Expression of cell type incongruent alpha-cardiac actin 1 subunit in medulloblastoma reveals a novel mechanism for cancer cell survival and control of migration

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Abstract

**Background:** Alterations in actin subunit expression have been reported in multiple cancers, but have not been investigated previously in medulloblastoma.

**Methods:** Bioinformatic analysis of multiple medulloblastoma tumor databases was performed to profile ACTC1 mRNA levels. Western blot was used to verify protein expression in established medulloblastoma cell lines. Immunofluorescence microscopy was performed to assess ACTC1 localization. Stable cell lines with ACTC1 overexpression were generated and shRNA knockdown of ACTC1 was accomplished. We used PARP1 cleavage by Western blot as a marker of apoptosis and cell survival was determined by FACS viability assay and colony formation. Cell migration with overexpression or knockdown of ACTC1 was determined by the scratch assay. Stress fiber length distribution was assessed by fluorescence microscopy.

**Results:** ACTC1 mRNA expression is highest in SHH and WNT medulloblastoma among all subgroups. ACTC1 protein was confirmed by Western blot in SHH subgroup and Group 3 subgroup cell lines with the lowest expression in Group 3 cells. Microscopy demonstrated ACTC1 co-localization with F-actin. Overexpression of ACTC1 in Group 3 cells abolished the apoptotic response to Aurora kinase B inhibition. Knockdown of ACTC1 in SHH cells and in Myc overexpressing SHH cells induced apoptosis impaired colony formation, and inhibited migration. Changes in stress fiber length distribution in medulloblastoma cells are induced by alterations in ACTC1 abundance.

**Conclusions:** Alpha-cardiac actin (ACTC1) is expressed in SHH medulloblastoma. Expression of this protein in medulloblastoma modifies stress fiber composition and functions in promoting resistance to apoptosis induced by mitotic inhibition, enhancing cell survival, and controlling migration.

**Keywords**

medulloblastoma, cardiac actin, migration, apoptosis, chemo resistance
Key points

- Alpha cardiac actin (ACTC1) is expressed in SHH and WNT medulloblastoma.
- ACTC1 incorporates into F-actin and confers resistance to apoptosis.
- ACTC1 is involved in control of survival and migration in SHH medulloblastoma.

Importance of the study

Alpha-cardiac actin 1 (ACTC1), which is normally expressed in cardiac muscle, was found to be aberrantly expressed in medulloblastoma with higher levels expressed in WNT and SHH subtypes compared to Group 3 and Group 4. We demonstrate that ACTC1 incorporates into F-Actin and renders medulloblastoma cells resistant to apoptosis induced by mitotic inhibitor. Additionally, ACTC1 is involved in control of survival and migration in SHH cells with or without MYC overexpression. Our study shows for the first time that actin subunit composition alters stress fiber dynamics and contributes to tumorigenicity in medulloblastoma. This may have important implications in other primary brain cancers that demonstrate aberrant actin subunit expression such as glioblastoma.
Introduction

The actin cytoskeleton and actin binding proteins play an important role in mechanisms of apoptosis in both normal cells and neoplastic cells.\(^1\) Apoptosis can be stimulated by changes in actin polymerization;\(^2,3\) by alteration of apoptosis regulating actin binding protein interaction; or by disruption of tropomyosin interactions with F-actin.\(^4,6\) Alterations in actin subunit composition can also have effects on cell morphology, migration, and growth.\(^3,7\) It has been previously reported that sonic hedgehog (SHH) medulloblastoma cells are resistant to apoptosis when mitosis is inhibited with a specific inhibitor of Aurora kinase B; whereas Group 3 cells demonstrated marked apoptosis.\(^8-10\) The contribution of actin subunit expression to resistance to mitotic inhibition and migration in medulloblastoma cells is not known. We do know that F-actin plays a critical role in granule cell precursor migration\(^11\) and SHH medulloblastoma cells show gene expression profiles of developing granule cell precursors.\(^12\) Abnormal expression of the alpha cardiac actin (ACTC1) actin subunit has been documented in multiple cancer types including head and neck,\(^13\) urothelial,\(^14\) prostate,\(^15\) and glioblastoma.\(^16\) Furthermore, \(ACTC1\) expression is upregulated in multi-drug resistant breast cancer cells\(^17\) and in lung cancer cells that survive exposure to paclitaxel.\(^18\)

We hypothesized that ACTC1 expression in medulloblastoma could alter F-actin composition, thereby conferring an anti-apoptotic function in response to mitotic inhibition in medulloblastoma cells. Identification of resistance mechanisms in medulloblastoma subgroups is crucial since it is often difficult to treat at recurrence. The identification of pro-tumorigenic factors is essential for the development of more effective targeted therapy that will impair tumor growth, reduce migration and ultimately enhance patient survival with minimal toxic side effects.
MATERIALS AND METHODS

Details for each method are provided in Supplementary Methods.

**Cell culture**

SHH cells were (DAOY, UW426, UW228) and their derivatives were grown as adherent cultures. Group 3 cells (D425, D458) were grown as suspension cultures. The UW426-Myc and UW228-Myc cell lines were provided by Dr. Annie Huang, The Hospital for Sick Children, and were originally derived by retroviral transduction to express higher levels of Myc protein compared to parental.19 The D425 and D458 cells were kindly provided by Dr. Darell D. Bigner, Duke University.

**Gene expression profiling in cell culture**

Total RNA was isolated from UW426 and UW426-Myc cells exposed to vehicle control (0.01% v/v DMSO) or 100 nM AZD1152-HQPA Aurora kinase B inhibitor. The cRNAs generated were hybridized onto the Human HT-12 v4.0 BeadChip. The BeadChip was stained as per Illumina protocol and scanned on the iScan (Illumina). The data files were quantified in GenomeStudio Version 2011.1 (Illumina). To identify differentially expressed genes, linear models were fitted using the limma package (v3.12). The t-statistics derived from pre-specified contrasts were moderated by empirical Bayes shrinkage. The moderated t-statistics were subsequently used for hypothesis testing and for Gene Set Enrichment Analyses. Microarray data, methods, and analysis have been uploaded to GEO (GSE159385). Gene enrichment maps were generated as described by Merico et al.20,21 using conservative thresholds.
ACTC1 analysis in published Medulloblastoma Tumor datasets

Previously published gene expression data was analyzed using R2 platform (https://hgserver1.amc.nl/cgi-bin/r2/main.cgi, Academic Medical Center (AMC) Amsterdam, Netherlands)). ACTC1 was analyzed in multiple Medulloblastoma datasets (Heidelberg (PMID: 21911727), Pomeroy (PMID: 21098324), Cavalli/Remke (PMID: 28609654), Kool (PMID: 18769486) and Gilbertson (PMID: 22722829)) by subgroups that were assigned in the original publication. Additionally, a multi-cancer RNA sequencing dataset in R2 (Mixed Pediatric Pan Cancer - Pfister - 272 - FPKM - informp3) was analyzed to show ACTC1 in Medulloblastoma subgroups as well as across other CNS and non-CNS tumors. Neuroblastoma and T-ALL with a single datapoint were removed. ANOVA was used to calculate significance. Pearson correlation was used to determine the correlation between the expression and methylation of ACTC1 in the Cavalli/Remke dataset. Chromosome 15 copy number information was provided in the original publication for Cavalli/Remke and significance was calculated using ANOVA.

Quantitative RT-PCR

MB cell lines (UW426, UW426-Myc) were exposed to 0.01% DMSO or 100 nM AZD1152-HQPA for 48 hr (Selleckchem, Houston, TX, USA). Total RNA was isolated and reverse transcription was performed on 100 ng of total RNA using random hexamer priming. Quantitative PCR using ACTC1 specific primers was performed and data analysis was performed using the Livak method. The experiment was conducted in three biological replicates and four technical replicates per run for each cell line.
Creation of stable ACTC1-overexpression lines

SHH (UW426, UW426-Myc) cells were transfected plasmid DNA using Lipofectamine 2000 (ThermoFisher Scientific, Waltham, MA, USA). For D458 cells, transfections were performed via electroporation instead. 48 hours after transfection, selection was done for all cell lines using hygromycin. A single clone was then picked and expanded. The cells were transfected with either a human ACTC1 overexpression plasmid containing a C-terminal FLAG-tag (Ref.# HG10960-CF, Sino Biological, Wayne, PA, USA), or an empty vector control. Expression of the plasmid transgene was confirmed by Western blot (WB) analysis for the FLAG tag.

Transient shRNA ACTC1 knockdown

Transfection of shRNA expression plasmids were performed using Lipofectamine 2000 (ThermoFisher Scientific) reagent. The cells were transfected with either an ACTC1 knockdown DNA plasmid (GeneCopoeia, Rockville, MD, USA; Cat# HSH018117-CH1) or a scrambled control (GeneCopoeia; Cat# CSHCTR001-CH1). The knockdown was confirmed by WB analysis.

Western blot

Cell lysates were obtained using RIPA lysis buffer that was modified to contain a protease inhibitor. Lysates from control liver and heart were obtained from mouse samples flash frozen in liquid nitrogen. Proteins were separated by SDS-PAGE on 10% gels and transferred to nitrocellulose membranes using a semi-dry transfer apparatus (Bio-Rad, Hercules, CA, USA). The membranes were incubated with primary antibody at 4°C overnight followed by incubation with horseradish peroxidase conjugated secondary antibody for 1 hr. The following antibodies and dilutions were used: anti-ACTC1
(Cell Signaling Technologies, Danvers, MA, USA) 1:1000, anti-PARPi (Cell Signaling) 1:1000, anti-GAPDH (Cell Signaling) 1:1000, anti-Aurora Kinase B (Cell Signaling) 1:1000, anti-beta Actin (abcam, Cambridge, UK; ab115777) 1:1000. Secondary antibodies used were anti-rabbit or anti-mouse IgG conjugated to horseradish peroxidase (Cell Signaling) at a 1:2000 to 1:30,000 dilution.

**FACS cell viability assay**

Cells were transfected with either a mock transfection, scrambled control, or ACTC1 knockdown plasmid 4 days prior to exposure to 100 nM AZD1152-HQPA (Selleck Chemicals LLC, Houston, TX, USA) or 0.01% v/v DMSO (Sigma, St. Louis, MO, USA) for 48 h. Cells were then stained with LIVE/DEAD® Fixable Aqua Dead Cell Kit (ThermoFisher Scientific, Waltham, MA, USA) according to the manufacturer’s protocol and analyzed on flow cytometer. For each experimental condition, 20,000-150,000 cells were analyzed in three independent trials. An Attune NxT flow cytometer (ThermoFisher Scientific, Waltham, MA, USA) was used with Attune NxT software. Data analysis was conducted with FCS Express 7 (De Novo Software, Pasadena, CA, USA).

**Colony formation assay**

The cells were allowed to grow in standard culture media with the media being changed every 3 days. After 5 days (UW426-Myc) or 7 days (UW426) of growth, the colonies were stained with crystal violet solution. The number of colonies per well was counted using a dissecting microscope with a minimum of 50 cells necessary to constitute a colony.
**Wound closure assay**

Cells were cultured in a confluent monolayer. The media was changed to a serum-free media and a scratch was created in the well with a 200 μL pipette tip. Phase contrast imaging was done at 0, 12 and 24 hours, and the diameter of the wound was measured using ImageJ software. Seven different points were measured along the scratch.

**Immunofluorescence**

Cells were seeded on a sterile glass coverslip and fixed with 4% paraformaldehyde. After blocking, incubations were done with primary antibodies overnight at 4°C, and with secondary antibodies for 1 hour. The following antibodies and dilutions were used: anti-ACTC1 (Cell Signaling Technologies) 1:1000, anti-FLAG (Sigma Aldrich Canada Ltd.) 1:500. Secondary antibodies used were Alexa-Fluor 488 nm anti-rabbit IgG (Life Technologies Inc., Burlington, ON, Canada) or Alexa-Fluor 488 anti-mouse IgG (Life Technologies Inc.) at a 1:500 dilution. Each coverslip was incubated in 1U of Alexa-Fluor 647 nm fluorescent Phalloidin (ThermoFisher Scientific) and in Hoechst nuclear stain (ThermoFisher Scientific) at a 1:4000 dilution. The coverslips were then imaged using a Zeiss LSM700 confocal microscope with Zen Black 2010 acquisition software (Carl Zeiss, Oberkochen, Germany).

**Stress fiber analysis**

Cells were stained with Alexa-Fluor 647 nm fluorescent Phalloidin (ThermoFisher Scientific) and Hoechst 177 nuclear stain (ThermoFisher Scientific). Images were acquired using a Zeiss LSM700 confocal microscope with Zen Black 2010 acquisition software (Carl Zeiss, Oberkochen, Germany). Stress fibers were analyzed using FSegment (Universitätsmedizin Greifswald, Greifswald, Germany) software in MATLAB Runtime (MathWorks, USA) according to the protocol by Rogge et al.²³ and using
the parameters listed in Supplementary Table 1. Images of at least 150 cells were taken for each cell line in three independent trials. 1 pixel length was equivalent to 0.1 μm.

Statistics

Prism 8 (GraphPad Software, Inc., CA, USA) was used for statistical analysis. All measures are reported as mean +/- standard error. Means were compared by independent samples Student's t-test. A P-value less than 0.05 was selected for significance. Unless otherwise indicated, all experiments were averaged for 3 independent trials.

RESULTS

ACTC1 expression in medulloblastoma varies with subgroup type

Alpha-cardiac actin 1 (ACTC1) is an actin isoform that is primarily found in cardiac tissue. Here we demonstrate that ACTC1 is also expressed in medulloblastoma. Analysis of five independent medulloblastoma tumor gene expression data sets demonstrated higher ACTC1 mRNA expression in SHH and WNT tumors compared to Group 3 or Group 4 tumors (Figure 1A, Supplementary Figure 1 A-D). ACTC1 mRNA expression was correlated with methylation status at the ACTC1 promoter region (Fig. 1B, Supplementary Figure 1E) and was independent of chromosome 15 copy number (Supplementary Figure 1F). Among childhood tumours for which gene expression has been previously assessed by RNA sequencing, ACTC1 mRNA was found to be highly expressed in SHH and WNT medulloblastoma as well as Rhabdomyosarcoma (Figure 1C). Western blot analysis confirmed the expression of ACTC1 in SHH cells (DAOY, UW228, UW426), isogenic SHH cells with Myc overexpression (UW228-Myc, UW426-
Myc) and Group 3 cells (D425, D458) (Figure 1E). The highest levels of ACTC1 protein is found in the SHH subgroup cells, and the lowest in Group 3 cells. UW228 and UW426 cells express much higher levels of ACTC1 than the D425 and D458 cells (P<0.01, n=3; Figure 1D-E). Myc overexpression does not alter the ACTC1 levels in SHH medulloblastoma (Figure 1D-E).

**ACTC1 incorporates into F-actin in SHH medulloblastoma cells**

Immunofluorescence imaging was performed on UW426 cells to determine if ACTC1 expressed in SHH medulloblastoma cells incorporates into polymerized actin filaments. Extensive overlap was observed between the polymerized F-actin which was stained with Phalloidin and the immunolabeled ACTC1 protein (Figure 1G). To verify this finding, imaging was also performed on UW426 with stable ACTC1 overexpression (UW426-ACTC1 OE), but this time probing for the FLAG-tag of the overexpressed recombinant ACTC1 protein. Immunolabeled FLAG-tagged ACTC1 co-localized with Phalloidin stained F-actin in UW426-ACTC OE cells. (Supplementary Figure 2). The results confirmed the finding that ACTC1 is a constituent of polymerized actin filaments in SHH medulloblastoma.

**ACTC1 mRNA levels are upregulated during Aurora kinase B inhibition in SHH cells.**

RNA expression profiling was performed in UW426 and UW426-Myc cells with or without exposure to Aurora kinase B specific inhibitor AZD1152-HQPA for 48 hr. We sought to determine the transcriptomic changes that take place in medulloblastoma cells upon Myc overexpression, and to determine if Myc regulated gene networks are differentially expressed in response to Aurora B inhibition. Furthermore, transcriptional profiling would enable us to define the gene networks that are differentially expressed upon Aurora B inhibition in Myc overexpressing cells compared to wild-type cells. Myc overexpression in UW426 cells resulted in upregulation of gene networks involved in ribosomal
biosynthesis (Figure 2A). Aurora B inhibition did not alter the transcriptional networks upregulated by Myc overexpression (Figure 2B). DNA binding genes, negative regulators of DNA binding, and regulators of multicellular organism processes (RMOP) were significantly and uniquely downregulated in Myc overexpressing cells in response to Aurora kinase B inhibition (Figure 2B). Gene networks for hypoxia response and skeletal development were identified as shared response genes to Aurora B inhibition in wild-type versus Myc overexpressing UW426 cells. The differentially expressed genes in each network with > 1 or < -1 log fold change (LogFC) and P-value <0.05, which did not overlap with shared response genes were as follows: DNA binding (ID3, -1.74 LogFC) and RMOP (ACTC1, -1.95 LogFC; MYL9, - 1.08 LogFC; PTGDS, -1.01 LogFC). Quantitative RT-PCR analysis of ACTC1 mRNA expression in UW426 versus UW426-Myc cells after 48 hours of Aurora kinase B inhibition revealed an increase in ACTC1 mRNA abundance in UW426 cells compared to UW426-Myc, D458, and D425 cells (Figure 2C).

Overexpression of ACTC1 protects against apoptosis induced by Aurora kinase B inhibitor in Group 3 cells, but not in SHH cells overexpressing Myc.

Given the association between increased ACTC1 mRNA abundance and resistance to apoptosis induced by Aurora kinase B inhibition in SHH cells, we tested whether Group 3 cells which have low endogenous ACTC1 levels and undergo apoptosis in response to Aurora kinase B inhibition are protected from apoptosis when ACTC1 is overexpressed. ACTC1 was stably overexpressed in Group 3 cells (D458) (Figure 3A). Upon Aurora kinase B inhibition for 48 hours, the ACTC1 levels remained unchanged in control and ACTC1 overexpressing cells (Figure 3A). PARP1 cleavage to the 89 kDa fragment was observed with Aurora kinase B inhibition in the control cells (D458), but was significantly reduced in the ACTC1 overexpressing cells (D458-ACTC1 OE) [Figure 3B]. Aurora kinase B protein
levels were not altered by ACTC1 overexpression or by Aurora kinase B inhibition in D458 cells (Figure 3B, Supplementary Figure 3A). Furthermore, overexpression of ACTC1 did not alter the protein levels of β-actin (ACTB) (Supplementary Figure 3A). Overexpression of ACTC1 in D458 cells enhanced cell viability under control conditions (P<0.0001) and in the presence of Aurora kinase B inhibitor (P<0.0001, Figure 3C). Thus, ACTC1 overexpression enhanced cell viability and suppressed intrinsic apoptosis that is induced by Aurora kinase B inhibition in Group 3 cells. When ACTC1 was overexpressed in SHH cells with wild-type background (UW426), there was no induction of PARP1 cleavage and the resistance to PARP1 cleavage induced by Aurora kinase B inhibition was unchanged (Figure 3D). PARP1 cleavage was induced by Aurora kinase B inhibition in SHH cells with Myc overexpression (UW426-Myc). Increased expression of ACTC1 in UW426-Myc cells did not result in suppression of PARP1 cleavage induced by Aurora kinase B inhibition (Figure 3D). ACTB and Aurora kinase B protein levels were not altered by ACTC1 overexpression or Aurora kinase B inhibition in UW426 and UW426-Myc cells (Supplementary Figure 3B). Thus, increased Myc expression bypasses the anti-apoptotic effects of ACTC1 expression in SHH medulloblastoma subjected to mitotic inhibition with Aurora kinase B specific inhibitor AZD1152-HQPA.

**Reduction of ACTC1 levels in SHH medulloblastoma cells induces apoptosis and is synergistic with Aurora kinase B inhibition.**

Knockdown of ACTC1 in UW426 and UW426-Myc cells was accomplished by transfection with plasmid DNA encoding a targeting shRNA. Out of the three shRNA clones tested, clone A showed the greatest knockdown in ACTC1 level and was therefore selected to be used in further experiments (Supplementary Figure 4A). The scrambled control transfected cells (UW426-ACTC1 KD sCTL and UW426-Myc-ACTC1 KD sCTL) expressed similar levels of ACTC1 as the non-transfected cells (Supplementary Figure 4A). Furthermore, the protein levels of ACTB were not changed by ACTC1
knockdown (Supplementary Figure 4B-C). ACTC1 knockdown resulted in a marked increase in PARP1 cleavage indicating activation of the intrinsic apoptosis pathway in wild-type and Myc-overexpressing UW426 cells (Figure 4A-B). The percentage of non-viable cells increased when ACTC1 was knocked down in the wild-type and Myc-overexpressing background (Figure 4 C-D). Additionally, there was a greater percentage of non-viable cells when ACTC1 knockdown was combined with Aurora kinase B inhibition in both wild-type and Myc overexpressing backgrounds (Figure 4C-D).

ACTC1 levels control cell survival as measured by colony formation in SHH medulloblastoma

To assess the role of ACTC1 in promoting cell survival, a colony formation assay was performed in SHH cells (UW426) with wild-type or Myc overexpression background in which ACTC1 was knocked down by shRNA or overexpressed from a transgene. UW426 cells in which ACTC1 expression was decreased by shRNA knockdwon (UW426-ACTC1 KD) showed a 69% ± 4.0% reduction in colony formation compared to the scrambled control (P<0.000001, Figure 5A). ACTC1 overexpression in UW426 cells had the effect of increasing colony formation by 70% ± 8.2% relative to control (P<0.001, Figure 5A). ACTC1 knockdown impaired the colony formation capability of UW426 cells overexpressing Myc (UW426-Myc) by 51% ± 1.7% compared to scrambled control (P<0.0001, Figure 5B). However, overexpression of ACTC1 in UW426-Myc cells did not augment colony formation (Figure 5B).
ACTC1 knockdown inhibits migration in SHH cells.

Given the important role of actin in cell migration, we sought to determine the effect of ACTC1 levels on SHH cell migration. Phase contrast images were taken of a scratch assay at 0, 12 and 24-hour time points with a brightfield microscope and quantitative analysis of the scratch diameter was performed. We found that the same distance was traversed by UW426 and UW426-ACTC1 KD sCTL scrambled control. However, knocking down ACTC1 resulted in a marked reduction in the distance migrated in 24 hours (from a total distance of 525 ± 40 μm by UW426-ACTC1 KD sCTL compared to 210 ± 32 μm by UW426-ACTC1 KD, P<0.001, n=3; Figure 6A-B).

In order to assess if knocking down ACTC1 had a similar impact on migration in SHH cells with Myc overexpression, a scratch assay was also performed with UW426-Myc and its derivatives. Knocking down ACTC1 resulted in a marked reduction in the distance migrated over the 24-hour period (from 598 ± 85 μm by UW426-Myc-ACTC1 KD sCTL compared to 167 ± 33 μm by UW426-Myc-ACTC1 KD, P<0.05, n=3; Supplementary Figure 5). The overexpression of ACTC1 in both the UW426 and UW426-Myc cells, which already have elevated levels of ACTC1, did not significantly increase wound closure rate compared to their controls (Figure 6A-B, Supplementary Figure 5). These findings suggest an essential role for ACTC1 in permitting the migration of SHH cells.

ACTC1 expression levels modify stress fiber composition in Group 3 and SHH cells.

Stress fibers are contractile structures found in non-muscle cells that are composed of filamentous actin (F-actin) bundles, α-actinin, and and non-muscle myosin II. Alterations in stress fiber composition in a cell can be used to monitor changes in F-actin dynamics induced by drugs. To test if ACTC1 expression has an effect on F-actin dynamics, we measured the distributions of stress fiber length in Group 3 cells (D458) with and without overexpression of ACTC1 and in SHH (UW426) with knockdown or
overexpression of ACTC1. Notably, the histogram for stress fiber length showed a pronounced increase in
the frequency of the dominant fiber length at 3 μm when ACTC1 is overexpressed in Group 3 cells 
(P<0.01, Fig. 7A, left). This is in keeping with an increase in total fiber length in the D458-ACTC1 OE
line as compared to the wild-type (Fig. 7A, right). Conversely, in SHH cells which show high levels of
endogenous ACTC1 expression, no significant increase in fiber length was observed when ACTC1 was
overexpressed (Fig. 7B, left). However, when ACTC1 was knocked down, there was a decrease in the
longer fiber lengths, with a shift in the dominating fibre length from 6 to 3 μm (P<0.01). Interestingly,
there was an overall increase in total fiber length when ACTC1 was knocked down, which is explained by
an increase in the number of the shorter stress fibers compared to the control (P<0.01, Fig. 7B right).
Thus, ACTC1 expression levels can modulate the length of actin stress fibers in medulloblastoma.

Discussion

Actin is a key structural protein in all cells and consists of Beta cytoactin and Gamma cytoactin
subunits in non-muscle cells of the human body. The finding of alpha cardiac actin (ACTC1) expression
in medulloblastoma is intriguing given evidence for alternative actin isoform expression during
mammalian neurodevelopment and the known developmental gene expression profiles found in these
tumors. Furthermore, our findings of a negative correlation between ACTC1 promoter methylation and
ACTC1 mRNA expression in medulloblastoma tumors is consistent with a report of reduced ACTC1 gene
expression in early adult mouse skeletal muscle when increased methylation is found around the
transcriptional start site. The analysis of ACTC1 mRNA expression across multiple pediatric cancers
revealed an association between high ACTC1 expression and SHH pathway driven tumours (SHH
medulloblastoma, WNT medulloblastoma, and Rhabdomyosarcoma). SHH medulloblastoma and
Rhabdomyosarcoma are tumours that share Gli1 activation as a common tumorigenic pathway as
evidenced by formation of both of these tumours in Ptc +/- knockout mice.
constitutively active Smoothened mutant in mouse endothelial progenitor cells results in aberrant expression of myogenic specification factors, including MyoD, and formation of fusion-negative Rhabdomyosarcoma.\(^{30-33}\) Since MyoD is a known positive regulator of \(ACTC1\),\(^ {34}\) these observations raise the possibility that \(ACTC1\) expression may be downstream of SHH pathway activation in both SHH medulloblastoma and Rhabdomyosarcoma.

The actin cytoskeleton undergoes dramatic changes during cellular transformation resulting in altered morphology, migration ability, anchorage-independent growth, and resistance to apoptosis.\(^ {35,36}\) Proteins that bind to actin subunits regulate the function of actin.\(^ {37,38}\) For example, in neurons, actin dynamics regulated by actin binding proteins gelsolin and coflin are linked to apoptosis signaling.\(^ {39}\) Actin has been actively studied as a central component of both extrinsic and intrinsic apoptotic signaling pathways.\(^ {40,41}\) However, the role of subunit composition and how it may alter apoptosis signaling in cancer cells is not known.

Distinct functionality of actin subunits has been observed in transgenic mouse models.\(^ {2,37}\) Three different theories for the functional uniqueness of actin subunits have been proposed based on experimental observation: 1) differing affinity for actin binding proteins, 2) difference in subcellular localization, and 3) altered F-actin function depending on the subunit mix.\(^ {26}\) The functional consequences of alpha-cardiac actin expression in medulloblastoma were not known prior to the work presented here. However, some recent observations in glioblastoma had suggested an important role in sensitivity to radiation and chemotherapy as well as regulation of invasion. Ohtaki \textit{et al.}\(^ {26}\) reported that patients with glioblastoma expressing high levels of \(ACTC1\) have shorter survival despite receiving radiation and alkylating chemotherapy. Additionally, GBMs with high \(ACTC1\) expression show a more invasive phenotype on neuroimaging,\(^ {17}\) and in-vitro siRNA knockdown of \(ACTC1\) in the U87 GBM cell line inhibited migration.\(^ {17}\)
We have demonstrated that medulloblastoma cells from the SHH subgroup show high ACTC1 expression and are resistant to apoptosis induced by inhibition of Aurora kinase B. The small molecule inhibitor of Aurora kinase B, AZD1152, causes G2/M arrest in medulloblastoma cells.\textsuperscript{42} When ACTC1 is overexpressed in Group 3 cells, which normally show low ACTC1 expression, the apoptotic response to Aurora kinase B inhibition is blocked. Furthermore, knockdown of ACTC1 in SHH MB cells increases cell death and strongly inhibits colony formation, indicating a clear functional role for ACTC1 in SHH cell survival. Interestingly, ACTC1 overexpression does not protect against apoptosis induced by Aurora kinase B inhibition in SHH cells that have been engineered to overexpress Myc. This could be explained by gene expression changes in Myc overexpressing SHH cells, that lead to alterations in actin binding proteins needed for ACTC1 mediated anti-apoptotic signaling.\textsuperscript{11,23} Alternatively, the increased levels of ACTC1 may not be sufficient to overcome the apoptotic signaling triggered by increased entry into G2/M and excessive endoreplication in Myc-overexpressing cells.\textsuperscript{43}

The actin cytoskeleton and actin binding proteins play an important role in mechanisms of apoptosis in both normal cells and neoplastic cells.\textsuperscript{11} Apoptosis can be stimulated by changes in actin polymerization;\textsuperscript{2,3} by alteration of apoptosis regulating actin binding protein interaction; or by disruption of tropomyosin interactions with F-actin.\textsuperscript{4-6} Minor changes in the actin isoform forming F-actin can affect tropomyosin positioning and thereby influence interactions between F-actin and regulatory binding proteins.\textsuperscript{3,7} Our observations of a marked reduction of PARP1 cleavage to the 89 kDa fragment when ACTC1 is expressed in Group 3 cells undergoing mitotic inhibition is indicative of decreased Caspase3/7 protease activity. Since Caspase 3/7 activation is downstream of loss of mitochondrial membrane potential and mitochondrial permeability transition pore formation in the intrinsic mitochondrial apoptosis pathway\textsuperscript{44}, ACTC1 interactions with F-actin may be regulating mitochondrial membrane potential in medulloblastoma. For example, in the yeast \textit{Neurospora crassa}, the conductance of mitochondrial voltage-dependent anion channels is differentially regulated depending on the type of actin subunit expressed.\textsuperscript{37}
Understanding the mechanism for anti-apoptotic and migration permissive signaling mediated by ACTC1 in medulloblastoma is the key next step in elucidating the functional role of ACTC1. Immunofluorescence imaging of ACTC1 co-localizing with F-actin suggests the possibility for co-polymerization of alpha-cardiac actin with other actin subunits in medulloblastoma cells. Actin subunit co-polymerization has been previously reported *ex-vivo* for alpha-cardiac actin and skeletal muscle actin. Additionally, changes in subunit composition of actin co-polymers modifies filament stability. Filament stability may be important for cell survival during mitosis since cells undergoing mitosis experience increased plasma membrane tension and actively assemble a stiff actin cortex. Localization of ACTC1 to the actin cortex in prophase (**Supplementary Figure 7**) further suggests a possible functional role for ACTC1 in maintaining cell integrity during mitosis. Alterations in the stability of actin filaments required for plasma membrane protrusions that function in migration could explain the significant inhibition of migration caused by ACTC1 knockdown in SHH cells. Here we have demonstrated that alteration in ACTC1 levels in medulloblastoma cells has a direct effect on stress fiber formation. The effects are cell-type dependent with overexpression of ACTC1 in Group 3 medulloblastoma resulting in a greater number of stress fibers with no change in the fiber length distribution; while knockdown of ACTC1 in SHH cells results in a shift towards shorter stress fibers and coupled with an increase total fiber formation. Stress fiber composition is known to vary with cell type and the rigidity of the culture substrate. In keeping with the low adhesion phenotype of D458 cells in culture, we observe shorter stress fibers in this cell line compared to UW426 cells, which adhere robustly to plastic substrate. The differences in actin subunit expression and associated stress fiber formation between Group 3 cells and SHH cells could explain the greater tendency of Group 3 tumours to show CSF dissemination at diagnosis compared to SHH tumours. Further experiments to measure the polymerization properties and actin binding protein interactions of ACTC1 F-actin and as well as those of co-polymers with Beta and Gamma cytoactin subunits will help to define the cytoskeletal changes that occur when ACTC1 is expressed.
The finding that non-cell type specific expression of ACTC1 is present in SHH medulloblastoma cells and plays a functional role in survival and migration adds to our understanding of the dynamic changes which occur in actin composition in cancer cells and their contribution to neoplastic transformation. Ultimately, this altered actin subunit expression may be a reflection of the neurodevelopmental transcriptional program that is a hallmark of SHH medulloblastoma, which is composed of cells at varying stages of granule cell precursor differentiation. Interestingly, genes that contribute to structural heart development are linked to neurogenesis, including granule cell progenitor survival and differentiation. The findings presented here suggest that high ACTC1 levels may serve as a biomarker for resistance to apoptosis induced by Aurora kinase B inhibition. This may extend to resistance to other mitotic inhibitors and radiation therapy, which induces G2/M arrest similar to Aurora kinase B inhibition.

Further work to understand the transcriptional regulation of ACTC1 in SHH medulloblastoma may provide insight into combination therapy that could be ultimately useful for treatment of cancers with high ACTC1 expression. Such a strategy would involve downregulation of ACTC1 and activation of apoptosis signaling through a mitotic inhibitor or radiation. ACTC1 expression is suppressed through promoter methylation and activated by histone H3/H4 acetylation in non-cancer cells. If a similar mechanism is involved in regulating ACTC1 expression in cancer cells, a combination of a DNA demethylase inhibitor such as C35 or a HDAC1 activator such as a Exifone with Aurora kinase B inhibition or radiation could be a potent therapy.
Conclusion

We demonstrate for the first time that alpha-cardiac actin is expressed in medulloblastoma cells. We show that alpha-cardiac actin subunit expression alters stress fiber composition and promotes SHH medulloblastoma cell survival, resistance to apoptosis, and migration. Our findings have important implications in the understanding of SHH medulloblastoma tumorigenesis and provide a novel target for therapeutic development.
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Figure Legends:

**Figure 1:** A) ACTC1 mRNA expression across tumor subgroups in the Cavalli/Remke dataset (n=763) showing increased expression in SHH and WNT tumors compared to Group 3 and 4 tumours. B) Correlation between ACTC1 mRNA expression and promoter methylation in medulloblastoma tumors in the Cavalli/Remke dataset. C) ACTC1 mRNA expression by RNA sequencing across multiple pediatric tumors (Pfister data set). D) Western blot demonstrating ACTC1 and Myc expression across multiple human medulloblastoma cell lines. Positive control (+) is mouse cardiac muscle lysate and negative control (-) is mouse liver lysate. 10 ug total protein loaded. E) Relative ACTC1 protein abundance in relation to GAPDH across human medulloblastoma cell lines. Error bars indicate standard error of the mean. ** P<0.005. F) Immunofluorescence imaging of UW426 cells labeled with phalloidin (F-Actin, red) and anti-ACTC1 antibody (green). Merged image shows the red-green overlap as yellow. Scale bar is 50 μm.

**Figure 2:** A) GO gene-set enrichment results for Myc overexpressing UW426 cells versus isogenic wild-type control. Nodes represent gene-sets and edges represent GO defined relations (Is-a, Part-of, Regulates). Gene-sets that did not pass the conservative enrichment significance thresholds (P<0.001, FDR-Q = 0.05, Overlap coefficient = 0.5) are not shown. Nodes are coloured according to enrichment results: red represents enrichment in Myc overexpressing cells (i.e. up-regulation due to Myc overexpression), whereas blue represents enrichment in wild-type cells (i.e. down-regulation due to Myc overexpression). Colour intensity is proportional to enrichment significance. Since conservative
thresholds were used to select gene-sets, most of the node colours are intense, corresponding to highly significant gene-sets. The results indicate an overall increase in Ribosomal gene expression in Myc overexpressing cells. B) Enrichment map displaying the enriched gene-sets in AZD1152-treated UW426 cells with endogenous Myc expression (UW426) versus Myc overexpression (UW426-Myc). Cells were treated with 100 nM AZD1152-HQPA for 48 hr prior to mRNA isolation. Enrichments were mapped to the inner node area and to the node borders, respectively. Red represents enrichment in AZD1152-treated cells (i.e. up-regulation after AZD1152 treatment), whereas blue represents enrichment in untreated cells (i.e. down-regulation after AZD1152 treatment); colour intensity is proportional to enrichment significance. Conservative threshold parameters (P<0.001, FDR-Q = 0.05, Overlap coefficient = 0.5) were used and gene sets not meeting these thresholds are not displayed. Gene clusters that are differentially expressed uniquely in Myc overexpressing cells or shared by both wild-type and Myc overexpressing cells are outlined. Genes involved in regulation of multicellular organism process and DNA binding are uniquely downregulated in UW426 cells with Myc overexpression exposed to AZD1152 compared to UW426 cells. C) ACTC1 mRNA expression changes in UW426, UW426-Myc, D458 and D425 cells treated with 100 nM AZD1152-HQPA for 48 hr. Fold change relative to untreated cells. Bar graphs show mean fold change from 4 technical replicates per sample. Error bars are standard error of the mean, *P<0.0001.

**Figure 3:** A) ACTC1 and cleaved PARP1 protein expression in Group 3 cells (D458 and D458-ACTC1 OE) exposed to 100 nM AZD1152 [MI] or DMSO control [C] for 48 hours, 10 ug total protein loaded. Positive control (+) is mouse cardiac muscle lysate and negative control (-) is mouse liver lysate. Data represents mean of triplicates ± SEM, *P<0.05, **P<0.005, ns- P>0.05. B) Aurora kinase B (AurkB)
protein expression in Group 3 cells (D458 and D458-ACTC1 OE) exposed to 100 nM AZD1152 [MI] or DMSO control [C] for 48 hours, 10 ug total protein loaded. Positive control (+) is U251 glioblastoma cell line lysate. Data represents mean of triplicates ± SEM, ns- P>0.05. C) Percentage of cell death determined by FACS in D458 wild-type (WT) and ACTC1-overexpressing (OE) cells after 48 hr of exposure to 0.01% DMSO (-), or 100 nM AZD1152 (+). Graphs represent mean of three biological replicates ± SEM.*P<0.05, ***P<0.001, ****P<0.0001. D) ACTC1 and cleaved PARP1 protein expression in SHH cells (UW426, UW426-ACTC1 OE, UW426-Myc and UW426-Myc-ACTC1 OE) exposed to 100 nM AZD1152 [MI] or DMSO control [C] for 48 hours, 10 ug total protein loaded. Positive control for ACTC1 (+) is mouse cardiac muscle lysate and negative control (-) is mouse liver lysate loaded at 1 ug total protein. The PARP band in the positive control lane is a technical artefact. GAPDH is low in the control lysates due to the lower amount of protein loaded. Data represents mean of triplicates ± SEM, *P<0.05, ** P<0.005, ns- P>0.05.

Figure 4: A) Western blot for ACTC1 and cleaved PARP1 protein in UW426 and UW426 cells overexpressing ACTC1 (UW426 ACTC1) 48 hrs after transfection with shRNA plasmid. KD - knockdown, sCTL - scrambled control. 10 ug total protein loaded. Positive control (+) is mouse cardiac muscle lysate and negative control (-) is mouse liver lysate. Graphs represent mean of three biological replicates quantified by densitometry. *P<0.05, ns - P>0.05. B) ACTC1 and cleaved PARP1 protein expression in UW426 cells overexpressing Myc (UW426-Myc) and overexpressing both Myc and ACTC1 (UW426-Myc-ACTC1) 48 hrs after transfection with shRNA plasmid. KD - knockdown, sCTL - scrambled control. 10 ug total protein loaded. Positive control (+) is mouse cardiac muscle lysate and negative control (-) is mouse liver lysate. Graphs represent mean of three biological replicates quantified by densitometry. *P<0.05, ns - P>0.05. C) Percentage of cell death determined by FACS in UW426 cells
under mock (Mock), scrambled control (sCTL) and ACTC1 knockdown (KD) shRNA transfected conditions exposed to 48 hours of 100 nM AZD1152 [MI+] or 0.01% DMSO [MI-]. Graphs represent mean of three biological replicates ± SEM. *P<0.05, **P<0.01, ***P<0.001, ****P<0.0001, ns - P>0.05.

D) Percentage of cell death determined by FACS in UW426-Myc cells under mock (Mock), scrambled control (sCTL) and ACTC1 knockdown (KD) shRNA transfected conditions exposed to 48 hours of 100 nM AZD1152 [MI+] or 0.01% DMSO [MI-]. Graphs represent mean of three biological replicates ± SEM. **P<0.01, ***P<0.001, ****P<0.0001, ns - P>0.05.

Figure 5: A) [Left] Photographs of crystal violet stained colonies of UW426 cells with ACTC1 knockdown (KD) or overexpression (OE). sCTL - scrambled control shRNA. [Right] Quantification of colony formation in UW426 cells with ACTC1 knockdown or overexpression Graphs represent mean of four biological replicates ± SEM. *P<0.05, *** P< 0.001, ****P< 0.0001, ns - P>0.05.

B) [Left] Photographs of crystal violet stained colonies of UW426 cells overexpressing Myc (UW426-Myc) with ACTC1 knockdown (KD) or overexpression (OE). sCTL - scrambled control shRNA. [Right] Quantification of colony formation in UW426-Myc cells with ACTC1 knockdown or overexpression. Graphs represent mean of four biological replicates ± SEM. *P<0.05, *** P< 0.001, ****P< 0.0001, ns - P>0.05
**Figure 6:** A) Phase contrast images of a scratch migration assay of UW426 cells (10x magnification) with knockdown (KD) or overexpression (OE) of ACTC1. sCTL - scrambled control shRNA. Scale bar 400 μm. B) Graph demonstrates the wound distance over time in UW426 cells with knockdown (KD) or overexpression (OE) of ACTC1. Data represents mean of 7 technical and 3 biological replicates ± SEM, ***P<0.001. C) Distribution of stress fiber lengths in wild-type (WT), ACTC1 knockdown (ACTC1 KD), ACTC1 overexpressing (ACTC1 OE) UW426 cells. Counts were summed in bins of 3 μm. D) Total stress fiber length in wild-type (WT), ACTC1 knockdown (ACTC1 KD), ACTC1 overexpressing (ACTC1 OE) UW426 cells. E) Distribution of stress fiber lengths in wild-type (WT) and ACTC1 overexpressing (ACTC1 OE) D458 cells. Counts were summed in bins of 3 μm. F) Total stress fiber length in wild-type (WT) and ACTC1 overexpressing (ACTC1 OE) D458 cells. Graphs C to F represent a mean of three biological replicates ± SEM. *P<0.05, **P<0.01, ns - P>0.05.
A  

0 hrs

24 hrs

B

UW426 Migration

C

UW426 Stress Fiber Length Distribution

D

UW426 Total Fiber Length

E

D458 Stress Fiber Length Distribution

F

D458 Total Fiber Length