Beare-Stevenson cutis gyrata syndrome (BSS) is a human genetic disorder characterized by skin and skull abnormalities. BSS is caused by mutations in the FGF receptor 2 (FGFR2), but the molecular mechanisms that induce skin and skull abnormalities are unclear. We developed a mouse model of BSS harboring a FGFR2 Y394C mutation and identified p38 MAPK as an important signaling pathway mediating these abnormalities. 

FGF receptor (FGFR) mutations that underlie the genetic basis of BSS are present in more than 100 human skeletal syndromes (10–13). The FGFR2 IIIb (NP_000132.3) of the juxtamembrane domain (4, 14–19). Expression of FGFs and FGFRs, not only in keratinocytes during skin development and homeostasis (26–31), but also in osteoblasts during calvarial development (32–35). FGFR2 IIIb transgenic mice expressing a dominant-negative Fgfr2+/Y394C mice exhibited epidermal hyperplasia and premature closure of cranial sutures (craniosynostosis) due to abnormal cell proliferation and differentiation. We found ligand-independent phosphorylation of FGFR2 and activation of p38 signaling in mutant skin and calvarial tissues. Treating Fgfr2+/Y394C mice with a p38 kinase inhibitor attenuated skin abnormalities by reversing cell proliferation and differentiation to near normal levels. This study reveals the pleiotropic effects of the FGFR2 Y394C mutation evidenced by cutis gyrata, acanthosis nigricans, and craniosynostosis and provides a useful model for investigating the molecular mechanisms of skin and skull development. The demonstration of a pathogenic role for p38 activation may lead to the development of therapeutic strategies for BSS and related conditions, such as acanthosis nigricans or craniosynostosis.

Introduction

Beare-Stevenson cutis gyrata syndrome (BSS) (MIM #123709) is an autosomal dominant disorder characterized by both skin and skull abnormalities, including cutis gyrata, acanthosis nigricans (AN), craniosynostosis, craniofacial dysmorphism, including choanal atresia, a prominent umbilical stump, and anogenital anomalies (1–3). Patients can be born with respiratory distress and may die within 50 days after birth. Survivors have significant developmental delay (1, 4). Skin abnormalities such as cutis gyrata and AN are common characteristics of this genetic disease (3). Cutis gyrata is characterized by furrowed skin with a corrugated appearance. The skin may exhibit hyperplasia of connective tissue histologically (3). AN presents as a brown-to-black, poorly defined, velvety hyperpigmentation of the skin, with a prevalence of 7% in unselected populations (5–7). Histologic evaluation of AN is characterized by hyperkeratosis and papillomatosis, with a thinned epidermis overlying the papillae. Acanthosis is usually confined to the troughs of the epidermal papillae, and hyperpigmentation is not always present (8, 9). Craniosynostosis, a common isolated congenital disorder, is characterized by premature fusion of sutures and abnormal cranial vault shape. It can also be associated with midfacial hypoplasia as well as increased intracranial pressure. Craniosynostosis occurs in 1 in 2,500 newborns across all ethnicities and is

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manifested keratinocyte hyperproliferation with the onset of inflammation (30, 31). FGFR2 IIIc is expressed in preosteoblasts and osteoblasts in both endochondral and intramembranous ossification (37). These studies suggest that FGFR2 plays an important role in the regulation of both epidermal maintenance and bone development.

FGFs bind to the linker region between the extracellular immunoglobulin-like domains, IgII and IgIII, of FGFR2. When the receptor is dimerized and phosphorylated, it activates downstream signaling pathways to control the balance among different cellular activities, including migration, proliferation, differentiation, and survival of cells (32, 38–40). The 2 main FGFR2 isoforms, epithelial FGFR2 IIIb and mesenchymal FGFR2 IIIc, show distinct ligand specificity for different FGF ligands (41–43). To date, no functional studies on the BSS mutant FGFR2 have been performed. FGFR2 is known to signal by several downstream pathways including the MAPKs ERK1/2 and p38, PI3K/AKT, PLCγ pathways, and others, depending on cell type, tissue-specific expression, and developmental processes (22, 39, 44, 45). The MAPK pathways are critical in normal epidermal development (46). Although studies have suggested that alteration of FGFR2 and its downstream pathways contribute to craniosynostosis conditions (22, 33, 47–51), the mechanism by which cranial vault and skin abnormalities, especially cutis gyrata and acanthosis, are induced remains unclear.

To understand the cellular and molecular pathogenesis of the skin and skull malformations in BSS and to provide information relevant to possible molecular approaches for treatment of the skin and skull abnormalities, we created the first mouse model for BSS with cutis gyrata, acanthosis, skull synostosis, and other abnormalities. (A–D) Note skin furrows and hyperplasia in the mutant mice at P0 (A and B). With age, the skin phenotype becomes more obvious at P3 (C and D). (E–H) Surface reconstructions of HRCT images show midfacial hypoplasia of the mutant mice at P0. Note premature fusion of the coronal suture (E–H; black arrows), zygomatic suture (E–H; white arrows), and squamosal suture (E–H; red arrow) (Supplemental Table 1). (I and J) Alizarin red S and Alcian blue staining of the chest show abnormal bony fusion of sternum in mutant at P0. (K and L) Gross observations reveal protruding and enlarged umbilical stump in mutant (arrow). A, C, E, G, I, and K are from littermate controls. B, D, F, H, J, and L show the corresponding organs and tissues in mutant mice.

**Results**

*Fgr2*+/Y394C mice show epidural abnormalities similar to those in human BSS. We created a mouse model for BSS by designing a gene-targeting construct to site-specifically knock in the Fgr2 Y394C mutation (mouse FGFR2 IIIc protein NP_034337.2, analogous to the Y375C substitution in humans) into the mouse genome using a knockin approach. Our analysis showed that mutant mice exhibit epidermal hyperplasia and craniosynostosis of the calvarial sutures, which are accompanied by abnormal proliferation and differentiation of keratinocytes and osteoblasts, respectively. We found constitutive activation of the FGFR2 receptor and the activation of MAPK p38 signaling in the mutant epidermis and skull. Our data demonstrate that pharmacological blockade of p38 activation partially suppressed epidermal hyperplasia and, to a lesser degree, premature closure of the coronal suture in *Fgfr2*+/Y394C mice, correlating with attenuated hyperproliferation and normal expression of differentiation markers. Our findings provide opportunities for dissecting the molecular mechanisms of BSS and developing novel therapeutic strategies related to similar, more common conditions of acanthosis nigricans or craniosynostosis.
mutant Fgfr2 allele generates a number of transcripts similar to that of the WT allele in both skull and skin tissues (Supplemental Figure 1B). At early postnatal stage P0, the Fgfr2+/Y394C mice could be distinguished macroscopically from their littermate controls by skin furrowing and dome-shaped heads (Supplemental Figure 1C), and with age the mutants showed reduced growth and early death, usually within 2 weeks (Supplemental Figure 1, D–F). Necropsy of P0 Fgfr2+/Y394C mice revealed multiple malformations in the skin, skull, sternum, and umbilicus, which resemble those of the human condition (refs. 1–3 and Figure 1). Each of these different defects showed 100% penetrance in mutant mouse. The mutant skin displayed typical furrowing and thickening, similar to the cutis gyrata and acanthosis observed in BSS patients (Figure 1, A–D). High-resolution micro-CT scans of Fgfr2+/Y394C mice revealed midfacial hypoplasia, brachycephalic-shaped skull, and premature fusion of coronal and other sutures (Figure 1, E–H, and Supplemental Figure 2). Morphometric analysis of 3D craniofacial landmarks showed comparable statistically significant differences between Fgfr2+/Y394C and controls at P0 and at P8, with the dysmorphology becoming more pronounced by P8 (Supplemental Table 1 and Supplemental Figure 2). Other abnormalities included fusion of bones of the sternum (Figure 1, I and J) and an enlarged umbilical stump (Figure 1, K and L).

Detailed characterization of skin and skull abnormalities was performed, as these 2 phenotypes are the major features in BSS patients (3). Histologically, the skin of E16.5, P0, and P5 Fgfr2+/Y394C mice displayed cutis gyrata, epidermal acanthosis, and papillomatosis (Figure 2, A–D). The thickness of the epidermis was significantly different between the mutant and WT (+/Y394C [n = 12] versus +/+ [n = 12] at E16.5: 35.9 ± 11.6 μm versus 25.9 ± 1.6 μm, P < 0.006; +/Y394C [n = 12] versus +/+ [n = 12] at P0: 48.8 ± 13.9 μm versus 27.1 ± 3.3 μm, P < 0.000007; +/Y394C [n = 6] versus +/+ [n = 6] at P5: 44.6 ± 4.3 μm versus 26.7 ± 2.6 μm, P < 0.000005) (Figure 2, C, D).
Figure 3

Fgf2<sup>-/-Y394C</sup> mice show abnormal histology, proliferation, and differentiation in the skull. (A and B) H&E staining shows the abnormal development of the mutant coronal suture at E17.5, with presynostosis and osteoid deposition between the osteogenic fronts (arrows, osteogenic fronts; arrowhead, presynostosis/synostosis). (C and D) Immunohistochemical staining of BrdU shows decreased numbers and abnormal distribution of positive cells in mutants at E17.5. (E and F) ALP staining shows broad and expanded ALP expression into the coronal suture of the mutants at E17.5. (G and H) Immunohistochemical staining of osteonectin shows accelerated bone formation in mutants at E17.5. (I and J) In situ hybridization of osteonectin shows accelerated bone formation in mutants at E17.5, with presynostosis and osteoid deposition between the osteogenic fronts (arrows, osteogenic fronts; arrowhead, presynostosis). (K and L) TUNEL staining shows no obvious change of apoptosis in the mutant. A, C, E, G, I, and K are from littermate controls. B, D, F, H, J, and L show the corresponding organs and tissues in mutant mice. Scale bars: 50 μm.

D, and O, and Figure 3, A and B). The height measurements of the top and bottom of the troughs of the papillomatosis showed significant differences in mutants from E16.5 to P0 (E16.5: top, mean 46.8 ± 4.5 μm versus bottom, mean 25.1 ± 2.5 μm; P0: 59.8 ± 9.3 μm to 37.7 ± 7.0 μm); thus the averages of the overall heights were significantly different and displayed greater statistical variation as compared with those of the controls (+/Y394C [n = 12] versus +/+ [n = 12] at E16.5: 35.9 ± 11.6 μm versus 25.6 ± 1.6 μm; P = 9 × 10<sup>-13</sup>; +/Y394C [n = 12] versus +/+ [n = 12] at P0: 48.8 ± 13.9 μm to 27.1 ± 3.3 μm, P < 0.05; +/Y394C [n = 6] versus +/+ [n = 6] at P5: 53.3 ± 51.6 μm; P > 0.07). WT and mutant epidermis were stained with Masson-Fontana from P0 to P7, and only patchy melanin and dendritic melanocytes were present in the stratum basale. Most of the melanin and melanocytes were in the matrix of the hair follicles. No clear difference could be discerned between the 2 groups (Supplemental Figure 3, A and B).

The basal layer of the epidermis was particularly affected by proliferative changes as compared with that of the WT littermates. Quantitation of cell proliferation in the epidermis showed that Fgf2<sup>-/-Y394C</sup> mice showed a 2-fold increase in Ki67-positive cell numbers compared with controls (E16.5: +/Y394C [n = 4] versus +/+ [n = 4]: 107.7 ± 12.0 versus 62.8 ± 6.7, P < 0.01; P0: +/Y394C [n = 4] versus +/+ [n = 4]: 38.2 ± 5.7 versus 19.8 ± 3.5, P < 0.01) (Figure 2, E, F, and P, E16.5). The cell proliferation in the dermis was the same in the mutant and WT (+/Y394C [n = 6] versus +/+ [n = 6] at P0: 94 ± 13.9 versus 83 ± 5.2, P > 0.09).

Immunohistochemical analysis of epidermal differentiation markers revealed increased expression due to hyperplastic cells in each layer, including keratin 14 (K14) in the basal layer (Figure 2, G and H), K10 in the spinous layer (Figure 2, I and J), and loricrin in the granular as well as cornified layers (Figure 2, K and L) in mutant mice, but none of these markers was ectopically expressed beyond the layer of regular detection. There were slightly, but not significantly, fewer apoptotic cells in the epidermis of Fgf2<sup>-/-Y394C</sup> mice relative to controls as determined by quantitation of TUNEL staining (+/Y394C [n = 2] versus +/+ [n = 2]: 29.8 ± 7.4 versus 27.7 ± 5.8, P > 0.05) (Figure 2, M, N, and Q). We also examined the development of hair follicles and did not find obvious differences in developmental timing, numbers, distribution, or structure between mutants and controls (Supplemental Figure 3). Permeability assays suggested a normal epidermal barrier in mutant mice as compared with controls from E14.5 to E18.5 (data not shown).

Fgf2<sup>-/-Y394C</sup> mice have skull synostosis with abnormal proliferation and differentiation. The coronal suture of the skull was examined microscopically at different developmental stages. In Fgf2<sup>-/-Y394C</sup> mice at E17.5, the coronal sutures exhibited synostosis and presynostosis, a condition characterized by proximate osteogenic fronts, showing deposition of osteoid as compared with the clearly separated fronts in controls (Figure 3, A and B). Cell proliferation was analyzed by calculating the ratios of the number of BrdU-positive cells to total cells between and including the osteogenic fronts in mutants and littermate controls. At E17.5, the ratio was obviously decreased in mutants as compared with controls (+/Y394C [n = 3] versus +/+ [n = 3]: 0.05 ± 0.02 versus 0.13 ± 0.02, P = 0.004) (Figure 3, C and D). Differentiation of the coronal suture was studied...
from E17.5 to P0 by using alkaline phosphatase (ALP), Runx2, osteopontin, and osteonectin markers. Apparent differences in the expression of these markers were found between mutants and controls. In mutants, the coronal suture exhibited ALP expression in the region between the 2 osteogenic fronts, whereas there was no ALP expression in the same region of controls (Figure 3, E and F). The staining of Runx2 also showed increased expression in the area between the 2 osteogenic fronts in mutants as compared with controls (Figure 3, G and H). Expression of osteonectin showed more overlap of the osteogenic fronts (Figure 3, I and J). These data indicated increased osteogenic differentiation as well as abnormal bone growth at the coronal suture in Fgfr2+/Y394C mice. There was no obvious difference in apoptosis between mutants and controls at E17.5 and P0 (Figure 3, K and L). These results showed that the epidermal hyperplasia as well as premature fusion of the coronal sutures of Fgfr2+/Y394C mice are associated with aberrant proliferation and differentiation.

The Y394C mutation causes constitutive activation of FGFR2 in rat chondrosarcoma chondrocytes and keratinocytes of Fgfr2+/Y394C mice. To learn about the mechanisms underlying the FGFR2 Y394C mutation in BSS, we analyzed the effect of this mutation on the signaling pathways downstream of FGFR2, including ERK1/2, p38, JNK, and PI3K/AKT pathways further investigated the signaling pathways downstream of the FGFR2 receptor and FRS2, a major FGFR downstream mediator of receptor function. Our results showed constitutive activation of the receptor and the activation of the downstream pathway p38 MAPK in both the epidermis and skull. To further study the relationship between p38 signaling and the skin and skull abnormalities in the mutant mice, we inhibited p38 activity by intraperitoneally injecting either SB203580, a p38 kinase inhibitor (53), or a PBS-vehicle control into pregnant females from E12.5 to E18.5. We used SB203580 because it is a pyridinyl imidazole inhibitor that specifically inhibits p38 kinase activity in an ATP-competitive manner. It has been extensively investigated in targeting p38 signaling in preclinical mouse models with different conditions, including inflammatory disease, cancer, pulmonary disease, and bone disorders (52, 53). We examined the phosphorylation of p38 in the epidermis of offspring after drug administration at P0. The PBS-treated mutant epidermis showed increased phosphorylated p38 as compared with WT controls (Figure 4, B–D). None of the other pathways showed obvious changes of phosphorylation in both tissues; however, we noted phosphorylated ERK1/2 was increased in mutant skull tissues (Supplemental Figure 5). These results indicate that the FGFR2 Y394C mutation contributes to the constitutive activation of the receptor and the activation of the downstream pathway p38 MAPK in both the epidermis and skull. In utero systemic p38 inhibition attenuates epidermal and skull abnormalities as well as abnormal proliferation and differentiation in Fgfr2+/Y394C mice. To further study the relationship between p38 signaling and the skin and skull abnormalities in the mutant mice, we inhibited p38 activity in vivo by intraperitoneally injecting either SB203580, p38 kinase inhibitor (53), or a PBS-vehicle control into pregnant females from E12.5 to E18.5. We used SB203580 because it is a pyridinyl imidazole inhibitor that specifically inhibits p38 kinase activity in an ATP-competitive manner. It has been extensively investigated in targeting p38 signaling in preclinical mouse models with different conditions, including inflammatory disease, cancer, pulmonary disease, and bone disorders (53, 54). We examined the phosphorylation of p38 in the epidermis of offspring after drug administration at P0. The PBS-treated mutant epidermis had high levels of phosphorylated p38, which was dramatically reduced in mutants treated with SB203580, indicating that this drug inhibited p38 activity successfully in vivo (Figure 5, A and B). Macroscopically, the skin showed slight improvement in the SB203580-treated mutant mice compared with the PBS-treated and untreated mutants. Histological examination of PBS-treated Fgfr2+/Y394C mice revealed typical skin phenotypes with epidermal hyperplasia and thickening at P0 (+/Y394C [n = 3] versus +/+ [n = 3]: 43.9 ± 1.7 μm versus 31.0 ± 0.7 μm, P < 0.01), whereas mutant mice treated with SB203580, showed that 59% of the microscopic views of the epidermis (31 out of 52 serial histological views of 9 mutant pups from 3 different litters) were histologically normal and the hyperplasia-driven thickening was decreased as compared with those of the PBS-treated mice (+/Y394C [n = 3] versus +/+ [n = 3]: 28.3 ± 2.4 μm versus 26.8 ± 0.7 μm, P > 0.05) (Figure 5, C–F, and W).
We further analyzed the skin of PBS- or SB203580-treated pups for Ki67 staining and expression of differentiation markers. Upon PBS treatment, Fgfr2\(^{+/Y394C}\) mice displayed 1.9 times as many proliferating cells in the basal layer as littermate controls (+/Y394C \(n=4\) versus +/+ \(n=4\): 76.3 ± 13.6 versus 47.0 ± 6.7, \(P<0.05\)) (Figure 5, G, H, and X) as well as increased expression of K14 in the basal layer (Figure 5, K, L, S, and T) and K10 in the spinous layer (Figure 5, O, P, S, and T), similar to the results without treatment. In contrast, upon SB203580 treatment, Fgfr2\(^{+/Y394C}\) mutant skin displayed only 1.1 times as many proliferating cells in the basal layer compared with controls (B). Lanes were run on the same gel, but were non-contiguous. (C–F) H&E staining shows epidermal hyperplasia in Fgfr2\(^{+/Y394C}\) mice treated with PBS (C and D), and SB203580 treatment rescued skin phenotype in mutants (E and F). (G–J) Ki67 staining shows increased number of Ki67-positive cells in the basal layer of PBS-treated mutants (G and H); hyperproliferation in the mutant epidermis was reduced after SB203580 treatment (I and J). (K–V) Staining of differentiation markers of epidermis shows increased expression of the markers in the basal (K14, K, L, S, and T) and spinous layers (K10, O, P, S, and T) of PBS-treated mutant mice; SB203580 treatment reduced the expression of these markers (K14, M, N, U, and V; K10, Q, R, U, and V). Scale bars: 50 μm. (W and X) Bar graph shows that the epidermal thickening (W) and the number of Ki67-positive cells (X) of mutants after the SB203580 treatment compared with those of the vehicle-treated mice. Results are presented as mean ± SEM. **\(P<0.01\).
with WT controls (+/+Y394C [n = 4] versus +/+ [n = 4]; 46.3 ± 6.5 versus 43.1 ± 3.8, P > 0.05) (Figure 5, I, J, and X). The expression of K14 and K10 also decreased to a level and distribution similar to that of the WT controls (Figure 5, M, N, U, and V [K14]; Figure 5, Q, R, U, and V [K10]).

Initial qualitative examination of the specimens showed that about half of the Fgfr2+/-Y394C mice (5 in 9 pups from 3 litters) treated with SB203580 had a relatively normal head shape at P0 (Supplemental Figure 6, A–C), but subsequent landmark-based quantification of skull morphology did not show significant differences between treated and untreated mutants (data not shown). The histological evaluation at comparable levels of the skull did not reveal obvious improvement in most of the mutant mice with drug treatment, as compared with the fused sutures of untreated or PBS-treated mutants. Only a small portion of drug-treated mutant pups (20% of mutants from 2 litters) showed slight improvement and increased patency in coronal sutures. These findings were further confirmed by ALP staining (Supplemental Figure 6, D–I). We also examined the phosphorylation of p38 in the skull of pups after drug administration at P0 by immunohistochemical staining. Our results showed that the treatment of SB203580 reduced the level of phosphorylated p38.
in coronal sutures of mutants compared with those untreated mutants, indicating that the drug inhibited p38 activity successfully in the skull (Supplemental Figure 6, J–L). Although systemic administration showed varying results, these data indicated that this SB203580 dosage treatment did ameliorate skin abnormalities and had certain cellular, but not appreciable, morphologic effect on skull abnormalities in the Fgfr2\(^{+/Y394C}\) mice.

Previous studies have shown that ERK1/2 and p38 are important pathways that may mediate the premature closure of sutures in Apert syndrome Fgfr2\(^{+/S252W}\) and Fgfr2\(^{+/P253R}\) mice (47, 49, 50). We treated our BSS Fgfr2\(^{+/S252W}\) mice with MEK/ERK inhibitor U0126 at previously published dosages (47), but the inhibition did not reveal obvious rescue of the skin and skull phenotypes (data not shown). We also treated Apert Fgfr2\(^{S252W/}\) mice on the same C57BL/6J background and same dosage of SB203580 as our BSS mice to target the p38 kinase with no obvious improvement in skull morphology (data not shown). These results also suggested that p38 modulation is more effective for treatment of skin phenotypes and to a lesser degree for cranial vault phenotypes in these craniosynostosis mouse models. The SB203580 treatment did not improve the survival rate of Fgfr2\(^{+/Y394C}\) mice, suggesting that other or multiple mutations cause the neonatal lethality and that refined or alternative treatment strategies are required.

Topical p38 inhibition attenuates skin abnormalities as well as abnormal epidermal proliferation and differentiation of Fgfr2\(^{+/Y394C}\) mice. To use a more direct approach and avoid undue systemic effects, we used topical treatment to downregulate p38 kinase specifically in the skin. The administration of SB203580 obviously repressed the epidermal hyperplasia and thickening in the mutants (+/+Y394C \([n = 3]\) versus +/+ [ \(n = 3\)]: 24.6 ± 0.3 \(\mu\)m versus 27.8 ± 1.5 \(\mu\)m, \(P < 0.05\)) compared with those of the vehicle-treated mice (+/+Y394C [ \(n = 3\) ] versus +/+ [ \(n = 3\) ]: 40.3 ± 3.7 \(\mu\)m versus 26.6 ± 1.3 \(\mu\)m, \(P < 0.01\) (Figure 6, A–D, and U). In drug-treated Fgfr2\(^{+/Y394C}\) mice, the number of Ki67-positive cells (+/+Y394C [ \(n = 4\) ] versus +/+ [ \(n = 4\) ]: 50.5 ± 5.0 versus 50.3 ± 4.7, \(P > 0.05\) ) (Figure 6, E–H, and V) and expression of K14 and K10 (Figure 6, I–T) were similar to those of WT littermate controls, as opposed to those in vehicle-treated control mice (+/+Y394C [ \(n = 4\) ] versus +/+ [ \(n = 4\) ]: 79.2 ± 17.2 versus 54.7 ± 7.6, \(P < 0.01\) ). Thus, p38 signaling exerts an important influence in skin development in Fgfr2\(^{+/Y394C}\) mice.

Discussion

Mice carrying the heterozygous Y394C mutation of the FGFR2 gene serve as an important mouse model showing both skin and skeletal defects. In our in vivo mouse model, we observed consistent skin furrowing/corrugation and hyperplasia as well as synostosis of the coronal sutures. High penetration of affected organs in this mouse model indicates that FGFR2 and its downstream pathways are crucial for the pleiotropic effects on skin and skull. The skin lesions observed in our Fgfr2\(^{+/Y394C}\) mouse model are consistent with human cutis gyrata and AN. The epidermis showed epidermal acanthosis and papillomatosis, as has been described for the human counterpart. The absence of increased pigmentation is not surprising, as that is not always present with AN, and other mice reported with acanthosis in the context of diseases such as benign tumors or psoriasis also do not have hyperpigmentation (55, 56), possibly due to strain or species differences. In the different layers of the epidermis, the Fgfr2\(^{+/Y394C}\) mutant mice had an increased number of proliferating cells in the basal layer and abnormal differentiation. Our studies and the work of others have provided evidence that FGFR signaling is also involved in the proliferation and differentiation of calvarial osteoblasts from patients and sutural development in Fgfr2\(^{+/Y394C}\) and other craniosynostosis mouse models (22, 33, 35, 48, 49). Collectively, these findings demonstrate that FGFR2 signaling regulates the disturbances of proliferation and differentiation in both the developing skin and skeleton.

Besides BSS, cutaneous phenotypes have been found in other congenital skeletal disorders caused by FGFR mutations. For example, there is AN in Crouzonodermoskeletal syndrome caused by the FGFR3 A391E mutation, severe achondroplasia with developmental delay caused by the FGFR3 K650M mutation, and thanatophoric dysplasia caused by the FGFR2 R248C mutation (57–61). Acne is a skin feature of Apert syndrome caused by the FGFR2 S252W or P253R mutation (23, 62). The mechanism by which only certain mutations of the FGFRs lead to cutaneous features is not well understood, but it is clear that FGFRs and their receptors are involved in many early developmental processes (63) and can have effects on many tissue types (35, 64–66). It is possible that certain FGFR mutations differentially activate certain downstream signaling pathways, whose effects are targeted at specific tissues or organs. It is also possible that there is a threshold effect and that skin abnormalities will occur if there is activation of the FGFR signaling caused by different mechanisms reaching the threshold.

In our BSS mouse model, the phenotype of BSS is consistent with the effect of the Y394C mutation on both FGFR isoforms. FGFR2 IIIb is mainly expressed in epithelial-derived skin tissue; FGFR2 IIIc is primarily expressed in mesenchymal-derived calvarial tissue (37, 41–43). The Y394C mutation involves the gain of a cysteine in the juxtamembrane domain of FGFR2. It has been shown by molecular modeling that the unpaired cysteines promote intermolecular disulfide bond formation, leading to dimerization of the mutant receptors and ligand-independent or constitutive FGFR activation (67, 68). Our results, which demonstrate that FGFR2 is constitutively activated in RCS chondrocytes transfected with the human FGFR2 Y394C mutation as well as in epidermal keratinocytes of Fgfr2\(^{+/Y394C}\) mice, are consistent with this model. Interestingly, constitutive activation of other FGFR2 or FGFR3 mutations located in different protein domains causes both skeletal and skin phenotypes, including FGFR3 A391E mutation in the transmembrane domain, K650M mutation in the tyrosine kinase domain, and R248C mutation in the extracellular domain (61, 69), suggesting that the mechanisms of constitutive activation converge to downstream signaling molecules, causing similar skin or skull phenotypes in these conditions.

We have provided evidence that constitutively activated FGFR2 controls abnormal skin and skull development by regulating p38 MAPK. We found that phosphorylated p38 signaling is increased in both the epidermis and calvarial tissues of Fgfr2\(^{+/Y394C}\) mice. It is known that p38 is one of the critical components within the downstream signaling pathways of different tyrosine kinase receptors, which are associated with skin development and disorders, including FGFR, EGFR, and IGF1 receptor (IGF1R) (70). Also, p38 signaling is involved in skin homeostasis or epidermal differentiation (71) and plays an important role in osteoblast differentiation (72, 73). Activation of the p38 pathway has been associated with craniosynostosis in Apert syndrome mice with the FGFR2 P253R mutation (49). Other intracellular signaling pathways important to skin hyperplasia or craniosynostosis include ERK and PKC\(\alpha\) pathways in craniosynostosis of Apert syndrome Fgfr2\(^{+/S252W}\) mice and osteoblasts of patients with Apert syndrome (33, 47, 49–51) as well as
ERK and AKT pathways in keratinocytes of human benign or malignant AN (74–76). Therefore, although we provide strong evidence for the role of the p38 pathway in skin and skeleton development, other parallel or downstream pathways may modulate the effects.

p38 Signaling is the first intracellular pathway involved in skin abnormalities related to craniosynostosis that has been demonstrated in an in vivo animal model. We further confirmed our findings by showing obvious suppression of the skin anomalies by selective inhibition of p38 signaling systemically. Our initial assessments suggest that the p38 inhibition ameliorates the skull abnormalities in BSS to a lesser degree. Because p38 plays a crucial role in the signaling cascades of diverse cellular processes, systemic p38 inhibition could result in undesired pharmacological activities in different organs, as suggested by previous studies (70). Thus, we performed topical administration on the skin and found obvious improvement of skin abnormalities. The effect of p38 inhibition on skin abnormalities in our mice suggests that FGFR2-mediated p38 activation is an important molecular pathogenesis of the skin phenotype in BSS and that the p38 pathway could be a potential pharmacologic target for the treatment of the skin abnormalities.

Interestingly, we saw activation of another important MAPK pathway, ERK1/2, in the skull of Fgfr2+/Y394C mice, whereas the activation of this pathway was not striking in the skin of Fgfr2+/Y394C mice. Further, the ERK1/2 pathway was not inhibited at the drug dosage used and did not contribute to rescuing skin and skull phenotypes. Similarly, the treatment of Apert syndrome Fgfr2+/S252W mice with a p38 inhibitor did not show obvious improvement of the craniosynostosis. These results may reflect the different functional effects of the FGFR2 Y394C and Fgfr2 S252W mutations on the receptors and downstream signaling pathways in different tissues. In our BSS mouse model, we speculate that FGFR2 Y394C mutation causes differential activation of the downstream intracellular pathways of FGFR2 in the skin and skull, with greater p38 and less ERK1/2 activation in keratinoctyes and greater ERK1/2 and less p38 activation in osteoblasts.

We now know that these different FGFR2 mutations result in different mechanisms. The FGFR2 receptor with the Y394C mutation is constitutively activated, triggering abnormal p38 signaling pathways without the presence of ligand, whereas in Apert syndrome, the receptors with S252W and P253R mutations and ERK1/2 pathways are activated in a ligand-dependent manner (77, 78). Also, p38 signaling appears to have a stronger effect on skin hyperplasia than ERK1/2 signaling, which is important in the etiology of FGFR-related craniosynostosis. Thus, inhibition with p38 or other components of the FGFR network may need to be tailored to optimize the therapeutic effects on diverse organs such as the treatment of skin and skull anomalies in BSS and potentially other common conditions of skin hyperplasia or craniosynostosis.

Methods

Generation of targeting construct and mutant allele genotyping. See Supplemental Methods.

Skeletal staining and histological analysis. Skeletal staining with Alizarin red S and Alcian blue was performed according to published methods (79). Histological sections (5 μm) were prepared from selected tissues that had been fixed in 4% paraformaldehyde for 24–48 hours and embedded in paraffin. Sections were stained with HE and histology. We studied the coronal sutures by sectioning the head with a sagittal plane using the anterior-posterior axis from the nose to vertebra. The sections analyzed were selected from the middle third portion of the unilateral coronal suture.

Epidermal and dermal thickness measurements and melanin staining. Epidermal thickness was assessed using 2 methods. Initially, ImageJ (http://rsb.info.nih.gov/ij/index.html) analysis of H&E-stained sections was used to quantify epidermal thickness. In each image, 5 different areas were analyzed, and these measurements were repeated on 2 to 4 different pictures for each mouse. Results were obtained from control mice at P0 (n = 3), treated mice at P0 (n = 2), and both control and treated mice at P5 (n = 2 each). Epidermal thickness measurement was further validated using an ocular micrometer in a Nikon Labophot 2 microscope by 2 observers (X. Zhou and R. Phelps). From 6 to 12 measurements were taken from the top of the stratum granulosum to the stratum basale. Dermal thickness measurements were taken at the same time at P0 and P5 from the base of the stratum basale to the top of the panniculus carnosus. Melanin was stained using the Masson-Fontana method. The areas chosen for study of the epidermis and melanin were from the upper dorsal back in all slides.

High-resolution micro-CT image acquisition and landmark data analysis. P0 and P8 mice were sacrificed and fixed in 4% paraformaldehyde. High-resolution microcomputed tomography (HRCT) images of all skulls were acquired at the Center for Quantitative X-Ray Imaging at Pennsylvania State University using an OMNI-X Universal HD600 Industrial X-ray CT system (Bio-Imaging Research Inc). Images of 21 P0 mice (n = 12+/Y394C; 9+/+) and 24 P8 mice (10+/Y394C; 14+/+) were acquired at 16- to 21-μm slice thickness and 15- to 20-μm pixel resolution. A standard minimum threshold of 92 mg/cm2 partial density of hydroxyapatite was used to create bone surfaces for morphometric analysis. The 3D coordinates of 40 biologically relevant cranial landmarks at P0 and 39 landmarks at P8 were recorded for all of the mice using AVIZO (Visualization Sciences Group). Previous analyses have demonstrated the accuracy and precision of this data collection method for CT scans (80, 81).

Differences in craniofacial shape between Fgfr2+/Y394C and nonmutant littermates were assessed separately for P8 and P0 mice using Euclidian Distance Matrix Analysis (EDMA) (82). Linear distances between all unique pairs of landmarks were calculated and were scaled to the geometric mean of all linear distances to account for differences in scale; the ratios between average linear distances for mutants and unaffected littermates were produced. The null hypothesis of similarity in form between mutant and nonmutant crania was evaluated for each linear distance by bootstrap-based 90% CIs, enabling localization of differences to specific anatomic loci.

Immunohistochemical, TUNEL, and ALP assays. Immunohistochemical assays were performed using the VECTOR M.O.M. Immunodetection Kit (Vector Laboratories) for BrdU (Sigma-Aldrich), Ki67 (Vector Laboratories), K14, K10, and loricrin (Novocastra). For BrdU labeling, pregnant female mice were injected with a 10 mg/ml solution of BrdU (Sigma-Aldrich) at 100 μg/g body weight 2 hours before sacrifice. The sections were counterstained with hematoxylin. Cell proliferation of the skin was analyzed by counting the number of Ki67-positive cells in a defined area at the back of the neck where AN occurs more severely in both humans (5) and, according to our observations, in mice. Cell proliferation of the skull was analyzed by calculating the ratios of the number of BrdU-positive cells to total cells between and including the osteogenic fronts. The counts for Ki67- and BrdU-positive cells in mutant and WT embryos were compared by using the t test. The TUNEL assay was done using the In Situ Cell Death Detection Kit, POD (Roche Applied Science) to detect apoptotic cell death by light microscopy. ALP staining was carried out as described (48). For each staining, at least 2 litters of mice were examined.

In situ hybridization. In situ hybridization was performed on sections as described by Wilkinson (83) with modifications. Mouse osteonectin cDNA fragments were each cloned into the pCRII-TOPO Vector (Invitrogen). The plasmids were linearized and antisense and sense single-stranded RNA probes were generated using T7, T3, or SP6 RNA polymerase with digoxigenin (Roche). At least 2 litters of mice were examined.
**In vitro FGFR2 functional study.** RCS chondrocytes were transfected using FuGENE 6 reagent according to the manufacturer’s protocol (Roche Applied Science). Vector-expressing WT FGFR2 was generated by cloning full-length human FGFR2 ORF into the pcDNA3.1 vector, according to the manufacturer’s protocol (Invitrogen). Y375C FGFR2 mutant was created by site-directed mutagenesis, according to manufacturer’s protocol (QuickChange II Site-Directed Mutagenesis Kit; Agilent Technologies). Empty plasmid was pcDNA3.1. At 36 hours after transfection, cells were lysed in ice-cold lysis buffer supplemented with proteinase inhibitors. Protein samples were resolved by SDS-PAGE, transferred onto a PVDF membrane (Bio-Rad), and visualized by luminescence (Amersham). The following antibodies were used: P-FGFR (Y653/Y654), P-FRS2 (Y196/Y436) (Cell Signaling Technology); FGFR2, ACTIN, and FRS2 (Santa Cruz Biotechnology Inc.). Keratinocyte culture and in vivo FGFR2 functional studies are included in the Supplemental Methods.

**Protein preparation and Western blotting.** Keratinocyte, epidermis, and calvarial tissues of skull vault were separately isolated by microdissection from P0 pups or E17.5 embryos from at least 2 different litters each included in the Supplemental Methods.

Keratinocyte culture and in vivo FGFR2 functional studies are included in the Supplemental Methods.

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