Pharmacological targeting of KDM6A and KDM6B, as a novel therapeutic strategy for treating craniosynostosis in Saethre-Chotzen syndrome

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Abstract

Background: During development, excessive osteogenic differentiation of mesenchymal progenitor cells (MPC) within the cranial sutures can lead to premature suture fusion or craniosynostosis, leading to craniofacial and cognitive issues. Saethre-Chotzen syndrome (SCS) is a common form of craniosynostosis, caused by TWIST-1 gene mutations. Currently, the only treatment option for craniosynostosis involves multiple invasive cranial surgeries, which can lead to serious complications.

Methods: The present study utilized Twist-1 haploinsufficient (Twist-1 del/+ ) mice as SCS mouse model to investigate the inhibition of Kdm6a and Kdm6b activity using the pharmacological inhibitor, GSK-J4, on calvarial cell osteogenic potential.

Results: This study showed that the histone methyltransferase EZH2, an osteogenesis inhibitor, is downregulated in calvarial cells derived from Twist-1 del/+ mice, whereas the counter histone demethylases, Kdm6a and Kdm6b, known promoters of osteogenesis, were upregulated. In vitro studies confirmed that siRNA-mediated inhibition of Kdm6a and Kdm6b expression suppressed osteogenic differentiation of Twist-1 del/+ calvarial cells. Moreover, pharmacological targeting of Kdm6a and Kdm6b activity, with the inhibitor, GSK-J4, caused a dose-dependent suppression of osteogenic differentiation by Twist-1 del/+ calvarial cells in vitro and reduced mineralized bone formation in Twist-1 del/+ calvarial explant cultures. Chromatin immunoprecipitation and Western blot analyses found that GSK-J4 treatment elevated the levels of the Kdm6a and Kdm6b epigenetic target, the repressive mark of trimethylated lysine 27 on histone 3, on osteogenic genes leading to repression of Runx2 and Alkaline Phosphatase expression. Pre-clinical in vivo studies showed that local administration of GSK-J4 to the calvaria of Twist-1 del/+ mice prevented premature suture fusion and kept the sutures open up to postnatal day 20.

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Background

Calvarial sutures are comprised of active mesenchyme forming the osteogenic fronts at the edges of the flat calvarial bones [1, 2]. Within the suture mesenchyme, a reservoir of TWIST-1+/Gli-1+ mesenchymal progenitor cells (MPC) exists which can either remain undifferentiated or differentiate to mature bone forming osteoblasts [3–7]. During embryonic and postnatal development, the sutures remain open up to adulthood, providing flexibility to the calvaria and allowing the cranium to accommodate for the growing brain [8–11]. However, dysregulation of MPC differentiation within these sutures results in excessive intramembranous ossification and premature fusion of the suture space or called craniosynostosis.

Craniosynostosis occurs in 1 in 2500 live births and can result in an unusual head shape, facial asymmetry, and most importantly, pre-fusion of the cranial sutures, causing increased pressure on the developing brain leading to neurological deficits [12, 13]. Currently, the only treatment for craniosynostosis is invasive cranial surgery, mainly involving the removal of the affected sutures and remodeling of the skull [14, 15]. These procedures could negatively impact the quality of life of children with craniosynostosis, leading to serious complications and enforcing the need for invasive surgical procedures [16–18]. The most prevalent syndromic craniosynostosis, Saethre-Chotzen syndrome (SCS), involves unilateral and bilateral coronal synostosis, facial asymmetry, occasional cleft palate, droopy eyelid, and mild limb deformities such as the shortened and united fingers and toes [19, 20]. SCS is induced by the deletion or non-sense mutations resulting in the loss of function or haploinsufficiency of the TWIST-1 gene [21]. There are more than 100 different TWIST-1 gene mutations related to SCS, resulting in a range of phenotypes from a simple unilateral coronal synostosis to a complex multiple suture synostosis [22]. Recently, it has been revealed that epigenetic mechanisms play a significant role in craniosynostosis where studies of genetically identical twins reported that one twin displayed craniosynostosis, whereas the other displayed normal skull development [23, 24]. These observations suggest that the development of craniosynostosis in only one identical twin is most likely due to epigenetic changes. However, until now, no study has thoroughly examined the role of epigenetics in SCS.

TWIST-1, a basic helix-loop-helix transcription factor, has been shown to mediate skeletal and head tissue development [9, 25, 26]. Its expression in MPC within the calvarial sutures is essential in maintaining its stemness characteristics, such as proliferation activity, and negatively regulating osteogenic differentiation by directly inhibiting major osteogenic genes [9, 27–30]. Furthermore, previous studies found that Twist-1 expression is required for correct establishment of the coronal sutures in mice [5, 31]. Haploinsufficiency of the TWIST-1 gene in SCS-derived calvarial cells results in a decrease in proliferation and increased osteogenic differentiation, leading to premature suture fusion [28, 32, 33]. TWIST-1 expression and function have been correlated with the epigenetic regulator Enhance of Zeste Homolog 2 (EZH2) in mediating SCS cranial bone cell growth and differentiation [32], where Ezh2 knockout in the mesenchymal lineage leads to craniosynostosis and other skeletal deformities [34]. EZH2 is a member of the Polycomb Repressive Complex 2 and acts as a methyltransferase which tri-methylates lysine-27 of the histone-3 tail (H3K27me3), to repress gene activation [35]. The counter demethylases, UTX (lysine demethylase 6A, KDM6A) and JMJD3 (lysine demethylase 6B, KDM6B), remove the tri-methylation mark on H3K27me3 to promote gene activation [36–39]. The enzymatic demethylase activity of these epigenetic modifiers is carried out by the Jumonji C catalytic domain, through a dioxygenase reaction that requires Fe (II) and α-ketoglutarate as co-substrates [40]. Previous studies have reported that KDM6A and KDM6B promote osteogenic differentiation of mesenchymal stem cells [41, 42], whereas EZH2 represses bone gene activation and mesenchymal stem cell osteogenic differentiation [41]. Additionally, loss-of-function mutation of KDM6A has been previously identified to be associated with a congenital skeletal tissue disorder, called Kabuki syndrome, with characteristics including malformed cranial bones [43–45]. Similarly, loss of KDM6B results in a severe delay of osteogenic differentiation in mice [46, 47] and lowered expression of Runx2 and Osterix, as a result of increased levels of H3K27me3 [48]. These observations provide affirmation that both KDM6A and B play important roles in promoting osteogenic differentiation of MPC.

To further examine epigenetic changes in SCS, we utilized Twist-1 heterozygous mutant mice (Twist-1<sup>+/del</sup>),
which display craniofacial defects including unilateral or bilateral coronal synostosis and limb abnormalities similar to the characteristic abnormalities described for SCS human patients [49]. The present study investigated the expression levels and role of Kdm6a and Kdm6b in the osteogenic potential of calvarial cells and calvarial explants derived from Twist-I(del/) mice. Furthermore, we assessed a potential drug therapy approach to reverse aberrant osteogenic differentiation in the sutures of Twist-I(del/) mice using the small-molecule cell-permeable selective inhibitor, GSK-J4 targeting Kdm6a and Kdm6b activity in vivo.

Methods

Isolation of mouse calvarial cells

Mouse calvarial stromal cells were derived from the calvaria of 15-day-old Twist-I(del/) heterozygous mice and wildtype mice in accordance with South Australia Health and Medical Institute (SAHMRI) Animal Ethics Committee approval # SAM347. The mice were humanely killed by CO2 inhalation followed by cervical dislocation. The calvarial bones were retrieved then digested twice with Collagenase I (3 mg/mL) and DNAase I (50 U/mL) in PBS for 40 min each time. Bone chips were cultured under hypoxia condition (5% O2) with growth media, 10 mM HEPES buffer, 1 × 10−3 M β-glycerol phosphate, 2 mM L-glutamine, 100 μg/mL streptomycin, 50 U/mL penicillin, and 50 μg/mL streptomycin for up to 7 days. Cells were disassociated with Collagenase I (3 mg/mL) and Dispase (3 mg/mL) solution, cultured with a seeding density of 8 × 10^3 cells/cm² and grown until confluent.

Osteogenic differentiation assays

Cells were cultured in osteogenic inductive media (αMEM supplemented with 10% fetal calf serum, 100 μg/mL L-ascorbate-2-phosphate, 10 mM β-glycerol phosphate, 2 mM L-glutamine, 1 mM sodium pyruvate, 10 mM HEPES buffer, 1 × 10−3 M dexamethasone, 50 U/mL penicillin, and 50 μg/mL streptomycin) for 1 week or 2 weeks. GSK-J4 at 1 μM and 2 μM or 0.1% dimethyl sulfoxide (DMSO) in osteogenic-inductive media were refreshed every 24 h. Alkaline phosphatase staining was performed using Leukocyte Alkaline Phosphatase Kit (Cat# 86R-1KT, Sigma-Aldrich Inc., North Ryde, NSW, AU) following manufacturer’s protocols. The activity of alkaline phosphatase was quantitated in triplicate and normalized to total protein level per well using Alkaline phosphatase assay kit (Cat#00231R, Sigma-Aldrich Inc., North Ryde, NSW, AU) following manufacturer’s instructions. Bone mineral deposits were stained with Alizarin red S (Cat# A5223, Sigma-Aldrich Inc., Melbourne, VIC, AU), following manufacturer instructions. Bone mineral deposits were stained with Alizarin red S (Cat# A5223, Sigma-Aldrich Inc., Melbourne, VIC, AU) following manufacturer’s instructions. Bone mineral deposits were stained with Alizarin red S (Cat# A5223, Sigma-Aldrich Inc., Melbourne, VIC, AU), following manufacturer’s instructions.

siRNA gene knockdown studies

Cells were seeded at 3 × 10^4 cells per well in 24-well plate the day before siRNA transfections to achieve approximately 70% confluency. Sequence-specific siRNA against Kdm6a (s75838 and s75839) and Kdm6b (s103747 and s103746) or negative siRNA#1 control (Ambion/Life Technologies, Scoresby, VIC, AU) were transfected into the cells at concentration of 20 pmol in transfection medium (αMEM with 10% fetal calf serum) with Lipofectamine RNAiMAX reagent (Thermo-Fisher Scientific, Scoresby, VIC, AU) as previously described [41]. The incubation period for the transfection to achieve at least a 50% knockdown of transcript levels was 72 h before changing the media to osteogenic inductive media.

GSK-J4 treatment

GSK-J4 (Cat# 12073, Cayman Chemical, Ann Arbor, MI, US) was reconstituted in DMSO and stored at −80°C. Cells were seeded at 4.2 × 10^4 cells per well into 24-well plate. GSK-J4 at 0.1 μM, 0.25 μM, 0.5 μM, 1 μM, 2 μM, 5 μM, and 10 μM or DMSO (0.1%) only were added to the cells in the presence of either growth or osteogenic inductive media.

Gene expression studies

Total RNA from cultured Twist-I(del/) calvarial cells was isolated using TRIzol reagent (Cat# 15596026, Invitrogen/Thermo Fisher Scientific, Waltham, MA, USA), according to manufacturer’s instructions. Synthesis of cDNA and real-time polymerase chain reaction (PCR) analysis were performed in triplicate as previously described [50]. Primer sets (GeneWorks Pty Ltd., Thebarton, SA, AU) used in this study were mouse β-Actin (Fwd: 5′-TTCTCTGGACAGATGCGAAG-3′; Rev.: 5′-AAGGTGTGTTTAACCGGAG-3′); mouse Kdm6a (Fwd: 5′-GGCTACGGGGTTGTTTGAAG-3′; Rev.: 5′-TCCAGGTGCCTGTAATATA-3′); mouse Kdm6b (Fwd: 5′-CCGGCTTTGCTGACTAA-3′; Rev.: 5′-CTGGACCAAGGGTGTGTTT-3′); mouse Ezh2 (Fwd: 5′-ACTTCGCGACACGCTCAGTG-3′; Rev.: 5′-TCTCCGAGAATATACTCTCCCCCAG-3′); mouse Twist-1 (Fwd: 5′-CAGCGGTCTGACTAC-3′; Rev.: 5′-TCCCTGAGAATATACTCTCCCCCAG-3′); mouse Alkaline Phosphatase (Fwd: 5′-CGCCAAGACACTCTTCTTGTGC-3′; Rev.: 5′-GGCTACCTTGGTGTTAGCTT-3′); mouse Runx2 (Fwd: 5′-CCCTGACTTCTGCGCTCG-3′; Rev.: 5′-TTATGGAGTGCTGCTTGG-3′).
Cell proliferation assay
Cells were cultured at $9 \times 10^3$ cells/well in 96-well plates in the presence of DMSO (0.1%) or a range of GSK-J4 concentrations (0.1 μM, 0.25 μM, 0.5 μM, 1 μM, 2 μM, 5 μM, and 10 μM) in growth inductive media (αMEM supplemented with 20% fetal calf serum, pyruvate, L-glutamine, P/S) for 7 days. The rate of cell proliferation was measured using cell proliferation ELISA, bromodeoxyuridine (BrdU) colorimetric kit (Cat# 11647229001, Roche Products Pty Limited, Sydney, NSW, AU), following manufacturer’s directions. Absorbance was read at 450 nm on an iMark microplate reader (Bio-Rad Laboratories, Hercules, CA, USA).

Cell viability assay
Cells were seeded at $2.6 \times 10^5$ cells/well into 6-well plates in growth inductive media and in the presence of 0.1% DMSO or GSK-J4 concentration range (0.1 μM–10 μM) for 7 days. The rate of apoptosis was measured using Annexin V and 7AAD staining procedure. For positive controls, apoptosis and necrosis were induced by adding 100% DMSO overnight and 70% Ethanol, respectively. Prior to reading, 5 μL of Annexin V-488 (Cat# A13202, Invitrogen/Thermo Fisher Scientific) and 20 μL of 7-amino-actinomycin (7AAD; Cat# A1310, Invitrogen/Thermo Fisher Scientific) were added to ~1 × $10^6$ cells as previously described [33]. Samples were analyzed immediately on LSRForessa X20 Analyzer (BD Biosciences, North Ryde, NSW, Australia).

Calvaria organ explant cultures
Whole calvaria organ explants were dissected from 4-day-old Twist-1del/+ mice and cultured at 8 × 10^3 cells/cm² in T75 flasks until confluent and then treated with GSK-J4 concentration range or 0.1% DMSO as vehicle control for 24 h. Histone extraction protocol was adapted from Abcam (Abcam, Melbourne, VIC, Australia). Briefly, 5 × $10^6$ cells were re-suspended in 1 ml of Triton Extraction Buffer (0.5% Triton X 100 (v/v), 2 mM phenylmethylsulfonyl fluoride (PMSF), 0.02% NaN₃ (w/v)). The nuclei lysates were incubated on ice for 10 min with gentle stirring and centrifuged for 10 min at 6500 g at 4 °C. Histone acid extraction was performed using 0.2 M HCl at a density of 2 × $10^7$ nuclei/mL overnight at 4 °C. Histone protein was collected on ice for 10 min with gentle stirring and centrifuged for 10 min at 6500 g at 4 °C. Histone protein was collected in the supernatant following centrifuge spin as described before and neutralized with 2 M NaOH at 1/10 of supernatant volume. Protein concentrations were measured using the Pierce Detergent Compatible Bradford Assay Kit (Cat# 1863028, Thermo Fisher Scientific). Western blot analysis was performed as described [50]. The membranes were blocked with 5% BSA blocking solution and probed overnight at 4 °C with an anti-mouse H3K27me3 rabbit polyclonal antibody (Cat# 07-449, Millipore, Bayswater, VIC, Australia). Rabbit IgG (Cat# I5006, Sigma–Aldrich) replaced the primary antibody as negative control, which showed no immunoreactivity. The percentage of H3K27me3-positive nuclei (brown) to total number of nuclei within the white box was quantitated using ImageJ software.

In vivo administration of GSK-J4 to calvaria of Twist-1del/+ mice
Two 3 mm² CollaCote sponges (Cat# 0101, Integra Life Sciences Services, Saint Priest, FRA) soaked in 0.1% DMSO as vehicle control or in GSK-J4 in the concentration of 2 μM was placed subcutaneously onto each side of the coronal sutures of 8-day-old Twist-1del/+ mice up to 20 days of age, in accordance with the SAHMRI Animal Ethics Approval (Ethics# SAM347). The calvaria of treated mice fixed in 10% formalin was analyzed using Masson’s trichrome staining. Mineralized calvarial bone formation relative to the length of bone analyzed was quantitated using OsteoMeasure software.

Immunohistochemical analysis
Calvaria was isolated from 10-day-old Twist-1del/+ and wildtype mice and then fixed in 10% formalin for 24 h, decalciﬁed with 14% EDTA (pH 7.2) overnight, and embedded in parafﬁn. The samples were cut transversely with thickness of 5 μm and processed as previously described [33]. The primary antibody used was an anti-mouse H3K27me3 rabbit polyclonal antibody (Cat# 07-449, Millipore, Bayswater, VIC, Australia). Rabbit IgG (Cat# I5006, Sigma–Aldrich) replaced the primary antibody as negative control, which showed no immunoreactivity. The percentage of H3K27me3-positive nuclei (brown) to total number of nuclei within the white box was quantitated using ImageJ software.

Western blot analysis
Calvarial cells were cultured at $8 \times 10^3$ cells/cm² in T75 flask until confluent and then treated with GSK-J4 concentration range or 0.1% DMSO as vehicle control for 24 h. Histone extraction protocol was adapted from Abcam (Abcam, Melbourne, VIC, Australia). Briefly, 5 × $10^6$ cells were re-suspended in 1 ml of Triton Extraction Buffer (0.5% Triton X 100 (v/v), 2 mM phenylmethylsulfonyl fluoride (PMSF), 0.02% NaNO₃ (w/v)). The nuclei lysates were incubated on ice for 10 min with gentle stirring and centrifuged for 10 min at 6500 g at 4 °C. Histone acid extraction was performed using 0.2 M HCl at a density of 2 × $10^7$ nuclei/mL overnight at 4 °C. Histone protein was collected in the supernatant following centrifuge spin as described before and neutralized with 2 M NaOH at 1/10 of supernatant volume. Protein concentrations were measured using the Pierce Detergent Compatible Bradford Assay Kit (Cat# 1863028, Thermo Fisher Scientific). Western blot analysis was performed as described [50]. The membranes were blocked with 5% BSA blocking solution and probed overnight at 4 °C with an anti-mouse H3K27me3 rabbit polyclonal antibody (Cat# 07-449, Millipore Corporation, North Ryde, NSW, Australia) and a rabbit anti-H4 antibody (Cat#ab10158, Abcam). Following two washes with TBS/0.1% Tween 20, the blots were incubated for 1 h in room temperature with fluorescence secondary antibody (anti-rabbit 800 nm or 680 nm, Li-Cor Biosciences, VIC, Australia). Blots were washed two more times and then scanned on Odyssey CLX Near-Infrared Fluorescence Imaging.
Calvarial cells derived from 15-day-old Twist-1

Results

Statistical differences (*) of \( p \) also used for the generation of graphs which showed 0.02798, http://www.graphpad.com/). The software was of statistical significance was carried out using GraphPad Experiments were performed in triplicates. Calculation of statistical significance was carried out using GraphPad PRISM 8 (GraphPad Software, La Jolla, CA, RRID: CR_002798, http://www.graphpad.com/). The software was also used for the generation of graphs which showed statistical differences (*) of \( p < 0.05 \) between samples.

Chromatin immunoprecipitation (ChIP) analysis

Calvarial cells were seeded at density of 8 \( \times 10^3 \) cells/cm\(^2\) in T75 flasks. Once the cells were confluent, the cells were cultured in growth or osteogenic-inductive media in the presence of GSK-J4 at 1 \( \mu \text{M} \) or vehicle control (0.1% DMSO) for 24 h. ChIP protocol was adapted from Abcam. Chromatin was crosslinked with a final of 0.75% formaldehyde for 10 min at room temperature with gentle rocking. Glycine at a final concentration of 125 mM was added and incubated whilst shaking for 5 min. After two washes with PBS, adherent cells were detached using 1x trypsin, and the remaining cells were scraped. Cells were lysed with FA lysis buffer (50 mM HEPES KOH pH 7.5, 140 mM NaCl, 1 mM EDTA pH 8, 1% Triton X-100, 0.1% Sodium deoxycholate, 0.1% SDS, and protease inhibitors) at 400 ul per one million cells. DNA was sheared with a probe sonicator (Diagenode Bioruptor Inc., Denville, NJ, USA) on ice and then used for immunoprecipitation as previously described [53]. Primary antibodies that were used for immunoprecipitation were anti-mouse H3K27me3 rabbit polyclonal (1 mg/ml; Cat# 07-449, Millipore) and IgG rabbit polyclonal control (1 mg/ml Millipore). Transcription start site (TSS) primer sets (GeneWorks Pty Ltd) used in this study: mouse Runx2 TSS (Fwd: 5′-AGGGAATACGACCCGCTCCTCATC-3′; Rev.: 5′-GGGACTGATCCCGCTCTGGTGC-3′), mouse Alkaline Phosphatase TSS (Fwd: 5′-AGGGAAGAG AGAGGCAGGG-3′, Rev.: 5′-TTCCCTATCTGCAGGC ACTC-3′).

Statistics

Experiments were performed in triplicates. Calculation of statistical significance was carried out using GraphPad System (Li-Cor Biosciences). Analysis and measurements were performed on Image Studio Lite software (Li-cor Biosciences).

Kdm6a and Kdm6b promoter the osteogenic differentiation capacity of Twist-1\(^{del/+}\) calvarial cells

The role of Kdm6a and Kdm6b during osteogenic differentiation in calvarial cells from Twist-1\(^{del/+}\) mice was assessed using two specific siRNA molecules targeting either Kdm6a or Kdm6b. Reduced gene expression levels of Kdm6a or Kdm6b and the mature osteoblast marker, Alkaline Phosphatase was confirmed by qRT-PCR analysis following siRNA treatment (Fig. 2a). Furthermore, the data showed a decrease in alkaline phosphatase enzymatic activity in siRNA Kdm6a or Kdm6b transfected Twist-1\(^{del/+}\) calvarial cells under osteogenic conditions, compared with scrambled siRNA controls (Fig. 2b, c). Parallel studies found that the level of Alizarin red-positive mineralized deposits was significantly reduced in cultures of siRNA Kdm6a or Kdm6b knockdown Twist-1\(^{del/+}\) calvarial cells under osteogenic conditions, compared with scrambled siRNA-treated cells (Fig. 2d). This was confirmed by reduced amounts of extracellular calcium levels observed in replicate cultures of siRNA Kdm6a- or Kdm6b-treated Twist-1\(^{del/+}\) calvarial cells, compared to the scrambled siRNA controls (Fig. 2e).

Kdm6a and Kdm6b inhibitor, GSK-J4, shows minimal toxicity in Twist-1\(^{del/+}\) calvarial cells

Twist-1\(^{del/+}\) calvarial cells were cultured with increasing concentrations of GSK-J4 to assess potential cytotoxic effects. Observable differences in cell density occurred in the presence of GSK-J4 between 2 and 10 \( \mu \text{M} \) (Fig. 3a). Quantitative analysis found that the proliferation rate was not affected in the presence of 0.1–0.5 \( \mu \text{M} \) GSK-J4, but cell proliferation was significantly reduced between 1 and 10 \( \mu \text{M} \) GSK-J4, as assessed by BrdU incorporation (Fig. 3b). Flow cytometric analysis of Twist-1\(^{del/+}\) calvarial cells showed that the percentage of early apoptotic (Annexin V positive), necrotic (7AAD positive), and late-stage apoptotic (Annexin V + 7AAD-positive) cells significantly increased with GSK-J4 treatment at the higher doses of 5 \( \mu \text{M} \) and 10 \( \mu \text{M} \) (Fig. 3c, d). Therefore, concentrations higher than 2 \( \mu \text{M} \) were eliminated from further studies.

Inhibition of Kdm6a and Kdm6b activity by GSK-J4 suppresses the osteogenic differentiation of Twist-1\(^{del/+}\) calvarial cells in vitro

We next assessed whether inhibition of Kdm6a and Kdm6b activity could suppress the osteogenic differentiation capacity of Twist-1\(^{del/+}\) calvarial cells. Western blot analysis showed an increase in H3K27me3 levels in histone lysates extracted from Twist-1\(^{del/+}\) calvarial cells treated with 1 \( \mu \text{M} \) or 2 \( \mu \text{M} \) GSK-J4, compared to 0.1%
DMSO (Fig. 4a), confirming the specificity of GSK-J4. Furthermore, Twist-1del/+ calvarial cells exhibited a reduction in Runx2 and Alkaline Phosphatase gene expression levels in the presence of 1 or 2 μM of GSK-J4 compared to vehicle alone controls, when cultured under osteogenic inductive conditions (Fig. 4b). Supportive studies showed that alkaline phosphatase activity was significantly suppressed in Twist-1del/+ calvarial cells treated with 2 μM of GSK-J4, compared to vehicle controls (Fig. 4c, d). Chromatin collected from replicate experiments was used to analyze levels of the inhibitory mark, H3K27me3, present on the Runx2 and Alkaline Phosphatase promoter transcription start sites (TSS), using ChIP analysis. The results demonstrated that H3K27me3 levels decreased dramatically on the Runx2 and Alkaline Phosphatase TSS under osteogenic inductive conditions compared to normal growth conditions (Fig. 4e, f). However, treatment with GSK-J4 resulted in increased levels of H3K27me3 on the Runx2 and Alkaline Phosphatase TSS, correlating with the suppression of these genes following GSK-J4 treatment (Fig. 4e, f). These findings suggested that the addition of GSK-J4 to Twist-1del/+ calvarial cell cultures increased the amount of H3K27me3 found on the promotors of osteogenic...
genes by inhibiting the activity of Kdm6a and Kdm6b during osteogenesis.

The effect of GSK-J4 on osteogenic differentiation was further examined using a murine calvarial organotypic explant model. Calvaria derived from Twist-1del/+ mice were cultured in media containing BMP2, to initiate bone formation, in the presence or absence of GSK-J4. The calvarial explants were then stained with Masson’s trichrome stain to identify newly mineralized bone (Fig. 5a).

Histomorphometric analysis revealed a reduction in total bone formation and thickness in calvarial explants treated with 1 μM and 2 μM of GSK-J4, compared to 0.1% DMSO vehicle alone treated explants (Fig. 5b, c).

**GSK-J4 treatment prevents craniosynostosis in Twist-1del/+ mice**

The ability of GSK-J4 treatment to prevent fusion of the coronal sutures in vivo was assessed using 3 mm²...
Collacote™ sponges containing either DMSO alone or GSK-J4 placed subcutaneously on top of the coronal sutures in pre-fusion from 8-day-old Twist-1del/+ mice. Sections of coronal sutures derived from 20-day-old Twist-1del/+ mice were stained with Masson’s trichrome stain and examined by histomorphometric analysis. The data showed that 80% of Twist-1del/+ mice treated with 2 μM GSK-J4 exhibited open coronal sutures at postnatal day 20, whereas unilateral or bilateral coronal craniosynostosis was observed in 83% of the Twist-1del/+ mice treated with DMSO alone (Table 1 and Fig. 5d). Furthermore, there was a significant reduction in the total mineralized bone formed and bone thickness in the coronal sutures of GSK-J4-treated Twist-1del/+ mice compared to DMSO treated control mice (Fig. 5e, f). These findings...
demonstrate that local administration of GSK-J4 underneath the skull cap can prevent premature coronal suture fusion that occurs in Twist-1 del/+ mice between postnatal days 9–20. Table 1 shows the number of Twist-1 del/+ mice with open coronal sutures, unilateral or bilateral coronal craniosynostosis following treatment of either DMSO (0.1%) or 2 μM GSK-J4 (n = 5–6 Twist-1 del/+ mice/treatment group).

**Fig. 4** Inhibition of Kdm6a and Kdm6b activity suppresses osteogenic differentiation of Twist-1 del/+ calvarial cells. a Western blot analysis of nuclear extracts isolated from Twist-1 del/+ calvaria cells treated with GSK-J4 (1 μM, 2 μM) or 0.1% DMSO vehicle control for 24 h to assess H3K27me3 levels relative to histone 4 (H4). b Real-time qPCR analysis of Runx2 and Alkaline Phosphatase (Alk Phos) transcript levels in Twist-1 del/+ calvaria cells under osteogenic inductions for 24 h. Data represent mean gene expression levels normalized to β-actin ± S.E. expression, *p < 0.05, two-tailed, not-paired, non-parametric student’s t test, n = 3 Twist-1 del/+ mice. c Representative images of alkaline phosphatase staining and d quantitation of alkaline phosphatase activity relative to total protein for Twist-1 del/+ calvaria cells treated with either 1 μM or 2 μM GSK-J4 or 0.1% DMSO, following 1 week of osteogenic induction. Data represent mean ± S.E., *p < 0.05, one-way ANOVA with Tukey’s multiple comparisons, n = 4 Twist-1 del/+ mice. Chromatin immunoprecipitation (ChIP) analysis of H3K27me3 levels on the transcriptional start sites of e Runx2 and f Alk Phos for Twist-1 del/+ calvarial cells cultured under normal growth media (NGM) or osteogenic inductive conditions (Osteo) for 1 week in the presence of either GSK-J4 (1 μM) or 0.1% DMSO. ChIP was performed using either IgG control antibody (IgG) or H3K27me3-specific antibody (K27me). Enriched genomic DNA was used to amplify the transcription start site of target genes. Data represent mean fold enrichment relative to input DNA ± S.E., *p < 0.05, one-way ANOVA with Tukey’s multiple comparisons, n = 4 Twist-1 del/+ mice.
In this study, we demonstrate for the first time that calvarial cells extracted from Twist-1del/+ mice express elevated levels of the H3K27me3 demethylases, Kdm6a and Kdm6b, whereas the gene expression levels of the counter histone methyltransferase, Ezh2, was reduced, compared to cells derived from littermate wildtype mice. This correlated to a reduction in the amount of H3K27me3 within the coronal sutures of Twist-1del/+ mice. These findings suggest that the altered balance in epigenetic enzymes that deposit or remove H3K27me3 are pivotal in driving the craniosynostosis phenotype. Both histone demethylases have been previously reported to promote osteogenic differentiation by removing the repressive mark, H3K27me3, on the promoter of osteogenic-promoting genes [41]. Similarly, we found that inhibition of Kdm6A and Kdm6b activity in Twist-1del/+ calvarial cells reduced gene transcript levels of Runx2 and Alkaline Phosphatase, following treatment with GSK-J4, correlating to lower levels of H3K27me3 on the respective promoters. Previous studies have shown that TWIST-1 induces EZH2 in cultured human mesenchymal stem cells increasing levels of EZH2 and H3K27me3 along the Ink4A locus and bone gene promoters to increase proliferation but suppress osteogenesis, which was diminished in cranial bone cells derived from SCS patients [32, 33, 41]. These observations support our findings that TWIST-1 mutations lead to aberrant EZH2 and KDM6A/B expression levels, suggesting that a balance of histone demethylases and methyltransferase is essential in maintaining the correct fate determination of cranial MPC. The epigenetic dysregulation seen in the Twist-1 haploinsufficient cells may therefore

|                  | DMSO control | 2 µM GSK-J4 |
|------------------|--------------|-------------|
| Open coronal sutures | 1            | 4           |
| Unilateral coronal craniosynostosis | 2            | 1           |
| Bilateral coronal craniosynostosis   | 3            | 0           |
mediate the premature maturation of bone cells as a result of Twist-1 mutation within the suture mesenchyme of the SCS mouse model. The aberrant osteogenesis in Twist-1 mutant cells has been reported previously in other studies and further confirmed by the present study with the increased expression of Runx2 and Alkaline Phosphatase when compared to wildtype cells [28, 33, 54]. Both of these osteogenic genes have been previously shown to be expressed within mouse calvarial cells at the osteogenic fronts and within the suture mesenchyme and thus have essential roles in the development of mouse calvaria [55, 56].

Functional studies using Twist-1 del/+ calvarial cells determined that the suppression of the expression of Kdm6a and Kdm6b led to the inhibition of early osteogenic differentiation shown in the reduced activity and expressions of Alkaline Phosphatase, and late osteogenic differentiation as seen in the reduced amount of mineral deposition and reduced calcium production. This is in agreement with previous studies reporting that Kdm6b-null mice display open calvarial sutures and less mineralized calvarial bones [46], whereas Kdm6a-null mice exhibit defects in neural crest formation [44]. Notably, the defects in Kdm6a knockout mice are more severe in female mice than in males [57, 58]. This suggested that Uty/Kdm6c, an enzymatically inactive paralog of Kdm6a located on the Y-chromosome, is able to compensate for the loss of Kdm6a in males [59, 60]. However, comparison between the sexes showed similar effects on osteogenesis following Kdm6a and Kdm6b inhibition (data not shown). This indicated that the enzymatic activity is essential in the regulation of H3K27me3 levels during calvarial osteogenic differentiation, and thus, Kdm6c activity was not able to compensate during Kdm6a and Kdm6b inhibition in this instance. Collectively, our findings provided evidence that Kdm6a and Kdm6b are putative targets in treating craniostenosis in Twist-1 del/+ mutant mice and confirmed previous studies that deregulated epigenetic patterns play significant roles in the development of craniostenosis [23, 24].

Our study further explored the utility of a chemical inhibitor, GSK-J4, in suppressing the osteogenic potential of Twist-1 del/+ calvarial cells. This inhibitor was designed to target KDM6A and KDM6B enzymatic activities by competitively binding with their active sites, responsible for the interaction between the co-substrate, α-ketoglutarate, and a histone-3 peptide [61]. Our study showed that the treatment of GSK-J4 using concentrations of 1 μM and 2 μM on Twist-1 del/+ calvarial cells resulted in a reduction in Runx2 and Alkaline Phosphatase gene expression and activity. Notably, the dosage used had little or no effect in the cell viability rate of the Twist-1 del/+ calvarial cells; however, higher doses significantly reduced the proliferation rate. This anti-proliferative effect of GSK-J4 has been previously described for embryonic bodies and tumor cells such as bone sarcoma [62], pediatric brain glioma [63], and acute lymphoblastic leukemia [64]. Whilst the reduction of proliferation was found to be caused by accumulation of cells at S-phase inhibiting cell cycle progression, our study showed that Twist-1 del/+ calvarial cells undergo apoptosis in the presence of 5 μM GSK-J4.

Functional studies showed that GSK-J4 treatment reduced the development of the total bone on whole calvarial organotypic explant cultures derived from Twist-1 del/+ mice. In a pre-clinical model of SCS, local administration of GSK-J4 to the calvaria of Twist-1 del/+ mice prevented premature coronal suture fusion and reduced the amount of mineralized calvarial bone formation. The local delivery approach of GSK-J4 administration was performed because Kdm6a is ubiquitously expressed and both Kdm6a and Kdm6b were shown to be essential in the correct skeletal patterning, brain and heart development [43, 46, 57, 65, 66]. Collectively, the present study demonstrated that GSK-J4 treatment effectively suppressed osteogenic differentiation of Twist-1 del/+ calvarial cells in both sexes and whole calvaria explants in vitro, and prevented coronal suture craniostenosis of Twist-1 del/+ mice in vivo by inhibiting the enzymatic activity of aberrant Kdm6a and Kdm6b levels and thus recovering the level of H3K27me3 marks on osteogenic genes.

Previous studies have employed GSK-J4 to understand the roles of the KDM6 subfamily members in regulating differentiation of embryonic stem cells and bone marrow-derived mesenchymal stem cells [65]. Moreover, GSK-J4 has been highly utilized in the studies of novel therapeutic strategies against various types of diseases, including acute lymphoblastic leukemia [64], myeloid leukemia [67], osteoarthritis [65] and breast [68], prostate [69], ovarian cancers [70], and brainstem glioma [63]. Of note, the later study also showed that normal brain cells from healthy children are unaffected by GSK-J4 treatment. This implies that the use of localized GSK-J4 treatment to reverse craniostenosis may have little or no adverse impact on brain development in children with SCS. However, further studies are required to perform pathological assessments of any potential GSK-J4 toxicity issues for various tissues and organs, as well as cognitive evaluations, using Twist-1 del/+ mice in the absence of pre-clinical large animal models of SCS.

Currently, the main treatment for craniostenosis involves an open calvaria remodeling surgery. This type of surgery might lead to serious complications such as cerebral contusions, cerebrospinal fluid leaks, hematomas, infections, and wound breakdowns [17, 18]. Additionally, in severe cases of craniostenosis, there is often a need for a follow-up treatment with repeated
surgery procedures and substantial hospitalization, as there is the possibility that sutures might fuse before the cranium has had the opportunity to expand appropriately to accommodate for the growing brain [71]. Despite the negative impacts on a patient’s wellbeing and health providers, therapies which do not involve invasive surgery have yet to be developed.

Conclusions
In this study, we demonstrate for the first time that the histone demethylases, Kdm6a, and Kdm6b have the potential to be used as novel therapeutic targets for the prevention of craniosynostosis in an SCS mouse model, where the Twist-1 gene is mutated. Therefore, this study could potentially minimize the need for invasive approaches by using a local drug therapy such as GSK-J4 to slow down the rate of premature suture fusion in children with SCS, which is irreversible following cessation of treatment.

Abbreviations
MPC: Mesenchymal progenitor cells; SCS: Saethre-Chotzen syndrome; H3K27me3: Tri-methylation on the lysine-27 of the histone-3 tail; EZH2: Enhance of Zeste Homolog 2; KDM6A: Lysine demethylase 6A; KDM6B: Lysine demethylase 6B; oMEM: α-Modification of Eagle’s medium; DMSO: Dimethyl sulfoxide; BrdU: Bromodeoxyuridine; PCR: Polymerase chain reaction; PMSF: Phenylmethylsulfonyl fluoride; ChIP: Chromatin immunoprecipitation

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Authors’ contributions
CP collected, analyzed, and interpreted all of the study data and was a major contributor in writing the manuscript. EC provided some of the cranial bone cells and skull sections, analyzed and interpreted the data, and was a contributor in writing the manuscript. DC analyzed and interpreted the data and was a contributor in writing the manuscript. PA analyzed and interpreted the data regarding the cranial suture development and the preclinical study. CG provided some of the study material. 3G analyzed and interpreted the data, provided funding for the study and was a contributor in writing the manuscript. All authors contributed in the creation of the study’s concept and design and read and approved the final manuscript.

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Availability of data and materials
The datasets used and/or analyzed during the current study are available from the corresponding author on reasonable request.

Ethics approval and consent to participate
The use of animals in this study is in accordance with the South Australia Health and Medical Institute (SAHMI) Animal Ethics Committee approval # SAM347.

Consent for publication
Not applicable

Competing interests
The authors declare that they have no competing interests.

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