The antiproliferative effect of indomethacin-loaded lipid-core nanocapsules in glioma cells is mediated by cell cycle regulation, differentiation, and the inhibition of survival pathways

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Abstract: Despite recent advances in radiotherapy, chemotherapy, and surgical techniques, glioblastoma multiforme (GBM) prognosis remains dismal. There is an urgent need for new therapeutic strategies. Nanoparticles of biodegradable polymers for anticancer drug delivery have attracted intense interest in recent years because they can provide sustained, controlled, and targeted delivery. Here, we investigate the mechanisms involved in the antiproliferative effect of indomethacin-loaded lipid-core nanocapsules (IndOH-LNC) in glioma cells. IndOH-LNC were able to reduce cell viability by inducing apoptotic cell death in C6 and U138-MG glioma cell lines. Interestingly, IndOH-LNC did not affect the viability of primary astrocytes, suggesting that this formulation selectively targeted transformed cells. Mechanistically, IndOH-LNC induced inhibition of cell growth and cell-cycle arrest to be correlated with the inactivation of AKT and β-catenin and the activation of GSK-3β. IndOH-LNC also induced G0/G1 and/or G2/M phase arrest, which was accompanied by a decrease in the levels of cyclin D1, cyclin B1, pRb, and pcdc2 and an increase in the levels of Wee1 CDK inhibitor p21WAF1. Additionally, IndOH-LNC promoted GBM cell differentiation, observed as upregulation of glial fibrillary acidic protein (GFAP) protein and downregulation of nestin and CD133. Taken together, the crosstalk among antiproliferative effects, cell-cycle arrest, apoptosis, and cell differentiation should be considered when tailoring pharmacological interventions aimed at reducing glioma growth by using formulations with multiples targets, such as IndOH-LNC.

Keywords: glioblastoma multiforme, PI3K/AKT, cell differentiation, indomethacin-loaded lipid core nanocapsules, cell cycle regulation

Introduction

Gliomas are the most common central nervous system malignancy. Their highly invasive and diffuse nature leaves little opportunity for a cure.¹ The infiltrative growth pattern of these tumors precludes curative neurosurgery, and tumor cells are usually resistant to irradiation, chemotherapy, and/or immunotherapy, leading to a low survival rate for patients with glioma.² Furthermore, the blood–brain barrier presents a huge challenge for the effective entry of a majority of drugs, and thus severely restricts therapy for gliomas. Treatment options are very limited and, in most situations, ineffective. Therefore, there is an urgent need for novel and effective therapies to treat these tumors.

Growth inhibition and the induction of cell death are among the major objectives of anticancer therapies. Apoptosis has particular relevance for the therapeutic treatment of...
tumors because the process favors the phagocytic elimination of dead tumor cells in the absence of an inflammatory response, which is normally associated with necrotic cell death. However, gliomas frequently develop resistance to apoptotic cell death. A number of deregulated signaling cascades mediate part of this resistance, including the phosphoinositide-3-kinase (PI3K/AKT) pathway. Despite the molecular heterogeneity of malignant gliomas, PI3K/AKT is a common signal transduction pathway altered in many of these tumors by the loss of phosphatase and tensin homologue (PTEN), the overexpression of epidermal growth factor receptor, and/or an increase of PI3K/AKT activity due to mutations in its regulators. Thus, inhibiting these pathways may be a way to decrease intrinsic and drug-induced resistance in glioblastoma multiforme (GBM), sensitizing GBM cells to apoptotic cell death.

The loss of the regulatory control of the cell cycle, which leads to unrestrained cell proliferation, is a hallmark of cancer. The normal progression of the cell cycle requires the sequential expression of cyclins, which activate cyclin-dependent kinases (Cdk) to phosphorylate target proteins required for cell-cycle progression. Cyclin D1 is one of the positive cell-cycle regulators necessary for the transition of cells from G1 to S phase. Its overexpression is one of the most commonly observed alterations in cancer. Recent studies have shown the role of cyclins not only as components of the cell cycle core machinery but also as cofactors for several transcription factors. Understanding deregulated cell-cycle pathways has provided a basis for designing molecular targeted therapies as new combination therapies and drug-delivery systems. Additional efforts focus on the development of novel therapeutic approaches using agents that promote cancer cell differentiation. Differentiation therapy changes the destructive behavior of cancer cells by reducing their proliferative capacity and inducing terminal differentiation and/or apoptosis. Recently, differentiation-inducing therapy has been proposed as a new potential strategy to treat malignant gliomas.

The development of delivery systems able to alter the biological profiles of active compounds is of the utmost importance and requires a multidisciplinary approach. Nanotechnology is at the core of biotechnology and is likely to play a significant role in this regard. One of the most important nanotechnology applications developed over the past decade is the nanovehicle, a nanoscale compound used as a therapeutic tool and designed to accumulate at specific sites in the body in order to improve pharmacotherapeutic outcomes. Hence, nanodrugs and nanodiagnostics have been developed to increase bioavailability profiles, enabling the administration of lower doses. This minimizes the adverse reactions found with conventional drugs administered in clinical practices and improves patient health. In the field of cancer therapy, there are many clinical applications based on nanotechnologies with a major development in drug delivery systems, particularly in solid tumors. Moreover, it has been suggested that nanoparticles could be used to carry drugs across the blood–brain barrier, an important requirement for the development of novel drugs against gliomas. The mechanisms underlying the encapsulation and release of several drugs from nanoparticulate systems have been extensively studied by our group. In a previous study, we demonstrated that polymeric nanocapsules could carry indomethacin, a powerful nonsteroidal anti-inflammatory drug, into brain tumors. Local delivery of indomethacin significantly reduced in-vivo GBM growth and improved animal survival in a rat glioma model. Taking into account the limited success of glioma therapy and these promising data, the aim of the present study was to investigate the mechanisms involved in the antiproliferative effect of indomethacin-loaded lipid core nanocapsules in glioma cells by evaluating PI3K/AKT survival pathways, cell-cycle regulatory molecules, and cell differentiation.

Materials and methods
Preparation of lipid-core nanocapsules
Lipid-core nanocapsule suspensions were prepared by interfacial deposition of polymer. At 40°C, indomethacin (0.010 g), poly(ε-caprolactone) (0.100 g), capric/caprylic triglycerides (0.33 mL), and sorbitan monostearate (0.077 g) were dissolved in acetone (27 mL). In a separate flask, polysorbate 80 (0.077 g) was added to 53 mL of water. The organic solution was injected into the aqueous phase under magnetic stirring at room temperature. After 10 minutes, the acetone was eliminated, and the suspensions were concentrated under reduced pressure. The final volume was adjusted to 10 mL for a drug concentration of 1 mg/mL. The indomethacin-loaded lipid-core nanocapsule formulation was named IndOH-LNC. Drug-unloaded nanocapsules (LNC) were prepared as described above, without adding indomethacin, and used as a control formulation.

Characterization of lipid-core nanocapsules
After preparation of the nanocapsule suspensions, the pH was determined using a potentiometer (B-474; Micronal, São Paulo, Brazil). The particle size, polydispersity index, and zeta potential of the suspensions were determined using a Zetasizer® Nano-ZS ZEN 3600 model (Malvern...
Instruments, Malvern, UK). The samples were diluted with pre-filtered water (MilliQ®) (particle size and polydispersity index) or a 10 mM NaCl aqueous solution (zeta potential). The measurements were made in triplicate. The total concentration of indomethacin in the formulations was measured by reverse phase high-performance liquid chromatography (PerkinElmer S-200, with injector S-200, detector UV-Vis, a guard-column [PerkinElmer, Waltham, MA, USA]; and a LiChrospher® 100 RP-18 column of 250 mm, 4 mm, and 5 µm [Merck KGaA, Darmstadt, Germany]). The mobile phase consisted of acetonitrile/water (70:30, v/v) adjusted to an apparent pH of 5.0 ± 0.5 with 10% (v/v) acetic acid. Each suspension (100 µL) was treated with acetonitrile (10 mL). The solution was filtered (Millipore 0.45 µm; Merck Millipore, Billerica, MA, USA) and injected (20 µL). The high-performance liquid chromatography method was previously validated. Linear calibration curves for indomethacin were obtained in the range of 1.00–25.00 µg/mL, presenting correlation coefficients higher than 0.9992. The encapsulation efficiency was determined by an ultrafiltration-centrifugation technique (Ultrafree-MC 10,000 MW, Millipore) at 15,300 g for 10 minutes. The encapsulated indomethacin in the nanocapsules was calculated from the difference between the total and the free-drug concentrations determined in the nanocapsule suspension and in the ultrafiltrate, respectively.

**Cell cultures**

Rat C6 glioma cells and human U138-MG glioma cells were obtained from American Type Culture Collection (ATCC) (Manassas, VA, USA). The cells were grown and maintained in Dulbecco’s modified Eagle’s medium (DMEM) low glucose containing 0.1% Fungizone® and 100 U/L gentamicin and supplemented with 5% (C6) or 15% (U138-MG) (v/v) fetal bovine serum. All culture materials were purchased from Gibco Laboratories (Grand Island, NY, USA). Cells were kept at a temperature of 37°C, a minimum relative humidity of 95%, and an atmosphere of 5% CO₂ in air.

Primary astrocyte cultures were prepared as previously described. Briefly, the cortices of newborn Wistar rats (1–2 days old) were removed and mechanically dissociated by sequential passage through a Pasteur pipette. The cell suspension was centrifuged at 1000×g for 5 minutes. The pellet was then resuspended in DMEM supplemented with 10% fetal bovine serum, 1% penicillin/streptomycin, and 0.1% amphotericin B. The cells (2 × 10⁵) were plated (1.5 × 10⁴ cells/cm²) in poly-L-lysine-coated 24-well plates. After 4 hours plating, plates were gently shaken and washed with phosphate buffered saline (PBS). The medium was changed, and cells were grown to confluence (21 days). Medium was replaced every 4 days.

**Cell treatments**

Indomethacin was purchased from Sigma-Aldrich (St Louis, MO, USA). Free indomethacin was dissolved in cell culture-grade dimethyl sulfoxide (DMSO) (Sigma-Aldrich). The control groups using this solution were named IndOH. The indomethacin-loaded lipid-core nanocapsules (IndOH-LNC) were prepared and characterized as described above. After reaching semiconfluence (glioma cell lines) or after 21 days (astrocytes), the cultures were exposed to IndOH or IndOH-LNC (5, 10, 25, 50, or 100 µM) for 1, 6, 24, 48, or 72 hours. The vehicle DMSO was present in the cells at a maximum concentration of 0.5%. Unloaded lipid-core nanocapsules (LNC) were considered the vehicle control of IndOH-LNC.

**Assessment of cell viability**

The 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide (MTT) assay and the trypan blue exclusion test were used to estimate glioma and astrocyte cell viability. The MTT method is based on the mitochondrial reduction of MTT to a formazan product. Cells were seeded (10⁴ per well) into 96-well plates and treated with 5, 10, 25, 50, or 100 µM of IndOH or IndOH-LNC for 24 hours. At the end of the treatment, the MTT assay was performed as described previously. In the trypan blue exclusion test, live cells with intact membranes exclude the dye, whereas dead cells do not stain. Cells were seeded (10⁴ per well) into 24-well plates. At the end of the treatment, cells were collected after trypsin digestion and were counted immediately in a hemocytometer.

**Annexin V staining**

Apoptotic cells were quantified using an Annexin V-FITC/Propidium Iodide (PI) double staining kit, according to the manufacturer’s instructions (Chemicon International, Temecula, CA, USA). Glioma cells were seeded (3 × 10⁴ per well) into 6-well plates and treated with 10, 25, or 50 µM of IndOH or IndOH-LNC for 24 hours. At the end of the treatment, cells were collected after trypsin digestion, and externalized phosphatidylserine was labeled (15 minutes, ice-cold) with 3 µL FITC-conjugated annexin V in 100 µL binding buffer in the dark. Next, PI (1 µg/mL) was added. Apoptotic and/or necrotic cells were quantified by a dual-color flow cytometry technique using a FACSCalibur flow cytometer (BD Biosciences, San Jose, CA, USA). Ten thousand events were analyzed per sample, and cells were classified as follows: live cells (Annexin−/PI−); early apoptotic cells (Annexin+/PI−); late
apototic (Annexin+/PI+); necrotic (Annexin–/PI+). The data obtained were analyzed with FCS Express 4 Flow Cytometry Software (De Novo Software, Los Angeles, CA, USA).

**Cell-cycle analysis**
Glioma cells were seeded (3 × 10⁴ per well) into 6-well plates and treated with 25 𝜇M IndOH-LNC for 24 and 48 hours. At the end of treatment, cells were trypsinized, centrifuged, and resuspended in a lysis buffer containing 3.5 mmol/L trisodium citrate, 0.1% v/v Nonidet P-40, 0.5 mmol/L Tris-HCl, 1.2 mg/mL spermine tetrahydrochloride, 5 μg/mL RNase, 5 mmol/L EDTA, and 1 μg/mL propidium iodide, pH 7.6. Afterwards, cells were incubated for at least 10 minutes on ice for cell lysis (10⁶ cells/mL lysis buffer).²⁷ The DNA content was determined by flow cytometry (FACSCalibur; BD Biosciences). Ten thousand events were counted per sample. FACS analyses were performed in the CellQuest Pro Software (BD Biosciences).

**Western blot analysis**
After the indicated treatments, glioma cells were homogenized in ice-cold lysis buffer (4% sodium dodecyl sulfate [SDS], 2 mM EDTA, and 50 mM Tris) containing a proteinase inhibitor cocktail, and the protein concentration was determined. Protein samples were prepared and Western blotted as described previously with minor modifications.²⁸ Equal amounts of proteins were resolved (50 μg per lane) on 8% or 10% SDS-PAGE. After electrophoresis, proteins were electrotransferred to nitrocellulose membranes using a semi-dry transfer apparatus (Trans-Blot SD; Bio-Rad Laboratories, Hercules, CA, USA). Membranes were incubated for 60 minutes at 4°C in blocking solution (Tris-buffered saline containing 5% nonfat milk and 0.1% Tween 20, pH 7.4) and further incubated with the appropriate primary antibody dissolved in the blocking solution overnight at 4°C.²⁸ The following primary antibodies were used: anti-phospho AKT (Ser473) (pAKT) (1:1000; Cell Signaling, Danvers, MA, USA), anti-AKT (AKT) (1:1000; Cell Signaling), anti-phospho GSK3β (Ser9) (pGSK3β) (1:2000; Cell Signaling), anti-GSK3β (GSK3β) (1:2000; Cell Signaling), anti-GFAP (glial fibrillary acidic protein) (1:2000; Chemicon International), anti-nestin (1:2000; Chemicon International), anti-cyclin D1 (1:1000; Cell Signaling), anti-CDK4 (1:1000; Cell Signaling), anti-CDK6 (1:1000; Cell Signaling), anti-pRb (Ser795) (1:500; Cell Signaling), anti-cyclin B1 (1:1000; Cell Signaling), anti-phospho cdc2 (cdc2) (1:500; Cell Signaling), anti-cdc2 (cdc2) (1:1000; Cell Signaling), anti-p21 (1:1000; Cell Signaling). After washing, the membranes were incubated with adjusted secondary antibodies coupled to horseradish peroxidase (1:1000) for 2 hours. The immunocomplexes were visualized using the ECL chemiluminescence detection system (Thermo Fischer Scientific, Waltham, MA, USA). Band density was measured using Optiquant software (Packard Instrument Company, Meriden, CT, USA). Uniform gel loading was confirmed with beta-actin (1:1000; Cell Signaling) as the standard.

**Immunocytochemistry**
Glioma cells were grown on coverslips and treated with 25 𝜇M IndOH-LNC for 24 hours (β-catenin experiment) or 72 hours (cell differentiation experiments). At the end of treatment, cells were fixed with 4% fresh paraformaldehyde in PBS for 30 minutes at room temperature. Samples were blocked for 1 hour with 1% normal goat serum in PBS containing 0.2% Triton X-100. The primary antibodies anti-GFAP (1:500; Chemicon), anti-nestin (1:500; Chemicon), CD133 (1:500; Abcam, Cambridge, MA, USA), and β-catenin (1:500; Sigma-Aldrich) were incubated overnight. For immunolabeling, the samples were incubated with species-specific fluorochrome-conjugated secondary antibodies for 90 minutes at room temperature. The secondary antibody used was anti-rabbit Alexa Fluor® 488 (green) (1:500; Life Technologies, Carlsbad, CA, USA) or anti-mouse Rhodamine (red) (1:500; Life Technologies). Nuclei were stained with DAPI (4′,6-diamidino-2-phenylindole) (1:1000; Sigma-Aldrich). After a final rinse with PBS, the slides were mounted using Pro-Long® Gold Antifade Reagent (Molecular Probes, Eugene, OR, USA). The slides were examined using an Olympus IX-70 fluorescence microscope (Olympus Corporation, Tokyo, Japan), and images were captured using a digital camera connected to the microscope (camera model U-LH100HGAP; Olympus Corporation) (magnification ×20).

**Statistical analysis**
All experiments were carried out at least three times. The results are presented as the mean ± standard deviation. The statistical comparisons of the data were performed by one-way or two-way analysis of variance followed by a Bonferroni post-hoc test using GraphPad Prism software version 5.01 (GraphPad Software Inc, La Jolla, CA, USA). P-values lower than 0.05 (P < 0.05) were considered significant.

**Results**
**Physicochemical characterization of IndOH-LNC**
The lipid-core nanocapsule formulations were prepared by interfacial deposition of poly(ε-caprolactone) without the need...
for any subsequent purification step. IndOH-LNC and LNC showed macroscopic homogeneous aspects, such as white bluish opalescent liquids. After preparation, the mean particle diameters determined by photon correlation spectroscopy (z-average diameters) were 231 ± 4 nm (IndOH-LNC) and 229 ± 5 nm (LNC). The suspensions showed monomodal size distributions and a polydispersity index of 0.12 ± 0.01 nm (IndOH-LNC) and 0.14 ± 0.02 (LNC), indicating the formulations were highly homogeneous with narrow size distributions. The pH values were 5.95 ± 0.1 (IndOH-LNC) and 6.1 ± 0.2 (LNC), and the zeta potential values were −7.0 ± 1.3 mV and −7.2 mV ± 1.8 mV, respectively. The indomethacin content was 0.998 ± 0.010 mg/mL, and the encapsulation efficiency was close to 100% for all batches.

**IndOH-LNC selectively decrease cell viability in glioma cells**

First, the MTT assay was used to evaluate whether IndOH and IndOH-LNC (5, 10, 25, 50, or 100 µM) affect the cell viability of gliomas after 24 hours of treatment. As shown in Figure 1, all concentrations of IndOH-LNC significantly reduced the cell viability of C6 and U138-MG cell lines (Figure 1A and B). In accordance with previous results using 48 hours of treatment, IndOH-LNC more potently reduced the cell viability when compared with respective concentrations of IndOH (Figure 1A and B). These results were confirmed by a trypan blue exclusion test (data not shown). In parallel, primary astrocyte cultures were used as a nontransformed model of glial cells in order to confirm the selectivity of IndOH-LNC. Whereas IndOH-LNC decreased the viability of the two GBM cell lines in a concentration-dependent manner (half-maximal inhibitory concentration [IC50] range: 25 µM), concentrations of IndOH-LNC up to 100 µM (IC50 > 500 µM) did not alter astrocytic viability significantly (Figure 1C). These results suggest that IndOH-LNC preferentially targets cancer cells.

**IndOH-LNC induce apoptotic cell death in glioma cells**

To characterize the cell death induced by IndOH-LNC, glioma cells were treated with 10, 25, or 50 µM of IndOH or IndOH-LNC for 24 hours, and annexin V-PI assays were carried out. The cytogram of the four quadrants in Figure 2 was used to distinguish the live (Annexin−/PI−), early apoptotic (Annexin+/PI−), late apoptotic (Annexin+/PI+), and necrotic (Annexin−/PI+) cells. In C6 glioma cells, 25 µM IndOH-LNC elicited externalization (flip-flop) of phosphatidylserine in approximately 25% of the cells (Annexin+/PI−). A low

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**Figure 1** Effect of IndOH and IndOH-LNC on the cell viability of gliomas and astrocytes. (A) C6 and (B) U138-MG glioma cell lines and (C) normal astrocytes were treated for 24 hours with different concentrations (5, 10, 25, 50, or 100 µM) of IndOH or IndOH-LNC, and MTT assays were carried out. Notes: The dashed line represents the IC50 values. Unloaded LNC were considered the vehicle control of IndOH-LNC. The cell viability is presented relative to that of control cells (100% cell viability). The values are presented as mean ± standard deviation for six independent experiments. Significant differences from control and between the respective concentrations of IndOH groups: ***P < 0.001 and **P < 0.01, as assessed by two-way analysis of variance followed by the Bonferroni post hoc test.

**Abbreviations:** IC50, half-maximal inhibitory concentration; IndOH, indomethacin; IndOH-LNC, indomethacin-loaded lipid-core nanocapsules; LNC, lipid-core nanocapsules; MTT, 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide.
percentage of cells (approximately 6%) was Annexin−/PI+ (necrosis), suggesting that IndOH-LNC induced cell death mainly by apoptosis (Figure 2A and C). The cell death profile was similar for all concentrations of IndOH-LNC (Figure 2A and C). Consistent with the cell viability results, IndOH-LNC was more potent in inducing apoptotic cell death than the respective concentrations of IndOH (Figure 2A and C). Similar results were obtained with U138-MG glioma cells. However, in these cells, IndOH-LNC treatment was even more effective (Figure 2B and D). In the U138-MG cells, our results showed that 25 μM IndOH-LNC induced early apoptosis in approximately 60% of the cells (Annexin+/PI−). A low percentage of cells (approximately 2%) was Annexin−/PI+ (necrosis) (Figure 2B and D), confirming that IndOH-LNC induced cell death mainly by apoptosis. Based on these results, the concentration of 25 μM (approximate IC50 value) was chosen for further experiments. Treatment of the C6 and U138-MG cells with LNC did not significantly alter cell death, indicating that the nanostructured architecture is not toxic for these models (Figure 2).

**IndOH-LNC modulate PI3K/AKT, GSK3β, and β-catenin survival pathways in glioma cells**

PI3K/AKT is over activated in a wide range of tumor types; it triggers a cascade of responses that include cell growth, proliferation, and survival.19 In an attempt to delineate the role of AKT signaling in IndOH-LNC-treated glioma cells, we determined the AKT phosphorylation status and the activation/inactivation profile of signaling targets downstream of AKT in C6 and U138-MG glioma cell lines. As shown in Figure 3, glioma cells expressed high levels of activated AKT, as determined by the phosphorylation of AKT at Ser473. IndOH-LNC treatment inhibited the constitutive activation of the PI3K/AKT pathway at all times evaluated (1, 6, and 24 hours), as demonstrated by the levels of the phosphorylated form of AKT in C6 and U138-MG cell lines (Figure 3). The total levels of AKT were not altered, indicating that the protein was modulated at the posttranslational level (Figure 3). We then evaluated whether the AKT downstream target GSK-3β was affected. IndOH-LNC treatment also decreased the phosphorylation of GSK-3β, thus activating this protein, in both cell lines without changing the total level of GSK-3β (Figure 3). This effect was observed at 6 and 24 hours of treatment for C6 cells and at 1, 6, and 24 hours for U138-MG cells. In the U138-MG glioma cells, the effect was more intense with 1 and 6 hours of treatment (Figure 4B). The treatment of the C6 and U138-MG cells with LNC did not significantly alter AKT or GSK-3β levels (Figures 3 and 4).

Considering the interplay between the Wnt/β-catenin and PI3K/AKT signaling cascades in gliomas,18 we evaluated whether IndOH-LNC treatment was able to modulate β-catenin expression in C6 glioma cells. β-catenin staining in cells treated with IndOH-LNC was lower than that in control cells, suggesting that the IndOH-LNC formulation reduces β-catenin expression (Figure 5).
Figure 2 IndOH and IndOH-LNC induce apoptosis in glioma cells. Flow cytometry experiments to quantify apoptotic and/or necrotic cells in (A) C6 and (B) U138-MG glioma cells treated with IndOH or IndOH-LNC (10, 25, or 50 µM) for 24 hours. Unloaded LNC were considered the vehicle control of IndOH-LNC. Values are expressed as the percentage of early apoptotic cells (%) (annexin V positive/PI negative) in (C) C6 and (D) U138-MG glioma cells treated with IndOH or IndOH-LNC.

Notes: Bars represent the mean ± standard deviation for three independent experiments. Significant differences from control and LNC groups and between the respective concentrations of IndOH group: *P < 0.05, and ***P < 0.001, as assessed by two-way analysis of variance followed by the Bonferroni post hoc test.

Abbreviations: FITC, fluorescein isothiocyanate; IndOH, indomethacin; IndOH-LNC, indomethacin-loaded lipid-core nanocapsules; LNC, lipid-core nanocapsules; PI, propidium iodide.
IndOH-LNC promote cell-cycle arrest and induce cell-cycle regulation

Cell-cycle analysis experiments were performed to evaluate the mechanisms underlying the antiproliferative effect of IndOH-LNC in glioma cells. The cell-cycle distribution indicated that 25 µM IndOH-LNC caused C6 glioma cells to arrest in G0/G1 phase after 24 hours treatment (Figure 6A). Specifically, at 24 hours of IndOH-LNC treatment, the cell population in the G0/G1 phase grew to 61% ± 0.8% compared with 50% ± 1.8% in control, while the proportion of cells in S phase decreased from 28% ± 0.9% to 18% ± 0.5% (Figure 6A). This cell-cycle distribution was maintained at 48 hours treatment in C6 glioma cells (Figure 6A). The G0/G1 arrest was accompanied by the formation of...
Figure 4 IndOH-LNC decrease the pGSK3β protein immunocontent in glioma cells. Western blotting for pGSK3β, GSK3β, and β-actin was performed at the indicated times in 25 µM IndOH-LNC-treated (A) C6 and (B) U138-MG glioma cells.

Notes: Unloaded LNC were considered the vehicle control of IndOH-LNC. The histogram presents the quantitative Western blot analysis of the GSK3β phosphorylation state. The densitometric values obtained for phospho- and total-GSK3β after treatments were normalized to their respective controls (control bar) (100%). Data are expressed as a ratio of the normalized percentages of pGSK3β and GSK3β. Bars represent the mean ± standard deviation for six independent experiments. Significant differences from control and LNC groups: ***P < 0.001, as assessed by one-way analysis of variance followed by the Bonferroni post hoc test.

Abbreviations: IndOH-LNC, indomethacin-loaded lipid-core nanocapsules; LNC, lipid-core nanocapsules.

A population sub-G1 (12% ± 1.5% of IndOH-LNC-treated cells compared to 0.9% ± 0.6% of control cells), suggesting that G0/G1 arrest preceded DNA fragmentation and, consequently, apoptosis (Figure 6A). The formation of sub-G1 apoptotic cells confirmed the result obtained with annexin V staining (Figure 2A and C). Similar results were obtained with U138-MG glioma cells at 24 hours of IndOH-LNC treatment (Figure 7A). At 48 hours of treatment, IndOH-LNC induced a pronounced arrest in G0/G1 phase (66% ± 0.9% compared with 54% ± 1.1% of control cells) and a modest arrest in G2/M (24% ± 0.9% compared with 19% ± 0.5% of control cells), while the proportion of cells in S phase decreased from 22% ± 0.4% to 7% ± 0.9% (Figure 7A). In the same way, the formation of sub-G1
apoptotic cells at 24 and 48 hours treatment was observed, suggesting that arrest in G0/G1 and G2/M phases preceded apoptosis (Figure 7A). The treatment of the C6 and U138-MG cells with LNC did not significantly alter the cell-cycle progression (Figures 6A and 7A).

To evaluate the mechanisms involved in cell-cycle regulation, we investigated the effect of IndOH-LNC on cell-cycle checkpoint signaling. Cyclin D1 is believed to be a cell-cycle checkpoint at the G0/G1 phase to S phase transition. In response IndOH-LNC, cyclin D1 protein levels significantly decreased in C6 and U138-MG cells after 24 hours treatment compared with levels in control cells (Figures 6B and 7B). Additionally, the phosphorylation of downstream pRb significantly decreased in both cell lines treated with IndOH-LNC (Figures 6C and 7C). On the other hand, cyclin-dependent kinases 4 and 6 (CDK4 and CDK6) proteins were unaltered after the treatment (Figures 6D, 6E, 7D, and 7E). The treatment of cells with LNC did not significantly alter the cell-cycle distribution or cell-cycle regulation (Figures 6 and 7).

In an attempt to find the signaling pathway that coordinates IndOH-LNC-induced G2/M arrest only in U138-MG glioma cells, we determined the activity of cyclin B1, pcd2, Wee1, and p21WAF1 in this cell line. Our results showed that IndOH-LNC treatment reduced cyclin B1 levels (Figure 8A) and the phosphorylation status of cdc2 (Figure 8B). On the other hand, IndOH-LNC treatment increased the levels of Wee1 (Figure 8C), a component of the G2/M checkpoint that inhibits cdc2. Furthermore, IndOH-LNC also promoted a high accumulation of the cyclin-dependent kinase inhibitor (CDKi) p21WAF1, a marker of cell-cycle arrest (Figure 8D). The treatment of the U138-MG cells with LNC did not significantly alter cyclin B1, pcd2, Wee1, or p21WAF1 (Figure 8).

**IndOH-LNC induce the differentiation of glioma cells**

The differentiation of glioma cells toward the astrocyte type is characterized by a morphological transformation and expression of differentiation markers. Microscopic observation of glioma cells treated with 25 µM IndOH-LNC for 72 hours revealed cells with altered morphology, such as mature astrocytes with smaller, round bodies and much longer, fine, tapering processes (data not shown). To confirm this observation, we examined the expression of GFAP, a well-established marker of mature astrocytes. Immunocytochemistry with anti-GFAP antibody showed that few untreated U138-MG glioma cells stained positive, but numerous cells treated with IndOH-LNC stained positive in the cytoplasm (Figure 9A). The GFAP immunoccontent was quantified by Western blot analysis (Figure 9B and C). Consistent with the immunocytochemistry analysis, the results confirmed that IndOH-LNC strongly upregulated GFAP protein expression in both cell lines (Figure 9B and C).

Another differentiation marker evaluated here was nestin. Nestin is used to identify neural stem cells and is expressed in almost all GBMs. Thus, the low staining for nestin in C6 glioma cells treated with IndOH-LNC indicated that this formulation was able to promote glioma cell differentiation (Figure 10A). This result was also confirmed and quantified by Western blot analysis (Figure 10B).

Additionally, U138-MG glioma cells were positive for the cell surface antigen CD133 (Figure 11), a marker of neural stem cells. The low staining for CD133 in cells treated with IndOH-LNC (Figure 11) confirms that IndOH-LNC promoted glioma cell differentiation.

**Discussion**

Biodegradable nanoparticles are emerging as novel and promising drug delivery systems. They offer an alternative
Figure 6 Cell-cycle arrest induced by IndOH-LNC in C6 glioma cells.

Notes: Flow cytometry was performed to determine the cell-cycle distribution (sub-G1, G1/G0, S, and G2/M) in 25 µM IndOH-LNC-treated (24 and 48 hours) cells. Western blotting for cell-cycle checkpoint proteins from G0/G1 phase to S phase (cyclin D1, pRb, CDK4, and CDK6) was performed. Unloaded LNC were considered the vehicle control of IndOH-LNC. The histogram presents the quantitative Western blot analysis for the protein indicated. The densitometric values obtained from treatments were normalized to their respective controls (control bar) (100%). Bars represent the mean ± standard deviation for at least six independent experiments. Significant differences from control and LNC groups: **P < 0.01, ***P < 0.001, as assessed by one-way analysis of variance followed by the Bonferroni post hoc test.

Abbreviations: IndOH-LNC, indomethacin-loaded lipid-core nanocapsules; LNC, lipid-core nanocapsules.

Figure 6

therapeutic approach for difficult-to-treat diseases such as brain tumors. In previous work, we showed that polymeric nanocapsules can successfully carry indomethacin into brain tumors.²⁰ Additionally, the local delivery of indomethacin reduces in-vivo GBM growth and improves animal survival. However, the mechanisms underlying the antiproliferative effect of indomethacin remained unclear. Here, we showed that IndOH-LNC modulates PI3K/AKT and GSK3β survival pathways, modulates cell-cycle regulation, and induces differentiation in glioma cells.
Figure 7 Cell-cycle arrest induced by IndOH-LNC in U138-MG glioma cells.

Notes: Flow cytometry was performed to determine the cell-cycle distribution (sub-G1, G1/G0, S, and G2/M) in 25 μM IndOH-LNC-treated (24 and 48 hours) cells. Western blotting for cell-cycle checkpoint proteins from G0/G1 phase to S phase (cyclin D1, pRb, CDK4, and CDK6) was performed. Unloaded LNC were considered the vehicle control of IndOH-LNC. The histogram presents the quantitative Western blot analysis for the protein indicated. The densitometric values obtained from treatments were normalized to their respective controls (control bar) (100%). Bars represent the mean ± standard deviation for at least six independent experiments. Significant differences from control and LNC groups: ***P < 0.001, as assessed by one-way analysis of variance followed by the Bonferroni post hoc test.

Abbreviations: IndOH-LNC, indomethacin-loaded lipid-core nanocapsules; LNC, lipid-core nanocapsules.

In agreement with a previous study,26 our results showed a potent and selective cytotoxic effect of IndOH-LNC in GBM cells. At 24 hours of treatment, IndOH-LNC was at least fourfold more cytotoxic than IndOH. Of great relevance, IndOH-LNC was selectively cytotoxic to glioma cells and spared normal glial cells, since astrocyte viability was not significantly affected. According to the literature, the higher cytotoxic effect of nanocapsules in glioma cells may be triggered by a higher rate of cell division or by increased endocytotic activity resulting in
enhanced nanocarrier uptake. Thus, in our experiment, the cellular uptake of drug-loaded nanocapsules was probably sufficient to achieve a high intracellular concentration of indomethacin-loaded nanocapsules, and the subsequent intracellular release of the drug led to glioma cell death. This result is relevant since tumor-specific targeting is an improvement over classical chemotherapeutics, which adversely affect both diseased and normal brain in a nonspecific manner.

Inhibition of apoptosis and cell-cycle deregulation are two mechanisms of tumor formation. Therefore, it is desired that chemopreventive agents act by inducing apoptosis and cell-cycle arrest to block the carcinogenic process. In addition, defects in apoptotic signaling pathways, such as p53 mutations and caspase downregulation, can promote chemoresistance and thereby limit the effectiveness of chemotherapeutics. Our results showed that, at all concentrations tested, IndOH-LNC induced cell death mostly by apoptosis in C6 and U138-MG glioma cells. Despite the distinct genetic differences between C6 (PTEN and p53 wild-type) and U138-MG (PTEN and p53 mutant) cell lines, we found that both lines were sensitive to cell death triggered by IndOH-LNC, although the U138-MG cell line was more sensitive. This result is relevant because only about 35% of GBMs have mutant p53; therefore, it is important to select a therapy that can function independently of tumors’ p53 mutational status.
Figure 9 IndOH-LNC increase the GFAP protein immunocontent in glioma cells. Cells were treated with 25 µM IndOH-LNC for 72 hours, and immunocytochemistry for anti-GFAP antibody (red) and DAPI (blue) was performed in (A) U138-MG glioma cells. GFAP protein immunocontent was quantified by Western blot analysis for (B) C6 and (C) U138-MG glioma cells.

Notes: Unloaded LNC were considered the vehicle control of IndOH-LNC. The densitometric values obtained from treatments were normalized to their respective controls (control bar) (100%). Immunocytochemistry experiments were performed at least three times. In the Western blot analysis, bars represent the mean ± standard deviation for six independent experiments. Significant differences from control and LNC groups: ***p < 0.001, as assessed by one-way analysis of variance followed by the Bonferroni post hoc test.

Abbreviations: DAPI, 4′,6-diamidino-2-phenylindole; GFAP, glial fibrillary acidic protein; IndOH-LNC, indomethacin-loaded lipid-core nanocapsules; LNC, lipid-core nanocapsules.
The PI3K/AKT signaling pathway has a key role in regulating cell-cycle progression and apoptosis. The ectopic expression of AKT induces cell survival and malignant transformation, whereas the inhibition of AKT activity stimulates apoptosis in a range of mammalian cells. Thus, AKT is an attractive target for chemotherapy. It has been postulated that the inhibition of AKT alone or in combination with standard cancer chemotherapy will reduce the apoptotic threshold in cancer cells. The PI3K/AKT pathway is frequently abnormally activated in GBM as a result of a mutation or the loss of the tumor suppressor PTEN. Given that AKT is activated in approximately 70% of gliomas, the deregulation of the PI3K signaling pathway is an attractive target for therapy. Our results showed that 25 µM IndOH-LNC (the IC₅₀ value, approximately) was effective in reducing the phosphorylation levels of AKT after 1, 6, and 24 hours of treatment in C6 and U138-MG glioma cell lines. AKT may exert one of its downstream effects via the phosphorylation/inactivation of GSK-3β, which in turn inhibits β-catenin ubiquitination and degradation, resulting in increased β-catenin nuclear translocation. The Wnt/β-catenin complex regulates the transcription of multiple genes involved in cellular proliferation, differentiation, survival, and apoptosis, such as Fra-1, c-myc, and cyclin D. Increasing evidence indicates that aberrant activation of the Wnt/β-catenin signaling pathway plays a key role in glioma development and progression. We therefore determined whether the IndOH-LNC–induced decrease in AKT phosphorylation changed GSK-3β phosphorylation and/or β-catenin levels. We found that the decrease in AKT phosphorylation (rendering it inactive) after treatment with 25 µM IndOH-LNC was accompanied by a decrease in GSK-3β phosphorylation (rendering it active). In addition, IndOH-LNC treatment reduced β-catenin levels. Taken together, these results suggest that AKT inactivation may lead to reduced nuclear levels of β-catenin, thereby

**Figure 10** IndOH-LNC increase the nestin protein immunocontent in C6 glioma cells. C6 glioma cells were treated with 25 µM of IndOH-LNC for 72 hours. (A) Immunostaining for anti-nestin antibody (red) and DAPI (blue). (B) Quantification of nestin protein immunocontent by Western blot analysis. Notes: Unloaded LNC were considered the vehicle control of IndOH-LNC. The densitometric values obtained from treatments were normalized to their respective controls (control bar) (100%). Immunocytochemistry experiments were performed at least three times. In the Western blot analysis, bars represent the mean ± standard deviation for six independent experiments. Significant differences from control and LNC groups: ***P < 0.001, as assessed by one-way analysis of variance followed by the Bonferroni post hoc test.

**Abbreviations:** DAPI, 4′,6-diamidino-2-phenylindole; IndOH-LNC, indomethacin-loaded lipid-core nanocapsules; LNC, lipid-core nanocapsules.

**Figure 11** IndOH-LNC reduce CD133-positive stem cells in the U138-MG glioma cell line. Notes: Cells were treated with 25 µM IndOH-LNC for 72 hours, and immunocytochemistry for CD133 (green) and DAPI (blue) was performed in U138-MG glioma cells. Images are representative of six independent experiments.

**Abbreviations:** DAPI, 4′,6-diamidino-2-phenylindole; IndOH-LNC, indomethacin-loaded lipid-core nanocapsules; LNC, lipid-core nanocapsules.
negatively modulating glioma cell growth. Moreover, in the same manner as AKT, GSK-3β and β-catenin are important in the generation of an anti-apoptotic response. For this reason, inactivation of AKT and β-catenin, as well as activation of GSK-3β, could account, at least in part, for the apoptosis induced by IndOH-LNC in GBM cells.

It is worth noting that IndOH-LNC treatment modulated the cell-cycle dynamics of GBM cells. Cell-cycle analysis indicated that 25 μM IndOH-LNC caused G0/G1 phase arrest in C6 and U138-MG glioma cells. This G0/G1 arrest was accompanied by the formation of a sub-G1 population of cells, suggesting that G0/G1 arrest preceded DNA fragmentation and consequent apoptosis. This confirms the result obtained with annexin V staining. IndOH-LNC induced G2/M phase arrest in U138-MG cells only, in agreement with the greater percentage of apoptotic cells observed among U138-MG cells than among C6 glioma cells. Additionally, G2/M arrest, as an early step in the apoptotic mechanism, could also contribute to the selectivity of IndOH-LNC, since cancer cells constantly progress through S to G2/M phase, in contrast to nonproliferative cells.

We observed down- or upregulation of a number of cell-cycle regulators in GBM cells treated with IndOH-LNC, which is consistent with the outcome of cell-cycle arrest. Cyclin D1, one of the checkpoints in the G1 to S phase transition, was downregulated by IndOH-LNC in glioma cells. The phosphorylation of pRb also decreased in both cell lines after IndOH-LNC treatment. On the other hand, the CDK4 and CDK6 proteins were unaltered, suggesting that the reduction of cyclin D1 and pRb proteins represses the cyclin D1-CDK4/6-Rb pathway. Furthermore, in agreement with the G2/M arrest observed in U138-MG glioma cells, IndOH-LNC treatment reduced cyclin B1 levels and the phosphorylation status of cdc2. We also observed an increase in Wee1 levels and an accumulation of p21WAF1, components of the G2/M checkpoint that inhibits cdc2. These results are in agreement with the AKT downregulation induced by IndOH-LNC treatment, since when AKT is phosphorylated (rendering it active) it inhibits Wee1 and p21WAF1, driving the inhibition of the cyclin B1/cdc2 complex and consequently the progression of G2/M.

The effects of cyclin D1 regulation on cell growth are clear. Cyclin D1 binds to CDK4/CDK6 and phosphorylates Rb, which then releases the transcription factor E2F1, allowing it to transcribe genes that promote the G1/S phase transition. Thus, downregulation of cyclin D1 reduces Rb phosphorylation, inhibits E2F1 release, and ultimately hampers the entry of cells into S phase. In the mechanism by which IndOH-LNC represses proliferation, elevated p21 could inactivate the cyclin D1-CDK4/6 complexes. Additionally, accumulating evidence demonstrates that cyclin D1 levels are tightly regulated through multiple mechanisms, including the PI3K/AKT/GSK-3β and/or Wnt/β-catenin pathways. Several studies reported that inhibition of the PI3K/AKT/GSK-3β pathway, which culminates in GSK-3β activation, enhanced cyclin D1 protein stability and caused cyclin D1 accumulation.46,47 GSK-3β negatively influences cyclin D1 expression by regulating mRNA transcription and protein degradation.47 When transformed to its active form, GSK-3β regulates gene expression by phosphorylating specific aminoterminal residues of β-catenin, such as Ser33, Ser37, and Thr41, which first increases the degradation of cytoplasmic β-catenin, then inactivates Wnt/β-catenin signaling, and finally reduces cyclin D1 promoter activity.47 Thus, in the genesis and development of cancer, cyclin D1 is considered a cellular proto-oncogene that may be a candidate target to inhibit cell proliferation and induce cell differentiation in cancer therapy. Taken together, our results showed that control of the cell cycle can be modulated by the inactivation of AKT and β-catenin and the activation of GSK-3β, as observed in GBM cells treated with IndOH-LNC.

Differentiation therapy, using agents that promote cancer cell differentiation, has been shown to be effective in vitro and in vivo treatments of several types of cancer cells.54 The most successful clinical application of differentiation therapy has been the introduction of retinoic acid for the treatment of acute promyelocytic leukemia.49 Furthermore, the induction of cellular differentiation has become a promising therapeutic strategy against solid tumors, including glioma.50 The tumorigenic potential of GBM cells is, in part, attributable to their undifferentiated (neural stem cell-like) phenotype. The differentiation of glioma cells toward an astrocytic type is characterized by morphological transformation and the expression of differentiation markers. Astrocytic differentiation of GBM cells is associated with the transcriptional induction of GFAP and repression of nestin, whereas the reciprocal transcription program operates in undifferentiated GBM cells. GFAP is a well known biomarker of astrocyte differentiation.51 Several reports demonstrated that transfection of astrocytoma cells with GFAP cDNA effectively inhibit proliferation and tumorigenicity and induce morphologic alteration. Furthermore, GFAP is not only a reliable marker for gliogenesis but also a tumor suppressor gene in malignant glioma pathogenesis.53,54 Nestin, an intermediate filament protein, is one marker that has been used to identify neural
stem cells and is expressed in almost all GBMs. Nestin is downregulated in mature cells and is expressed in astrocytes of the adult central nervous system in response to cellular stress, such as neoplastic transformation. A high level of nestin expression reportedly indicates significantly shorter survival of glioma patients. Nestin may therefore correlate with invasiveness, being expressed mainly in the tumor periphery in the human brain. Additionally, the coexpression of nestin with the CD133 cell surface antigen in tumor cells has been investigated in detail because this phenotype is considered a marker of cancer stem cells. In the present study, we found that IndOH-LNC treatment upregulated GFAP protein expression and downregulated nestin in GBM cells. In addition, IndOH-LNC treatment induced a low staining for CD133 in U138-MG glioma cells. Therefore, our findings suggest that IndOH-LNC might function as a potential differentiation-supporting agent in treatments against malignant gliomas. The molecular mechanisms underlying the regulation of cancer cell differentiation remain elusive. Cell-cycle progression and differentiation are opposing but tightly linked events. In particular, G1 exit is a critical stage of the cell cycle where cells decide either to re-enter the cell cycle or differentiate. Compared with the definite role of cyclin D1 in cell-cycle regulation, the action of cyclin D1 in cell differentiation is less clear. However, reports have demonstrated that inhibition of cyclin D1 induces the differentiation of several tumor cell lines. Considering that our results showed that downregulation of cyclin D1 was induced by IndOH-LNC treatment in GBM cells, this mechanism could contribute to the cell differentiation observed here.

**Conclusion**

Overall, the data presented herein clearly show that multiple molecular mechanisms are involved in the tumor-suppressive effects of IndOH-LNC in glioma cells. IndOH-LNC activated AKT and activated GSK-3β, thus inhibiting growth and inducing apoptotic cell death. In conjunction with these changes, IndOH-LNC modulated the cell-cycle dynamics of GBM cells. Cells arrested in G0/G1 and G2/M phases after IndOH-LNC treatment, with concomitant decreases in cyclin D1 and pRb phosphorylation and increases in the levels of Wee1 and CDK inhibitor p21WAF1, among others. Additionally, the IndOH-LNC formulation promoted GBM cell differentiation. These findings, together with our previous results using a preclinical model of glioma, suggest that IndOH-LNC may be a candidate for further testing in clinical studies of gliomas.

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**Disclosure**

The authors report no conflicts of interest in this work.

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