Battling Glioblastoma: A Novel Tyrosine Kinase Inhibitor with Multi-Dimensional Anti-Tumor Effect (Running Title: Cancer Cells Death Signalling Activation)

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Abstract: Glioblastoma (GB), a grade IV glioma, with high heterogeneity and chemoresistance, obligates a multidimensional antagonist to debilitate its competence. Considering the previous reports on thioesters as antitumor compounds, this paper investigates on use of this densely functionalized sulphur rich molecule as a potent anti-GB agent. Bio-evaluation of 12 novel compounds, containing α-thioether ketone and orthothioester functionalities, identified that five analogs exhibited better cytotoxic profile compared to standard drug cisplatin. Detailed toxicity studies of top compound were evaluated in two cell lines, using cell viability test, apoptotic activity, oxidative stress and caspase activation and RNA-sequencing analysis, to obtain a comprehensive molecular profile of drug activity. The most effective molecule presented half maximal inhibitory concentration (IC50) values of 27 µM and 23 µM against U87 and LN229 GB cells, respectively. Same compound effectively weakened various angiogenic pathways, mainly MAPK and JAK-STAT pathways, downregulating VEGF. Transcriptome analysis identified significant promotion of apoptotic genes, and genes involved in cell cycle arrest, with concurrent inhibition of various tyrosine kinase cascades and stress response genes. Docking and immunoblotting studies suggest EGFR as a strong target of the orthothioester identified. Therefore, orthothioesters can potentially serve as a multi-dimensional chemotherapeutic possessing strong cytotoxic, anti-angiogenic and chemo-sensitization activity, challenging glioblastoma pathogenesis.

Keywords: glioblastoma; thioester; anti-angiogenesis; tyrosine kinase inhibitor
1. Introduction

Glioblastoma (GB) is a type of brain tumor with almost 100% recurrence rate and high resistance to current treatment modalities. The high heterogeneity exhibited within GB tumors render it unresponsive to single-target cytotoxic/anti-angiogenic agents and demands a significant clinical need for new multi-dimensional oncogenic inhibitors for GB.

Angiogenesis is a significant feature of GB, attributed to the overexpression of vascular endothelial growth factor (VEGF). Few of the most significant proangiogenic regulators that stimulate angiogenesis indirectly by inducing VEGF mRNA expression include the growth factors epidermal growth factor (EGF), transforming growth factor (TGF-α and TGF-β), tumor necrosis factor α (TNF-α), keratinocyte growth factor, insulin-like growth factor 1 (IGF-I), fibroblast growth factor (FGF), platelet-derived growth factor (PDGF), and cytokines (interleukin (IL)-1α and IL-6) and amplifications of Ras/Raf genes [1]. Multiple strategies have been developed to target VEGF/VEGFR receptor (VEGFR)–mediated angiogenesis, including VEGF blockade, VEGF Trap, and suppression of VEGFR signaling via receptor tyrosine kinase inhibitors (TKIs) [2–4]. EGFR belongs to the ErbB family receptors of Class I receptor tyrosine kinases (RTKs). Almost 60% of glioblastoma patients have some kind of genomic alteration affecting EGFR pathway [5]. Downstream effects of EGFR is mediated through phosphoinositide 3-kinase (PI3K), mitogen-activated protein kinase (MAPK), signal transducer and activator of transcription 3 (STAT3) pathways, and Src family kinases [6]. A number of studies have focused on inhibiting both VEGFR and EGFR so as to improve drug efficiency, including monotherapy with a multi-targeted tyrosine kinase inhibitor (e.g., vandetanib, AEE788, BMS-690514) [7,8]. Other significant pathways regulating tumor angiogenesis directly or indirectly via VEGF includes MAPK pathway [9], JAK-STAT pathway [10,11], and PI3K-AKT [12] pathway. Thus, a multi-targeted chemoagent that can effectively sequester multiple pathways involved in VEGF regulation would be an effective solution to tackle tumor pathogenesis.

Some of us have recently reported the unprecedented autoxidative condensation of 2-aryl-2-lithio-1,3-dithianes (Scheme 1) [13]. The result of such transformation was a small library of highly functionalized molecules containing α-thioether ketones and orthothioesters functionalities, among others. Motivated by the desire to find new agents capable of multi-target inhibition as promising approaches in the development of glioblastoma cancer drugs [14], we have set to assess the antitumor properties of these intricate molecules.

![Scheme 1. Synthesis and structures of studied orthothioesters.](image-url)
2. Materials and Methods

2.1. Synthesis of Ortho-thioesters

Reactions were monitored through thin-layer chromatography (TLC) with commercial silica gel plates (Merck silica gel, 60 F254). Visualization of the developed plates was performed under UV lights at 254 nm and by staining with cerium ammonium molybdate, 2,4-dinitrophenylhydrazine and vanillin stains. Flash column chromatography was performed on silica gel 60 (40–63 μm) as a stationary phase. NMR spectra were recorded with Varian Mercury 300 MHz or Jeol ECZR 500 instruments using CDCl$_3$ as solvent and calibrated using tetramethylsilane as internal standard. Chemical shifts (δ) are reported in ppm referenced to the CDCl$_3$ residual peak (δ 7.26) or TMS peak (δ 0.00) for $^1$H-NMR and to CDCl$_3$ (δ 77.16) for $^{13}$C-NMR. The following abbreviations were used to describe peak splitting patterns: s = singlet, d = doublet, t = triplet, m = multiplet. Coupling constants, J, were reported in Hertz (Hz).

High-resolution mass spectra (HR-MS) were recorded on a Waters ESI-TOF MS spectrometer. Tetrahydrofuran (THF) was dried by distillation under argon with sodium metal and benzophenone as an indicator. Dichloromethane (DCM) was dried by distillation under argon with calcium hydride. Structural elucidation of all compounds tested in biological assays was performed by $^1$H and $^{13}$C-NMR, and HR-MS. All compounds of interest were isolated by chromatography and their purity (>97%) assessed by NMR.

Compounds 1–4 were prepared and characterized as reported elsewhere [13], through antioxidative condensation of 2-aryl-2-lithium-1,3-dithianes, prepared from treatment of 2-aryl-1,3-dithianes with n-BuLi followed by air exposure.

$5a$: (4-(1,3-Dithian-2-yl)-2-methoxyphenoxy)(tert-butyl)diphenylsilane (1.09 g, 2.27 mmol) was dissolved in dry THF (10 mL) in an argon purged round-bottom flask. The solution was cooled to −78 °C in an acetone/liquid nitrogen bath. n-BuLi (1.3 equivalents) was added dropwise to the reaction mixture at −78 °C. After addition, the solution was left stirring at −78 °C for 20 min, and then left to warm up to room temperature for 40 min. The argon balloon was replaced with an atmospheric air balloon and an additional needle was inserted in the septum as to allow air flow through the surface of the solution. After one minute, the solution was quenched with saturated aqueous NH$_4$Cl solution (20 mL). Et$_2$O (20 mL) was added and the layers separated. The organic phase was collected and the aqueous phase was extracted with Et$_2$O (2 × 20 mL). The organic phases were combined and dried over MgSO$_4$. The solvent was filtered and evaporated to yield crude 3b. The mixture was dissolved in dry methanol (8 mL) in an argon purged round-bottom flask. Then, ammonium fluoride (95 mg, 2.56 mmol, 1.1 equivalents) was added and the solution was stirred overnight at room temperature. The methanol was evaporated and water (10 mL) was added. The aqueous phase was extracted with DCM (3 × 10 mL) and the organic phases were combined and dried over MgSO$_4$. The solvent was evaporated and the product was purified by flash chromatography, eluent hexane:ethyl acetate (3:2), to give the benzoyl derivative 5b as an amorphous white solid (163 mg, 0.151 mmol) in 95% yield.
1H NMR (500 MHz, CDCl$_3$) δ ppm 9.02 (s, 3H), 8.51–8.48 (m, 6H), 7.76–7.67 (m, 6H), 7.62 (dd, $J = 8.4, 2.0$ Hz, 2H), 7.25–7.08 (m, 5H), 5.56 (s, 1H), 3.86 (s, 3H), 3.85 (s, 3H), 3.83 (s, 3H), 3.31 (dd, $J = 13.3, 11.2$ Hz, 2H), 2.79 (dd, $J = 14.3, 3.6$ Hz, 2H), 2.70–2.58 (m, 4H), 2.14–2.11 (m, 1H), 1.98–1.91 (m, 1H), 1.86–1.81 (m, 2H). 13C-NMR (125 MHz, CDCl$_3$) δ 193.8, 162.6, 162.6, 162.2, 151.7, 151.6, 151.0, 148.5, 148.5, 143.8, 140.9, 139.5, 139.4, 136.1, 135.9, 134.9, 131.2, 131.2, 130.8, 130.1, 130.0, 128.3, 128.1, 128.1, 125.4, 122.9, 122.4, 122.3, 121.4, 120.6, 112.9, 112.8, 64.1, 56.3, 56.2, 56.2, 55.1, 32.8, 30.9, 29.5, 28.5, 24.3. HR-MS (ESI) $m/z$ calculated for C$_{51}$H$_{43}$N$_3$O$_16$S$_4$Na$^+$ [M + Na]$^+$ 1104.1418, found 1104.1385.

2.2. Cell Culture

Human GB cell lines, U87 cells were grown in Minimum Essential Medium (MEM, Product# 51416C, Sigma-Aldrich, St. Louis, MO, USA) with 10% Fetal Bovine Serum (FBS) 2 mM sodium pyruvate (Product# S8636, Sigma-Aldrich, St. Louis, MO), 1% Penicillin-Streptomycin and 0.025 mg/mL Amphotericin B. LN229 and non-cancerous cell line, mouse embryonic fibroblast (MEF) cells were cultured in Dulbecco’s Modified Eagle Medium—high glucose (DMEM, Catalog# L0102, Biowest, Riverside, CA, USA) containing 5% FBS (Product # F1051, Sigma-Aldrich, St. Louis, MO), 1% Penicillin-Streptomycin (Product # P4333, Sigma-Aldrich, St. Louis, MO) and 0.025 mg/mL Amphotericin B (Sigma-Aldrich, St. Louis, MO). U87 and LN229 are the standard cell lines derived from malignant gliomas used commonly for cytotoxicity study. MEF Cells were maintained at 37 °C in a humidified incubator supplemented with 5% CO$_2$. Three biological and technical repeats were used for each condition.

2.3. In Vitro Cytotoxicity Assay

Cytotoxicity assay was performed to determine the cell growth inhibitory effect of the compounds following treatment for 48 h on the GB cell lines, U87, LN229. This assay was performed in two stages. At first, a high concentration, 100 µM, of compounds were used as well as for the positive control Cisplatin, CIS (Sigma-Aldrich, USA). Following this, the top compound was selected, and different concentrations (100 µM, 75 µM, 50 µM, 25 µM, and 10 µM) of the compound were tested to determine the IC$_{50}$. Treated cells were harvested by centrifugation at 1200 rpm for 10 min. Cell viability was determined using trypan blue staining. The number of live and dead cells were counted using a Countess II Automated Cell Counter (Thermo Fisher Scientific, Carlsbad, CA, USA). The proliferation inhibition percentage of each sample was determined using the following formula to determine the dose-response curve. From the dose-response curve, the IC$_{50}$ value of each compound was calculated. The cytotoxicity of the top compound and Cisplatin at a concentration of 10 µM was also evaluated in normal brain cells (MEF). Percentage of inhibition of cell proliferation was calculated using the following formula:

$$\text{Proliferation inhibition (\%) = } \left( \frac{\text{Mean No. of untreated cells (DMSO control) - Mean No. of treated cells x100}}{\text{Mean No. of untreated cells (DMSO control)}} \right)$$

2.4. Double Staining Assay

U87 and LN229 cell lines were grown as described previously, followed by cells treatment with IC$_{50}$ concentration of 5a and incubated for 48 h. Untreated (Negative) and Cisplatin (positive) controls were also maintained. Apoptosis/necrosis detection was carried out using Annexin V-FITC and PI (Thermo Fisher Scientific). The apoptosis determination was performed following the standard protocol suggest by the manufacturer. Briefly, the cells were cultured in 6-well plate with an initial cell density of 7 x 10$^5$ cells/well. The cells were incubated for 48 h with 5a, positive control and untreated cells conditions and then harvested and washed in cold phosphate buffered saline (PBS). The cell pellets were then resuspended in 1× Annexin-binding buffer provided in the kit. Consequently, 5 µL of FITC conjugated Annexin V and 1 µL of the 100 µg/mL PI working solutions were added to the 100 µL
of cell suspension. The cells were incubated in dark for 15 min, at room temperature, after which the stained cells were observed for fluorescence to distinguish apoptotic cells. The image acquisition was done by using an EVOS imaging system (ThermoFisher Scientific) with 20× objective magnification. More than 300 cells were used for each analysis. The percentage of apoptosis was quantified based on the cells stained with Annexin V-FITC positive and PI negative and both Annexin V-FITC and PI positive. The percentage of necrosis was calculated based on the cells with Annexin V-FITC negative and PI positive [15–17]. The fold change in apoptosis was calculated against the untreated cells.

2.5. Caspase Activity Assay

In-vitro caspase-3 and caspase-7 activity was determined using Caspase-Glo® 3/7 Assay Systems (Promega Corporation, Madison, WI, USA). The reagent was prepared as mentioned in the manufacturer protocol. The U87 and LN229 cells were grown overnight in a 96-well plate and were treated with an IC_{50} of 5A. Negative control, positive control and blank (medium+ Dimethyl sulfoxide (DMSO)+ dye) were also maintained. Cells were incubated at 37 °C in a humidified incubator supplemented with 5% CO_2 for 5 h and then equilibrated at room temperature for 30 min. 100 µL of Caspase-Glo 3/7 reagent was added to 100 µL of cells/well and was incubated in a dark-chamber. The luminescence signal was quantified (Chameleon Multi-label Detection Platform) at one hour after treatment. Magnitude of fold change in luminescence between treated and untreated cells were determined using the following formula:

\[
\text{Fold increase} = \frac{F_{\text{test}} - F_{\text{blank}}}{F_{\text{control}} - F_{\text{blank}}}
\]

2.6. Intracellular Redox Potential Test

To evaluate the redox potential of 5a, a comparative test, using H_2O_2 and standard drug against untreated cells, was performed using H2DCFDA (Catalog no.# D399 Life Technologies, Eugene, OR, USA). The U87 and LN229 cells were grown overnight in a 96-well plate and were treated with an IC_{50} of 5a for 5 h at 37 °C in a humidified incubator supplemented with 5% CO_2. Negative control, positive control and blank were maintained. Baseline effect due to solvent was determined as well. After 5 h of treatment, cells were harvested by centrifugation at 3000 rpm for 10 min and incubated with 200 µL of 2 µM H2DCFDA for 30 min at 37 °C in the CO_2 incubator. Cells were then washed with pre-warmed PBS and resuspended in 200 µL of pre-warmed medium. Next, 100 µL of suspension was transferred to each well, in a 96-well plate and incubated at room temperature for 20 min. Finally, fluorescence signal was measured using Chameleon Multi-label Detection Platform (Excitation 485 nm, Emission 535 nm).

2.7. RNA Isolation and Gene Expression Evaluation

Cell were incubated with IC_{50} of test compound and standard drug Cisplatin for 48 h at 37 °C in a humidified incubator supplemented with 5% CO_2. A negative control was maintained as well. All conditions were conducted in triplicated samples for the isolation of RNA. Total RNA of >9.25 ng/µL was isolated using GeneJET RNA purification kit (Catalog no #K0731), according to the manufacturer’s instructions. The yield was then measured spectrophotometrically using NanoDrop-1000 (Thermoscientific, Wilmington, NC, USA). After quantification, the cells were considered for quality assessment by TapeStation. The gene expression analysis was performed by using Illumina Next Seq High throughput profiling (Illumina NextSeq 500). The sequencing produced data in bcl format which was converted into FASTQ file format.

2.8. Transcriptome Analysis

Differential expression (DE) and statistical analysis were performed using DESeq2 [18] (release 3.3) in R (version 3.2.4) (https://bioconductor.org/packages/release/bioc/vignettes/DESeq2/inst/doc/DESeq2.html). p-values were adjusted for multiple testing using the Benjamini-Hochberg procedure [19]. A false discovery rate adjusted p-value (i.e., q-value) < 0.05 was set for the selection of DE genes.
2.9. Phosphorylation of MAPKs and Other Serine/Threonine Kinases

The U87 cells were treated with an IC$_{50}$ concentration of 5a and DMSO, maintained in CO$_2$ humidified temperature for 48 h. The total protein was extracted using RnD protein extraction kit for the immunoblotting experiment. The phosphorylation of three families of mitogen-activated protein kinases (MAPKs), including the extracellular signal-regulated kinases (ERK1/2), c-Jun N-terminal kinases (JNK1-3), and different p38 isoforms, was analyzed following the manufacturer protocol, RnD Human Phospho-MAPK array kit. In detail, cell lysates are mixed with a cocktail of biotinylated detection antibodies and incubated with Proteome Profiler Human Phospho-MAPK Array. Streptavidin-HRP and chemiluminescent detection reagents are added and the signals that are produced at each spot correspond to the amount of phosphorylated protein bound. The captured and control antibodies have been spotted in duplicate on nitrocellulose membranes. The signals are measured using ChemiPro Luminescence detection system. The relative level of phosphorylation was analysed for 26 MAPK proteins as described in the manufacturer’s protocol.

2.10. Gene Ontology (GO) and Pathway Analysis

Gene ontology [20] and ClusterProfiler package [21] was used for pathway analyses. We performed with the over-representation test for the GO biological process and KEGG pathways [22]. The package supports the human genome. We used the binominal test and Bonferroni correction for multiple testing and displayed z-scores to indicate whether a potential regulator was activated or inhibited. We used the default settings for statistical analysis in both the KEGG pathways and GO terms. In this analysis, pathways and GO terms only with a $p$-value $<$ 0.05 were included.

2.11. Evaluation of Structure-Activity Relationship

In order to make a comparative study on the binding of 5a with different receptors, sequences and structures of six receptors taken from Protein Data Bank (PDB)—Fibroblast Growth Factor receptor (FGFR, PDB ID: 2FDB), Epidermal growth Factor Receptor (EGFR, PDB ID: 4UIP), Platelet-Derived Growth Factor Receptor (PDGFR, PDB ID: 1PDG), c-MET Receptor (PDB ID: 3DKC), c-KIT Receptors (PDB ID: 6GQK) and Vascular Endothelial Growth Factor receptors (PDB ID: 3V2A) were obtained from the Protein Data bank. To study the binding efficiency and to identify the important amino acid residues contribute to the binding, Patchdock molecular docking program was used. The default parameters were used for the molecular docking in this study. Patchdock carries out rigid docking, with surface variability implicitly addressed through liberal intermolecular penetration.

2.12. Statistical Analysis

All experiments described in the present study were performed as with three biological and technical repeats. The data were presented as the mean ± standard error of mean. Statistical analyses between two groups were performed by Student’s $t$-test. Differences among multiple groups were tested by one-way analysis of variance following by a Dunnett’s multiple comparison test (GraphPad Prism 7.04, San Diego, CA, USA). $p < 0.05$ was considered to indicate a statistically significant difference.

3. Results

3.1. Synthesis of Orthothioesters

The orthothioesters 1-4 were synthesized through autoxidative condensation of 2-aryl-2-lithium-1,3-dithianes, prepared in situ from treatment of 2-aryl-1,3-dithianes with $n$-BuLi followed by air exposure. The presence of bromide when preparing 1c demanded for replacement of $n$-BuLi by lithium diisopropylamide (LDA) in order to avoid transmetalation. The compounds were obtained in reasonable yields and pure after quenching with NH$_4$Cl saturated aqueous solution, usual work-up and purification by silica chromatography. The silyl ether group in compound 3b
was cleaved by treatment with 1 equivalent of NH$_2$F to yield phenol 5a. Derivatization of 5a with m-nitrobenzoyl chloride delivered carboxylic ester 5b in excellent yield. Purity evaluation results of the novel compounds are given in the Supplementary file 1.

3.2. 5a Inhibited Proliferation of Glioma Cells

The cytotoxicity of the orthothioesters 1–5 against the growth of U87 GB cell line was performed with 100 μM concentration of the compounds as described in the method section. Upon evaluation of the 12 compounds depicted in Scheme 1, 5a was identified to be the most cytotoxic compound inducing 90% cell death and 5b was the least cytotoxic with 0% cell death. This was a 20-fold higher activity compared to the standard drug cisplatin and 31.8-fold compared to the negative control (Figure 1a). The analysis of cytotoxic data of all compounds tested shows that while the presence of halogens or nitrile in the 4-position of the aromatic rings is detrimental for cytotoxicity (as for compounds 1b–d), having a methoxy group in the 3-position has an opposite effect. Compounds 2c, 3a and 5a, decorated with a 3-methoxy substituent in the aromatic rings, showed cytotoxicity similar to or higher than 1a. Compounds 3b and 5b on the other hand, also decorated with the same 3-methoxy substituent but having bulky substituents in the vicinal 4-position, have residual or absent cytotoxicity. No improvement on cytotoxicity was observed when replacing the aryl groups by heteroaryl such as pyridyl in 4a. The cytotoxicity on normal brain cells, MEF, was also performed for the top compound 5a (Figure 1b) and cisplatin at a concentration of 10 μM. The results show that 5a have 11% of growth inhibition, whereas cisplatin have 48% of inhibition, suggesting that 5a is more toxic to the cancer cells than the normal cells (Figure 1c).

**Figure 1.** In vitro screening and evaluation of dose response of thioester compounds (a). The percentage of mortality rate on treating U87 cells with novel panel of 12 thioester derivatives along with CIS (positive control) 1% DMSO (negative control), at a 100 μM concentration (b). Chemical structure of the investigative drug 5a. (c) Cytotoxicity analysis of non-cancerous brain cells, MEF and cisplatin at 10 μM concentration. (d) Dose response curve of U87 cells (left) and (e) LN229 (right) cells treated with the top compound (5a), CIS and 1% DMSO was analyzed to determine IC$_{50}$ value. Treatment was carried out at 48h and cellular viability was measured by Trypan Blue exclusion method. Data points and error bars represent mean ± standard error of the mean (SEM) (n = 6 per group). Significant when p < 0.05, one-way ANOVA.

In order to investigate the in vitro anticancer activity of 5a and to determine its IC$_{50}$ value, we evaluated cytotoxicity against cell lines, U87 and LN229 in the presence of increasing concentrations of this compound for 24 h, ranging from 0 μM–150 μM. The viability of cells was determined according to cell counting based on Trypan Blue assay and the IC$_{50}$ value of 5a was determined as 27 μM in U87.
cells and 23 µM in LN229 cells (Figure 1d,e). The IC$_{50}$ value of cisplatin was identified as 53 µM in U87 cells and 115 µM in LN229.

### 3.3. 5a Induced Negligible Oxidative Stress and Promoted Caspase Activation

Reactive oxygen species (ROS) mediated caspase activation of tumor cells during stress and subsequent cell death has been repeatedly reported by various studies [23,24]. In the present study, both intracellular ROS and caspase in U87 cells were quantified to verify oxidative stress and cellular response upon 5a treatment. After exposure of U87 cells to 5a at IC$_{50}$ for 5 h, we detected an oxidative increase of 3.3% increase in the 5a treated cells when compared to untreated cells. The standard drug cisplatin and positive control H$_2$O$_2$ on the other hand, marked a 4.2% and 2.2% increase in oxidative response respectively (Figure 2a). Similarly, exposure of LN229 cells to 5a at IC$_{50}$ for 5 h demonstrated negligible change in the treated cells when compared to untreated cells (Figure 2a). The difference between treated and untreated conditions was confirmed to be statistically insignificant per ANOVA test ($p$-value $< 0.05$), establishing that the 5a does not increase ROS in the tested GB cells.

![Figure 2.](image)

**Figure 2.** Comparative evaluation of oxidative stress response and caspase activation by the 5a in two different cell lines (a) Effect of 5a on intracellular reactive oxygen species (ROS) production by GB cells as interpreted by fluorescence level. Evaluation suggested no significant increase in ROS level upon drug treatment ($\alpha = 0.05$). (b) Caspase activity displayed by GB cells analyzed using caspase 3/7 luminescence assay. Significant increase in caspase activity of cells was observed after treatment with IC$_{50}$ of 5a for 5h compared to control. The values are expressed as means ± standard error of the mean (SEM) of triplicate measurements of biological repeats (n = 3). Significance: **** $p < 0.0001$. 
Considering the role of Caspase activation, we determined the caspase activity of U87 cells using caspase3/7 assay after a treatment period of 5 h with 5a at IC₅₀. Interestingly, U87 5a-treated cells displayed an increase of caspase 3/7, displaying a 0.43-fold increase in comparison to untreated cancer cells, whereas positive control displayed no significant increase (0.003) in caspase activity (Figure 2b). Similarly, in LN229 cells 5a-treated cells demonstrated a 1.37-fold increase with respect to untreated cells, whereas the standard drug cisplatin exhibited a 0.05 fold increase in caspase levels. The difference between treated and untreated conditions was confirmed to be statistically significant per ANOVA test (p-value < 0.05).

3.4. 5a Enhanced Apoptosis and Cell Cycle Arrest

Defective apoptotic machinery enhances tumor pathogenesis by permitting survival of genetically unstable cells leading to treatment resistance. To understand if the decrease in viability was due to apoptosis, we treated U87 cells with IC₅₀ of 5a and we determined the apoptosis effect using double staining method.

The results revealed that 5a was much effective in inducing apoptosis exhibiting a 3.75-fold increase in programmed cell death, whereas the standard drug cisplatin induced a 1.7-fold increase when compared to untreated cells (Figure 3a,b). To explore in detail the effect of 5a transcriptomics, data analysis was carried out, and we identified 1148 differentially expressed genes (DEGs) when 5a is compared with untreated (negative control) samples (q-value < 0.05). We also compared the 5a and cisplatin samples as a positive control group, both individually and combined as a single “affected” group. In these comparisons, 3132 DEGs were identified, with the largest number of DEGs identified in comparison with the cisplatin samples. In total, 595 out of 4280 DEGs were common in both comparisons. The complete lists of DEGs from the cell line analysis and all pairs of comparisons are provided in Supplementary file 2. Gene expression analysis indicated that 5a has a multidimensional impact on various tumorogenic features, especially on tyrosine kinase signaling (Figures 4 and 5a) and was effective in increasing expression of agonists of cell death such as DKK1, ADM, HMGA2, and CAV1. In addition, downregulation of chemoprotective factors such as HMOX1, HSPs, and CYP1B1 were detected. Adding to its cytotoxic effects, 5a was also found to suppress mitochondrial membrane stability as evident by dysregulated levels of HSPA1A and BNIP3.

Gene expression analysis also points out that treatment of U87 cells with 5a leads to inhibition of G1/S transition leading to cell cycle arrest. Upregulation of cell cycle inhibitory genes involved in G1/S such as DACT1, SUSD2, and CTDSP1 signifies the efficiency of the drug as a cell cycle inhibitor (Figure 5c). Additionally, genes favoring G1/S transition such as PLRG1 and ADAM17 were effectively downregulated due to 5a treatment. The cell cycle interruption is further strengthened by suppression of CCND3 affecting the CDK4 activity associated with this cyclin, which is necessary for cell cycle progression. Several studies have proposed that agents that interfere with DNA repair can act as a therapeutic strategy targeting double-strand break repair pathways or abrogate cell cycle checkpoints. HMGA2 gene involved in negative regulation of double-strand break repair via non-homologous end joining was found to be risen upon the treatment with 5a demonstrating its genotoxicity leading to the apoptotic death of cancer cells. Additionally, notable chemo-sensitizing was visible in treated cells as evident from transcriptome levels of HSPA1A, CLU, and TXN (Table 1).
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ANOVA test (3.4. was due to apoptosis, we treated U87 cells with IC50 of 5a and we determined the apoptosis effect genetically unstable cells leading to treatment resistance. To understand if the decrease in viability difference between treated and untreated conditions was confirmed to be statistically significant per

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The results revealed that 5a was found to effectively induce apoptotic cell death when compared to untreated cells. (b) The histogram represents quantification of apoptosis after treating the cell lines with IC50 of 5a, for 48 h, in U87 cell line. Cisplatin was used as a positive control and untreated cells as negative control. The fold change of apoptosis has been calculated against untreated U87 cells. The values are expressed as means ± standard error of the mean (SEM) of triplicate measurements of biological repeats (n = 3). **** p < 0.0001 as produced by ANOVA test. (c) Key genes involved in Apoptotic process and their log2(fold-change), that are differentially expressed upon R114 treatment, compared to untreated. log 2(fold-change) cutoff = 1.5.

Figure 3. Evaluation of apoptotic activity induced by 5a. (a). Phase contrast image of non-apoptotic cells in untreated condition (left); fluorescence image of U87 cells exhibited upon PI (middle) and Annexin V (right) staining. 5a was found to effectively induce apoptotic cell death when compared to untreated cells. (b) The histogram represents quantification of apoptosis after treating the cell lines with IC50 of 5a, for 48 h, in U87 cell line. Cisplatin was used as a positive control and untreated cells as negative control. The fold change of apoptosis has been calculated against untreated U87 cells. The values are expressed as means ± standard error of the mean (SEM) of triplicate measurements of biological repeats (n = 3). **** p < 0.0001 as produced by ANOVA test. (c) Key genes involved in Apoptotic process and their log2(fold-change), that are differentially expressed upon R114 treatment, compared to untreated. log 2(fold-change) cutoff = 1.5.

Figure 4. Representation of DEGs in major signal transduction pathways, in 5a treated cells. The color code denotes over-representation of that pathway by DEGs. FDR represents the corrected over-representation probability. Color scale denotes the proportion of entities identified among the total entities enriched in a pathway (Image generated via the reactome pathway analysis tool; https://reactome.org/PathwayBrowser/).
Figure 5. Transcriptional profile of 5a compared to standard drug (a) Venn diagram showing 595 genes commonly differentially expressed in both treatments versus control, as assessed by RNA-seq data analysis of U87 cell line. (b) Visual representation of top 14 pathways associated with differentially expressed genes upon 5a treatment with reference to control. (c) Gene interaction network of cell cycle genes affected upon 5a treatment.

Table 1. Key genes affected by the top drug-like compound.

| Signalling Pathways | MAPK | TNF | PI3K-Akt | STAT | ErbB | NF-kB |
|---------------------|------|-----|----------|------|------|-------|
| AREG                | CASP3| AREG| CCND3    | CDKN1A| AREG| BCL10 |
| CASP3               | FOS  | CCND3| CDKN1A   | IL15 | CDKN1A| IL1R1 |
| CRK                 | IL15 | AREG| CDKN1A   | CDKN1A| CRK | PTGS2 |
| Downregulated genes | FGF2 | IL18R1| MCL1     | HBEFG| JUN | TNF   |
| FOS                 | JUN  | MCL1| STAT1    | JUN  | JUN |       |
| IL1A                | JUNB | PGF | SGK1     |       |     |       |
| IL1R1               | PTGS2| JUNB| THBS1    |       |     |       |
| JUN                 | RIPK1| TNF |         |       |     |       |
| LAMTOR3             | PGF  |     |          |       |     |       |
|                     | TNF  |     |          |       |     |       |
| Upregulated genes   | DUSP6| JAG1| BCL2L1   | BCL2L1| EGFR| BCL2L1|
| EGF                 | LIF  | EGFR| EGFR     | MYC  | EGFR|       |
| EPHA2               | RPS6KA5| EPHA2| LIF     | MYC  | MYC |       |
| MET                 | F2R  | GNG4| PDGFA    | PDGFB| PDGFB|       |
| MYC                 | IRS1 | PDGFA|       |       |     |       |
| PDGFA               | ITGA6| MET | VEGFA    |       |     |       |
| RPS6KA5             | STMN1| MYC | BMP4     | VEGFA| RASA3|       |
|                     |      |     |         |PDGFA |      |       |
|                     |      |     |         |PDGFB |      |       |
|                     |      |     |         |BMP4  |      |       |
|                     |      |     |         |TGFA  |      |       |
|                     |      |     |         |       |      |       |
3.5. 5a Is an Anti-Angiogenic Agent

To gain more insight on anti-angiogenic efficacy of 5a, we analyzed the modulation of genes in U87 glioma cells treated with the investigative drug. 5a was found to adversely affect the cytokine receptor pathway exerting its effect through interaction of multiple pathways, eventually downregulating VEGF expression. VEGFD itself exhibited a log2 (fold change) of -4.5. Other key genes (TNF, HMOX, IL1A, CYP1B1, and PTGS1) involved in positive regulation of VEGF were identified to be downregulated by more than 1.75-fold. Key genes involved in inhibition of angiogenesis, including NR2F1, SEM3A, ERRFI1, SPRY1, ADM, NRP1, etc., were found to be upregulated (log2 (fold change) > 1.5). Moreover, enhancers of migration and angiogenesis such as GPNMB, PTGS2, CD274, and ZNF703 were significantly downregulated suggesting suppression of tumor malignancy (Table 1 and Supplementary file 2).

5a advocates inhibition of angiogenesis targeting VEGF pathway, via multiple pro-angiogenic regulators. The mRNA levels of VEGF were markedly decreased in the U87 cells treated with the IC50 concentrations of 5a. Moreover, other angiogenic enhancers such as HMOX1, IL1A, CYP1B1 PGF, and FGF were effectively suppressed by 5a. Additionally, such a molecule potentially weakened angiogenesis by affecting pathways involved in the crosstalk by interrupting key regulators involved. The phosphorylation of multiple kinase proteins involved in the VEGF signaling pathway were quantified using immunoblotting. The upregulation of CREB, GSK-3αβ, GSK-3β, MSK2, p38α, p38γ, and p53 and down regulation of JNK1 validates that the VEGF pathway might be targeted by 5a (Figure 6a–c).

3.6. 5a Restricts MAPK Signaling Cascade and Downregulates TNF Expression

Activation of the MAPK pathway leads to the transcription of genes that encode proteins involved in the regulation of essential cellular functions, such as cell growth, cell proliferation, and cell differentiation. U87 glioma cells treated with IC50 concentrations of 5a exhibited downregulation of positive regulators of MAPK pathway such as GADDs, RIPK, CTNNB1 and PSAP, whereas, it enhanced the expression of inhibitors of MAPK signaling such as MYC, SPRY1, EZR, RASA3 and CAV1. Additionally, 5a reduced the expression of activators of ERK cascade (TNF, CTGF, FGF, JUN) and enhanced its negative modulators such as CIQL4 and TNIP1 (Figure 5b, Table 1 and Supplementary file 2). TNF exerts its biological functions by activating distinct signaling pathways such as nuclear factorκB (NF-κB) and c-Jun N-terminal kinase (JNK). We observed a significant reduction of TNF levels in 5a treated cells compared to untreated cells. Inhibitors of TNF production like ERRFI1, RARA, VSIR and AXIL were found to be overexpressed explaining the reduction in TNF level. Additionally, immunoblotting revealed the phosphorylation of CREB, p38, JNK proteins involved in MAPK/TNF signaling pathway. The relative higher-level expression of CREB, p38 and reduced expression of JNK protein, also validates the inhibition of MAPK signaling cascade by 5a in U87 cells (Figure 6a–c).

3.7. 5a Negatively Affects JAK-STAT Pathway

The direct and mediated mechanisms of JAK-STAT signaling in tumor cell survival, proliferation, and invasion have made the JAK-STAT pathway a feasible target for drug development and cancer therapy. Interactions of JAK/STAT pathway with the RTK/Ras/MAPK pathway, TGF-β signaling pathway and PI3K pathway amplifies the effect mediated through the regulation of JAK-STAT signaling. The current study revealed that 5a is a strong suppressor of JAK-STAT pathway causing notable downregulation of major and downstream genes such as STAT1, MCL1, CCND3, CDKN1A, and IL15 (Table 1 and Supplementary file 2). Upregulation of PTPN2, an inhibitor of JAK expression was observed, which is also a contributor of endoplasmic reticulum stress-induced intrinsic apoptotic signaling pathway. Additionally, upregulation of CAV1 and HMGA was identified explaining the downregulation of JAK-STAT pathway. Phosphorylation analysis of multiple kinases shows the upregulation of CREB and p38 proteins, suggesting a role of 5a in inhibiting JAK-STAT signaling pathway (Figure 6a–c).
Several cross-family interactions among tyrosine kinases may significantly alter angiogenic signaling, leading to anti-angiogenic drug resistance. Reports say FGF-driven angiogenesis is blocked by VEGF inhibition, which suggests that FGF controls angiogenesis upstream of VEGF by modulating VEGF function [27]. Additionally, in the case of glioma tumorigenesis, PDGF-expression is assumed to contribute to the expansion of an established tumor as well as the regulation of the angiogenic switch for initial tumor development [28]. Studies also testify that there are cross family interactions between VEGF and PDGFR [29]. In addition, cKIT and MET are another two important RTKs to be explored as angiogenic drivers [30]. c-KIT signaling promotes cell proliferation and survival, exerting its effect through Ras-Erk pathway as well as JAK/STAT pathway [31]. Chen et al. asserted that intracrine VEGF function can be regulated by MET signaling and plays a significant role in controlling VEGFR2 [32].
In this context, we sought to explore interactions between promising orthothioester 5a and six different tyrosine kinase receptors—FGFR, EGFR, PDGFR, c-MET, cKIT, and VEGFR-2, by in silico docking study. The results show that the Epidermal growth Factor Receptor shows good binding efficiency with thioester with high docking score (7256) (Figure 7a). The two-dimensional ligand interaction diagram of thioester with EGFR is shown in Figure 7b. Conclusively, our study evidences a multidimensional anti-tumor effect of the novel thioester drug 5a.

| S.No | RECEPTORS   | Patchdock Score |
|------|-------------|-----------------|
| 1    | FGFR        | 5712            |
| 2    | EGFR        | 7256            |
| 3    | PDGFR       | 5094            |
| 4    | c-MET       | 6194            |
| 5    | cKIT        | 6890            |
| 6    | VEGFR2      | 4700            |

**Figure 7.** Docking score of 5a with various kinase receptors (a) Docking analysis of various cytokine receptors (Fibroblast Growth Factor receptor—FGFR, PDB ID: 2FDB, Epidermal growth Factor Receptor—EGFR, PDB ID: 4UIP, Platelet-Derived Growth Factor Receptor—PDGFR, PDB ID: 1PDG, c-MET Receptor, PDB ID: 3DKC, c-KIT Receptor- PDB ID: 6GQK, and Vascular Endothelial Growth Factor receptors PDB ID: 3V2A) with 5a. Docking study was carried out using the Patchdock program. (b) 2-dimensional interaction diagram of EGFR with ligand.

### 3.8. 5a Induced Phosphorylation of MAPKs and Other Serine/Threonine Kinases

The effect of 5a on the phosphorylation levels of three families of MAPKs, such as ERK1/2, JNK1–3 and different p38 isoforms, was analyzed. The arrays used in the experiments are based on the analysis of kinase specific antibodies which are spotted in duplicates along with three reference spots and PBS as negative spots. The mixture of biotinylated anti-phospho-kinase antibodies and streptavidin-HRP conjugate differentiates the phosphorylated and non-phosphorylated protein between the control and 5a treated cells. The luminescence induced by the addition of chemiluminescence reagent was captured using the ChemiPro detection system (Figure 6a). In detail, the relative levels of phosphorylation of 26 kinases such as Akt1, Akt2, Akt3, Akt, CREB, ERK1, ERK2, GSK-3α/β, GSK-3β, HSP27, JNK1, JNK2, JNK3,
was observed in tumor cells upon treatment with 5α. FGF can act synergistically with VEGF. The present study validates potential of a novel orthothioester pathways as observed in our experimental results. The present research thus suggests 5α (Ser<sup>133</sup>) path as a candidate anti-GB agent. Additionally, 5α multi-targeted agent against GB signaling pathway and highlighting its significance as anti-tumor HSPs, CLU, SNAI1, PTGS2, stress proteins such as TNF-α/β, 5α. Furthermore, the investigative drug also exertively induced chemo-sensitization by suppressing other proangiogenic factors [35]. FGFs have been reported to have potent proangiogenic effects through the stimulation and release of other proangiogenic factors [35]. 5α was found to effectively reduce FGF levels, causing a synergistic anti-angiogenic effect as well as increased sensitization to VEGF suppression. Additionally, potent inhibition of TNF-α by 5α may also contribute to its anti-angiogenic impact. Though TNF is widely promoted as an antineoplastic agent, its dual role as a angiogenesis promoter and inhibitor has been discussed in multiple reports [36–38]. A study by Giraudo et al. discussed the VEGF mediated role of TNF-α in the initiation and maintenance of angiogenesis and increased vascular permeability [39]. Furthermore, the investigative drug also effectively induced chemo-sensitization by suppressing stress proteins such as HSPs, CLU, SNAI1, PTGS2, and MCL1. Taken together, we postulate 5α as a multi-targeted agent against GB signaling pathway and highlighting its significance as anti-tumor agent. Additionally, 5α exerts its cytostatic effect via key genes involved in regulation of cell cycle pathways as observed in our experimental results. The present research thus suggests 5α as a candidate GB chemotherapeutic with multiple anti-cancer properties.

4. Discussion

In order to challenge tumor recurrence and resistance of GB, agents that can synergistically confront multiple oncogenic pathways are gaining great importance in GB drug discovery. In a pursuit to identify a multi-targeted chemo-agent against glioblastoma, we assessed bioactivity of a panel of new orthothioester derivatives as anti-GB agents in vitro.

Cytotoxicity evaluation identified an orthothioester 5α, which has inhibited tumor cell growth in a dose-dependent manner in GB cell lines. In addition, it displayed a more effective anti-tumor activity than current standard drug cisplatin in both GB cell lines U87 and LN229. Gene expression profiling revealed that this effect is due to the suppression of multiple kinase signaling pathways regulating VEGF levels in GB cells suggesting the possible anti-angiogenic effect. Additionally, inhibition of HMOX1 abrogates VEGF-induced endothelial activation and subsequent angiogenesis [33]. A robust silencing of HMOX1 observed in our study convinces curbing of VEGF-induced pathogenic angiogenesis.

Virtual binding studies using bioinformatics tools suggests EGFR as a key target of 5α, exerting its effect through multiple pathways leading to the inhibition of angiogenesis, cell cycle arrest, and apoptosis. Considering previous reports that EGFR can activate β-catenin via receptor tyrosine kinase-PI3K/Akt pathway [34], inhibitory ligand binding on EGFR is expected to diminish WNT signaling. Our observation of weakened expression of CTNNB1 and other positive regulators of WNT pathway (DAB2, GSKIP) corresponds to this notion.

The candidate drug 5α seems to be an attractive anti-angiogenic agent acting beyond VEGF/VEGFR pathway. Notable downregulation of growth factors (VEGFD, FGF2, PGF, TNF-α) and cytokine IL-1A was observed in tumor cells upon treatment with 5α. FGF can act synergistically with VEGF to amplify tumor angiogenesis and are implicated in the emerging phenomenon of resistance to VEGF inhibition. FGFs have been reported to have potent proangiogenic effects through the stimulation and release of other proangiogenic factors [35]. 5α was found to effectively reduce FGF levels, causing a synergistic anti-angiogenic effect as well as increased sensitization to VEGF suppression. Additionally, potent inhibition of TNF-α by 5α may also contribute to its anti-angiogenic impact. Though TNF is widely promoted as an antineoplastic agent, its dual role as a angiogenesis promoter and inhibitor has been discussed in multiple reports [36–38]. A study by Giraudo et al. discussed the VEGF mediated role of TNF-α in the initiation and maintenance of angiogenesis and increased vascular permeability [39]. Furthermore, the investigative drug also effectively induced chemo-sensitization by suppressing stress proteins such as HSPs, CLU, SNAI1, PTGS2, and MCL1. Taken together, we postulate 5α as a multi-targeted agent against GB signaling pathway and highlighting its significance as anti-tumor agent. Additionally, 5α exerts its cytostatic effect via key genes involved in regulation of cell cycle pathways as observed in our experimental results. The present research thus suggests 5α as a candidate GB chemotherapeutic with multiple anti-cancer properties.

5. Conclusions

Drugs that can act as multi-targeted agents can enhance efficacy and confront chemoresistance exhibited by GB cells. Thioesters has been investigated as an antitumor agent in multiple studies [40,41]. The present study validates potential of a novel orthothioester 5α, as an excellent pharmacological scaffold possessing strong cytotoxic, anti-angiogenic, and chemo-sensitization activity. The compound 5α exerted extensive killing effects on two different glioma cell lines by effectively weakening resistance
pathways and enhancing apoptotic machinery. Additionally, 5a is a strong cytostatic agent acting on key genes involved in regulation of cell cycle pathways. In particular, the capability of 5a to impede various pathogenic signaling cascades leading to GB pathogenesis, by acting on multiple tyrosine kinase pathways, makes it an appealing anti-GB agent. Taken together, our report provides new insights on how underexplored thioester derivatives can act as a potent pharmacological scaffold against glioblastoma.

**Supplementary Materials:** The following are available online at http://www.mdpi.com/2073-4409/8/12/1624/s1: NMR spectroscopy results of previously unreported compounds are attached as Supplementary file 1. List of differentially expressed genes and pathway analysis are attached as Supplementary file 2.

**Author Contributions:** J.R.V., C.A.M.A., and N.R.C. synthesized and characterized the compounds and A.V. executed the experiments and data analysis. S.K.M. executed docking studies. A.M. (Akshaya Murugesan) executed protein studies. A.M. (Aliyu Musa) analyzed RNA-seq data. O.Y.-H. and M.K. conceived and managed all studies. All the authors contributed to writing the manuscript.

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**Abbreviations**

- **GB** Glioblastoma
- **TKI** Tyrosine Kinase Inhibitor
- **RTK** Receptor Tyrosine Kinases
- **ROS** Reactive Oxygen Species
- **GO** Gene Ontology
- **TLC** Thin-Layer Chromatography
- **DCM** Dichloromethane
- **TCF** Tetrahydrofuran
- **DE** Differentially expression
- **DEG** Differentially expressed gene

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