Allosteric Antagonist Modulation of TRPV2 by Piperlongumine Impairs Glioblastoma Progression

João Conde, Ruth A. Pumroy, Charlotte Baker, Tiago Rodrigues, Ana Guerreiro, Bárbara B. Sousa, Marta C. Marques, Bernardo P. de Almeida, Sohyon Lee, Elvira P. Leites, Daniel Picard, Amrita Samanta, Sandra H. Vaz, Florian Sieglitz, Maíke Langini, Marc Remke, Rafael Roque, Tobias Weiss, Michael Weller, Yuhang Liu, Seungil Han, Francisco Corzana, Vanessa A. Morais, Cláudia C. Faria, Tânia Carvalho, Panagis Filippakopoulos, Berend Snijder, Nuno L. Barbosa-Morais, Vera Y. Moiseenkova-Bell,* and Gonçalo J. L. Bernardes*

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ABSTRACT: The use of computational tools to identify biological targets of natural products with anticancer properties and unknown modes of action is gaining momentum. We employed self-organizing maps to deconvolute the phenotypic effects of piperlongumine (PL) and establish a link to modulation of the human transient receptor potential vanilloid 2 (hTRPV2) channel. The structure of the PL-bound full-length rat TRPV2 channel was determined by cryo-EM. PL binds to a transient allosteric pocket responsible for a new mode of anticancer activity against glioblastoma (GBM) in which hTRPV2 is overexpressed. Calcium imaging experiments revealed the importance of Arg539 and Thr522 residues on the antagonistic effect of PL and calcium influx modulation of the TRPV2 channel. Downregulation of hTRPV2 reduces sensitivity to PL and decreases ROS production. Analysis of GBM patient samples associates hTRPV2 overexpression with tumor grade, disease progression, and poor prognosis. Extensive tumor abrogation and long term survival was achieved in two murine models of orthotopic GBM by formulating PL in an implantable scaffold/hydrogel for sustained local therapy. Furthermore, in primary tumor samples derived from GBM patients, we observed a selective reduction of malignant cells in response to PL ex vivo. Our results establish a broadly applicable strategy, leveraging data-motivated research hypotheses for the discovery of novel means tackling cancer.

INTRODUCTION

Natural products provide ample opportunities to develop innovative medicines. However, understanding their mechanisms of action remains a bottleneck to unlock their promise in drug discovery. Chemoproteomics is a privileged approach to unveil new biology for molecules of therapeutic interest. However, such methods are laborious and time-consuming and unlikely identify membrane proteins and targets with only minute expression. Moreover, chemoproteomics relies on structural manipulation of the chemical entities under study, which may have unpredictable outcomes. Considering the large volume of publicly available bioactivity data, statistical learning of ligand−target relationships has become tractable in recent years and amenable to providing probabilistically motivated target binding hypotheses. One such ligand-based target-identification program (SPiDER: self-organizing map-based prediction of drug equivalence relationships) has been validated for the deorphanization of natural products and discovery of new biology, drug repurposing and the unravelling of primary and secondary pharmacology. Indeed, in silico technologies are a growing concept in drug discovery that may offer complementary/alternative solutions for not only target identification programs but also discovery chemistry in general.

Piperlongumine (PL) is an alkaloid that has attracted much attention as a selective anticancer agent despite its unclear pharmacology and mode of action. Thus, unveiling the underlying pharmacology can open new research avenues and leverage the development of PL as an efficacious cancer treatment. Here, we used SPiDER, as an alternative to
chemical proteomics, to identify potential biological targets for PL that could be pursued in downstream validation studies. We report on an unprecedented link between PL and the human transient receptor potential vanilloid 2 (hTRPV2) channel. To achieve this, we performed extensive experimental validation—including cryoelectron microscopy (cryo-EM)—to illuminate the PL−TRPV2 molecular recognition mechanism. Our studies reveal a new, transient binding site in TRPV2, whose recognition results in selective, potent allosteric antagonism by PL. Moreover, we found that the anticancer activity of PL is high in cell lines with high hTRPV2 expression, which is a marker of poor prognosis in glioblastoma multiforme (GBM). Most importantly, local (intracranial) and sustained delivery of PL in mice with GBM using a bespoke material produced a potent and selective antitumor effect and almost complete remission of the disease. Additionally, we observed a selective reduction of malignant cells in response to PL ex vivo in primary tumor samples derived from GBM patients. These results open up new vistas for successful GBM therapies.

**RESULTS AND DISCUSSION**

**Self-Organizing Maps Identify TRP Channels as Targets of Piperlongumine.** We used the SPIIDER method \(^{5,13}\) that employs self-organizing maps and topological descriptors to infer biological targets for small molecules. By correlating PL to the structure of reference ligands, targets were predicted (Table S1). We prioritized TRP channels for screening given the high prediction confidence and availability of assays for panel screening. Moreover, a privileged link
between several members of the TRP channel family and natural products had been established. We further motivated screening of PL against TRP channels by processing the SPiDER output with a data visualization algorithm. PL was projected onto a region of low TRP ligand density, which suggests that its pharmacophore has been scarcely employed to interrogate TRP-channel biology as opposed to molecules within other regions of chemical space (Figure 1a).

To validate the drug-target prediction, we tested PL in functional, cell-based fluorescence assays. PL showed antagonism of hTRPV2 (IC₅₀ = 4.6 μM ± 0.13 log units) and selectivity against a TRP channel panel (Figure 1b). Agonistic activity was not detected in any case. In calcium-imaging experiments with HEK293T cells transiently transfected with hTRPV2-RFP, we confirmed significant inhibition (p < 0.0001; one-way ANOVA and Dunnett post hoc test) of calcium influx evoked by cannabidiol—an hTRPV2 agonist (Figure 1c).
Similarly, PL inhibited calcium influx evoked by cannabidiol (IC50 = 1 μM ± 0.52 log units; Figure 1d, Figure S1) in a patch-clamp assay. The Michael acceptor motifs in PL have been previously implicated in covalent and irreversible modification of proteins.18 To confirm that PL reversibly inhibits hTRPV2, we performed wash-out assays. The functional hTRPV2 activity could be reinstated, which shows that binding is either noncovalent or that PL presents a fast off-rate after covalent modification of hTRPV2 (Figure 1e,f). This observation contrasts with a recent report on the covalent and stable modification of GSTO1 by PL.21 We next synthesized two PL analogues, as a racemic mixture, comprising a cyclopropane ring and without Michael acceptor motifs. The trans derivative displayed a perfect alignment with that of PL (Figure S2). No functional activity was found for both small molecules (Figure S3), suggesting a potential role of the conjugated motifs for molecular recognition, namely, for inducing a bioactive conformation. To further understand modulation of TRPV2 by PL, we probed whether PL and cannabidiol compete for the same binding pocket through a series of coincubation experiments. The maximal effect and apparent v_max value of cannabidiol decreased with increasing PL concentrations (Figure 1g,h). Both observations are consistent with noncompetitive (allosteric) antagonism.22 Neither autofluorescence nor SCAM behavior23 that could interfere with assay readouts was detected at relevant assay concentrations (Table S2).

**Molecular Mechanism of Piperlongumine Interaction with TRPV2.** We used cryo-EM to gain atomistic insight into the molecular mechanism of TRPV2 antagonism by PL. We reconstituted the full-length rat TRPV2 (rTRPV2, 77% identity and 90% similarity to hTRPV2) into nanodiscs and incubated with 50 μM PL (≈10 times the EC50) before preparing grids for cryo-EM. This yielded an rTRPV2−PL complex data set with a single state at 3.5 Å with C4 symmetry (Table S3 and Figures S4 and S5). This state was similar to the apo truncated rabbit TRPV2 in amphipols (PDB 5AN8) and the cannabidiol (CBD) state 2 rTRPV2 in nanodisc (PDB 6U88) (Figure 2a, Figures S4 and S5) structures determined previously.24−27 The reconstructed map has the highest quality in the transmembrane region, but no density is resolved for the pore turret (residues Ser564−Pro587).

Recent structures of rTRPV2 in nanodiscs show density for lipid tails in the vanilloid pocket;25 these densities are absent in the PL-bound cryo-EM map (Figure S6). Instead, we observed an elongated density starting at the vanilloid pocket and stretching along the top of the S4−S5 linker that we have attributed to PL. This density is wedged above the S4−S5 linker in a pocket formed by the S4 helix and the S5 and S6 helices of an adjacent TRPV2 monomer (Figure 2b, Figure S6). The trimethoxylphenyl moiety in PL is oriented toward the C-terminal end of the S4−S5 linker and the vanilloid pocket, whereas the dihydropyridinone group is oriented toward the N-terminal end (Figure 2c). PL is stabilized in this new allosteric pocket by hydrogen bridges between the two carbonyl groups of PL and the side-chains of Thr522 and Arg539 (Figure 2d). A density similar to that of PL is not
Figure 4. hTRPV2 expression is required by CNS cancer cell lines and associated with disease progression and poor prognosis in glioblastoma multiforme. (a) Cell lines with high hTRPV2 expression (as defined in Figure S12a) are sensitive to PL activity (−AUC). **** represents p < 0.0001 (Wilcoxon rank-sum test). (b) Cell lines from the Central Nervous System (CNS) lineage are the most sensitive to hTRPV2 loss. The effect of hTRPV2 loss-of-function was quantified with the CERES dependency score: a lower score represents a higher likelihood that the gene is essential in a given cell line. Left: smoothed histogram of distribution of hTRPV2 dependency scores across all cell lines. (c) hTRPV2 expression is a marker for poor prognosis in human gliomas. Overall survival Kaplan–Meier plots for patient stratification based on hTRPV2 median expression in low- and high-grade gliomas. p values for log-rank tests for differences in survival are shown. (d) hTRPV2 expression increases concomitantly with brain tumor stage. hTRPV2 expression across GTEx normal and TCGA tumor (grouped in LGG grade II, LGG grade III, and GBM) brain samples. Black points and lines represent the median ± upper/lower quartiles. ** represents p < 0.01 and **** p < 0.0001 (Wilcoxon rank sum test). (e) qRT-PCR of hTRPV2 in matched primary and recurrent GBM pairs. PGK1 was used as an internal control, and the Wilcoxon matched pairs signed rank test was used for statistics. (f) hTRPV2 is expressed in endothelial and microglial/macrophage brain tumor cells. hTRPV2 expression (log2 of counts per million, CPM) across 3589 single cells derived from four GBM human samples, grouped by cell type. Data are shown as mean ± standard error. (g) hTRPV2 is expressed in diffuse human gliomas, in glial (white arrowhead) and in endothelial cells (black arrowhead); and endothelial cells in GBM display an intense and diffuse staining of the cytoplasmic membrane in areas of microvascular proliferation, which contrasts with the weak or negative staining of the endothelial cells of grade II and III gliomas, and with the cytoplasmic and punctate staining pattern of glial cells.
present in any currently published TRPV2 X-ray or cryo-EM map.\textsuperscript{24–28} (Figure S6). rTRPV2 Arg539 (Arg537 in humans) is conserved as either Arg or Lys across the TRPV family in humans. Thus, while it may be involved in PL recognition, it is unlikely to account for TRPV2 selectivity. Moreover, Thr522 (Thr520 in humans) is highly conserved across TRPV2 in different species but is not conserved across the TRPV family in humans (Figure S7). While the Thr520/522 position is also a Thr in human TRPV4, it is a Met in TRPV3 and the otherwise highly homologous TRPV1 (Figure S7). Altogether, this may make Thr520/522 key for PL specificity. To challenge this hypothesis, validate the binding site, and test the contribution of these two residues to the effect of PL, we mutated Arg539 to Ala and Thr522 to Met in hTRPV2 and measured changes in calcium uptake. In calcium-imaging experiments using the wild type hTRPV2, we reconfirmed inhibition of calcium influx by PL ($p \leq 0.0001$; two-way ANOVA) when cannabidiol was evoked (Figure 2e,f, Figures S8–S10). The Arg539Ala mutation showed a loss of inhibition by PL ($p \leq 0.0001$; two-way ANOVA), consistent with its predicted role as a PL binding site. More interestingly, the Thr522Met mutation not only showed a loss of inhibition by PL ($p \leq 0.0001$; two-way ANOVA) but also overall increased calcium uptake both with and without the addition of PL. It is not possible to determine from these experiments whether this increase in calcium uptake was due to increased hTRPV2 expression and/or trafficking to the cell membrane or because the Thr520Met mutation increased channel activity in response to CBD. The Thr520Met mutation not only alters hydrogen bridging possibilities but also introduces a bulkier residue that could restrict access of PL to the binding site. Taken together, our data highlight the relevance of these two residues in the antagonism of TRPV2 by PL with changes in calcium influx.

Structurally, PL induces conformational changes in the TRPV2 selectivity filter and lower gate. Both remain closed when PL is bound; the conformation of the lower gate shifts slightly downward, and His651 rotates out of the pore relative to the recently published apo rat TRPV2 in nanodisc structure (PDB 6U84) (Figures 3a,b). The downward shift in the PL-bound structure seems to originate at the S4–S5 linker, which shifts toward S5 by 4.0 Å and downward by 2.9 Å, with the adjacent helices, including S6, shifting to compensate (Figure 3c). The shift of the S4–S5 linker toward S5 is reminiscent of the changes seen in the recent structure of rTRPV2 bound to CBD, state 2 (PDB 6U88) (Figure S11), though that structure did not have the additional downward shift of the S4–S5 linker seen in the PL-bound structure. On top of this, the PL-bound structure has the C-terminal domain wrapped around its $\beta$-sheet region paired with a downward pivot of the intracellular ankyrin repeat domains (Figure 3d), again as seen in the CBD state 2 structure (PDB 6U88) (Figure S11). Overall, the entire molecule shifts downward by 2.8 Å relative to the central pore, creating a transient atherosclerotic tunnel where PL sits (Figure 3d). The transmembrane domain of the PL-bound structure also aligns well to the apo truncated rabbit TRPV2 structure, which was predicted to be desensitized\textsuperscript{27} (Figure S11). Taken together, these changes suggest that PL may inhibit TRPV2 by immobilizing the S4–S5 linker and locking TRPV2 in a desensitized state.

hTRPV2 Expression Is Required by CNS Cancer Cell Lines and Associated with Disease Progression and Poor Prognosis in GBM. To probe the role of hTRPV2 on the antiproliferative effects of PL, we screened PL against the NCI-60 panel of cancer cell lines\textsuperscript{29} and analyzed the publicly available CTRP data.\textsuperscript{30} In both panels, cell lines expressing hTRPV2 were very sensitive to PL (Wilcoxon rank-sum test $p < 0.0001$ and 0.005, respectively, for the CTRP and NCI-60 panels; Figure 4a, Figures S12–S14), supporting the importance of hTRPV2 modulation as a mechanism that contributes to the anticancer activity by PL. To identify cancer types potentially more vulnerable to hTRPV2 targeting, we analyzed CRISPR-Cas9 hTRPV2 loss-of-function screens performed in 764 cancer cell lines within the Cancer Dependency Map (DepMap).\textsuperscript{31} Cell lines from the central nervous system (CNS) were the most sensitive to hTRPV2 loss (Figure 4b, Figure S15), suggesting CNS tumors as candidate disease models for hTRPV2 targeting.

To further investigate the importance of hTRPV2 in different cancer types, we tested its prognostic value across 9785 tumor samples belonging to 30 cohorts from The Cancer Genome Atlas (TCGA; Table S4). High hTRPV2 expression is strongly associated with poor survival in patients with gliomas ($p = 4.37 \times 10^{-15}$, log-rank test; Figure S16), also when stratified into the low-grade glioma (LGG)\textsuperscript{32} and GBM\textsuperscript{33} cohorts individually ($p = 0.028$ and 0.0087, respectively, log-rank test; Figure 4c). The prognostic value of hTRPV2 was independently validated in two other cohorts of gliomas and GBMs from the Chinese Glioma Genome Atlas (CGGA, http://www.cggga.cn/; Figure S17). We also found a significant progressive increase of hTRPV2 expression with tumor grade (Figure 4d) and from primary to matched recurrent GBM samples (Figure 4e). These results thus associate hTRPV2 overexpression with worse outcome in GBM.

We next analyzed the transcriptomes of human GBM single cells\textsuperscript{34} and found hTRPV2 expressed predominantly in endothelial and microglia/macrophage tumor cells but not in neoplastic tumor cells (Figure 4f, Figure S18) or normal brain samples\textsuperscript{35} (Figure S19). Concordantly, hTRPV2 expression is negatively associated with tumor purity, i.e., with the number of neoplastic cells vs non-neoplastic cells present in the LGG and GBM samples (Spearman’s correlation coefficient, $\rho = −0.45$ and −0.36, $p = 7.5 \times 10^{-25}$ and $3.1 \times 10^{-14}$, respectively; Figure S20), showing that hTRPV2 is mainly expressed in the tumor microenvironment. This was validated in selected samples from patients diagnosed with grade II–IV glioma, where immunostaining for hTRPV2 showed expression in glial and endothelial cells (Figure 4g, Figure S21). Most significantly, in areas of microvascular proliferation, a characteristic pathologic feature of GBM, endothelial cells showed a significant overexpression of hTRPV2 (Figure S22), and a distinct staining distribution/pattern (Figure 4g). In glial cells, staining was cytoplasmic and with a punctate pattern, whereas endothelial cells showed an intense and diffuse staining of the cytoplasmatic membrane (Figure 4g). Higher-magnification imaging depicts a clear punctate staining in glial cells and diffuse membranous staining in the endothelial cells lining tumor vessels (Figure S22). This links with the fact that high hTRPV2 expression plays a role in angiogenesis\textsuperscript{26} and tumor progression and associates with poor prognosis (Figure 4c–e). When we tested the prognostic value of hTRPV2 independent of the endothelial and microglia/macrophage content, we confirmed that high hTRPV2 expression was still significantly associated with prognosis in GBM (Cox proportional-hazards regression $p < 0.05$) but not in low-grade glioma.
Altogether, our data and others' support GBM as a candidate disease for treatment with hTRPV2 ligands, including PL.

**hTRPV2 Knockdown in Glioblastoma Cells Reduces Sensitivity to PL and ROS Production.** We first tested the effect of PL in the viability of different GBM cell lines (U251, U87, YKG1, and D8TRG/06MG) which express significant amounts of hTRPV2 (Figure S23 and Table S5). These cell lines were sensitive to PL (with IC_{50} values of 3.0, 6.6, 4.7, and 6.6 μM, respectively) after 72 h. Endothelial brain cells, which also express hTRPV2, were sensitive to PL (IC_{50} = 1.6 μM), while both microglia and astrocytes were less sensitive (IC_{50} > 10 μM) (Figure S23 and Table S5). Despite the high hTRPV2 expression found in the tumor microenvironment, the high sensitivity to PL of cell lines expressing hTRPV2 and high hTRPV2 dependency of CNS cell lines support the use of GBM cell line models to study the relationship between hTRPV2 and PL treatment. Our data show that cells expressing hTRPV2 are sensitive to PL.
To investigate the relationship between hTRPV2 and PL treatment, we knocked down the expression of hTRPV2 (by 82 ± 12%, see Figure S24) in the U251 glioblastoma cells (TRPV2KD) using the CRISPRi technology with dCas9-KRAB-guide RNA plasmids.38 We found that the PL-induced effect on viability is diminished when hTRPV2 expression is reduced as shown in Figure S25a−c. TRPV2KD cells are 2-fold less sensitive to the antiproliferative effects of PL (p = 0.0002, unpaired t-test, Figure S25a). We then performed RNA-sequencing analysis of U251 cells (TRPV2WT or TRPV2KD).

Figure 6. PL-loaded hydrogel scaffolds treat glioblastoma multiforme in vivo. (a) Experiment design showing the development of a biocompatible, PL-loaded hydrogel suitable for local treatment of GBM. Experiment flow: U251 or U87-MG cells were injected intracranially (n = 5 per group). 8 days later, engraftment was confirmed with bioluminescent imaging. On day 9, PL-loaded or unloaded hydrogels were implanted. Tumor growth was assessed at regular intervals by using bioluminescent imaging. (b) Histological image of a full mouse brain showing the hydrogel implant. (c) Bioluminescent images of U251 and U87-MG xenograft mice treated with PL-loaded hydrogel (50 mg/kg) or unloaded hydrogel (control). Data show that PL treatment successfully reduces tumor volume over a period of 21 days relative to the control, in which the tumor grows exponentially, on both models of GBM. Tumor burden in mice treated with PL-loaded or unloaded hydrogel as measured by luciferase activity for (f) U251 and (g) U87-MG models. Data are expressed as luciferase intensity (ROI on tumor site) as an indication of average tumor load in the control group at each time point and represent group mean ± SEM. Data show a significant tumor burden difference between both groups (p ≤ 0.01, n = 5, Mann−Whitney test) for (f) U251 and (g) U87-MG models. The Kaplan−Meier survival curve shows that mice treated with PL-loaded hydrogels have significantly increased survival (p ≤ 0.001, log-rank Mantel-Cox test) relative to the control group for (h) U251 and (i) U87-MG models. The experiment cutoff criteria included an increase in tumor size of around 500% relative to the original size measured for each mouse prior to hydrogel implantation.
treated with PL for 48 h. Principal component analysis confirmed a stronger effect of PL-treatment in TRPV2WT cells, with little separation between control and treatment conditions in TRPV2KD cells (Figure S25d) confirming successful knockdown of hTRPV2. Indeed, comparison of the fold changes (PL-treated vs DMSO control) between TRPV2WT and TRPV2KD cells identified only 100 genes that were significantly (p < 0.05) differentially expressed in the knockdown as opposed to 7302 genes in the case of TRPV2WT cells (Figure 5a). This was further confirmed when comparing the fold changes introduced by PL in TRPV2WT vs TRPV2KD cells, which yielded high correlation to PL-treated TRPV2WT cells only (Figure 5b). This shows that hTRPV2 knockdown in glioblastoma cells reduces the sensitivity to PL and its induced transcriptic changes.

Inspection of significantly differentially expressed genes upon PL treatment (p < 0.05 and absolute fold-change >1.5) identified several candidates linked to the electron transport chain as well as in trafficking of ATP (including NDUFAB1, NDUFAF7, KDM1A, and SLC25A5, Figure 5c). Indeed, gene-set enrichment analysis (GSEA) highlighted the down-regulation of oxidative phosphorylation as well as that of several mitochondrial signatures, including that of the mitochondrial envelope (Figure 5d, Figure S25e,f). To validate this observation, we asked if the mitochondrial membrane potential was compromised following treatment with PL. Using the cationic dye JC-1 that changes its fluorescent signal according with the energy state of the mitochondria, we interrogated mitochondrial depolarization and found a decrease in TRPV2WT U251 cells treated with PL, which was not observed in TRPV2KD U251 cells (Figure 5e). In agreement with this observation, ATP content was reduced following PL treatment of U251 cells compared to a negative control (Figure 5f).

PL is known to cause an increase in ROS, which ultimately leads to cell death; however, the mechanism by which this occurs has not previously been associated with TRPV2.38 We found that U251-TRPV2-KD cells treated with PL accumulate significantly less ROS relative to the control cells with normal hTRPV2 expression levels (Figure 5g). This was further supported by rescue studies using N-acetyl-L-cysteine (NAC). NAC is commonly used to identify and test ROS inducers and to inhibit ROS. Indeed, NAC was able to rescue U251 and U87MG cell viability in the presence of PL (Figure S26). This shows that NAC interfered with the activity of PL, further supporting that PL affects mitochondrial dysfunction. This implies that hTRPV2 plays a key role in the formation of ROS in cells after PL treatment.

Taken together, these observations suggest that PL treatment results in a TRPV2-dependent decrease of mitochondrial membrane potential, which ultimately leads to the loss of cellular ATP content. A recent study demonstrated that the increase of cellular ROS by defective mitochondria links to loss of translation in agreement with our transcriptomic analysis, which identified strong downregulation of translational signatures (Figure 5d).

Treatment of GBM via Sustained and Local Implanted PL-Scaffolds. Having obtained promising in vitro functional assay data and established a mode of action in a preclinical scenario, we next explored the therapeutic potential of PL in a murine orthotopic xenograft model of GBM (Figure 6a,b). In our experience, PL was poorly soluble in biocompatible formulations, and subcutaneous injections of 50 mg/kg resulted in toxicity with limited effects on intracranial tumor growth (Figure S27). After considering these results, we sought out a more effective mode for in vivo drug delivery. Thus, we encapsulated PL in β-cyclodextrin (1:2 ratio), which dramatically improved its aqueous solubility. Furthermore, to enhance therapeutic efficacy, we aspired to eliminate the challenge of crossing the blood-brain barrier and create a therapy suitable for local and sustained drug administration. To this end, we devised implantable dextran-dendrimer hydrogel scaffolds that were doped with PL encapsulated in β-cyclodextrin. The cyclodextrin was used as a ‘carrier’ molecule to facilitate the dissolution of PL. β-Cyclodextrin is a cyclic oligosaccharide, which consists of a macrocyclic ring of 7 glucose subunits with a hydrophobic interior and hydrophilic exterior that forms a complex with the hydrophobic PL compound. This new material allowed us to obtain a high concentration of PL at the tumor site. Moreover, we minimized leakage of PL to healthy tissue by decorating the hydrogel with aldehyde groups from oxidized dextran to interact with cancer tissue amines and form adhesive bonds, as previously reported.41–44 Profiling of our drug delivery system showed a significant discharge of PL in the first 4 h followed by steady release (3% of total) for at least 192 h under physiological conditions (Figure S28). These data provide a rationale to the in vivo toxicity experiments in which PL-doped hydrogels were able to induce near complete cell death after incubation with U251 cells for 24 h. The empty hydrogels did not affect cell viability (Figure S29).

We proceeded to study the in vivo therapeutic efficacy of the PL-doped hydrogels in two murine models of GBM (U251 and U87-MG). The U251 malignant glioma cell line is known to mimic the salient features of human GBM and displays similarities at the genetic level.45,46 The U87 GBM model is commonly used for assessing tumor angiogenesis and antiangiogenic therapies in GBM.46,47 Tumors were induced through direct injection of 2.5 × 105 U251-GFP-luc or 2.5 × 104 U87-MG-luc cells into the brain of athymic nude mice (Crl:NU(Ncr)-Foxn1nu) via a burr hole 2.5 mm lateral and 1.5 mm posterior to bregma, above the right cerebrum. Nine days post-tumor-injection, hydrogel scaffolds loaded with PL (50 mg/kg) were implanted in the supratentorial region directly overlaying the area where the tumor was implanted in the right hemisphere of the brain (Figure 6a,b). Empty hydrogel implants, i.e., lacking the PL component, were used as controls. Tumor volume was measured over time by bioluminescence with Xenogen IVIS Lumina for 21 days after hydrogel implantation (Figure 6c). The difference in survival between the two groups was highly significant (p < 0.001, log-rank Mantel–Cox test) for both U251 (Figure 6d) and U87 (Figure 6e) models. The mice exposed to PL-doped hydrogel showed significant tumor size reduction at the end of the study (n = 5, p < 0.01 Mann–Whitney test) for both U251 (Figure 6f) and U87 (Figure 6g) models. To further demonstrate the superiority of the local PL-hydrogel scaffold, the efficacy of the local delivery was compared with systemic (subcutaneous) injection of PL alone, both at a final concentration of 50 mg/kg PL (Figure S27). We found that although local administration of PL by means of a hydrogel scaffold resulted in almost complete remission of the GBM tumor, no effect on tumor reduction was attained following systemic administration of the same concentration of PL. These results provide substantial evidence that the hydrogel scaffold is essential for higher therapeutic performance. In
addition, no apparent signs of disease (not related to local tumor growth) were seen in any of the animals for 21 days after hydrogel implantation as indicated by the preservation of steady body weight (Figure S30), pointing at both drug and hydrogel biocompatibility. Histopathology of the liver, kidney, lung, spleen, gastrointestinal tract, and heart showed no significant changes or evidence of toxicity of PL. No evidence of necrosis or other lesions (e.g., inflammation) was seen in the brain, at the site of implantation (Figure S31).

Monitoring of tumor growth in the brain of PL-treated mice showed delayed tumor progression (Figure 6d–g). Leptomeningeal dissemination was seen both through in vivo imaging, as a second primary and more intense bioluminescence signal, dorsal and caudal to the primary tumor and overlaying the posterior part of the brain and spinal cord (Figure 6c), and also microscopically as tumor cells infiltrating and expanding the leptomeninges. Meningeal bioluminescence was seen in 60% (for U251 model) and 80% (for U87 model) of the mice in the control group. Meningeal bioluminescence was seen both through imaging, in vivo (TMZ), or DMSO as a negative control for 48 h. Then, to identify potentially malignant cells, dissociated cells were fluorescently labeled with the astrocyte marker S100B (Figure S31). For the U251 model used to identify the best treatment option for patients, ultimately leading to better outcomes such as increased progression-free-survival.48 Here, we tested PL ex vivo drug responses in five surgically derived GBM patient samples at three different drug concentrations. Freshly dissociated cells that were initially cryopreserved before the assay were incubated with PL, standard-of-care temozolomide (TMZ), or DMSO as a negative control for 48 h. Then, to identify potentially malignant cells, dissociated cells were immunofluorescently labeled with the astrocyte marker S100B as well as RESTIN (Figure 7), a marker for proliferating neural progenitors’ cells, while CD45 (Figure S34) was included as a counterstain to identify nonmalignant immune cells. In four out of five tested patient samples (80%), PL induced a significant decrease in the fraction of glioblastoma cells (defined inclusively as either S100B or RESTIN positive cells) at the highest tested concentration (50 μM) compared to the negative control. Among these four responders, two patient samples also responded to PL at lower concentrations. The observed effect was “on-target”—preferentially affecting the putatively malignant cells and not the immune populations, which was also reflected in our relative fraction readout. Furthermore, when compared to TMZ, which is usually used at higher in vitro concentrations than the ones tested, PL

**Figure 7.** Ex vivo drug response of GBM cells derived from five patients. In this experiment, the ZH prefix is followed by the unique patient ID (n = 5). Colored lines represent the tested drugs; piperlongumine (PL, red) and temozolomide (TMZ, gray), where the x-axis corresponds to the concentrations tested (2, 20, 50 μM). The y-axis denotes fraction reduction of S100B+CD45− or RESTIN+CD45− cells identified by immunofluorescence relative to the DMSO control (50 μM). Values plotted as mean ± SEM. Two-way ANOVA followed by Dunnett’s multiple comparisons test was performed to calculate statistical significance (n = 4–5 replicate wells per treatment condition, n = 15 wells for DMSO control). * p_adj < 0.05, ** p_adj < 0.01, *** p_adj < 0.001, **** p_adj < 0.0001.

**PL Induces Selective ex Vivo Response in GBM Patient-Derived Cells.** Next, we evaluated the clinical potential of PL using Pharmacoscopy, a technology that utilizes high content automated microscopy to measure ex vivo drug response directly in patient-derived cells.48,49 The relevance of Pharmacoscopy has been previously demonstrated in a clinical trial for aggressive hematological malignancies, where it was used to identify the best treatment option for patients, ultimately leading to better outcomes such as increased progression-free-survival.48 Here, we tested PL ex vivo drug responses in five surgically derived GBM patient samples at three different drug concentrations. Freshly dissociated cells that were initially cryopreserved before the assay were incubated with PL, standard-of-care temozolomide (TMZ), or DMSO as a negative control for 48 h. Then, to identify potentially malignant cells, dissociated cells were immunofluorescently labeled with the astrocyte marker S100B as well as RESTIN (Figure 7), a marker for proliferating neural progenitors’ cells, while CD45 (Figure S34) was included as a counterstain to identify nonmalignant immune cells. In four out of five tested patient samples (80%), PL induced a significant decrease in the fraction of glioblastoma cells (defined inclusively as either S100B or RESTIN positive cells) at the highest tested concentration (50 μM) compared to the negative control. Among these four responders, two patient samples also responded to PL at lower concentrations. The observed effect was “on-target”—preferentially affecting the putatively malignant cells and not the immune populations, which was also reflected in our relative fraction readout. Furthermore, when compared to TMZ, which is usually used at higher in vitro concentrations than the ones tested, PL
exerted a stronger effect within this range of low doses (Figure 7, Figure S34). Taken together, we provide promising evidence of PL-induced selective drug response in tumor samples derived from GBM patients. As our tested cohort is relatively small, we hope future studies will further investigate the clinical potential that PL might hold as a putative GBM therapy.

**CONCLUSIONS**

Statistical learning may be used to accelerate drug discovery programs by prioritizing biochemical screens; such methods ought to be generally applicable to molecules of synthetic and natural origins. Under such a concept, our computer-driven and disease agnostic approach constitutes also a viable alternative to swiftly repurpose chemical matter and motivate the development of innovative therapeutics. Here, by dissecting the chemical features of PL, a hitherto unknown relationship to hTRPV2 emerged. The mechanistic insights for hTRPV2 have been rare, and its (patho)biological relevance is largely unknown.30,51 To further our understanding of the hTRPV2 mechanism of modulation by exogenous compounds, we determined the PL-bound full-length rat TRPV2 structure at an overall resolution of 3.5 Å and identified a transient, allosteric binding site for PL along the S4−S5 linker of the channel. A high-quality open state of TRPV2 has not yet been resolved, so it is difficult to predict the exact mechanism of PL inhibition of channel opening. However, the S4−S5 linker plays a key role in channel opening across the entire TRP channel family,52 which suggests that small molecule ligands for this region modulate function. Calcium imaging data of hTRPV2 mutants with the addition of PL further established the importance of the Arg539 and Thr522 residues in TRPV2 for this region modulate function. Calcium imaging data of hTRPV2 channel’s calcium influx modulation and highlighted the role of Arg539 against PL antagonism for TRPV2, consistent with the cryo-EM structure.

Importantly, we establish that modulation of hTRPV2 by PL affords a biologically relevant phenotype, in particular against GBM in which hTRPV2 overexpression correlates with tumor grade, and that the anticancer activity of PL is high in cell lines with high hTRPV2 gene expression, which is required by CNS cancer cell lines. Moreover, the CRISPRi-induced down-regulation of hTRPV2 showed that GBM cells treated with PL accumulate significantly less ROS relative to the control cells and that knockdown of hTRPV2 also decreases sensitivity to PL, which suggests the importance of TRPV2 for the anticancer activity of PL. Indeed, with the presence of damaged mitochondria (decreased mitochondrial membrane potential and ATP content) in hTRPV2WT treated with PL, increased ROS fold is expected in these cells. These increased ROS levels impinge upon nuclear transcription of mitochondrial targeted proteins, as observed in the RNaseq analysis. Finally, we also showed that hTRPV2 is a marker of poor prognosis for GBM patients, and immunohistochemistry of GBM tissues samples showed a strong correlation between hTRPV2 overexpression and aggressiveness of the disease.

GBM remains an unmet medical need, with newly diagnosed patients having a life expectancy of only 11−15 months (which has not improved in the last 30 years), and new therapeutics are urgently sought after. To that end, formulation of PL in a hydrogel for sustained intracranial release afforded an efficacious GBM treatment in mice using two orthotopic GBM models to harness an innovative mechanism of action. This treatment resulted in a highly significant decrease in disease progression and increase in survival, which greatly contributes to the avoidance of relapses of GBM. Furthermore, using Pharmacoscopy, we have demonstrated that PL shows promising on-target ex vivo responses in surgically derived GBM patient samples, further strengthening the clinical potential and applicability of our findings. Altogether, supported by functional and structural data, we showed the power of computer perception and decision making from complex data patterns and expect it to play an increasing role in future drug discovery and in the success of GBM therapy.

**ASSOCIATED CONTENT**

*Supporting Information*

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acscentsci.1c00070.

- Materials and methods; detailed characterization of small molecules; and additional data and figures (PDF)
- NMR, HPLC, and mass spectrometry data for the cis analogue (PDF)
- NMR, HPLC, and mass spectrometry data for the trans analogue (PDF)
- Model validation report (PDF)
- GBM patient clinical annotations (XLSX)

**Accession Codes**

The cryo-EM density map and atomic coordinates of PL-bound rat TRPV2 are deposited into the Electron Microscopy Data Bank and Protein Data Bank under accession codes EMD-21705 and PDB 6WKN.

**AUTHOR INFORMATION**

*Corresponding Authors*

Vera Y. Moiseenkova-Bell — Department of Systems Pharmacology and Translational Therapeutics, Perelman School of Medicine, University of Pennsylvania, Philadelphia, Pennsylvania 19104, United States; Email: vbmb@pennmedicine.upenn.edu

Gonçalo J. L. Bernardes — Instituto de Medicina Molecular João Lobo Antunes, Faculdade de Medicina, Universidade de Lisboa, 1649-028 Lisboa, Portugal; Yusuf Hamied Department of Chemistry, University of Cambridge, CB2 1EW Cambridge, United Kingdom; orcid.org/0000-0001-6594-8917; Email: gb453@cam.ac.uk, gbernardes@medicina.ulisboa.pt

*Authors*

João Conde — Instituto de Medicina Molecular João Lobo Antunes, Faculdade de Medicina, Universidade de Lisboa, 1649-028 Lisboa, Portugal; orcid.org/0000-0001-8422-6792

Ruth A. Pumroy — Department of Systems Pharmacology and Translational Therapeutics, Perelman School of Medicine, University of Pennsylvania, Philadelphia, Pennsylvania 19104, United States

Charlotte Baker — Instituto de Medicina Molecular João Lobo Antunes, Faculdade de Medicina, Universidade de Lisboa, 1649-028 Lisboa, Portugal

Tiago Rodrigues — Instituto de Medicina Molecular João Lobo Antunes, Faculdade de Medicina, Universidade de Lisboa, 1649-028 Lisboa, Portugal

Ana Guerreiro — Instituto de Medicina Molecular João Lobo Antunes, Faculdade de Medicina, Universidade de Lisboa, 1649-028 Lisboa, Portugal

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Author Contributions

△J.C., R.A.P., C.B., and T.R. contributed equally. J.C. designed the hydrogel scaffold and in vivo delivery strategy, and C.B. and J.C. performed in vivo experiments. T.R. designed and performed cheminformatics studies and analyzed data. C.B., A.G., B.B.S., M.C.M., F.S., E.P.L., A.S., and M.C.M. conducted biochemical and cell biology studies, which included Ca^{2+} flux assay measurements with the assistance of S.H.V., protein purification, sample preparation, and data analysis and interpretation. R.A.P., A.S., and V.Y.M.-B. conducted cryo-EM studies and analyzed and interpreted data with the assistance of Y.L. and S.H. B.P.d.A. and N.L.B.-M. designed and performed bioinformatics analyses of TCGA, CTRP, NCI-60, DepMap, and single-cell transcriptomic data. M.L. acquired mass spectrometry data. M.L. and D.P. interpreted and analyzed mass spectrometry, and M.R. supervised the work performed by M.L. and D.P. F.C. performed preliminary molecular dynamic simulations and DFT minimizations. P.F. interpreted and analyzed RNA sequencing data. T.C. performed the histopathological analysis of mouse xenographs and TRPV2 immunohistochemistry scoring. R.R. performed histopathological analysis of human tumors and TRPV2 immunohistochemistry scoring. S.L. and B.S. performed Pharmacoscopy experiments, and human tumor samples used were surgically removed and analyzed by T.W. and M.W. J.C., R.A.P., C.B., T.R., and G.J.L.B. are coinventors on a patent application (WO2019/054891 A2) that incorporates discoveries described in this manuscript. R.R., J.C., and G.J.L.B. conceived the research. V.Y.M.-B. and G.J.L.B. wrote the manuscript with contributions from the remaining authors. G.J.L.B. conceived the research. V.Y.M.-B. and G.J.L.B. supervised the research. All authors agreed on the final version of the manuscript.

Notes

The authors declare the following competing financial interest(s): J.C., C.B., T.R., and G.J.L.B. are coinventors on a patent application (WO2019/054891 A2) that incorporates discoveries described in this manuscript. T.R., J.C., and G.J.L.B. are cofounders and shareholders of Targ.Tex S.A., a company that develops TRPV2 antagonists for the treatment of cancer.
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