High-throughput screening identifies candidate drugs for the treatment of recurrent respiratory papillomatosis

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A B S T R A C T

Recurrent respiratory papillomatosis (RRP) is a benign neoplasm of the larynx caused mainly by human papillomavirus type 6 or 11 and its standard treatment involves repeated surgical debulking of the laryngeal tumors. However, significant morbidity and occasional mortality due to multiple recurrences occur. Conditional reprogramming (CR) was used to establish a HPV-6 positive culture from an RRP patient, named GUMC-403. High-throughput screening was performed at the National Center for Advanced Technology (NCATS) to identify potential drugs to treat this rare but morbid disease. GUMC-403 cells were screened against the NPC library of > 2800 approved drugs and the MIPE library of > 1900 investigational drugs to identify new uses for FDA-approved drugs or drugs that have undergone significant research and development. From the two libraries, we identified a total of 13 drugs that induced significant cytotoxicity in RRP cells at IC50 values that were clinically achievable. We validated the efficacy of the drugs in vitro using CR 2D and 3D models and further refined our list of drugs to panobinostat, dinaciclib and forskolin as potential therapeutics for RRP patients.

1. Introduction

Recurrent respiratory papillomatosis (RRP) is a rare disease with an actual incidence of approximately 20,000 cases in the United States [1]. RRP is characterized by the growth of tumors in the respiratory tract caused by the human papillomavirus type 6 or 11 (HPV-6 or -11) and generally classified into two subtypes: juvenile-onset RRP (JORRP) and adult-onset RRP (AORRP) [2,3]. JORRP cases, which develop before the age of 14, are more recurrent and aggressive than their AORRP counterpart [4]. Despite the fact that RRP primarily occurs on and around the laryngeal vocal cords, these growths can spread downward and affect the bronchi, the trachea and intermittently the lung parenchyma [2,3]. Life-threatening breathing complications such as acute respiratory distress can result from untreated papilloma. When progression to the lung occurs there are limited treatment options, the disease is fatal [5].

Presently, there is no “cure” for RRP, and no single adjuvant treatment has reliably been shown to be effective in eliminating RRP [6]. The backbone of RRP therapy is surgical excision to debulk the papilloma without injuring the normal tissues [7]. A characteristic feature of this disease is the tendency for the papilloma to reappear after surgical excision, causing JORRP patients to have an average 4.4 surgeries per year [1,6,7].

One of the limitations for investigating RRP treatment has been the lack of a suitable cell culture system. As reported previously, Conditional reprogramming (CR) allows rapid and efficient isolation and propagation of primary tumor cells [8,9]. In contrast to existing conventional cancer cell lines, the conditionally reprogrammed (CR) tumor cells maintain the cancer-specific mutations and phenotypic heterogeneity typically seen in the primary tumor [10–12]. Therefore, CR cells represent an advanced cancer model for preclinical drug development. In the past, we isolated and propagated CR cultures from an HPV-11-positive RRP patient. We utilized the generated cell line to detect an HPV-11 mutation that may have been responsible for the
observed aggressive clinical phenotype. We also used the patient’s cells for a limited drug screening and identified vorinostat, an HDAC inhibitor, as a potential treatment and subsequently showed that vorinostat was effective for arresting tumor growth in the patient [13].

To extend this line of work, we have used high-throughput screening to identify potential new drugs effective against RRP. The RRP CR cells, which contained episomal HPV-6 DNA, were screened by the National Center for Advanced Translational Sciences against the NPC library of 2816 approved drugs and the MIPE library of 1912 investigational drugs. From approximately 4700 drugs we have identified 3 that might be clinically actionable.

2. Methods

2.1. Cell isolation and propagation of normal and tumor tissue samples

Lung tissue (right lower lobe) was obtained at surgery with the patient’s written consent according to the Georgetown University Hospital IRB. The sample was digested with collagenase and trypsin, and the obtained cells were propagated using the Conditionally Reprogrammed (CR) system (Fig. 1), that includes the use of a bed of irradiated murine fibroblasts and a F medium supplemented with Y-27632 Rho-kinase inhibitor (Enzo Life Sciences) as described previously [8,9]. The cells, named as GUMC-403, were used for HTS, 2D and 3D cell viability assays. Previously a laryngeal sample from another patient was used to generate normal cells, named GUMC-228 (HPV negative), which used as a control in the 2D cell viability assay.

2.2. DNA isolation, cloning, and sequencing

DNA was isolated and purified from the cultured cells, GUMC-403, or directly from the tissues using the DNeasy Blood and Tissue Kit (Qiagen) and was amplified with specific primers for HPV-6, HPV-11 or with the Rolling Circle Amplification (RCA) kit (Illustra TempliPhi, GE Healthcare Life Sciences), as published previously [13]. The products of RCA were digested with EcoR I, BamHI, Hind III, Nde I and EcoR V. Two DNA fragments from the digestion were isolated from agarose gel, and cloned into pUC19 separately and sequenced from two directions with the use of Primer Walking Services (Genewiz).

2.3. HPV-6 viral copy number assay and E6:E2 ratio assay

Genomic DNA was extracted from RLL-403 and GUMC-403 and diluted with nuclease-free water to 10 μg/ml then used for all ddPCR experiments. ddPCR, probes were designed to amplify products of 100-300bp as recommended. The reaction mixtures contained ddPCR Probe Supermix (Bio-Rad Laboratories, Hercules, CA, USA), template DNA (1 μl) and primers (final concentration, 25 nM) in a final volume of 20 μl. Each reaction was then loaded into a sample well of an eight-well disposable cartridge (DG8™; Bio-Rad Laboratories) along with 70 μl of droplet generation oil (Bio-Rad Laboratories). A QX200™ Droplet Generator (Bio-Rad Laboratories) was used to generate the droplets as per the manufacturer’s directions. Droplets were then transferred to a 96-well PCR plate, heat-sealed with foil and amplified using a conventional thermal cycler to the end point (95 °C × 5 min (1 cycle), ...
read. The experiment was paused and the plate removed from the upper chamber followed 1 h incubation. Next, the background was lower chambers while 50

The Tagman assay used is forward primer 5′-TATCCA-3′, (reverse primer) 5′-TCGTGATTGTTAGTGATG) -3′, and probe 56-FAM/AC CAC ACG C/ZEN/A GTA CCA ACA TGA CA/-(CCACGTCTGCAACGACCATA) -3′. A 2.5 μM of HPV-6 L1 copy number reference assays were purchased commercially (Applied Biosystems). The HPV6-L1 Tagman assay used is forward primer 5′-(TGG AAG ATG TAG TTA CGG (GCTGG AAG ATG TAG TTA CGG GACC ) -3′, (reverse primer) 5′-(TTCCATGA AATTCTAGGCAGCA) -3′, and probe 56-FAM/AC CAC ACG C/ZEN/A GTA CCA ACA TGA CA/-TGGACCCGTGGAC TTTCGAGGCGGC -3′ normalization was done using DMSO as 100% cell viability and empty wells. For 2D drug culture treatments, 7 diﬀerent concentrations of Panobinostat, Dinaciclib or Forskolin were added in 50 μl to make up the volume to 100 μl. Controls included bortezomib at 9.2 μM, DMSO only, and empty wells. Plates were then returned to the incubator for 48 h. To assess cell viability, 3 μl of CellTiter-Glo reagent (Promega) was added to each well of the plates using a solenoid value dispenser. Plates were then incubated at room temperature for 15 min and then, luminescence signal was read using a ViewLux (PerkinElmer) with a 2 s exposure time. Data normalization was done using DMSO as 100% cell viability and empty wells as 0% viability.

2.6. High throughput screen (HTS) assay

GUMC-403 cells were plated at a density of 700 cells/well in 5 μl of (F + Y medium) into white tissue culture treated polystyrene plates (Corning Cat. 7464) using a MultiDrop Combi dispenser with small volume cassette. All plates were covered with a stainless steel gasketed lid and placed into an incubator at 37 °C/95% RH/5% CO2 overnight. The next day, 23 μl of MIPE 4.0 library, NPC library, and control compounds were added to each plate using a pin tool dispenser [17,18]. Controls included bortezomib at 9.2 μM, DMSO only, and empty wells. Plates were then returned to the incubator for 48 h. To assess cell viability, 3 μl of CellTiter-Glo reagent (Promega) was added to each well of the plates using a solenoid value dispenser. Plates were then incubated at room temperature for 15 min and then, luminescence signal was read using a ViewLux (PerkinElmer) with a 2 s exposure time. Data normalization was done using DMSO as 100% cell viability and empty wells as 0% viability.

2.7. Cell viability assay for 2D and 3D

GUMC-403 cells (5.0 × 10^3 cells/well in 100 μl of F + Y medium) were seeded in a 96 wells plate for 2D monolayer (WVR, Radnor, PA) or (5.0 × 10^3 cells/well in 50 μl of F + Y medium) in a 96-well ULA round-bottomed plates for 3D spheres culture (CLS3474, corning). Seeded cells were incubated overnight at 37 °C in a cell culture incubator with 5% CO2 levels. For 2D drug culture treatments, the medium was replaced with fresh medium that have 7 diﬀerent concentrations of the 13 drugs (drugs are listed in Table 1). For the 3D drug culture treatments, 7 diﬀerent concentrations of Panobinostat, Dinaciclib or Forskolin were added in 50 ul to make up the volume to 100 ul. Cell viability assay was conducted using The Veritas microplate luminometer turner Biosystems. The CellTiter-Glo® Luminescence Cell Viability Assay (G7570, Promega, Madison, WI) kit for 2D culture or CellTiter-Glo® 3D Cell Viability Assay (G9681, Promega, Madison, WI) kit for 3D culture and GloMax®-96 Microplate Luminometer Software (Promega) were used for data analysis according to the manufacturer’s protocol. The cell viability reading was measured after 3 days for 2D culture or 5 days for 3D culture and the treated cells luminescence reading was normalized to that of vehicle (DMSO) treated cells.

2.4. Amplification of papillomavirus oncogene transcribes PCR (APOT-PCR) for HPV-6

For detection of the physical status of HPV6 (episomal vs. integrated), a 3′-RACE APOT assay was used which was based on Huebers et al. [15]. After reverse transcription of RNA, a nested PCR with a set of 5′-Primer 5′-GGAGCCGACAAGATTCCA ACC -3′; 2nd 5′-primer: 5′-CTCTGTTGCTTGATGTGGATGTGGACAGC-3′ both located in the E7 open reading frame of HPV-6 and a 3′-Frohman primer (for both nested PCR-setups) was used. PCR products were separated on a 1% agarose gel. Visible bands were cut out of the gel, purified (Gel extraction kit, QIAGEN) and sequenced. Sequences were compared with NCBI and UCSC database entries to determine viral sequences or virus-human fusion sequences indicating viral integration.

2.5. Chemotaxis/cell migration assay

Cell migration experiments were performed using the xCELLigence RTCA (ACEA Biosciences Inc.) system as described before [16]. xCELLigence allows the examination of the cell migration process in real time by measuring electrical impedance. Experiments were carried out in 16-well plates (CIM-16, ACEA Biosciences Inc.). Briefly, 160 μl of cell culture medium with and without serum were dispensed in the lower chambers while 50 μl serum-free medium were added to the upper chamber following 1 h incubation. Next, the background was read. The experiment was paused and the plate removed from the RTCA-DP device. Then, 1.0 × 10^6 of GUMC-403 cells or Human Fore-skinned Keratinocytes (HKF) cells in a total volume of 50-μl serum-free medium/well were seeded on the upper chamber. Plate was placed back in the RTCA system and incubated for 27 h, performing measurements every 10 min. The electrical impedance is reflection of cell number and software was used to generate the migration activity of each condition. The graph represents the average of triplicates.

Table 1

| Drugs | Name | Mode of Action | Diseases Treated | Library |
|-------|------|----------------|-----------------|---------|
| Drug 1 | Panobinostat | HDAC inhibitor | Multiple myeloma(approved), HIV-1AART combination (in trial) | MIPE |
| Drug 2 | Dinaciclib | CDK2, CDK5, CDK1 and CDK9 inhibitor | In clinical trials for various cancer | MIPE |
| Drug 3 | Forskolin | ubiquitously activating of eukaryotic adenylyl cyclase | Glaucoma | NPC |
| Drug 4 | Veretoprin | Photo-sensitizing agent derived from porphyrin in endothelial cells. | Photodynamic therapy for abnormal blood vessels | MIPE |
| Drug 5 | Fomipinole | Inhibitor of the enzyme alcohol dehydrogenase. | Antidote for methanol or ethyl alcohol poisoning | MIPE & NPC |
| Drug 6 | Carfilzomib | Selective proteasome inhibitor | Multiple myeloma | MIPE |
statistical significance, the experiment including was carried in a 3 technical replicates and conducted at three independent times.

2.8. Xenograft assay

To determine in vivo tumorigenicity for the GUMC-403, 1 \times 10^6 cells were suspended in 200 μL of Matrigel HC (BD-growing Biosciences). The Matrigel-suspended cells were injected subcutaneously into the left and right flanks of 6-week-old male mice with severe combined immunodeficiency (Taconic, Germantown, NY). The growth of xenografts was measured weekly with callipers. Animals were housed at the Georgetown University animal care facility according to institutional guidelines. Animal protocol #14-033-100171 was approved by Institutional Animal Care and Use Committee (IACUC) at Georgetown University. All experiments were performed in accordance with the protocol relevant guidelines and regulations.

2.9. Statistical analysis

Unpaired student’s t-test was used to compare drug treatment response in primary cells. Data (mean ± s.e.m.) were calculated and plotted using GraphPad Prism 6.0 (La Jolla, CA).

3. Results

3.1. Generation and characterization of HPV-6 positive CR cultures from RRP patient

The patient was a 29 year old female with more than a 26-year history of recurrent respiratory papillomatosis. She had undergone more than 90 laryngeal ablation surgeries to control viral-induced tumors and had been additionally treated with intralesional cidofovir. However, the treatment was not able to slow tumor growth or its progression into the lung. Computed Tomographic (CT) scanning revealed that there were multiple pulmonary nodules that had accelerated in growth. Bronchoscopy was performed and papillomas from the right lower lobe (RLL) were excised and submitted for both pathology examination and cell isolation. Histology of the tumor revealed squamous papillary proliferation (Fig. 2A, Left) and intraepithelial mucocytes (Fig. 2A, Right). To facilitate the analysis of molecular alterations and drug screening in RRP, we established a cell culture (GUMC-403) from a right lower lobe papilloma biopsy using conditional reprogramming [9]. The cells could be observed as early as 2 days after isolation, and the primary culture reached confluence in 10 days (Fig. 2B). The cells were maintained for more than 28 population doublings (52 days) with an average growth rate of 45 h/doubling.

HPV typing using specific primers for HPV-6 or HPV-11 or general primers for HPVs was conducted. Only bands with primers for HPV-6 and general HPV were detected in the biopsy and GUMC-403 (Fig. 2C). The amplified bands were isolated and cloned. Sequencing of the PCR products showed that all the products matched HPV-6 DNA. HPVs can exist in two forms in the infected cells either episomal form or integrated form [19]. To confirm the anticipated episomal form of the HPV-6 DNA in GUMC-403, we performed rolling circle amplification (RCA) followed by digestion by a set of restriction enzymes [20]. The expected number and size of bands for HPV-6 were detected in the viral genome [13]. This cell line represents an HPV-6 positive cell culture containing episomal viral DNA.

Quantitative PCR was used for viral copy number assessment. The average viral copy number is 1.08 copy per cell in GUMC-403 at passage 2. Viral genome copy number gradually decreased during passages and was undetectable at passage 6. This is consistent with earlier studies that these HPV positive cells lack the maintenance of the episomal viral genome. All the characterization assays and drug-screening experiments were done between passages 2 and 3. Quantitative PCR assays were performed to verify the presence of HPV-6 genome at the time.

3.2. GUMC-403 cells maintain RRP characteristics

The capacity of tumor cells to migrate is a critical property for cancer metastasis [21]. Therefore, a transwell migration assay was used to measure the chemotaxis of GUMC-403 cells by electric impedance in response to a chemoattractant fetal bovine serum (FBS). GUMC-403 cells showed higher migration potential compared to human foreskin keratinocytes (HFK, negative control) in the presence or in the absence of fetal bovine serum (FBS) during a 27 h period (Fig. 2A). We further analyze GUMC-403 for in vivo tumorigenicity, cells were trypano- sinized from a culture at passage 5 and injected subcutaneously into immunodeficient mice. The tumors were measurable as early as 8 weeks post-injection. Xenograft experiments were performed three times independently, with a total of 11 out 12 xenograft sites producing tumors. Similar to the primary tumor (Fig. 2A), the xenografts were composed of atypia squamous papillary proliferation with focal koilocytic (Fig. 3B, Left) and intraepithelial mucocytes (Fig. 3B, Right). These results demonstrate that the cell line GUMC-403 maintains the tumorigenic phenotype in vitro and in vivo and mimic the original tumor.

3.3. High Throughput Screening assay for RRP cells

Drug screening was performed at the National Center for Advancing Translational Sciences (NCATS) using 1536 well plates. Drugs from the NPC library (8 point dilutions, 1:5 dilution) and MIFE library (11 point dilutions, 1:3 dilution) were evaluated at concentrations ranging from 0.5 nM to 50 μM [15,16]. Survival curves with an area under the curve (AUC) less than 425 for MIFE or less than 460 for NPC and Curve Class of −1.1 were selected [22]. Out of 4728 drugs, 45 drugs matched the criteria, and 13 were selected for further validation in 2D and 3D cultures. Priority was given to compounds based on their degree of cell killing, clinical status and FDA approval.

3.4. 2D and 3D drug sensitivity validation

The 13 potential candidates from NCATS screening (Table 1) were further validated using 2 and 3-dimensional culture systems. For the validation assays and to further narrow down the potential candidates, we used normal laryngeal cells isolated from a second RRP patient, named GUMC-228 (HPV negative), since the GUMC-403 patient did not provide normal tissue. 5.0 × 10^5 cells/well of GUMC-403 and GUMC-
228 were seeded in 96 well plates, allowed to attach for 24 h. The cell monolayer was then treated for 72 h with (Fig. 4A, I) Verteporfin (Fig. 4A, II), Fomepizole (Fig. 4A, III), Carlifozimib (Fig. 4A, IV), Flavopiridol (Fig. 4A, V), AT-7519 (Fig. 4A, VI), SNS-032 (Fig. 4A, VII), Romidepsin (Fig. 4A, VIII), PF-04691502 (Fig. 4A, IX), Sertindole and (Fig. 4A, X) Crenolanib in concentrations ranging from 50 μM to .5 nM. The cell viability assays revealed higher selective cytotoxicity toward GUMC-403 compared with GUMC-228 when treated with (Fig. 4B, I) Panobinostat (Fig. 4B, II), Dinaciclib and (Fig. 4B, III) Forskolin, depicted by an IC50 of 0.035 μM for Panobinostat, 0.010 μM for Dinaciclib and 5.157 μM for Forskolin (Table 2). The Cytotoxic effect of these drugs induced morphological changes within 3 days of treatment and ultimately a decreased cell viability compared to DMSO treated control (Fig. 4C I, II, III and IV).

In recent years, the use of 3D cell culture systems has been increasingly used as an in vitro model for drug discovery. Therefore testing drug candidates for efficacy and tissue distribution can be enhanced through 3D culture such as multicellular tumor spheroids (MCTS) due to the in vivo like microenvironment [23]. 5.0 × 10^3 cells/well of GUMC-403 and GUMC-228 were seeded in low attachment plates and incubated overnight. GUMC-403 cells formed spheres around the sizes of 200–400 μM (Fig. 5A left) while GUMC-228 cells failed to form spheres (Fig. 5A right). The RRP cells were treated for 5 days with Panobinostat, Dinaciclib or Forskolin. In the 3D culture, the IC50 of Panobinostat, Dinaciclib or Forskolin were 0.030 μM, 0.010 μM and 1.920 μM respectively, as calculated from dose-response curves (Fig. 5B, Table 2). Thus, we found that Panobinostat, Dinaciclib, and Forskolin have a similar cytotoxicity for RRP Cells in 2D and 3D in vitro models.

4. Discussion

Recurrent respiratory papillomatosis is a fatal disease once it has metastasized to the lung and there are limited therapeutic options since no single adjuvant therapy has been shown to be effective in eliminating RRP. A major limitation to study RRP tumor progression and treatment is the lack of an appropriate cell culture system. Recently CR culture has shown to maintain and imitate the ordinary biology of their primary tissue such as tracheal epithelium, ectocervical epithelium or breast tumors [10,11]. Additionally, intra-tumoral heterogeneity was maintained in CR cells suggesting oligoclonality of these cultures [12]. In an earlier study, we isolated and established a continuous CR cell cultures from a patient with recurrent respiratory papillomatosis. The HPV-11 positive CR cell culture helped us to detect a unique and important mutation in the viral genome. More importantly, the primary patient’s RRP cells enabled us to run a limited-scale drug screening and identified an effective therapy for the patient in less than two weeks [13]. In this study, we successfully established a RRP cell culture system that contains episomal HPV-6. Unlike in the earlier HPV-11 positive RRP cells, we did not detect any significant mutations of the HPV6 viral genome in this cell line. In xenograft assays, measurable tumors were observed as early as eight-weeks post injection into immunodeficient mice. The original cellular characteristics of the RRP tumor, as well as HPV genome, were maintained in the xenografts.

Recently, CR cells have been used in translational research for drug discovery. Using small scaled high-throughput drug screening with 306 clinical and emerging cancer drugs on CR cells, Saeed et al. (2017) have identified Bcl-2 family inhibitor navitoclax as a potential treatment for castration-resistant prostate cancer (currently is being tested in a
clinical trial) \[24\]. Chen et al. (2017) have used CR to grow cells from rare salivary gland cancers and identify regorafenib as a potential therapeutic drug \[25\]. Recently, Alamri et al. (2018) have shown that allosteric AKT inhibitor MK2206 can inhibit the growth of Mucoepidermoid carcinoma (MEC) cells in 2d and 3d CR culture \[26\]. Formerly, we have used CR method to identify vorinostat as an effective treatment for an HPV-11 positive RRP case; the appropriate therapy was identified in less than two weeks \[13\]. In contrast of the limited, small-scale drug screening in our earlier HPV-11 case using 96-well plates, the drug repurposing study in this HPV6 case was done in a high throughput format with thousands of drugs using 1536-well plates. The rapid expansion of 35 million cells of the RRP CR cells met the demand for a large number of testing cells.

Drug repurposing has emerged a novel approach in finding new treatments for unmet health conditions due to the well-defined side-effect profiles and the established bioavailabilities of drugs that led to FDA approval \[27,28\]. In this study, we identified three drugs cytotoxic to RRP cells and all of them have IC 50s that are within the safe range of therapeutic use and covering different classes of drugs such as CDK inhibitors, HDAC inhibitors, and non cancer drugs such as Forskolin. Panobinostat has been approved for multiple myeloma and cutaneous T-cell lymphoma \[29,30\]. Panobinostat (Farydac®, LBH589) is under clinical investigation for a range of hematological and solid tumors worldwide in both oral and intravenous formulations \[29,31\]. Panobinostat represses tumor cell growth by interacting with nonhistone and histones proteins as well as autophagy-mediated targets, apoptotic and tumorigenesis pathways involved in the development of cancer \[32\]. Recently, Vorinostat and Panobinostat were shown to inhibit HPV-18 E6 and E7 activities, leading to stabilize host cell tumor suppressors as well as abolishing the viral DNA amplification \[33\].

Our drug screening also found CDK inhibitors that had potent cytotoxicity against RRP cells. For instance, dinaciclib is an inhibitor of CDK1, CDK2, CDK5, and CDK9, and is active in a broad range of cancer cell lines originating from leukemia, melanoma, osteosarcoma and pancreatic cancer \[34–37\]. The mechanism by which dinaciclib inhibits RRP cells is unknown but previous study has shown that it inhibits RB phosphorylation in cancer cells at concentrations between 12 and 500 nM \[38,39\]. The IC50 from this study is 10 nm from 2D and 3D cell validation assays. Interestingly, the combination of HDAC and CDK inhibitors is a new leukemic and melanoma strategy since in combination they activate caspase; induce mitochondrial damage, and alter cell cycle regulation \[40,41\]. Therefore, we will explore the potential synergy of HADCi and CDKi present for RRP treatment.

Unlike panobinostat and dinaciclib, forskolin is a natural product that has been isolated from the roots of the plant Coleus Forskohlii \[42\]. Forskolin exhibits a wide range of pharmacological properties such as anti-obesity, asthma, and glaucoma by the stimulation of adenylyl cyclase activity and increases intracellular levels of cyclic AMP \[43–45\]. It has also been reported forskolin has anti-cancer activity. Treatment of colon cancer line KM12C with the adenylyl cyclase activator Forskolin completely inhibits their growth at the concentration of 50uM \[46,47\].

![Fig. 3. GUMC-403 cells maintain invasive and tumorigenic properties. (A) GUMC-403 cells exhibit a higher migration capacity in either the presence or absence of FBS compared with HFK (The graph is the average of triplicates). (B) Xenograft H&E staining is displaying koilocytes (Left, black arrow) and intraepithelial mucocytes (Right, black arrow). (40× magnification. Size bars = 200 μm).](image-url)
In the present study, we demonstrated that forskolin was cytotoxic to RRP cells at concentrations as low as 1.67 μM.

The use of three-dimensional (3D) cellular systems for drug discovery has been explored in drug discovery. A recent study, using the same HTS screen platform, showed similar drug responses between cancer cell lines growing as 2D monolayers and 3D spheres [48]. In this study, we developed the first 3D tumor model of RRP using patient's primary CR cells. Interestingly, we were able to validate drug sensibility in our 3D sphere system, and IC50s for the three drugs were similar in 2D and 3D systems. However, the 3D RRP culture does have the potential to discover novel mechanisms and targets and to accelerate target identification and validation, given that the gene expression patterns found in 3D models can better mimic physiological conditions [23].

Early studies have shown that low-risk HPV is known to be difficult to maintain in cultured cells [49]. In GUMC-403 cells, the average viral copy number is 1.08 copy per cell at passage 2. Viral genome copy number gradually decreased during passage, and was undetectable at passage 6. All the characterization assays and drug-screening experiments were done between passages 2 and 3. Quantitative PCR assays were performed to verify the presence of HPV-6 genome at the time.

Interestingly the xenograft assay with a relative late passage cells (passage 5) produced 11 tumors out of the 12 xenograft sites. More importantly the xenografts were composed of atypia squamous papillary proliferation with focal koliocytotic and intraepithelial mucocytes. These results suggested that even in the late passage the cell line GUMC-403 maintains the tumorigenic phenotype. CR cells, like GUMC-403, can be very useful for drug screening assay for RRP. However, the importance of validation of the HPV presence as well as the maintenance of tumorigenic phenotype should be performed at the same time.

5. Conclusion

In this study, by using conditional reprogramming and high throughput screen platforms, we identified and validate Panobinostat, Dinaciclib or Forskolin as effective drugs for recurrent respiratory papillomatosis therapy.

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Author contributions

Conception and design: F.A., S·P., F·W., H.Y. and R.S. Development of methodology: F.A., F·W., D.Z., S·P., X.Z., K·W., R.G., M.F., C.T and H.Y. Performed the high-throughput assay:., X.Z., R.G., M.F., C.T and H.Y. Analysis and interpretation of data: F.A., D.Z., F·W., K·W., L.A., H.Y. and R.S.; Writing, Review and/or revision of the manuscript: F.A., H·Y and R.S.; Administrative, technical, or material Support: F.A., D.Z., S·P., N.G., and H.Y. Study supervision: F.A., H·Y., T.C., and R.S.

Conflicts of interest

The authors declare no competing financial interests.

List of abbreviations

CR Conditional Reprogramming
PCR polymerase chain reaction
3D three-dimensional
2D two-dimensional
ROCK Rho-associated protein kinase
HFK human foreskin keratinocyte;
PDs population doublings.

Appendix A. Supplementary data

Supplementary data to this article can be found online at https://doi.org/10.1016/j.pvr.2019.100181.
Fig. 5. Validation of cytotoxicity of Panobinostat, Dinaciclib and Forskolin on the GUMC-403 3D culture. (A) Morphology of GUMC-403 spheres in ULA round-bottomed plates (left) and GUMC-228 (right). (B) Dose–response curves for (I) Panobinostat, (II) Dinaciclib and (III) Forskolin show similar cytotoxicity to 2D on 3D culture of GUMC-403. (10× magnification. Size bars = 400 μm). Data represents mean ± s.d. from 3 independent measurements, each in triplicate.

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