Gli2 mediates the development of castration-resistant prostate cancer

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Abstract. Glioma-associated oncogene family zinc finger 2 (Gli2), a key component of the hedgehog signaling pathway, has been previously demonstrated to promote the malignant properties of prostate cancer in vitro. However, the role of Gli2 in the development of castration-resistant prostate cancer (CRPC) has yet to be fully elucidated. In the present study, Gli2 expression was knocked down in androgen-responsive prostate cancer cells using an inducible Gli2 short hairpin RNA. Suppression of Gli2 expression resulted in significant reduction of cell viability, increased the proportion of cells in the G1 phase of the cell cycle and reduced the expression of genes associated with cell cycle progression. Gli2 knockdown sensitized both androgen-dependent and -independent prostate cancer cells to the antiandrogen drug Casodex and prevented the outgrowth of LNCaP prostate cancer cells. In addition, Gli2 knockdown significantly suppressed the development of CRPC in a LNCaP xenograft mouse model, which was reversed by the re-expression of Gli2. In conclusion, to the best of our knowledge, the present study was the first occasion in which the essential role of Gli2 expression resulted in significant reduction of cell viability, increased the proportion of cells in the G1 phase of the cell cycle and reduced the expression of genes associated with cell cycle progression. Gli2 knockdown sensitized both androgen-dependent and -independent prostate cancer cells to the antiandrogen drug Casodex and prevented the outgrowth of LNCaP prostate cancer cells. In addition, Gli2 knockdown significantly suppressed the development of CRPC in a LNCaP xenograft mouse model, which was reversed by the re-expression of Gli2. In conclusion, to the best of our knowledge, the present study was the first occasion in which the essential role of Gli2 in the development of CRPC was demonstrated, providing a potential therapeutic target for the intervention of CRPC.

Introduction

Prostate cancer is one of the most frequently diagnosed malignancies in men. According to a previous report, prostate cancer is the third most common cause of death associated with cancer in men in the United States (1). Hormone therapy is frequently applied to treat advanced metastatic prostate cancer (2,3). Although androgen deprivation therapies are effective on the short term, relapse regularly occur shortly following clinical remission due to highly aggressive tumors that are resistant to hormone therapy; this is referred to as castration-resistant prostate cancer (CRPC). The development of CRPC is a complex process that involves reactivation of androgen receptor (AR) signaling (4). Therefore, to improve overall survival, novel strategies in inhibiting the emergence of this phenotype of prostate cancer are urgently needed.

The hedgehog signaling pathway is pivotal to the growth, patterning and morphogenesis of a number of tissues during embryogenesis (5-7). Canonical hedgehog signaling in vertebrates is initiated by the engagement of ligands, including Sonic (Shh), Indian (Ihh) or Desert (Dhh) hedgehog to the Patched (Ptc) receptor on corresponding target cells (8,9), resulting in the dissociation of smoothened (Smo) from the membrane and its activation. The accumulation of active Smo occurs in the primary cilium and induces the processing, re-localization and finally the activation of the glioma-associated oncogene family zinc finger 2 transcription factor (Gli) (10,11).

Although accumulating evidence supports the notion that the hedgehog pathway is involved in the tumorigenesis of a variety of malignancies, including skin, breast, lung, brain, pancreatic, liver, stomach and prostate cancer (12-28), the role of this signaling pathway in prostate cancer remains controversial. Previous evidence supporting this association includes reported findings that prostate cancer cell growth was significantly inhibited by the Smo inhibitor cyclopamine or Gli knockdown (25,28,29). In addition, the expression of components of the hedgehog signaling pathway such as Gli was also found to be is enhanced in the malignant prostate epithelium compared with their benign counterparts (26,29,30). However, by contrast, there remains a lack of evidence for elevated canonical hedgehog signaling activity in commonly used prostate cancer cell lines (31,32).
Despite its controversial role in the tumorigenesis of prostate cancer, evidence suggests that the canonical autocrine-like hedgehog signaling is reactivated in prostate cancer cells following androgen deprivation (26,30,33,34). Additionally, Gli expression and activity can also be regulated by noncanonical signaling pathways, including Ras- or TGF-β-mediated signaling (35). In the present study, investigative focus was placed on the role of Gli2, an intracellular mediator of the hedgehog pathway, on CRPC physiology. To the best of our knowledge, the present study was the first in which Gli2 expression was demonstrated to be necessary for the transition of prostate cancer cells from the androgen-dependent to androgen-independent phenotype in vitro and in vivo, suggesting that Gli2 serves as a driver in the progression of prostate cancer to the more severe CRPC.

Materials and methods

Cell culture. The prostate cancer cell lines LNCaP, 22RV1 and VCaP were originally obtained from American Type Culture Collection (ATCC). LNCaP and 22RV1 were cultured in McCoy's 5A medium supplemented with pyruvate, amino acids, nutrients and 10% FBS (Gibco; Thermo Fisher Scientific, Inc.) as previously described (34). VCaP cells was grown in RPMI-1640 medium (Invitrogen; Thermo Fisher Scientific, Inc.) supplemented with 10% FBS and 0.1% penicillin. Cultures were maintained in 5% CO₂ at 37°C. These cell lines were authenticated using the DNA markers of short tandem repeats used by ATCC.

Construction of the Tet-inducible Gli2-specific short hairpin RNA (shRNA) expression plasmid. The Gli2 shRNA was cloned into the Tet-pLKO-puro vector for the inducible expression of Gli2 shRNA by doxycycline (DOX). The Tet-pLKO-puro vector was a gift from Dmitri Wiederschain (Addgene plasmid no. 21915; http://n2t.net/addgene:21915; RRID:Addgene_21915; Addgene, Inc.) (36). In the absence of doxycycline (DOX-), the Gli2 shRNA expression is repressed by the constitutively expressed TetR protein (36). Upon addition of doxycycline (DOX+) to the culture medium, this repression is removed and Gli2 shRNA expression is triggered, resulting in Gli2 knockdown.

A number of distinct Gli2-targeting shRNA constructs were designed and cloned into lentiviral vectors, which were then used to transfect LNCaP cells. The efficiency of Gli2 shRNA transfection was determined by measuring Gli2 mRNA levels using reverse transcription-quantitative PCR (RT-qPCR) in transfected LNCaP cells. The most efficient Gli2 shRNA-coding sequence in knocking down Gli2 was chosen and its sequence was as follows: 5'-CCTGGCATGACTACCACTATGCTCGAGCATAGTGGTAGT-3' and reverse, 5'-AAATTAAAAACC TGGGCATGACTACCACTATGCTCGAGCATAGTGGTAGT-3'. These two complementary oligos were annealed and ligated into the Tet-pLKO-puro vector between the AgeI and EcoRI restriction sites. This recombinant plasmid was named pTet-Gli2shR thereafter. To construct the control vector, the Tet-pLKO-puro vector was first digested with AgeI and EcoRI I to release the stuffer segment, following which the 5' overhang ends were blunted using DNA Polymerase I, Large (Klenow) Fragment (New England BioLabs, Inc.) where the two blunted ends were ligated using T4 ligase (New England Biolabs, Inc.). This vector was named pTet-ctl thereafter.

Construction of the Gli2ΔN expression plasmid. The cDNA sequence for Gli2, obtained from the plasmid pcS2-MT GLI2 FL plasmid, which was a gift from Erich Roessler (Addgene plasmid no. 17648; http://n2t.net/addgene:17648; RRID:Addgene_17648) (37), was used for generating Gli2ΔN cDNA, an isoform of Gli2 lacking the 32 amino acid residues in the N-terminal repressor domain. Gli2ΔN cDNA was subsequently cloned into the pSDM101 vector as described previously (38). Briefly, Gli2ΔN expression plasmid was produced by cloning two fragments into the pSDM101 vector using the In-Fusion HD Cloning kit according to manufacturer's protocol (cat. no. 011614; Clontech Laboratories, Inc.). The homemade pSDM101 plasmid is a lentiviral vector pWPI backbone, which was a gift from Didier Trono (Addgene plasmid no. 12254; http://n2t.net/addgene:12254; RRID:Addgene_12254), with a linker inserted at the Pmel1 site to introduce BamHI, Mlu1 and NdeI restriction sites. The pSDM101 backbone was digested using the Pmel1 and BstBI restriction enzymes, where the larger fragment was purified using the EZ DNA Gel Extraction kit (Enzymax, LLC) for use as the vector. The first fragment for the In-Fusion HD Cloning kit, which was part of Gli2ΔN cDNA sequence at the 5'-terminal end, was amplified by PCR using the pcS2-MT GLI2 FL plasmid as template and CloneAmp™ HiFi PCR Premix according to manufacturer's protocols (cat. no. 639298; Clontech Laboratories, Inc.). The second fragment, which forms the remaining part of Gli2ΔN cDNA sequence and part of the pSDM101 backbone sequence, was amplified using the pSDM101-Gli2 plasmid as template, which was constructed by cleaving pcS2-MT GLI2 FL using EcoRI and XbaI restriction enzymes to form the Gli2 cDNA sequence, which was then cloned into the pSDM101 vector using the Pmel1 site. Primers used for PCR amplification was as follows: Fragment 1 forward, 5'-CTAGGCCCTCGAGTATTTTTAAACATGGCGCCTCA CCTCCATCA-3' and reverse, 5'-ATAGTATGTTGTCATGCC AGGACACTGGCTGCC-3'; Fragment 2 forward, 5'-CAT GCCACTACTATGAGCCCCCATGCCTGCTA-3' and reverse, 5'-TTGATATCGAATTCGAAGCTTGAGCTCGA-3'. The thermocycling conditions for both PCR reactions were as follows: 35 cycles of 98°C for 10 sec, 55°C for 5 sec and 72°C for 15 sec. All restriction enzymes mentioned above were purchased from New England Biolabs, Inc.

Stable transfection of plasmids and selection. In total, 2.5 µg pTet-Gli2shR or pTet-ctl plasmids were co-transfected with two packaging vectors, 1.9 µg psPAX2 (Addgene, Inc.) and 0.75 µg pMD2.G (Addgene, Inc.) into 1.5x10⁶ 293T cells using Lipofectamine® 2000 (Thermo Fisher Scientific, Inc.) according to manufacturer's protocol, to produce lentiviral particles. The lentiviral particle-containing medium were collected 48 h after transfection and subsequently added to the culture of the prostate cancer cells at a ratio of 1 ml viral particle-containing medium supplemented with 8 µg/ml
polybrene to 2 ml DMEM-based culture medium without antibiotics to create prostate cancer cell lines expressing the inducible Gli2 shRNA (LNCaP Gli2shR and 22RV1 Gli2shR) and corresponding controls (LNCaP Ctrl and 22RV1 Ctrl). The transduced cells were selected using 1 µg/ml puromycin following infection with lentiviral particles. pSDM101 vectors encoding either Gli2AN or control (CV) were similarly introduced into LNCaP Gli2shR cells by lentiviral transduction to generate cell lines of LNCaP Gli2shR/Gli2AN and LNCaP Gli2shR/CV genotypes, which were selected for using GFP fluorescence.

The lentiviral vector pLKO.1-Gli2 shRNA (Gli2 shR1) and corresponding control vector pLKO.1 were purchased from Sigma-Aldrich; Merck KGaA. These two vectors (2.5 µg) were subsequently used to produce lentiviral particles using 293T lentiviral packaging cells in protocols aforementioned and filtered prior to prostate cancer cell transduction. The sequence of Gli2 shR1 used is as follows: 5'-CCGGCTGGACAGGGA TGACTGTAAGCTCGAGCTTACAGTCATCCCTGTCCAG TTGGTT-3'.

**MMT cell viability assays.** LNCaP, 22RV-1 and VCaP cells were seeded in triplicate at a density of 5,000 cells/well into 96-well tissue culture plates, where they were allowed to adhere for 24 h and treated with/without 1 µM DOX and/or 30 µM Casodex® (Sigma-Aldrich; Merck KGaA) at 37˚C for various indicated time periods. Relative cell viability was quantified by MTT analysis using a protocol described previously (34).

**Whole transcriptome sequencing.** Prostate cancer cell lines LNCaP Gli2shR and LNCaP Ctrl were treated with either DOX or vehicle to generate four conditions with two replicates for each type: i) Ctrl DOX; ii) Ctrl DOX+; iii) Gli2sh R DOX; and iv) Gli2sh R DOX+. Total RNA was isolated from the cells using the RNeasy Mini kit according to manufacturer’s protocol (Qiagen GmbH), following which sequencing libraries were constructed using Truseq RNA Library Prep Kit 2 set B according to manufacturer’s protocol (cat. no. RS-122-2002; Illumina, Inc.) and sequenced in a HiSeq® 2000 system (Illumina, Inc.) using a single-read 50-bp sequencing protocol. The reads were aligned to the reference human genome (hg19 assembly; University of California Santa Cruz) using TopHat, a spliced read mapper for RNA sequencing (version 2.0.8b) (39). Only ±2 mismatches were permitted during in the alignment and HTSeq version 0.6.0-count was used to count the gene expression reads (40). R/DESeq (version 1.26.0) (41), DESeq2 (version 1.14.1) (42) and limma (version 3.30.13) (43) were used to identify differentially expressed genes (DEGs) by comparing the two samples from the GLI2shR DX+ condition with the other six samples. Significant DEGs were identified using a filter of Bonferroni’s correction-adjusted P≤0.01 for each package and sequence read abundance per gene >40% within each sample/library. The DEGs were defined as the intersected overlapping genes found by all three packages. DEGs were analyzed further for the functional enrichment of gene ontology using the Database for Annotation, Visualization and Integrated Discovery (DAVID) platform (44,45) and QIAGEN’s Ingenuity® Pathway Analysis (Ingenuity Systems; Qiagen Sciences, Inc.).

**Soft agar assay.** LNCaP Gli2shR, LNCaP Ctrl, 22RV1 Gli2shR and 22RV1 Ctrl cells were first resuspended in 250 µl 0.4% agarose supplemented with 5% FBS and seeded onto a layer of 0.8% agarose supplemented with 5% FBS, such that the final cell density was 2,000 cells/well in 24-well tissue culture plates. DOX (0.1 or 1 µM) and/or Casodex (30 µM) or DMSO were added five days after seeding, when small colonies could be observed, following which the plates were maintained in an incubator for a further 10-15 days. Colonies were subsequently stained using 1 mg/ml p-iodonitrotetrazolium violet staining solution (Sigma-Aldrich; Merck KGaA) for 1-2 days in an incubator. All incubations were performed at 37˚C. The plates were scanned using a scanner (Epson Perfection 2450 PHOTO; Seiko Epson Corporation). The colony number in the scanned image of each well with no magnification was quantified using ImageJ software (version 1.8.0; National Institutes of Health).

**Analysis of androgen-independent progression of LNCaP cells in vitro.** On day 0, 1x10⁶ LNCaP Ctrl cells and LNCaP Gli2shR cells were first seeded into 6-well plates in medium supplemented with 10% FBS. Following overnight incubation, the medium was changed to an androgen-depleted medium containing 10% charcoal-stripped FBS (Gibco; Thermo Fisher Scientific, Inc.), where the cells were treated with 1 µM DOX or vehicle control. When the Ctrl cells reached 90% confluence, cells were passaged simultaneously using the same ratio (1:3-1:4). All incubations were performed at 37˚C. Phase contrast images of the cells were taken on days 1, 10 and 25 following culture using a light microscope (magnification, x40). Cell numbers were counted from five fields of view per condition.

To investigate the association of Gli2 expression with androgen-independent progression in vitro, 1x10⁶ LNCaP cells were first seeded into 6-well plates in DMEM supplemented with 10% FBS. Following overnight incubation, the medium was changed to an androgen-depleted medium containing 10% charcoal-stripped FBS supplemented with or without 10 pM synthetic androgen R1881 (cat. no. NLP005; PerkinElmer, Inc.) on day 0. Gli2 expression levels in LNCaP cells were subsequently assessed by reverse transcription-quantitative PCR (RT-qPCR) analysis on days 1, 3, 7, 14 and 21.

**Western blot analysis.** LNCaP, 22RV1 and VCaP cells were harvested and lysed using a lysis buffer containing 50 mM Tris-HCl (pH 7.4), 150 mM NaCl and 0.5% NP-40 supplemented with a protease inhibitor (Thermo Fisher Scientific, Inc.) and a phosphatase inhibitor cocktail (1 mM NaVO₃, 5 mM NaF and 20 mM B-glycerophosphate). Total protein concentration was quantified using bicinchoninic acid protein assay (Thermo Fisher Scientific, Inc.) and equal amounts of protein (20-50 µg) were separated by 5% SDS-PAGE, transferred to a nitrocellulose membrane under constant voltage and blocked with 5% non-fat dried milk dissolved in TBS containing 0.05% Tween-20 (TBST) at room temperature for 30 min. The membranes were subsequently incubated with primary antibodies at 4˚C overnight before incubation with secondary antibodies at 4˚C for 1 h. All primary and secondary antibodies were diluted in TBST supplemented with 3% milk or BSA (Sigma-Aldrich; Merck KGaA).
Proteins were visualized by chemiluminescence using Immobilon Western Chemiluminescent HRP Substrate (cat. no. WBKLS0050; EMD Millipore). Primary antibodies against Gli2 (1:200; cat. no. NB600-874; Novus Biologicals, LLC; and cat. no. SC-20291; Santa Cruz Biotechnology, Inc.) and GAPDH (1:5,000; cat. no. 80602-840; Calbiochem; EMD Millipore) and horseradish peroxidase-conjugated secondary antibodies (Goat anti-mouse IgG; cat. no. 115-035-003; 1:4,000; Goat anti-rabbit IgG, cat. no. 111-035-003; 1:3,000; Donkey anti-goat IgG, cat. no. 705-035-003; 1:4,000; all from Jackson ImmunoResearch Laboratories, Inc.) were used for western blotting, in procedures that were described previously (34). The Western blot images were obtained using the FluorChem™ E imager with AlphaView software version 4.1.4 (ProteinSimple).

Reverse transcription-quantitative PCR (RT-qPCR). RNA was extracted from prostate cancer cells and xenograft tumors using RLT Plus buffer (Qiagen China Co., Ltd.) and RNA binding columns (cat. no. EZCR101; Enzymax, LLC), where the extracted RNA was shredded using a EZshredder column (Enzymax LLC) to remove genomic DNA. Total RNA (2 µg) was reverse-transcribed into cDNA using a high-capacity cDNA reverse transcription kit (Invitrogen; Thermo Fisher Scientific, Inc.). The temperature protocol used for reverse transcription was as follows: 25°C for 10 min, 37°C for 60 min and 70°C for 15 min. Subsequent qPCR was performed using Power SYBR® Green PCR Mix from Life Technologies, Thermo Fisher Scientific, Inc. The thermocycling conditions were as follows: Initial denaturation at 95°C for 10 min, followed by 40 cycles of denaturation at 95°C for 15 sec, annealing at 60°C for 15 sec and extension at 70°C for 20 sec. All primers used in this study were designed using Primer Blast of NCBI and synthesized by Integrated DNA Technologies, Inc. Primer pair specificity was determined by the generation of a single peak on a dissociation curve (melting curve) at the end of RT-PCR cycling program. The relative expression level for each gene was calculated according to the 2^-ΔΔCq method (46), where GAPDH was used as the reference gene. The primer sequences for Gli2 and Gli2AN are forward, 5'-GGGTTCAACCCAGGGTGTCCTCCAGCTGT-3' and reverse, 5'-GATGGAGGGCAGGGTCAAGGAGTTT-3' and GAPDH forward, 5'-GGGTCAACCAGGTGTCCAGCACTGT-3' and reverse, 5'-GGCCCAATAAGCACCACCAAATCCGT-3'.

Cell cycle analysis. Cell cycle analysis was performed with LNCaP Gli2shR and LNCaP Ctl cells. Cells were cultured in medium supplemented with 10% FBS and treated with 1 µM DOX or vehicle for 48 h before being harvested and incubated with 70% ethanol overnight at 4°C for fixation. The fixed cells were pelleted at 376 x g for 10 min at room temperature, washed twice with PBS and a DNA staining solution ([PBS, 0.5% Triton X-100, 50 µg/ml propidium iodide (Sigma-Aldrich; Merck KGaA) and 500 µg/ml RNase A (Worthington Biochemical Corporation)] was added to the cell pellets for resuspension (cell density, 1x10^6 cells/ml). Cells were then incubated at 37°C in the dark for 45 min followed by storage at 4°C until analyzed using a BD™ LSR II flow cytometer (Becton, Dickinson and Company). FlowJo software (version 7.0; FlowJo LLC) was used to analyze flow cytometry data. Each assay was performed in triplicate.

Animal studies. All mouse experiments were performed in accordance with the appropriate guidelines (47) and were approved by the Institutional Animal Care and Use Committee of the Department of Laboratory Animal Resources at the University of Texas Health Science Center (San Antonio, USA). Animal health and behavior were monitored daily. All xenograft-bearing mice had a single tumor and were euthanized by cervical dislocation after being anesthetized with 2% isoflurane inhalation before the largest xenograft tumor reached 2 cm³, with a maximal diameter <2 cm. In total, 40 male (age, 6-8 weeks; weight, 16-20 g) severe combined immunodeficient (SCID) mice were obtained from Taconic Farms Inc. Animals were housed in polycarbonate standard mouse cages (35x20x15 cm) with wood shavings (Lignocel) as bedding material. Housing conditions were kept constant (temperature, 18-21°C; humidity, 40-70%) with a 12-h light/dark cycle. Pelleted mouse food and water was provided ad libitum. LNCaP Gli2shR cells or LNCaP Ctl cells (6x10^6 cells in 120 µl), suspended in 50% Matrigel (Corning Life Sciences) in cold PBS were injected subcutaneously into the right flank of each mouse (n=20 mice per group). After 2-3 weeks, 26 mice with tumor burdens >100 mm³ (volume = 0.5 x length x width²) in volume (16 mice from theCtl group and 10 mice from the Gli2shR group) were recruited. In total, six Ctl cell-injected mice were used for measuring Gli2 expression in the xenograft tumors with IHC and the rest of the mice were subdivided into four groups (n=5 mice per group). All 26 mice were surgically castrated to mimic androgen deprivation therapy under anesthesia with 2% isoflurane inhalation. The mice were kept warm on a heating pad during the surgery. Triple Antibiotic/pain relieving ointment (Mupirocin 2%; NDC 51672-1354-0; Taro Pharmaceutical Industries, Ltd.) was applied at the surgical site following surgery and during recovery. Buprenorphine (0.05-0.1 mg/kg) was injected subcutaneously twice a day for two days following surgery for analgesia. A total of 3 mice injected with Ctl cells were euthanized on day 1 whilst the other 3 mice injected with Ctl cells were euthanized on day 15 following castration. Tumor volume gradually reduced for the following four weeks before DOX treatment was initiated prior to tumor relapse. Castrated mice with tumors were initially administered with drinking water containing 5% sucrose (for palatability purposes) with or without 1 mg/ml DOX, which was then reduced to 1.25% sucrose after six days of treatment for the groups not treated with DOX as the control mice tended to consume too much water, whilst the DOX treatment groups continued receiving 5% sucrose with 1 mg/ml DOX. Water was replenished every four days. Tumor size and body weight were measured once every 2-4 days after surgical castration. The development of androgen independence was monitored by the growth of tumors. On day 60 following castration, when a clear difference was observed between the Gli2shR DOX⁺ and Gli2shR DOX⁻ groups, in addition to the average tumor volume of mice in the Gli2shR DOX⁺ group reaching 1,000 mm³, the experimental endpoint was reached. The mice were euthanized by cervical dislocation following anesthesia with 2% isoflurane inhalation, following which the tumors were excised. Parts of tumor tissue was then used to for the extraction of RNA for RT-qPCR whilst another part was used for mRNA expression analysis.
fixed with 10% formalin for 24 h at room temperature, which was then replaced with 70% ethanol until paraffin embedding. Detection of Gli2 and Ki67 expression was performed using immunohistochemistry (IHC) assay.

To investigate tumor response to DOX withdrawal, 6 mice from the LNCaP Gli2shR group (n=20) aforementioned developed tumors later. They were castrated when tumors reached an average volume of ~270 mm³ 2 months after the inoculation of tumor cells. In total, 8 mice failed to develop tumors or exhibited small tumors (4 in the Ctl group and 4 in the Gli2shR group), which were euthanized at this time. The 6 mice were divided into two groups (n=3 per group) with one group receiving 1.25% sucrose-containing drinking water without DOX and the other group receiving 5% sucrose-containing drinking water with 1 mg/ml of DOX starting 3 days after castration. Water was replenished every 4 days. On day 51 following castration, all drinking water was changed to regular water. On day 63 after castration, when the maximum tumor volume reached 1,966 mm³, all three mice in the control group were euthanized, whilst the three DOX-treated mice were euthanized 81 days after castration when tumor growth began to accelerate.

**IHC assay.** Tissue sections of formalin-fixed and paraffin-embedded tumor tissues were cut into 4.5-µm sections using a LEICA RM2255 microtome (Leica Microsystems GmbH) and dried at room temperature for 24 h. The sections were then heated to 75°C for 15 min and rehydrated using xylene followed by a descending ethanol series. Antigen retrieval was performed by incubation with a sodium citrate solution (10 mM, pH 6.0) or an EDTA solution (1 mM, pH 8.0) at 95°C for 15 or 30 min, respectively. Sections were then blocked for endogenous peroxidase using 3% hydrogen peroxide (Thermo Fisher Scientific, Inc.) for 15 min at room temperature, permeabilized by incubation in 0.3% TBST for 5 min followed by another incubation in 0.025% TBST for 5 min at room temperature and blocked in 10% goat (Equitech-Bio, Inc.) or donkey (Equitech-Bio, Inc.) serum for 1 h at room temperature. Primary antibodies against Gli2 (1:200; cat. no. NB600-874; Novus Biologicals, LLC) or anti-Ki67 (1:50; cat. no. 550609; BD Biosciences), diluted in 1X PBS with 0.025% Triton X-100 and 5% serum from the species in which secondary antibody was raised were added to the tissue sections followed by incubation at 4°C overnight. The sections were then incubated with biotinylated secondary antibodies at room temperature for 1 h (1:200; cat. nos. NB600-874; Novus Biologicals, LLC) or anti-Ki67 (1:50; cat. no. 550609; BD Biosciences), diluted in 1X PBS with 0.025% Triton X-100 and 5% serum from the species in which secondary antibody was raised were added to the tissue sections followed by incubation at 4°C overnight. The sections were then incubated with biotinylated secondary antibodies at room temperature for 1 h (1:200; cat. nos. 550337 and 550338; BD Pharmingen; BD Biosciences). Streptavidin-Horseradish Peroxidase and DAB Substrate kit (BD Pharmingen; BD Biosciences) were used for the detection of staining under a light microscope (magnification, x40), whilst counterstaining was performed using hematoxylin at room temperature for 2 min.

**Statistical analysis.** Student’s t-test was performed to compare the means of two groups, whereas one-way or two-way ANOVA followed by Tukey’s or Sidak’s multiple comparisons test was used for analyzing data when the means of ≥2 groups were compared. Results are presented as the mean ± SEM. P<0.05 was considered to indicate a statistically significant difference. All the statistical analyses
Results

Gli2 knockdown inhibits prostate cancer cell proliferation and induces cell cycle arrest at the G₀/G₁ phase. LNCaP Gli2shR and 22RV1 Gli2shR cell lines were generated by transfection with the pTet-Gli2shR plasmid. Preliminary data in LNCaP Gli2shR cells demonstrated that 1 µM DOX treatment induced a marked reduction in Gli2 protein expression (Fig. S1). LNCaP Gli2shR, 22RV1 Gli2shR and corresponding control cells were treated with 1 µM DOX for 48 h. Gli2 mRNA and protein expression were both observed to be markedly lower in LNCaP Gli2shR and 22RV1 Gli2shR cells following DOX treatment compared with corresponding control cells (Fig. 1A and B). In addition, treatment of LNCaP Gli2shR and 22RV1 Gli2shR cells with 1 µM DOX resulted in significant reductions in cell viability (Fig. 1C). By contrast, cell viability in corresponding control cells was not affected by 1 µM DOX treatment (Fig. 1C).

To verify the aforementioned observations, a separate Gli2 shRNA (Gli2 shR1) lentiviral vector was stably transfection into the LNCaP, 22RV1 and VCaP cell lines, which also resulted in marked reductions in Gli2 expression and significantly reduced cell viability (Fig. S2).

To investigate the effects of Gli2 knockdown on cell cycle progression, LNCaP Gli2shR and LNCaP Ctl cells were treated with 1 µM DOX or vehicle for 48 h, following which the cells were harvested for cell cycle analysis. Gli2 knockdown resulted in a significantly increased proportion of LNCaP cells in the G₀/G₁ phase (Fig. 1D) and a significant reduction of cells in S phase (Figs. 1D and S3). To clarify that this observed growth inhibitory effect is Gli2 specific, a rescue experiment was performed by first stably transfecting LNCaP Gli2shR cells with the Gli2ΔN expression vector or control pSDM101 vector (CV), followed by treatment with or without DOX and measurement of cell viability using the MTT assay. Compared with cells transfected with the control vector, Gli2 expression as found to be significantly higher in LNCaP Gli2shR cells transfected with Gli2ΔN (Fig. S4A and B), which also significantly enhanced cell viability in the
absence of DOX (Fig. S4C). Following treatment with DOX, LNCaP Gli2shR/Gli2ΔN cell viability was significantly higher compared with cells transfected with the control vector. Therefore, restoration of Gli2 expression reversed the inhibitory effects of Gli2 knockdown on LNCaP cell viability (Fig. S4C). These findings suggest that the cell growth inhibition following Gli2 shRNA transfection was specifically due to Gli2 knockdown.

Transcriptome analysis consolidates the role of Gli2 in cell cycle regulation. Prostate cancer cell lines LNCaP Gli2shR and LNCaP Ctl were treated with either DOX (0.1 µM or 1.0 µM) and/or 30 µM Casodex or equivalent control vehicle, DMSO. Cell viability was measured on days 2, 4, 6 and 8 by measuring absorbance at 595 nm. Data is presented as the mean ± SEM of absorbance of 3 wells per group. ****P<0.0001, Casodex vs. Casodex +0.1 DOX. Soft agar assay was performed with LNCaP Gli2shR, LNCaP Ctl, 22RV1 Ctl and 22RV1 Gli2shR cells following treatment with DOX (0.1 µM or 1.0 µM) and/or 30 µM Casodex. The number of colonies formed were then counted 15-20 days later. Data is presented as the mean ± SEM of three experimental replicates. *P<0.05, **P<0.001 and ***P<0.0001. ns, not significant; OD, optical density; CAS, Casodex; DOX, doxycycline; Gli2, Glioma-associated oncogene family zinc finger 2; Ctl, control vector not encoding the shRNA; Gli2shR, vector encoding Gli2 short hairpin RNA.
Gli2 knockdown (Fig. 2C), suggesting that Gli2 is required for driving cell cycle progression, consistent with the aforementioned findings that Gli2 knockdown resulted in cell cycle arrest (Fig. 1).

**Gli2 knockdown exerts additive effects with Casodex in the inhibition of anchorage-dependent and -independent growth of prostate cancer cells.** Following observations that Gli2 may serve a significant role in prostate cancer cell cycle regulation, the effect of Gli2 downregulation on the inhibitory properties of the antiandrogen Casodex was next assessed. LNCaP Ctl, LNCaP Gli2shR, 22RV1 Ctl, and 22RV1 Gli2shR cells were first treated with DOX (0.1 or 1.0 µM) and/or 30 µM Casodex or DMSO, following which MTT assay was performed on days 2, 4, 6 and 8 to measure cell viability. All cells exhibited significant reductions in cell viability in response to Casodex (Fig. 3A), suggesting the growth-promoting properties of AR in these cell types. Interestingly, induction of Gli2 knockdown by DOX treatment in combination with Casodex resulted in further significant inhibitions of cell viability in Gli2shR cells, but not in control cells, seeded onto a plastic surface (Fig. 3A) and on soft agar (Fig. 3B-E). Taken together, these data suggest that downregulation of Gli2 was sufficient to enhance the sensitivity of LNCaP and 22RV1 cells to the anti-proliferative effects of the androgen antagonist Casodex. Gli2 expression is associated with androgen-independent progression. The association of Gli2 expression with androgen-independent progression was next examined in vitro and in vivo. LNCaP cells were cultured in an androgen-depleted medium containing 10% charcoal-stripped serum supplemented with or without 10 pM synthetic androgen R1881. Gli2 expression levels of LNCaP cells were subsequently assessed by RT-qPCR analysis.

The role of Gli2 on androgen-independent prostate cancer tumor growth was next investigated. LNCaP Ctl cells and LNCaP Gli2shR cells were first cultured in an androgen-depleted medium, following which they were treated with 1 µM DOX or vehicle. The number of cells in the LNCaP Gli2shR DOX+ group was significantly lower compared with that in the LNCaP Gli2shR DOX- and LNCaP Ctl DOX+ cells. Taken together, these results suggest that Gli2 is required for driving cell cycle progression.

**Figure 4.** Gli2 knockdown prevents the outgrowth of LNCaP cells cultured in an androgen-depleted medium. (A) LNCaP Ctl and LNCaP Gli2shR cells were maintained in androgen-depleted medium and chronically treated with or without 1 µM DOX. Phase contrast images of the cells were obtained after 1, 10 and 25 days. Magnification, x40. (B) Cell numbers were quantified on day 25. Data are presented as the mean ± SEM of three experimental repeats. ***P<0.001, ns, not significant; DOX, doxycycline; Gli2, Glioma-associated oncogene family zinc finger 2; Ctl, control vector not encoding the shRNA; Gli2shR, vector encoding Gli2 short hairpin RNA; hpf, high powered field.
groups after 10 days (Fig. 4A and B), with the difference more notable after 25 days. These observations suggest that androgen deprivation-induced Gli2 upregulation is necessary for prostate cancer transition into androgen-independence.
For in vivo studies, 16 SCID mice bearing xenograft tumors formed by injection with LNCaP Ctl cells were castrated when the tumors reached an average size of 200-300 mm³. A total of 3 mice were euthanized each on days 1 and 15 after castration, whereas the remaining 10 mice used for with or without DOX treatment with 5 mice in each group, which were euthanized on day 60 (Fig. 5B). Gli2 expression in tumor tissues were measured using immunohistochemical staining. When cultured in an androgen-depleted environment, Gli2 expression in LNCaP cells was upregulated in a time dependent manner, but not when in the presence of the synthetic androgen R1881 (Fig. S5A), consistent with previous observations (33,34). Gli2 expression was also found to be upregulated in a time dependent manner in LNCaP xenograft tumors following castration, where they became castration resistant (Fig. S5B).

Gli2 knockdown inhibits the growth of castration-resistant tumors in vivo. The effects of Gli2 knockdown on the androgen independence of LNCaP xenografts following castration was next examined in vivo. Male SCID mice were inoculated with either LNCaP Gli2shR or LNCaP Ctl cells mixed with Matrigel, following which they were castrated when tumors reached an average volume of 200-300 mm³ two weeks after injection (Fig. 5A).

Following castration, mice were subdivided into two groups, where one group were provided with drinking water supplemented with DOX, whilst the other was provided with drinking water without DOX. Mice from all groups exhibited tumor relapse 40 days after castration (Fig. 5B), but mice in the LNCaP Gli2shR DOX⁺ group displayed continuous tumor regression (Fig. 5B). Differences between LNCaP Gli2shR DOX⁺ and LNCaP Gli2shR DOX⁻ were notable shortly after DOX treatment, which became significant from day 45 onwards (Fig. 5B). Consistent with the final tumor volume, tumor weights of mice from the LNCaP Gli2shR DOX⁺ group was significantly reduced compared with those in the LNCaP Gli2shR DOX⁻ group, whilst no significant differences were observed between the two LNCaP Ctl groups (Fig. 5C). Mice from the LNCaP Gli2shR DOX⁺ group also exhibited significantly increased loss in body weight compared with the LNCaP DOX⁻ group (Fig. 5D). Analysis of Gli2 expression in tumor tissues revealed that Gli2 mRNA and protein expression was markedly lower in the LNCaP Gli2shR DOX⁺ group compared with the LNCaP Gli2shR DOX⁻ group (Fig. 5E and F). Immunohistochemistry also revealed reduced Ki67 expression in tumor tissues isolated from mice in the LNCaP Gli2shR DOX⁺ group, suggesting reduced LNCaP xenograft cell proliferation in vivo (Fig. 5G).

To investigate tumor response to DOX withdrawal, six mice bearing LNCaP Gli2shR tumors were castrated and divided into two groups three days following castration, with one group receiving DOX and the other without DOX. After a short response to surgical castration, LNCaP Gli2shR DOX tumors quickly relapsed in the 47 days following treatment, but not in LNCaP Gli2shR DOX⁻ tumors (Fig. 5H); significant differences were observed in the tumor volumes between these groups from day 10 after DOX treatment onwards (Fig. 5H). DOX treatment was withdrawn after 47 days, where tumor relapse was observed in both LNCaP Gli2shR groups. In conclusion, these data suggest that the suppression of Gli2 expression can sensitize LNCaP tumors to androgen deprivation, resulting in significant regression of LNCaP tumors and preventing the progression of androgen-sensitive LNCaP tumors to castration-resistant tumors in SCID mice.

Discussion

Accumulating evidence suggest that the re-activation of canonical hedgehog signaling occurs in prostate cancer cells during androgen-deprivation (27,34). In addition, Gli2 expression and activity can be regulated by alternative signaling pathways, including Ras and TGF-β signaling (35). Therefore, in the present study, the role of Gli2, a critical component of the hedgehog signaling pathway, in the progression of hormone-naïve prostate cancer to CRPC was studied. Analysis of Gli2 expression in LNCaP tumors in castrated SCID mice showed that castration was associated with Gli2 upregulation. This was consistent with a previous study, which showed that androgen deprivation resulted in increased Shh, Gli2 and Ptc expression in LNCaP and other androgen-responsive prostate cancer cell lines in vitro (33). In addition, Narita et al. (26) previously compared the Gli2 expression profiles of benign prostate hyperplasia, prostate cancer treated with neoadjuvant hormonal therapy and androgen-independent prostate cancer using a tissue microarray and found that Gli2 expression was significantly higher in prostate cancer compared with benign prostate hyperplasia, which was reduced following androgen ablation in a time-dependent manner; by contrast, Gli2 expression was found to be reactivated in androgen-independent prostate cancer. However, it should be noted that increases Gli2 mRNA expression was not observed when compared between untreated and hormone deprivation therapy-treated prostate cancers in a limited number of gene expression profiling studies (48,49). Given the heterogeneity of gene expression among prostate cancers in humans, the ≤20 samples tested in these two previous paired studies of prostate cancer pre- and post-hormone deprivation therapy is likely to be insufficient, where a larger sample size is required to verify the regulation of Gli2 expression in prostate cancer in humans during hormone deprivation therapy.

One of the novel findings in the present study was that LNCaP tumors with reduced Gli2 expression failed to progress to CRPC following castration-induced androgen deprivation. A previous study targeted Smo using either cyclopamine or siRNA demonstrated in vitro that Hedgehog/Gli signaling supported androgen-independent growth of prostate cancer cells in a low androgen environment (27). However, the role of Gli transcription factors in CRPC progression remains to be fully elucidated. In another previous study, which used PC-3 xenografts as an advanced model of CRPC, found that targeting Gli2 using an antisense oligonucleotide induced CRPC apoptosis (26). An important distinction in the present study is that tumors from LNCaP cells were used in SCID mice as a preclinical prostate cancer model. LNCaP xenografts exhibit similar behavior compared with clinical prostate cancer tumors, since both typically relapse following a temporary remission due to androgen deprivation (50). Therefore, LNCaP xenograft represent a superior model in mimicking the CRPC progression process. A number of studies have previously demonstrated...
increased hedgehog/Smo expression and subsequent signaling activity in hormone deprivation therapy-treated and/or metastatic human prostate cancers, where Gli2 was revealed as an important mediator of hedgehog/Smo signaling (25,30,34). In the present study, Gli2 knockout prevented the formation of CRPC, suggesting that the targeting of Gli2 expression in the early stages following hormonal therapy may be beneficial in prolonging relapse-free survival, even though the upregulation of Gli2 mRNA expression after castration in prostate cancer cell lines and xenografts has not been corroborated in human samples.

Despite being androgen-independent, AR signaling remains a pivotal driver in mediating disease progression in CRPC (51,52). Mechanistically, elevated expression of AR signaling coactivators in tumor cells facilitates AR activity in a low androgen environment (53). Previous studies showed that Gli2 can potentially serve as a coactivator of AR, where it interacts with the Tau5/AF-5 N-terminal activation domain of AR, referred to as the androgen-independent activation domain (27,54). Gli2 may also function by supporting androgen signaling in CRPC cells through the coactivation of the full-length AR and its splice variants (54).

Another plausible explanation for the sustained AR signaling may be the prevalence of AR splice variants, which encode truncated versions of the AR protein that lacks the C-terminal ligand binding domain (4,55). These versions of AR are transcription factors that are androgen-independent and are constitutively activated. The molecular mechanism by which AR splice variants are generated remains unclear. A previous study demonstrated that androgen deprivation therapy increased the rate of AR gene transcription and the recruitment of splicing factors to the AR pre-mRNA, in turn contributing to elevated AR-V7 levels in prostate cancer cells (56). Two RNA splicing enhancers and associated binding proteins (U2 Small Nuclear RNA Auxiliary Factor 65 and Serine and Arginine Rich Splicing Factor 1) were identified to serve critical roles in the splicing of the AR pre-mRNA into AR-V7 (56). Therefore, future studies will be focused on investigating the effects of Gli2 on the expression of AR splice variants under androgen-deprived conditions and associated mechanism in the development of CRPC.

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Availability of data and materials

The analyzed datasets generated during this study are available from the corresponding author on reasonable request.

Authors' contributions

LZS made substantial contributions to the conception and design of the study. LX, HB, CZ, TQ, BZW, YHW, JHY and HYZ contributed to the experimental design and performed the experiments, in addition to data analysis and interpretation. XG, CZ and YZ performed the whole transcriptome sequencing analysis. LX and LZS wrote and revised the manuscript. PJH provided the mice for the xenograft experiment and performed the acquisition of data. WZ performed analysis and interpretation of the data. All authors have read and approved the final version of manuscript.

Ethics approval and consent to participate

All animal experiments were conducted following appropriate guidelines. They were approved by the Institutional Animal Care and Use Committee and monitored by the Department of Laboratory Animal Resources at the University of Texas Health Science Center (San Antonio, USA).

Patient consent for publication

Not applicable.

Competing interests

The authors declare that they have no competing interests.

References


