubulin levels in glioblastoma cell lines.** A western blot analysis for acetylated α-tubulin was performed to ensure the effect of tubastatin A as an inhibitor of HDAC6 (Fig. 1) in glioblastoma cell lines. As expected, acetylated α-tubulin protein levels increased in the groups treated with tubastatin A, both alone and together with cyclopamine or temozolomide, compared with the control group, confirming the indicated mechanism of action of this drug. In the samples treated with cyclopamine or temozolomide alone, no significant differences were observed.

**Tubastatin A, alone and combined with temozolomide, reduces clonogenicity of glioblastoma cell lines.** To evaluate the clonogenic capacity of T98G and LN405 glioblastoma cells after treatment with cyclopamine, temozolomide, tubastatin A, combination of cyclopamine with tubastatin A, combination of temozolomide with tubastatin A, and DMSO for 72 h, two different colony formation experiments were performed: attachment-dependent (2D colonies); and attachment-independent conditions (3D colonies) (Fig. 2).
Tubastatin A single treatment, and its combination with cyclopamine and temozolomide, significantly reduced the number of colonies counted in both experiments. The combination of temozolomide with tubastatin A was the most efficient strategy for reducing clonogenicity of glioblastoma cells, compared with the untreated group or cells treated with temozolomide alone.

**Tubastatin A decreases the migration capacity of glioblastoma cell lines.** A wound healing or scratching assay was then conducted to analyze the migration capacity of T98G and LN405 cells following treatment with cyclopamine, temozolomide, tubastatin A, combination of cyclopamine with tubastatin A, and combination of temozolomide with tubastatin A (Fig. 3). Even if all groups exhibited a closure of the scratch after 48 h, differences were evident among different treatments. Cyclopamine alone had no significant effect on reducing cell migration, compared with the control group. Tubastatin A induced a reduced migration rate; however, a notable inhibition was observed when both drugs were added together. The single treatment with temozolomide reduced cell migration more, compared with the individual treatment with cyclopamine. Tubastatin A produced different results in the two cell lines. When tubastatin A was combined with temozolomide, inhibition upon migration was enhanced, as demonstrated by the inability of these cells to close the gap.

**Tubastatin A downregulates the sonic Hh pathway in glioblastoma cell lines.** To observe whether tubastatin A treatment had any effect on the regulation of the sonic Hh pathway, in LN405 (Fig. 4) and T98G (Fig. 5) glioblastoma cell lines, total RNA was extracted from each experimental group and RT-qPCR (Table III) was performed following reverse transcription for GLI1 and Patched 1 (PTCH1) genes. Ribosomal 18S was used as a reference gene for the relative quantification of these two genes using the comparative Cq method. Treatment with cyclopamine had a significant effect on GLI1 and PTCH1, both significantly reducing the expression following treatment with
Figure 3. Changes in cell migration after treatments of glioblastoma cells. Single treatment with TubA, and combination with Cyp and TMZ, reduced cell migration with respect to Cyp or TMZ alone in glioblastoma cells. (A) LN405 cells. Images of the gap were captured at 0, 8, 24, 32 and 48 h following scratching in every treatment condition. (B) LN405 cells. Graphs representing the percentage of the migrated area in every condition are incorporated. (C) T98G cells. Images of the gap were captured at 0, 8, 24, 32 and 48 h following scratching in every treatment condition. (D) T98G cells. Graphs representing the percentage of the migrated area in every condition are incorporated. TubA, tubastatin A; Cyp, cyclopamine; TMZ, temozolomide.

Figure 4. Changes in the expression of sonic hedgehog and EMT markers after treatments of LN405 glioblastoma cells. mRNA fold expression of sonic hedgehog markers (GLI1 and PTCH1) and EMT markers (Slug and Snail), compared with 18S in LN405 glioblastoma cells. Data are presented as the mean ± standard deviation. *P<0.05, **P<0.01 and ****P<0.0001, compared with control; ººººP<0.0001, compared with TMZ single treatment. TubA, tubastatin A; Cyp, cyclopamine; TMZ, temozolomide; PTCH1, Patched 1; Slug, Snail family transcription repressor 2.
these drugs. Nevertheless, in both cases, the most significant decrease of PTCH1 expression was observed when cells were treated both with cyclopamine and tubastatin A, indicating that acetylation of α-tubulin and the possible restoration of primary cilia may result in a downregulation of the sonic Hh pathway.

Additionally, the combined treatment of temozolomide and...
tubastatin A decreased GLI and PTCH1 expression more, compared with temozolomide alone.

**Tubastatin A downregulates the expression of mesenchymal markers in glioblastoma cell lines.** EMT was investigated by RT-qPCR (Table III) for the mesenchymal markers Snail and Snail family transcriptional repressor 2 (Slug) in LN405 (Fig. 4) and T98G (Fig. 5) glioblastoma cell lines. The greatest reduction in expression of those markers was observed in the tubastatin A single treatment. However, when tubastatin A did not produce the greatest decay in expression of the markers, the double treatments exhibited the greatest decay.

**Tubastatin A accelerates temozolomide action on apoptosis in glioblastoma cell lines.** In order to investigate the effect of cyclopamine, tubastatin A and temozolomide on apoptosis in T98G and LN405 glioblastoma cells, two different experiments were performed: Cell death detection ELISAPLUS (Fig. 6); and Caspase-Glo 3/7 assay (Fig. 7). Evolution of both apoptosis processes was monitored at 24, 48 and 72 h after treatment. Both experiments demonstrated similar results, strengthening the validation of these assays.

The treatment with cyclopamine had different effects when inducing apoptosis in the two cell lines. In LN405 cells, cyclopamine continued to induce apoptosis after 72 h of treatment, whereas it did not have any effect in T98G cells after 48 h, when the values were equalized with respect to its control. The combination of cyclopamine with tubastatin A increased apoptosis at 24 h in both cell lines more than cyclopamine alone.

Tubastatin A notably induced apoptosis at 24 and 48 h with respect to the control, but this induction was attenuated after reaching 72 h. The combination of tubastatin A with temozolomide improved the increase of apoptosis induced by temozolomide alone, at 24 and 48 h. However, at 72 h, temozolomide alone produced the greatest increase in apoptosis, indicating that the combination with tubastatin A accelerates the action of temozolomide on apoptosis.

**Discussion**

Glioblastoma is the most common and the most malignant form of brain tumor (1). Despite the therapeutic effort conducted to treat this type of glioma, the majority of patients develop resistance and have a poor prognosis (2,34). The aim of the present study was to investigate the effect of tubastatin A, a selective HDAC6 inhibitor, on glioblastoma cell lines. The cells used in the present study, which lack the primary cilium (14), may restore it under treatment with tubastatin A, and consequently, the cells may also reverse their malignant phenotype and become sensitive to chemotherapy. The present results demonstrated statistical differences encountered following treating the glioblastoma cell lines with cyclopamine, temozolomide, tubastatin A, combination of cyclopamine and tubastatin A, combination of temozolomide with tubastatin A, or DMSO, as a vehicle control.

With western blot analysis, an increase in acetylated α-tubulin levels following treatment with tubastatin A, alone or combined with cyclopamine or temozolomide, was demonstrated (Fig. 1). This result was expected, as inhibition of HDAC6 inhibits α-tubulin deacetylation (15,35). The increase in acetylated α-tubulin was greater when tubastatin A and temozolomide was used in the LN405 cell line, while this
increase with the combination treatment was not observed in the T98G cell line. Therefore, it should be considered that LN405 and T98G may produce different results, as although they are glioblastoma cell lines, they are different at the molecular level, as LN405 cells exhibit mutations in TP53 and phosphatase and tensin homolog (PTEN), while T98G cell exhibit mutations in TP53, but not in PTEN.

Both in vitro cell tumorigenicity experiments, the 2D colony formation assay and the 3D soft agar colony formation assay (Fig. 2), revealed a significant decrease in clonogenicity of glioblastoma cell lines following inhibition of HDAC6. A similar effect was observed when tubastatin A was combined with cyclopamine in the 3D assay. However, the combined therapy with temozolomide was the most efficient treatment when it comes to reducing tumor growth, as determined by the number of colonies counted. This indicates that HDAC6 inhibition successfully reduces glioblastoma cell growth, as occurs in other tumor types, including cholangiocarcinoma (36), and may sensitize them to chemotherapy with temozolomide.

This observation was reinforced following analyzing the results for the cell migration test in the wound healing assay (Fig. 3). As occurs in the colony formation assays, single treatment with cyclopamine or temozolomide did not produce any significant difference, compared with the control group. However, when tubastatin A was added, alone or combined with any of the two other drugs, a significant inhibition of the cell migration capacity was observed, which indicates tubastatin A as a potential adjuvant drug to be administered together with temozolomide.

Modulation of the sonic Hh pathway by tubastatin A was also investigated. As depicted in Figs. 4 and 5, tubastatin A was the only drug that achieved a decrease in the mRNA levels of GLI1 and PTCH1, and therefore a downregulation of the sonic Hh pathway, when administered alone. However, in the present study, a greater decrease was expected when tubastatin A was combined with cyclopamine, as restoration of the primary cilium caused by tubastatin A would allow the action of cyclopamine on Smo inhibition. This combined treatment also reduced GLI1 expression, compared with untreated cells. These results may confirm the role of the primary cilium in sonic Hh regulation and the efficiency of the cyclopamine and tubastatin A treatment in modulating the pathway when disrupted.

Finally, regarding the two genes associated with EMT, the RT-qPCR experiments demonstrated a notable decrease in expression of the mesenchymal genes Snail and Slug in all treated groups, compared with the control group, even though these differences were not statistically significant (Figs. 4 and 5). These observations indicated a reversion of the EMT transition in these cells, as these markers belong to a mesenchymal phenotype (37,38). However, a report published by Stepanenko et al (39) demonstrated that a number of glioblastoma cell lines treated with temozolomide increase vimentin and Slug levels, although it was not the case for T98G cells. Additionally, it was documented (40) that the loss of α-tubulin acetylation acts as a mesenchymal marker for EMT; therefore, increased expression of HDAC6 induces EMT. It can be considered that treatment with tubastatin A is not only inducing the reversion of EMT to a MET phenotype in glioblastoma cells, as it has been demonstrated to increase acetylated α-tubulin levels, but also represses TGF-1 induced EMT in cultured peritoneal mesothelial cells, preventing peritoneal fibrosis (30).

In the apoptosis assays (Fig. 6 and 7), cyclopamine increased cell death during the first 24 h after treatment, but cannot be considered an efficient drug when it comes to inducing apoptosis. Additionally, the cyclopamine and tubastatin A combination treatment for 24 h produced an increased effect, compared with cyclopamine alone. Simultaneously, temozolomide single treatment did not produce increased apoptosis, compared with the untreated group, which is a poor result. Nevertheless, the most notable result from this experiment is the highlighted effect of the inhibition of HDAC6 on inducing apoptosis. Tubastatin A has been demonstrated to promote cell death at a high rate not only in glioblastoma cells, but also in other tumor types, including gastric cancer (41). Additionally, tubastatin A accelerates temozolomide action, as the effect of single treatment with temozolomide was already achieved 24 h earlier with the combined treatment. This may be due to the fact that the inhibition of HDAC6 favors the sensitization of the cells to temozolomide, since tumors with overexpression of HDAC6 have an increased resistance to temozolomide (42). Furthermore, a novel mechanism for a possible explanation of the increase of apoptosis by HDAC6 inhibition has recently been determined (43), demonstrating that HDAC6-selective inhibition is a novel epigenetic anticancer therapeutic strategy targeting the p53-Hsp90 complex that can be applied to wild-type and mutated p53-bearing cancer, with similar efficacy.

The present results indicated that the primary cilium acts as a tumor suppressor in these glioblastoma cells, as well as in other glioblastoma cell lines, including U87-MG, U-373G, U-138MG or U-251MG (44), and other tumor types, including cholangiocarcinoma (36), breast cancer (45), melanoma (46), sporadic clear cell renal cell carcinoma (47), prostate cancer (48) and lung cancer (49). However, there are studies that indicate that cilia may function as an oncogene in other cases, including pancreatic ductal carcinoma (50) and medulloblastoma (51), and other different glioblastoma cell lines, due to the complicated heterogeneity of glioblastoma attributed to the different responses of different cell lines or patients (52). For this reason, characterization of the patients is notable in order to classify them as cilia positive or cilia negative, prior to choosing which therapeutic strategy fits them best. However, a number of other cellular tests and molecular experiments with more markers are required with different glioblastoma cell lines prior to determine the result, as the observations up to now are preliminary.

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Authors' contributions
AU, EE, BM, JAR, MAI and JSC conceived and designed the experiments. AU and EE performed the experiments. AU, EE and JSC analyzed the data. EE and JSC drafted the paper. All authors contributed to refine analysis of data and writing of the manuscript. All authors read and approved the final manuscript.

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References


