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RESEARCH COMMUNICATION

Pediatric glioma-associated KIAA1549:BRAF expression regulates neuroglial cell growth in a cell type-specific and mTOR-dependent manner

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Tandem duplications involving the BRAF kinase gene have recently been identified as the most frequent genetic alteration in sporadic pediatric glioma, creating a novel fusion protein [f-BRAF] with increased BRAF activity. To define the role of f-BRAF in gliomagenesis, we demonstrate that f-BRAF regulates neural stem cell (NSC), not astrocyte, proliferation and is sufficient to induce glioma-like lesions in mice. Moreover, f-BRAF-driven NSC proliferation results from tuberin/Rheb-mediated mammalian target of rapamycin (mTOR) hyperactivation, leading to S6-kinase-dependent degradation of p27. Collectively, these results establish mTOR pathway activation as a key growth regulatory mechanism common to both sporadic and familial low-grade gliomas in children.

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In the pediatric population, the most common solid tumors originate in the brain, where they can be associated with significant morbidity. Among the histological subtypes, low-grade pilocytic astrocytomas [PAs] are the most frequently encountered brain tumors [Central Brain Tumor Registry of the United States 2012]. PAs arise either sporadically or in the context of the Neurofibromatosis type 1 [NF1] inherited cancer disposition syndrome. Tumors arising in children with NF1 typically develop along the optic pathway, following biallelic inactivation of the NF1 tumor suppressor gene in neuroglial cells. The NF1 protein [neurofibromin] is a negative regulator of the RAS proto-oncogene such that neurofibromin loss in neuroglial protein (neurofibromin) is a negative regulator of the RAS tumor suppressor gene in neuroglial cells. The optic pathway, following biallelic inactivation of the NF1 gene, defines the role of this signature genetic alteration in neuroglial cell function and glioma formation. We demonstrate that f-BRAF regulates neuroglial cell growth in a cell type-specific manner via tuberous sclerosis complex (TSC)/Rheb activation of mTOR and is sufficient to induce glioma-like lesions in vivo.

Results and Discussion

The discovery of the signature KIAA1549:BRAF genetic alteration represents a major advance in our understanding of the molecular pathogenesis of sporadic PA. However, subsequent studies have raised important questions regarding the role of this potential driver mutation in neuroglial cell proliferation and PA tumorigenesis. Moreover, prior studies have examined the impact of wild-type or oncogenic BRAF [BRAFV600E] expression on astrocyte or astroglial progenitor function, rather than the signature PA-associated BRAF fusion protein. To determine whether f-BRAF expression is sufficient to confer a growth advantage to neuroglial progenitor cells, we expressed f-BRAF in cerebellar NSCs, since the majority of sporadic PAs arise in the cerebellum [Ohgaki and Kleihues 2005]. Primary NSC cultures generated from postnatal day 1 [P1] mouse cerebella expressed the Sox2 stem cell and Olig2 glial progenitor markers, but not glial fibrillary acidic protein [GFAP; astrocyte] [Supplemental Fig. S1A], and were capable of multilineage differentiation [Supplemental Fig. S1B]. Cerebellar NSCs were transduced with either empty pBABE-puro vector [pBABE] or f-BRAF. Since BRAF activates MAPK in a MEK-dependent manner, a constitutively active MEK mutant [MEKQ65E; caMEK] was included. Following caMEK or f-BRAF expression in cerebellar NSCs, there was increased MAPK activation [1.5-fold and twofold increased Thr202/Tyr204 phosphorylation, respectively] [Fig. 1A] and increased NSC proliferation, as assessed by secondary neurosphere diameters [28% and 46% increases, respectively], direct cell counting [1.5-fold increase] [Fig. 1B] and limiting dilution analysis [37 ± 7 pBABE-transduced vs. 2 ± 1 f-BRAF-transduced cells] [Supplemental Fig. S1C].

Oncogenic BRAF has been reported to induce activation of the mTOR pathway in melanoma, thyroid, and breast carcinomas [Bachhia et al. 2010; Duong et al. 2012; Faustino et al. 2012]. In addition, previous studies from our laboratory have demonstrated that the NF1 gene product neurofibromin regulates murine astrocyte, NSC, and glioma growth in an mTOR-dependent manner.
mTOR inhibition attenuated the f-BRAF-induced increases in S6 activation [Fig. 1E] and proliferation [Fig. 1F,G], similar to that observed following PD901 treatment. Finally, whereas MEK inhibition blocked f-BRAF-induced mTOR activation, rapamycin had no effect on f-BRAF-induced MAPK activation [Supplemental Fig. S1G]. Collectively, these results demonstrate that f-BRAF regulates NSC proliferation by activating the mTOR pathway in a MEK-dependent manner.

Several studies have shown that ectopic expression of either RAF or BRAF alone is not sufficient to induce gliomas in mice [Lyustikman et al. 2008; Robinson et al. 2010; Gronych et al. 2011], suggesting that BRAF may not be a driver mutation. However, BRAF expression was targeted to progenitor cells in the cerebral hemispheres rather than the more typical locations (cerebellum, optic pathway, and brainstem) where f-BRAF-containing PAs arise in children. To establish a causative role for the signature BRAF alteration in gliomagenesis, we show that f-BRAF expression resulted in a fivefold increase in cerebellar NSC soft agar colony formation [Supplemental Fig. S2A]. Next, mCherry-expressing cerebellar NSCs infected with control [pBABE] or f-BRAF were injected into the cerebella of 3-wk-old wild-type mice and analyzed at 2.5 and 6 mo post-transplantation. At 2.5 mo, we observed threefold increases in the numbers of Ki67+ and Iba1+ cells at the injection sites in mice transplanted with f-BRAF-expressing NSCs (n = 3) relative to control virus-infected NSCs (n = 3). In addition, there was only a modest increase in cellularity and GFAP immunoreactivity and no nuclear atypia or cells with piloid morphologies [Supplemental Fig. S2B,C]. However, at 6 mo post-injection, mice transplanted with f-BRAF-expressing NSCs (n = 5) harbored clear areas of increased cellularity near the injection sites, consistent with glioma-like lesions, including abnormal cell clusters with pale nuclei [Fig. 2A, H&E, inset], increased GFAP+ cells with piloid morphologies and elongated hair-like cytoplasmic processes [Fig. 2A, inset], increased numbers of proliferating [Ki67] cells [Fig. 2A,B], increased microglial [Iba1] infiltration [Fig. 2A,B], and increased numbers of endothelial cells (data not shown) relative to pBABE-containing NSC-injected control mice (n = 4). Whereas mCherry+ cells were identified at the injection sites from both pBABE- and f-BRAF-transduced NSC-transplanted mice [Fig. 2A], both NSC-derived astrocytes [GFAP and mCherry double-positive cells] [Supplemental Fig. S2E] and host-derived astrocytes [GFAP+ only] [Supplemental Fig. S2E] were detected in the f-BRAF-expressing lesions. Consistent with the in vitro results, there was also increased pMAPK and pS6 immunostaining in f-BRAF-expressing NSC-engrafted mice at both 2.5 and 6 mo of age [Fig. 2C; Supplemental Fig. S2D]. While no gross neurological deficits or changes in survival were observed, body weight was reduced by 28% in mice injected with f-BRAF-expressing NSCs.

The finding that f-BRAF expression in cerebellar NSCs generated glioma-like lesions in the cerebellum following engraftment likely reflects differences in the innate capabilities of these brain region-specific NSCs. In this regard, previous studies from our laboratory have revealed NSC regional specificity in response to f-BRAF expression and NF1 gene loss. Whereas lateral ventricle NSCs exhibit marginal or no increase in NSC proliferation following NF1 inactivation or f-BRAF expression, respectively, third ventricle and brainstem NSCs demonstrate increased proliferation and...
Fig. 2. f-BRAF induced glioma-like lesions in mice, and mTOR activation was observed in human PAs. (A) Injection sites [dotted lines] contain mCherry+pBABE- or f-BRAF-expressing cells. Abnormal cell clusters with pale nuclei [insets, H&E], increased number of GFAP-immunoreactive cells with piloid morphologies [insets], increased Ki67+ cells [arrows and inset; see also B, top], and increased Iba1+ cells [arrows and inset; see also B, bottom] were observed in f-BRAF-induced lesions. (C) Increased numbers of pMAPK+ and pS6+ cells [arrows] were found in f-BRAF-induced lesions. (D) Representative images show increased pMAPK and pS6 staining, along with increased GFAP and Sox2 immunoreactivity, in f-BRAF-containing human PAs. A normal human brain was used as a reference control. Error bars denote mean ± SD. *(P < 0.05; **P < 0.01. Bars, 50 μm.

BRAF regulation of neural growth

To determine whether sporadic human PAs likewise exhibit increased mTOR activation, we used a tissue microarray [TMA] containing 54 sporadic PAs. Consistent with the in vitro findings linking MAPK and mTOR activation in murine NSCs, a strong correlation was observed between mTOR activity (pS6 immunostaining) and MAPK activation [pMAPK immunostaining]. Since the BRAF genotyping was not available for all the cells contained in the TMA, we analyzed 14 sporadic PA cases in which the precise f-BRAF mutation was known and found that all f-BRAF-containing PA tumors examined had increased pMAPK and pS6 immunoreactivity relative to normal brains [Fig. 2D, Supplemental Fig. S2H]. In addition, these tumors exhibited diffuse GFAP expression and contained a smaller population of cells with Sox2 expression [Fig. 2D, Supplemental Fig. S2H].

To define the mechanism by which f-BRAF activates mTOR in cerebellar NSCs, we performed several experiments based on prior studies in other cell types, demonstrating that MAPK can activate mTOR by several mechanisms, including 90-kDa ribosomal protein S6 kinase (RSK) activation and inactivation of TSC protein function [Ma et al. 2005; Carriere et al. 2008]. First, we showed that f-BRAF expression does not increase RSK phosphorylation [Thr573], excluding RSK-mediated TORC1 activation as the mechanism underlying f-BRAF growth regulation [Fig. 3A]. Second, we observed a 2.6-fold increase in tuberin phosphorylation [Ser939] following f-BRAF expression in NSCs [Fig. 3A]. Similarly, there was a 2.9-fold and 1.6-fold increase in tuberin phosphorylation in NSCs after caMEK and BRAFV600 expression, respectively [Supplemental Fig. S3A,B]. Moreover, treatment of f-BRAF-expressing NSCs with PD901 reduced the observed increases in tuberin and S6 phosphorylation [Fig. 3B], demonstrating that f-BRAF-induced tuberin phosphorylation was MEK-dependent.

Since prior reports on BRAF growth regulation in astrocytes have yielded conflicting results, we transduced primary GFAP+ but Sox2/Olig2-negative astrocytes from P1–P2 mouse cerebella [Supplemental Fig. S3C] with pBABE, BRAFV600, or f-BRAF. Consistent with the known effects of BRAF on MAPK activation, both BRAFV600 and f-BRAF expression increased MAPK phosphorylation [6.5-fold and 4.4-fold, respectively], but neither BRAF molecule increased astrocyte proliferation [Fig. 3C]. The inability of BRAF to increase astrocyte proliferation was not the result of induced cellular senescence or programmed cell death [TUNEL labeling] [Supplemental Fig. S3D]. Similarly, this failure to increase cell proliferation was also observed in another astrocyte population [forebrain] despite increased MAPK activation [Supplemental Fig. S3E]. Moreover, the inability of BRAF to increase astrocyte proliferation was not dependent on mitogen [EGF, PDGF, or PACAP] availability [Supplemental Fig. S3F]. Consistent with the finding that BRAF-induced MAPK activation is dependent on MEK activity [Supplemental Fig. S3G], caMEK expression also did not increase primary astrocyte proliferation [Supplemental Fig. S3H]. Together, these results demonstrate that f-BRAF increases neuroglial cell proliferation in a cell type-specific manner unrelated to its ability to activate MEK/MAPK signaling. This observed cell type specificity underscores the critical cellular context in which cancer-associated driver mutations must occur in order to facilitate proliferation, transformation, and tumorigenesis. The differential response of NSCs and astrocytes to f-BRAF expression is consistent with previously reported roles for neuroglial progenitors in gliomagenesis [Taylor et al. 2005; Alcantara Llaguno et al. 2009; Liu et al. 2011].

To elucidate the mechanism underlying this cell type-specific effect, we compared the status of mTOR pathway activation following f-BRAF expression in astrocytes and NSCs. While f-BRAF expression in NSCs resulted in an 8.6-fold increase in S6 phosphorylation, there was no change in S6 phosphorylation following f-BRAF expression in astrocytes [Fig. 3D]. Furthermore, there was no change in tuberin phosphorylation following f-BRAF expression in astrocytes [Fig. 3E], indicating that MAPK, although activated by f-BRAF, was unable to phosphorylate and inactivate tuberin and thus could not activate mTOR in astrocytes.

We next sought to define the potential mechanism underlying differential tuberin phosphorylation in NSCs relative to astrocytes. Following hypotonic lysis fractionation, there was ~13-fold more tuberin in the membrane fraction of astrocytes compared with NSCs following normalizaton to total tuberin in each fraction [Fig. 3F]. These results support a model in which cell type-specific BRAF-mediated mTOR activation reflects the subcellular localization of tuberin. Polycystin-1, an integral membrane protein, suppresses mTOR activation by directly interacting with...
phosphorylation, or increase f-BRAF-expressing astrocyte type astrocytes (Supplemental Fig. S3J), increase tuberin down as tuberin did not alter tuberin localization in wild-type astrocytes (Supplemental Fig. S3J), increase tuberin phosphorylation, or increase f-BRAF-expressing astrocyte proliferation (Supplemental Fig. S3K), excluding polycystin-1 as the molecular etiology responsible for the cell type-specific effects of f-BRAF in astrocytes and NSCs. It is likely that other mechanisms account for the differential subcellular localization of tuberin relevant to its inactivation by phosphorylation.

Tuberin phosphorylation leads to loss of tuberin/hamartin complex inhibition of Rheb and subsequent increased Rheb-mediated mTOR activation (Castro et al. 2003; Inoki et al. 2003). We next sought to determine whether f-BRAF regulates mTOR activation in a tuberin/Rheb-dependent manner in NSCs. First, to confirm that TSC complex inactivation was responsible for f-BRAF-mediated mTOR activation, we expressed f-BRAF in wild-type or Tsc1−/− NSCs. Compared with their wild-type counterparts, empty vector-expressing Tsc1−/− NSCs exhibited a 2.5-fold increase in S6 phosphorylation as well as increased NSC proliferation (1.5-fold increase in cell number and 23% increase in neurosphere diameter). However, f-BRAF expression did not further increase S6 phosphorylation or proliferation in Tsc1−/− NSCs (Fig. 4A,B).

Second, ectopic Rheb expression in NSCs using a conditional Rheb-expressing mouse strain (Banerjee et al. 2011) resulted in a 2.1-fold increase in S6 activation (Supplemental Fig. S4A) and increased NSC proliferation (1.4-fold in cell number; 23% increase in neurosphere diameter) (Supplemental Fig. S4B). Third, to determine whether f-BRAF regulation of mTOR required Rheb activation, Rheb expression was reduced by shRNA-mediated knockdown. Attenuated Rheb expression [70% reduction] using two different shRNA constructs in f-BRAF-transduced NSCs reduced S6 phosphorylation and proliferation to wild-type levels with no effect on wild-type NSCs (Fig. 4C,D; Supplemental Fig. S4C). Together, these results establish TSC/Rheb-mediated mTOR activation as the mechanism underlying f-BRAF-mediated growth regulation of cerebellar NSCs.

Since mTOR can regulate cell growth in either an mTORC1-dependent (4E-BP1 and STAT3) or mTORC2-dependent (AKT) manner, we examined the activation state of each of these downstream signaling molecules. No changes in 4E-BP1 (Thr37/46), STAT3 (Ser727), or AKT (Ser473) were observed following f-BRAF expression in cerebellar NSCs (data not shown), mTOR can also regulate proliferation and cell cycle progression by increasing p27 phosphorylation and reducing total p27 expression (Hong et al. 2008; Dalvai et al. 2010). Following f-BRAF expression, there was increased p27 phosphorylation (Thr187) and decreased total p27 expression (Supplemental Fig. S4D). Moreover, both MEK (PD901) and mTOR (rapamycin) inhibition reduced p27 phosphorylation and restored p27 protein expression to wild-type levels (Supplemental Fig. S4D), indicating that f-BRAF controls NSC proliferation via mTOR-dependent regulation of p27 levels.

To elucidate the mechanism underlying f-BRAF regulation of p27 expression in cerebellar NSCs, we examined the mTOR downstream target S6K. Human S6K1 [hS6K1] expression increased NSC proliferation as well as p27 phosphorylation and decreased p27 protein levels (Supplemental Fig. S4E,F). In addition, S6K1 knockdown using two different shRNA constructs abrogated the f-BRAF-mediated increase in cell proliferation, S6 activation, and p27 phosphorylation and restored p27 expression to wild-type levels (Fig. 4E,F; Supplemental Fig. S4G). To assess whether p27 phosphorylation required cyclin-dependent kinase
mTOR pathway inhibitors for these common childhood cancers. It is worth noting that while both f-BRAF activation and NF1 loss lead to mTOR activation, the mechanism underlying mTOR regulation in each case is distinct. Whereas neurofibromin loss activates mTOR and leads to TORC2-dependent AKT activation, resulting in p27 phosphorylation and degradation [Lee et al. 2010], f-BRAF expression causes MEK-dependent tuberin inactivation and Rheb-directed TORC1/S6K-mediated p27 phosphorylation and degradation. These divergent mechanisms underscore the innate heterogeneity at the tissue (brain region), cellular (astrocyte/NSC), and molecular levels relevant to gliomagenesis.

Materials and methods

Mice

Tsc1+/−/lox mice [Uhlmann et al. 2002] and conditional Rheb-expressing transgenic mice [Banerjee et al. 2011] were generated as previously described. All mice were maintained on a C57BL/6 background and used in accordance with an approved Animal Studies protocol at Washington University.

Human tumor samples

Tumor specimens were obtained from the Tissue Procurement Core Facility at Washington University under an approved Institutional Review Board protocol.

Primary astrocyte and NSC cultures

Cerebellar hemispheres were microdissected from the brains of C57BL/6 mouse pups to establish primary astrocyte and NSC cultures, and proliferation assays were performed as previously described [Dasgupta and Gutmann 2005; Lee et al. 2010; Banerjee et al. 2011]. Wild-type and Tsc1+/− mice were generated from pups following infection with LacZ and Cre adenovirus, respectively. Retroviral and lentiviral (Supplemental Table 1) transduction was performed [Lee et al. 2010] for overexpression and knockdown studies, respectively. Expression of the f-BRAF transgene was verified by RT–PCR [Lee et al. 2012]. All in vitro experiments were performed at least three times using primary NSC and astrocyte cultures generated using pups from different litters and were independently transduced with the respective constructs.

Subcellular fractionation

Following hypotonic (10 mM HEPES at pH 7.9, 1.5 mM MgCl2, 10 mM KCl, protease inhibitors) lysis and protein degradation without affecting S6 activation. α-Tubulin served as an internal loading control. (H) Proposed mechanism of f-BRAF NSC growth regulation. (Veh) Vehicle. Error bars denote mean ± SD. (*) $P < 0.01$; (**) $P < 0.05$; (NS) not significant.
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a hamster syringe 3 mm below the dura mater using a stereotaxic apparatus. Mice were euthanized 2.5 and 6 mo later.

**Immunostaining**

Paraffin or frozen sections were processed (Dasgupta and Gutmann 2005) prior to staining with appropriate antibodies (Supplemental Table 2).

**Statistical analysis**

All in vitro experiments were analyzed using the Student's t-test. Statistical significance was set at P < 0.05.

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**References**


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