Gli2 and MEF2C activate each other’s expression and function synergistically during cardiomyogenesis in vitro

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Received July 25, 2011; Revised November 7, 2011; Accepted November 24, 2011

ABSTRACT

The transcription factors Gli2 (glioma-associated factor 2), which is a transactivator of Sonic Hedgehog (Shh) signalling, and myocyte enhancer factor 2C (MEF2C) play important roles in the development of embryonic heart muscle and enhance cardiomyogenesis in stem cells. Although the physiological importance of Shh signalling and MEF2 factors in heart development is well known, the mechanistic understanding of their roles is unclear. Here, we demonstrate that Gli2 and MEF2C activated each other’s expression while enhancing cardiomyogenesis in differentiating P19 EC cells. Furthermore, dominant-negative mutant proteins of either Gli2 or MEF2C repressed each other’s expression, while impairing cardiomyogenesis in P19 EC cells. In addition, chromatin immunoprecipitation (ChIP) revealed association of Gli2 to the Mef2c gene, and of MEF2C to the Gli2 gene in differentiating P19 cells. Finally, co-immunoprecipitation studies showed that Gli2 and MEF2C proteins formed a complex, capable of synergizing on cardiomyogenesis-related promoters containing both Gli- and MEF2-binding elements. We propose a model whereby Gli2 and MEF2C bind each other’s regulatory elements, activate each other’s expression and form a protein complex that synergistically activates transcription, enhancing cardiac muscle development. This model links Shh signalling to MEF2C function during cardiomyogenesis and offers mechanistic insight into their in vivo functions.

INTRODUCTION

The mammalian heart is the first organ to develop and is essential for life. Perturbations in cardiogenesis can lead to congenital heart disease, the most prevalent birth defect worldwide. Heart development in vivo starts with the formation of the cardiac crescent, where the first heart field progenitor cells fuse to form the linear heart tube and give rise to the left ventricle. Second heart field progenitor cells then migrate to form pharyngeal and splanchnic mesoderm, which will form the right ventricle and the outflow tract (1,2). In order to properly define and maintain the cardiac identity, Sonic Hedgehog (Shh) signalling pathway members and myocyte enhancer factor 2 (MEF2) proteins are required as shown by various animal models [(3–10) and reviewed in ref. (1,2)].

In mammals, the Shh signal is transmitted into the cell by the patched1/smoothened (Ptch1/Smo) regulatory complex and is mediated by transcription factors glioma-associated factor (Gli) 1, 2, 3 [reviewed in refs (11,12)], which bind the TGGGTGTC DNA consensus sequence (13). Gli1 acts as a transcriptional activator, but is dependent on Gli2- and/or Gli3-mediated transcription. Gli3 is a transcriptional repressor (11). Using genetic inducible fate mapping, members of the Shh signalling pathway were shown to be expressed in murine myocardial progenitor cells starting from embryonic day (E) 7.0–8.0 (3). The expression of Gli1 in some atrial and ventricular myocytes was confirmed in another study when tamoxifen was administered to the R26R^{Gli1-CreERT2} embryos at E6.5 (10). Thus, embryonic cardiomyocytes and/or cardiac progenitors were exposed to Shh signalling during development.

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The Shh pathway participates in the establishment of a proper number of cardiac progenitor cells during early vertebrate heart development in zebrafish (3). Inhibition of the Shh signalling resulted in an early defect in myocardial progenitor specification leading to reduction of both ventricular and atrial cardiomyocytes (3). Additionally, activation of Shh signalling resulted in an increase of cardiomyocytes (3). The importance of the Shh signalling pathway in mammalian heart development was demonstrated by total and tissue-specific knockout studies. Smo<sup>−/−</sup> mice showed delayed formation of heart tube with delayed Nkx2-5 expression (4), whereas Ptc<sup>1−/−</sup> mice, where the negative regulation of Shh signalling was removed, demonstrated upregulated Nkx2-5 expression during heart development (4). Moreover, in Shh<sup>−/−</sup> mice there were atrial septal defects and aberrant development of the outflow tract (5). Additionally, Gli<sup>2−/−</sup>/Gli<sup>3−/−</sup> mice showed cardiac outflow tract anomalies (6,14). Tissue-specific removal of the Shh signalling pathway members in murine second heart field demonstrated their role in atrioventricular septation and the development of the outflow tract (8–10). In addition, Shh signalling was found to be important in proliferation of second heart field progenitors in chicken embryos (7). Therefore, Shh signalling via Gli2 is important for embryonic heart development.

In addition to Gli transcription factors, cardiomyogenesis is also regulated by MEF2 family members. The four vertebrate MEF2 proteins, MEF2A, MEF2B, MEF2C and MEF2D belong to the MADS box family (MCM1, Agamous, Deficiens, SRF) of transcription factors and bind A/T rich DNA sequence (T/C)TA(A/T)TA(G/A) (15). MEF2C is the first MEF2 family factor to be expressed in heart myocardium progenitors starting from E7.5 (16,17).

Loss-of-function mutations in the single Mef2 gene in Drosophila lead to a block of the development of all muscle cell types during embryogenesis (18). In mammalian embryogenesis, however, the four MEF2 family members play redundant roles and can rescue the phenotype resulting from a specific Mef2 gene knockout [reviewed in ref. (15)]. For example, Mef2c-null mice undergo cardiomyogenesis but die from a failure to undergo heart looping morphogenesis and proper development of the right ventricle, as well as the outflow tract (19,20). Cardiomyogenesis might be occurring due to a 7-fold increase in MEF2B protein observed in Mef2c-null mice, which can partially compensate for the phenotype. Overlapping functions of MEF2C and MEF2B are also supported by the evidence of normal development of Mef2b-null mice (19). The possibility of the phenotype rescue by other MEF2 family members was eliminated by expression of a dominant-negative fusion protein, of MEF2C with the engrailed repression (EnR) domain (MEF2C/EnR) under the control of the Nkx2-5 enhancer that resulted in a severe disruption of cardiomyogenesis in mice (21). Thus, both Shh signalling and MEF2 family proteins play an important role in cardiomyogenesis in vitro.

The results from in vitro experiments have largely been reproduced in vitro. First, members of the Hedgehog signalling pathway and MEF2 transcription factors are expressed during cardiomyogenesis in embryonic stem (ES) cells (22–25). During cardiomyogenesis in mouse ES (mES) cells, committed cardiac progenitor cells express MEF2C and GATA-4 transcription factors after the formation of the mesodermal cells as characterized by the expression of Brachyury T (BraT) and Oct-4. Cardiac progenitor cells terminally differentiate into cardiomyocytes and express contractile proteins such as myosin heavy chain (MHC) and cardiac Troponin T (22).

A similar pattern of gene expression to ES cells can be seen in studies using pluripotent P19 embryonal carcinoma (EC) cells (26–28). The P19 EC cells were originally isolated from an induced teratocarcinoma of male mouse testes by injection of immature mES cells (26). These cells can contribute to tissues in live-born chimeric mice (29) and can undergo controlled differentiation into cardiac and skeletal muscle cells (up to 15%) in vitro when treated with dimethylsulphoxide (DMSO) (27,28,30–34). We have previously shown that retinoic acid regulates skeletal myogenesis in P19 EC, mouse and human ES cells through upregulation of Pax3/7-positive progenitor cells (35,36), similar to a role for retinoic acid and Pax3/7 progenitors during embryogenesis (37,38). Moreover, Nkx-MEF2C/EnR was found to abrogate cardiac myogenesis both during murine embryogenesis and in P19 EC cells (21). Similarly, Nkx/EnR impaired cardiac muscle formation in vivo (39) and in P19 EC cells (40). More recently, we found that skeletal myosin light chain kinase (MLCK) is important for skeletal myogenesis in mES and P19 EC cells, as well as in the activation of satellite cells via interaction and subsequent phosphorylation of the MEF2C protein (25). For Gli2 and MEF2C expression, previous publications demonstrate that the members of the Shh signalling pathway and MEF2 family members are expressed during cardiomyogenesis in P19 EC cells (41,42) and P19CL6 cells (43,44), as well as in cardiomyocytes derived from mES cells stably expressing the neomycin-resistance gene under the regulation of the cardiac α-MHC promoter (23). Therefore, P19 EC cells use similar pathways as those identified in the embryo, mouse and human ES cells and are a suitable model to study cardiac and skeletal myogenesis in vitro.

Gain-of-function experiments in P19 EC cells revealed that both Gli2 and MEF2C are able to induce cardiomyogenesis through the upregulation of Nkx2-5, GATA-4 and BMP-4 factors (41,45). Inhibition of Shh signalling in vitro resulted in aberrant cardiomyogenesis in P19 and P19CL6 cells (42,43). Surprisingly, the ubiquitous expression of MEF2C/EnR, driven by the <sup>pgk</sup> promoter, enhanced cardiomyogenesis in P19 cells, likely by binding and inhibiting Class II histone deacetylases (HDACs), resulting in the recruitment of non-cardiac muscle cells into the lineage (46). However, consistent with the gain-of-function experiments, the cardiac-restricted expression of MEF2C/EnR, driven by an Nkx2-5 enhancer, ablated cardiomyogenesis in P19 EC cells (21), demonstrating that MEF2 factors, or genes regulated by MEF2, are essential for cardiomyogenesis. Therefore, Shh signalling members and MEF2 proteins are important for cardiomyogenesis in stem cells.
Although Gli2 and MEF2C activate myogenic differentiation programs in a similar fashion (41,45,47,48), and their physiological role in heart development is known ([3–6,18,21] and reviewed in ref. (15)), their detailed function and the mechanism of their regulation is unclear. In this article, we examined the relationship between Gli2 and MEF2C during mES and P19 EC cellular cardiomyogenesis. We have found that Gli2 and MEF2C regulate each other’s expression and associate with their respective gene elements, as well as form a protein complex capable of synergizing on gene promoters. We therefore link, for the first time, the Sonic Hedgehog signalling pathway to the function of MEF2C protein during cardiomyogenesis in vitro.

METHODS
Transgenic mice
Gli2+/− heterozygous mice [obtained from Dr A. Joyner, Sloan-Kettering Institute, New York, NY, USA; (49)] were maintained on CD1 background. Gli2+/−/− homozygous mice were generated by crossing two Gli2+/− heterozygous mice.

mES cell culture
D3 mES cells (ATCC, # CRL-1934) were maintained as described in ref. (25,35). For differentiation, cells were aggregated in hanging drops containing 800 cells each for the first 2 days. After culturing aggregates in suspension for an additional 5 days, they were plated on tissue-culture grade adherent plates or 0.1%-gelatin covered coverslips for the remaining 2 days of the 9-day differentiation protocol.

P19 EC cell culture
P19 EC cells (ATCC, #CRL-1825) were cultured and differentiated in the presence of 1% DMSO as previously described (50). Briefly, cells were aggregated in the presence of DMSO for 4 days and formed aggregates were plated onto 0.1%-gelatin covered coverslips or adherent tissue-culture grade dishes. P19 EC cells overexpressing various factors including P19[Gli2], P19[MEF2C], P19[Control], P19[Gli/EnR], P19[MEF2C-TAP] and P19[TAP] cells were described elsewhere (25,47). P19[Nkx-MEF2C/EnR] cells stably overexpressing MEF2C/EnR under the regulation of the Nkx2-5 enhancer (within the region from −9435 to −7353 nt relative to Nkx2-5 transcriptional start site) were described in ref. (21).

Immunofluorescence
On Day 6 of P19 EC and Day 9 of mES cell differentiation, cultures were fixed and incubated with MF20 monoclonal antibody supernatant to detect expression of MHC as previously described (50). Cy3-conjugated secondary antibodies (Jackson Immuno Research Laboratories, USA) were used to detect immunofluorescence (IF) as described in ref. (51). Hoechst dye was used as a nuclear marker. Indirect IF was captured using a Leica DMI6000B microscope (Leica Microsystems GmbH, Germany) or a Zeiss Axioscope microscope (Carl Zeiss MicroImaging GmbH, Germany). Images were collected at ×200 or ×400 magnification using Hamamatsu Orca AG camera (Hamamatsu Photonics, Germany) or Sony 3CCD camera (Sony Corporation, Japan) and processed using Velocity 4.3.2 (Perkin Elmer, Canada) or Axiovision (Carl Zeiss MicroImaging GmbH, Germany) software.

Quantitative PCR analysis
Total RNA from differentiating mES and P19 EC cells was harvested using RNasy Mini Kit (Qiagen, Canada) and analysed using real-time quantitative PCR (QPCR) as described in refs (35,52). Briefly, 1µg of RNA was reverse-transcribed (RT) to synthesize cDNA using Quantitect Reverse Transcription Kit (Qiagen, Canada). No-RT reaction was included. One-twentieth of RT reaction was used as a template for QPCR amplification using specific primers listed in Table 1 and the FastStart SYBR Green kit (Roche Applied Sciences, Canada) or Promega GoTaq QPCR Master Mix (Promega, WI, USA). Data were acquired using ABI7300 and ABI7500 QPCR (Applied Biosystems, CA, USA) or Eppendorf Realplex2 (Eppendorf, Canada) instruments. Data were normalized to β-actin, analysed as described in ref. (53), using P19 or P19 control cell line Day 0 (monolayer cultures) as a calibrator, and expressed as percent maximum. Data

Table 1. Oligonucleotide sequences of primers utilized for real-time QPCR

<table>
<thead>
<tr>
<th>Target</th>
<th>Forward primer</th>
<th>Reverse primer</th>
</tr>
</thead>
<tbody>
<tr>
<td>β-actin</td>
<td>AAATCGTGCGTGACATCAA</td>
<td>AAGGAAGGCTGGAAAGAGC</td>
</tr>
<tr>
<td>BraT</td>
<td>CTGGACTTGAGGACGCTGT</td>
<td>TGACCTTGCTGAAAGACACAG</td>
</tr>
<tr>
<td>GATA-4</td>
<td>AAAACCGGAAGCGCAAGTCT</td>
<td>TTAGTGAGGTGGAGAAGGCTGAGT</td>
</tr>
<tr>
<td>Gli1</td>
<td>CCAAGCCAATTTGTCAGGA</td>
<td>ACGATTCGATGTGACCAAAG</td>
</tr>
<tr>
<td>Gli/EnR</td>
<td>GGAGAGGTTGGAGCCAGTA</td>
<td>CAGGATTCGGTGATCTGCT</td>
</tr>
<tr>
<td>Gli2</td>
<td>CAACGCCAATTCCAGAGC</td>
<td>AGATGTTATCTGATGTTACCA</td>
</tr>
<tr>
<td>MEF2C</td>
<td>TCTGTCTGAGGAGGCACTAG</td>
<td>GGCTGTAGGTGAGGCTGAGT</td>
</tr>
<tr>
<td>MHC6</td>
<td>CAACAAACCATAGAAGCTAG</td>
<td>ACATACAAAGGAGCTAGT</td>
</tr>
<tr>
<td>Nkx2-5</td>
<td>AAGGAAGCGAACGCACTGTC</td>
<td>GCTCGAGAATGTTAAATGTCGA</td>
</tr>
<tr>
<td>Oct-4</td>
<td>CAGCCACGACCAATCGTCT</td>
<td>GTCATGCGTGCTGAGT</td>
</tr>
<tr>
<td>Pch1</td>
<td>AAGAAGAATCGCGCAAGTTT</td>
<td>CTGCTATCTATATGCTGAGT</td>
</tr>
<tr>
<td>Tbx1</td>
<td>CTTCTCGGTCGAGTATGAC</td>
<td>TGATGGTGAGGCTGAGT</td>
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represent mean ± standard error of mean (SEM) from at least two independent biological experiments and using two clonal populations per cell line.

**Northern blot analysis**

RNA from differentiating P19 and P19[NKx-MEF2C/EnR] cells was harvested using the LiCl/Urea method and analysed using northern blot procedure as described previously (50). Briefly, a total of 12 µg of RNA for each sample was separated by denaturing gel electrophoresis and transferred to Hybond-N nylon membranes (GE Life Sciences, Canada). The membranes were hybridized with α-32P deoxyctydine 5'-triphosphate labelled MEF2C/EnR and 18S cDNA fragments described previously (21). Results shown are representative of three separate clones for each cell line.

**Immunoblot analysis**

Total protein extracts from differentiating mES and P19 EC cells and whole eyes of E16 Gli2+/− heterozygous and Gli2−/− homozygous mice were harvested using radioimmunoprecipitation assay buffer. In total, 10–20 µg of protein was resolved using 4–12% gradient NUPAGE gels (Invitrogen, Canada) according to manufacturer’s protocol using 3-(N-morpholino)propanesulfonic acid/sodium dodecyl sulfate running buffer. Resolved proteins were transferred to polyvinylidene fluoride or nitrocellulose membranes, blocked in 5% milk and reacted with Gli2 (54), calmodulin binding protein (CBP) (Millipore, MA, USA), MEF2C (Santa Cruz, sc-13266), α-tubulin (Sigma, DM1A) or β-actin specific (Sigma, AC-74) antibodies. Signal was detected using HRP-conjugated secondary anti-rabbit (Millipore, MA, USA), anti-mouse (Cell Signalling, MA, USA) or anti-goat (Santa Cruz, sc-2020) antibodies.

**ChIP assays**

A quantity of 50 µg of chromatin from Day 5 differentiating P19[MEF2C] cells and 150 µg of chromatin from Day 4 differentiating P19[Gli2] cells was immunoprecipitated using 2 µg of MEF2C specific (Santa Cruz, sc-13266), Gli2 specific (Santa Cruz, G-20) or goat IgG non-specific antibodies (Invitrogen, Canada) as described in ref. (51). Briefly, cells were cross-linked with 4% formaldehyde (Fisher Scientific, Canada) and chromatin was sheared as described in ref. (51). Sheared chromatin was incubated with Gli2, MEF2C or IgG antibodies and the immune complexes were captured using protein G sepharose beads as described previously (51). Gli2, MEF2C or IgG-bound chromatin was quantified as a percent chromatin input using QPCR analysis as described above. To be considered a true association, each ChIP sample was examined for the enrichment of a chromatin locus immunoprecipitated with a specific antibody and compared with the same chromatin locus immunoprecipitated with a non-specific IgG (ANOVA with P < 0.05), and compared with a negative control locus, such as the Ascl1 (55) or Hdac4 locus (ANOVA followed by post hoc Tukey honestly significant difference (HSD) test with P < 0.05). Data represent mean ± SEM from three independent biological experiments. Primers are listed in Table 2.

**Co-immunoprecipitation assays**

HEK-293 cells (ATCC, #CRL-1573) were seeded at 1 million cells per 100 mm tissue-culture grade dish and 24 h later co-transfected with a total amount of 3 µg of

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**Table 2. Oligonucleotide sequences of primers utilized for ChIP-QPCR experiments**

<table>
<thead>
<tr>
<th>Target gene</th>
<th>Position of BS in mm9 genome</th>
<th>Strand Orientation</th>
<th>Forward primer</th>
<th>Reverse primer</th>
</tr>
</thead>
<tbody>
<tr>
<td>Me2c A</td>
<td>Chr13: 83556197 - 83556209</td>
<td>+</td>
<td>TGGAAAAGGGAAATATCCCACTTAGA</td>
<td>TGGCATGGGTTTCACACCTAAG</td>
</tr>
<tr>
<td>Me2c B</td>
<td>Chr13: 83589537 - 83589549</td>
<td>+</td>
<td>AGTGGGCTGACCCATCGTTC</td>
<td>TTCTCCGGAACGTACTTCC</td>
</tr>
<tr>
<td>Me2c C</td>
<td>Chr13: 83657044 - 83657056</td>
<td>+</td>
<td>TCTCCAGTTCCCTGGGAAGGAA</td>
<td>CCTTCGCGGTAAGGATCTTGG</td>
</tr>
<tr>
<td>Me2c D</td>
<td>Chr13: 83663947 - 83663935</td>
<td>+</td>
<td>ACACACGCACACTTCGTCT</td>
<td>GACCCACGACAGAATCTCAAA</td>
</tr>
<tr>
<td>Me2c E</td>
<td>Chr13: 83734849 - 83734861</td>
<td>+</td>
<td>TTCCCCATTGAGCCTACCC</td>
<td>ACCACGGAGCAGAAGC</td>
</tr>
<tr>
<td>Me2c F</td>
<td>Chr13: 83772819 - 83772831</td>
<td>+</td>
<td>AGGGCTACTTCCGACCC</td>
<td>AACGTTTCGTCAGGTGAG</td>
</tr>
<tr>
<td>Me2c G</td>
<td>Chr13: 83800000 - 83800012</td>
<td>+</td>
<td>GAGCCCCCTCTCTAAATTC</td>
<td>TTGGGAGCTATGTGTCTGG</td>
</tr>
<tr>
<td>Me2c H</td>
<td>Chr13: 83803968 - 83803980</td>
<td>+</td>
<td>CGACGAGGCTGCTTACGT</td>
<td>AGATGCACATTTGGGCTGCT</td>
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<tr>
<td>Me2c I</td>
<td>Chr13: 83879226 - 83879238</td>
<td>+</td>
<td>CTTCCAGTTCCCTGGGAAGGAA</td>
<td>CCTTAATTTCCCTAAGGGAAG</td>
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<tr>
<td>Ptc1 A</td>
<td>Chr13: 86667821 - 86667833</td>
<td>−</td>
<td>TATTGCAGTCGAGGAGGTT</td>
<td>GGAGGGCGAAATTTACTCACG</td>
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<tr>
<td>Gli1 A</td>
<td>Chr13: 126778677 - 126778689</td>
<td>−</td>
<td>GCCACCCCTCTCTACCTTCTATC</td>
<td>GCCACCCAGCGAGACGGCAAC</td>
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<tr>
<td>Ascl1 A</td>
<td>Chr13: 86936603 - 86936615</td>
<td>−</td>
<td>CCTAAGATCAATGGGCGCAA</td>
<td>CCCCAACCATGCTCAGAG</td>
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<tr>
<td>Gli2 A</td>
<td>Chr13: 120906302 - 120906324</td>
<td>−</td>
<td>AACAGGGGCTCTCTCCTCAGAAC</td>
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<td>Gli2 B</td>
<td>Chr13: 12088621 - 12088643</td>
<td>−</td>
<td>GCATTCTCTAAGCTTGGTG</td>
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<td>Gli2 C</td>
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<td>Gli2 F</td>
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<tr>
<td>Gata4 A</td>
<td>Chr13: 63864687 - 63864697</td>
<td>+</td>
<td>AAGGGAGCTCTTCTCTCCTC</td>
<td>GTTGGAGGCTACGGGAGGTA</td>
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<tr>
<td>MyoG A</td>
<td>Chr13: 136186098 - 136186104</td>
<td>+</td>
<td>ATTTGCCCTTTCGTTTCTCTC</td>
<td>GCCTACACGACCTTAAAC</td>
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<tr>
<td>Hdac4 A</td>
<td>Chr13: 93901621 - 93901643</td>
<td>+</td>
<td>ATCTCCACTGTTGGTCCTG</td>
<td>GGTGGTCTATTTGATGCTTG</td>
</tr>
</tbody>
</table>

Chr: chromosome; BS: binding site.
Gli2, MEF2C-Flag or Flag-vector (DYKDDDDK flag peptide) using Eugene (Roche, Canada). On completing 24 h post-transfection, cells were washed twice with ice-cold phosphate buffered saline and lysed with buffer containing 50 mM Tris–HCl, pH 7.4, 150 mM NaCl, 1 mM EDTA and 1% Triton X-100. Lysates were clarified by centrifugation for 15 min at 13 krpm. In total, 0.5 mg of protein was subjected to FLAG immunoprecipitation (IP) according to manufacturer’s protocol (Sigma-Aldrich, MO, USA). Bound proteins were eluted by boiling the beads in 1 × SDS–PAGE loading buffer for 10 min at 95°C. Transfection efficiency was monitored by co-transfecting Green fluorescent protein (GFP) and analysing its autofluorescence. All protein extraction and incubation procedures were carried out at 4°C in the presence of 1 × protease inhibitor cocktail (Roche, Canada) and 0.5 mM phenylmethanesulphonylfluoride (PMSF).

Eluted proteins from co-immunoprecipitation assays in HEK-293 cells were resolved using 4–12% gradient NuPAGE gels, transferred to polyvinylidene fluoride membranes and reacted with Gli2- or MEF2C-specific antibodies as described above. Gli2 purification efficiency was quantified by band densitometry analysis using ImageJ program (56).

**Reporter assays**

Gli2, MEF2C-Flag and GATA-4 expressing plasmids are described elsewhere (25,47,57). Gli-responsive promoter is described in ref. (58) and MEF2-responsive promoter is from Affymetrix, CA, USA. The Nkx2-5 promoter, comprising −3059 to +223 nt relative to the Nkx2-5 transcriptional start site, termed Nkx2-5-luc, is described in Ref. (59).

P19 EC cells were plated at a density of 150,000 cells/35 mm tissue-culture grade dish and transiently co-transfected, as described above, 24 h later with a total amount of 3 μg of DNA with or without Gli2, MEF2C-Flag and the Nkx2-5 promoter. Gli-responsive promoter or MEF2-responsive promoter driving luciferase gene in the ratio 2:1 relative to luciferase reporters. Transfection efficiency was monitored by transfecting Renilla as described previously (51). On completing 24 h after transfection, cells were washed twice with ice-cold phosphate buffered saline and lysed according to Dual Luciferase Kit protocol (Promega, WI, USA). Luciferase activity was assayed using 10–15 μl of lysate and LmaxII384 luminometer (Molecular Devices, USA). To remove the background, the normalized activity of the luciferase reporter plasmid alone was subtracted from its activity in the presence of Gli2 and/or MEF2C and GATA-4. The activity of the luciferase reporter in the presence of protein expression plasmids was normalized to the activity in the presence of Gli2 (arbitrarily set at 1). Synergy was calculated as described in ref. (60) according to the following formula:

\[
\text{Synergy} = \frac{\text{RLUs(MEF2C+Gli2)}}{\text{RLUs(MEF2C)} + \text{RLUs(Gli2)}}
\]

where relative luciferase units (RLUs) represent luciferase activity normalized to *Renilla* activity. P19 cells transfected with GATA-4 and Nkx2-5-luc in ratio 2:1 served as a positive control.

**Bioinformatics analysis**

Conserved Gli DNA binding sites and MEF2 DNA binding sites in the *Mef2c* and *Gli2* genes (±100 kb) were identified using Mulan (Multiple Sequence Local Alignment and Visualization tool) as described previously (61). The primers for identified binding sites were designed using Primer 3 software (62) (for primer sequences, see Table 2).

Putative genomic targets for both Gli and MEF2 transcription factors were identified using SynoR (Genome miner for synonymous regulation) as described previously (63). The distance between neighbouring DNA binding sites was set to (i) 4–100 bp; (ii) 100–200 bp and (iii) 200–300 bp. The mouse genome (mm9) was used as a base and was compared with the human genome (hg18). Identified genes are listed in Supplementary Table S1. Functional annotation analysis of the identified potential targets was performed using DAVID (Database for Annotation, Visualization, and Integrated Discovery) software as described in Refs. (64,65). Selected gene ontology biological processes are listed in Table 3.

**Statistical analysis**

ANOVA followed by post hoc Tukey HSD test was performed using XLSTAT11 software (Addinsoft, NY, USA) to determine statistical significance between mean values of two groups (*P < 0.05; **P < 0.01).

**RESULTS**

Gli2 and MEF2C are expressed during cardiomyogenesis in mES and P19 EC cells

Differentiation of mES cells into cardiac muscle is schematically depicted in Figure 1A, using a 7-day aggregation protocol (25), where cardiac muscle forms as early as Days 8–9 (Figure 1A). Formation of cardiac muscle from mES cells was confirmed by indirect IF with an anti-MHC antibody, MF20 (Figure 1B).

Changes in gene expression during cardiomyogenesis in mES cells were examined for several markers (Figure 1C). Downregulation of Oct-4, a marker of pluripotency in ES cells (66), confirmed that mES cells differentiated efficiently (Figure 1C, panel Oct-4). Notably, a slight upregulation of Oct-4 was observed on Day 3, in agreement with a previously reported role for Oct-4 in cardiac mesoderm formation (67). The mesodermal marker BrAT (68), was also upregulated on Day 3 of mES differentiation (Figure 1C, panel BrAT). The highest expression of the cardiac muscle precursor transcription factors Tbx5, Nkx2-5 and GATA-4 (69–71) was detected during Days 6–9 of mES differentiation (Figure 1C, panels Tbx5, Nkx2-5 and GATA-4). Finally, the highest expression of the cardiac muscle marker MHC6 was detected on Day 9 of mES differentiation (Figure 1C, panel MHC6). During
mesenchymal stem cells (mES) differentiation, MEF2C mRNA levels became highly upregulated starting from Day 6 (Figure 1D, MEF2C panel), whereas Gli2 mRNA levels were downregulated starting from Day 3 (Figure 1D, Gli2 panel). There was a trend in Gli1 mRNA downregulation on Days 3–6 of differentiation (Figure 1D, panel Gli1). In contrast, Ptch1

Table 3. Selected gene ontology biological processes significantly enriched among genes containing Gli and MEF2 conserved DNA binding clusters in UTR, promoter, intron and coding sequence regions

<table>
<thead>
<tr>
<th>Category</th>
<th>Targets</th>
<th>Fisher’s exact P-value</th>
<th>Example genes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cellular process</td>
<td>562</td>
<td>2.1E-17</td>
<td>E2F4, Glis1, CCNB2, Hdac9, RBBP4, TCF4</td>
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<tr>
<td>System development</td>
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<td>6.1E-19</td>
<td>Tbx15, Fgf10, GSK3b, VEGFC, Sox6, Bmp1, Ar</td>
</tr>
<tr>
<td>Regulation of gene expression</td>
<td>172</td>
<td>6.4E-7</td>
<td>DNM1T3A, POU4F2, Smarca2, FOXA2, FOXD3, HOXD10, Lhx2, Phrn1</td>
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<tr>
<td>Cell differentiation</td>
<td>143</td>
<td>4.5E-13</td>
<td>ANGPT1, Bmp7, Fgf1, PAX2, Ptc1, Hhip, Twist1</td>
</tr>
<tr>
<td>Nervous system development</td>
<td>99</td>
<td>3.3E-15</td>
<td>BDNF, NRXN1, FOXP2, Gfra1, HOXB3</td>
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<tr>
<td>Embryonic development</td>
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<td>Nkx2, Pdgfr, Pdgfr2, HOXD10, ITGA8, Nr4a3</td>
</tr>
<tr>
<td>Neurogenesis</td>
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<td>8.5E-9</td>
<td>Ntn11, NAV1, Lhx8, APP, Ctnn2</td>
</tr>
<tr>
<td>Skeletal system development</td>
<td>31</td>
<td>9.1E-5</td>
<td>Bmp5, Sox5, HOXB6, Tapt1</td>
</tr>
<tr>
<td>Chromatin organization</td>
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<td>9.2E-4</td>
<td>MYST4, Pht15, ASH11, ING2, Rnf2</td>
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<tr>
<td>Heart development</td>
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<td>MEF2C, FOXP1, TTN, Hdac5, ALPK3</td>
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<tr>
<td>Muscle organ development</td>
<td>17</td>
<td>1.5E-2</td>
<td>Dmd, Utrn, Rora, CACNG2</td>
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<tr>
<td>Vascular smooth muscle</td>
<td>16</td>
<td>5.1E-4</td>
<td>Cacna1C, Cacna1D, BRAF, Prkca</td>
</tr>
</tbody>
</table>

For complete list of genes, see Supplementary Data. Altogether, 957 potential target genes were analysed as described in ‘Materials and Methods’ section. The background set of genes used was the entire mouse genome.

Figure 1. Gli2 and MEF2C are expressed during mES differentiation. (A) Schematic representation of cardiomyogenesis in D3 mES cells, as described in ‘Materials and Methods’ section. (B–E) mES cells were differentiated and examined (B) for MHC expression using MF20 antibodies on Day 9 of differentiation, n = 3. Error bars represent ± SEM. Primers are listed in Table 1. (C) Values were compared with Day 0 values and statistical significance was determined using ANOVA (*P < 0.05 and **P < 0.01). (D) To determine statistical significance between all days analysed, ANOVA followed by post hoc Tukey HSD analysis was performed, *P < 0.05 and **P < 0.01. n.s. = not significant.; (E) Immunoblot analysis of total protein extracts using MEF2C- and Gli2-specific antibodies at the time shown. Arrows designate Gli2 or MEF2C protein band(s), and asterisk denotes non-specific binding of the Gli2 antibodies. α-Tubulin served as a loading control. Lanes were spliced out from the same autoradiogram as designated by vertical lines. Gli2 band densities were measured from one representative experiment using ImageJ program (56), normalized to α-tubulin and expressed as percent maximum; (F) Immunoblot analysis of eye total protein extracts from E16 Gli2+/C0 and Gli2/C0/C0 mice using Gli2-specific antibodies. B-ACTIN served as a loading control.

mES differentiation, MEF2C mRNA levels became highly upregulated starting from Day 6 (Figure 1D, MEF2C panel), whereas Gli2 mRNA levels were downregulated starting from Day 3 (Figure 1D, Gli2 panel). There was a trend in Gli1 mRNA downregulation on Days 3–6 of differentiation (Figure 1D, panel Gli1). In contrast, Ptch1
mRNA was upregulated on Days 6–9 of mES differentiation (Figure 1D, panel Ptc1). MEF2C and Gli2 protein expression levels correlated with their mRNA expression levels (Figure 1E). MEF2C protein expression was induced after Day 3 of differentiation and Gli2 protein expression was downregulated after Day 3, although its expression persisted until Day 9 (Figure 1E).

To confirm that the upper band on the Gli2 immunoblot did indeed correspond to Gli2 protein, we analysed whole eye total protein extracts from Gli2<sup>+/−</sup> heterozygous and Gli2<sup>−/−</sup> homozygous mice. The only band absent in the Gli2<sup>−/−</sup> protein sample was the upper band corresponding to ~180 kDa (Figure 1F). Thus, the upper band represents Gli2 that correlates well with previously published data (54,55) and with the theoretical molecular weight of 165 kDa for full-length mouse Gli2 protein (database accession number Q0VG1). Therefore, both Gli2 and MEF2C proteins were expressed during mES cell cardiomyogenesis (summarized in Table 4).

Since mES cells spontaneously differentiate into cell types of all three germ layers upon removal of leukemia inhibitory factor (77) and only a small proportion of cells differentiate into cardiac muscle using a standard embryoid body differentiation protocol, it is difficult to specifically correlate the expression profile of MEF2C and Gli2 with cardiac myogenic differentiation in a heterogeneous population of mES cells. P19 EC cells, in contrast, mainly differentiate into mesoderm and endothelial lineages upon addition of DMSO (27,30–34). Moreover, P19 EC myogenic differentiation is very similar to mES myogenic differentiation (25,35). For these reasons, we compared the expression pattern of Gli2 and MEF2C during mES differentiation with DMSO-induced differentiation of P19 EC cells.

Cardiomyogenic differentiation of P19 EC cells is schematically presented in Figure 2A. P19 EC cells treated with DMSO successfully differentiated into cardiac muscle on Day 6 as observed by indirect IF using MHC-specific antibodies (Figure 2B). Gene expression changes during cardiomyogenesis in P19 EC cells were followed by QPCR analysis of myogenic markers (Figure 2C). Downregulation of Oct-4 starting from Day 2 indicated the loss of pluripotency in EC cells (Figure 2C, panel Oct-4). Induction of mesoderm is indicated by the robust upregulation of BraT expression on Day 2 of differentiation (Figure 2C, panel BraT). Induction of cardiomyogenesis was confirmed by elevated expression of Nkx2-5, Tbx5 and GATA-4 during Days 5–6 of P19 EC differentiation (Figure 2C, panels Nkx2-5, Tbx5 and GATA4). The robust expression of MHC6 on Day 6 of differentiation confirmed the formation of cardiac muscle in EC cells (Figure 2C, panel MHC6). The expression of MEF2C mRNA was significantly elevated (<i>P</i> < 0.01) during Days 5–6 of DMSO-induced differentiation of P19 EC cells (Figure 2D, panel MEF2C), and resembled the temporal pattern observed in mES cells (Figure 1D, panel MEF2C). There was a significant increase (<i>P</i> < 0.05) in Gli2 mRNA expression during Days 5–6 of P19 EC cardiomyogenesis (Figure 2D, panel Gli2). Gli1 mRNA was upregulated on Day 4 of differentiation (Figure 2D, panel Gli1). This data correlates with previous reports showing upregulation of MEF2C and Gli2 mRNA expression during P19 EC DMSO-induced myogenesis using northern blot analysis (35,41,42,47,51,78). Therefore, MEF2C and Gli2 transcription factors are expressed during mES and P19 EC cell cardiomyogenesis (summarized in Table 4).

**Gli2 regulates expression of MEF2C during cardiomyogenesis in P19 EC cells**

To study the ability of Gli2 to regulate the expression of MEF2C during cardiomyogenesis in stem cells, we examined P19 cells that stably overexpressed Gli2 (Figure 3). Stable overexpression of Gli2 was confirmed in Table 4. Summary of gene expression changes in cell lines treated with or without DMSO

<table>
<thead>
<tr>
<th>Cell line</th>
<th>Treatment</th>
<th>BraT</th>
<th>Nkx2-5</th>
<th>GATA-4</th>
<th>Tbx5</th>
<th>MHC6</th>
<th>Gli2</th>
<th>MEF2C</th>
<th>References</th>
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</thead>
<tbody>
<tr>
<td>mES</td>
<td>–</td>
<td>+D3</td>
<td>+D6–9</td>
<td>+D6–9</td>
<td>+D6–9</td>
<td>+D9</td>
<td>+D6–9</td>
<td>+D9</td>
<td>+D6–9</td>
</tr>
<tr>
<td>P19</td>
<td>DMSO</td>
<td>+D1–3</td>
<td>+D4–6</td>
<td>+D3–6</td>
<td>+D4–6</td>
<td>+D5–6</td>
<td>+D4–6</td>
<td>+D4–6</td>
<td>+D4–6</td>
</tr>
<tr>
<td>P19[Gli2]</td>
<td>DMSO</td>
<td>±D2</td>
<td>+D6</td>
<td>++D6</td>
<td>+D6</td>
<td>+D6</td>
<td>+D6</td>
<td>+D6</td>
<td>+D6–9</td>
</tr>
<tr>
<td>P19[MEF2C-TAP]</td>
<td>DMSO</td>
<td>+D2</td>
<td>+D6</td>
<td>++D6</td>
<td>+D6</td>
<td>+D6</td>
<td>+D6</td>
<td>+D6</td>
<td>+D6–4</td>
</tr>
<tr>
<td>P19[Gli/EnR]</td>
<td>DMSO</td>
<td>+D0–9</td>
<td>±D4–6</td>
<td>±D4–6</td>
<td>±D4–6</td>
<td>±D4–6</td>
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<td>±D4–6</td>
<td>±D4–6</td>
</tr>
<tr>
<td>P19[Nkx-MEF2C/EnR]</td>
<td>DMSO</td>
<td>+D2–6</td>
<td>±D0–6</td>
<td>±D0–6</td>
<td>±D0–6</td>
<td>±D0–6</td>
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<tr>
<td>P19[β-caten/EnR]</td>
<td>DMSO</td>
<td>±D0–9</td>
<td>N.D.</td>
<td>N.D.</td>
<td>N.D.</td>
<td>N.D.</td>
<td>N.D.</td>
<td>N.D.</td>
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</tr>
<tr>
<td>P19[ Pax/EnR]</td>
<td>DMSO</td>
<td>±D0–9</td>
<td>N.D.</td>
<td>+D9</td>
<td>N.D.</td>
<td>N.D.</td>
<td>N.D.</td>
<td>N.D.</td>
<td>N.D.</td>
</tr>
<tr>
<td>P19[Mef2c/EnR]</td>
<td>DMSO</td>
<td>±D0–9</td>
<td>N.D.</td>
<td>+D6</td>
<td>N.D.</td>
<td>N.D.</td>
<td>N.D.</td>
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</tr>
<tr>
<td>P19[Myod1/EnR]</td>
<td>DMSO</td>
<td>N.D.</td>
<td>N.D.</td>
<td>±D9</td>
<td>N.D.</td>
<td>N.D.</td>
<td>±D9</td>
<td>–D9</td>
<td>(76)</td>
</tr>
<tr>
<td>P19[EnR]</td>
<td>DMSO</td>
<td>±D0–6</td>
<td>±D0–6</td>
<td>±D0–6</td>
<td>N.D.</td>
<td>N.D.</td>
<td>±D0–6</td>
<td>±D0–6</td>
<td>(40)</td>
</tr>
</tbody>
</table>

* The expression of Nkx2-5 and MEF2C was elevated during Days 0–3, whereas it was downregulated during Days 4–6 of P19[Nkx-MEF2C/EnR] differentiation as compared with P19 control cells [ref. (21) and Figure 7C].

Cell lines indicated on the right were aggregated under the conditions described in ‘Materials and Methods’ section and the induction of muscle marker gene expression was monitored. (+ + +: high expression; +: elevated expression; ±: normal expression; ±: downregulated expression; –: not expressed or basal level of expression; N.D.: not determined). The timing of temporal gene expression in parental mES and P19 EC cells is shown. For stable cell lines, the change in expression relative to the respective control cell lines and the day examined, were indicated (D = day).
by immunoblot analysis using Gli2-specific antibodies (Figure 3A). Antigenic analysis using MHC-specific antibodies revealed an enhancement in the formation of cardiomyocytes in Day 6 differentiated P19[Gli2] cultures (Figure 3B and C). This was confirmed by QPCR analysis, which showed increased levels of Nkx2-5, Tbx5, GATA-4 and MHC6 mRNA in P19[Gli2] cells as compared with P19[Control] cells (Figure 3D, panels Nkx2-5, Tbx5, GATA-4 and MHC6) (summarized in Table 4). Enhancement of cardiomyogenesis by Gli2 is consistent with previous findings (3,41). Notably, overexpression of Gli2 resulted in a slight downregulation (15 ± 4%) of BraT expression on Day 2 of differentiation (Figure 3D, panel BraT). Furthermore, P19[Gli2] cells overexpressing Gli2 protein showed a statistically significant \( P < 0.05 \) upregulation of MEF2C mRNA levels on Day 4 of DMSO-induced differentiation (Figure 3D, panel MEF2C), similar to results shown in the absence of DMSO (41). There was also an 11- to 20-fold upregulation of MEF2C protein levels in P19[Gli2] cells on Days 6–9 of DMSO-induced differentiation as compared to P19[Control] Day 4 (Figure 3A). The increases in MEF2C protein expression on Days 6–9 in P19 and P19[Gli2] cells (Figure 3A) correlate with the pattern of MEF2C mRNA expression in P19 cells (compare with Figure 2D), which peaks by Day 5. The increase in MEF2C mRNA on Day 4 in P19[Gli2] (Figure 3D) cells did not correlate with an obvious increase in MEF2C protein on that day (Figure 3A), possibly due to post-transcriptional regulation of the MEF2C protein [reviewed in ref. (15)]. Overall, overexpression of Gli2 upregulates MEF2C mRNA and protein expression while enhancing cardiomyogenesis (summarized in Table 4).

Since Gli transcription factors have overlapping functions both in vivo and in vitro (79–82), the use of a dominant negative fusion protein of Gli2 with the EnR domain (Gli/EnR) would result in repression of all Gli2-bound regulatory elements and the inability of Gli1 or Gli3 to rescue the phenotype. This approach has successfully been used previously both in vitro and in vivo (21,74,75,83,84). Furthermore, overexpression of Engrailed does not interfere with cardiomyogenesis in P19 EC cells (40). Stable overexpression of Gli/EnR (Figure 4C, panel Gli/EnR) resulted in the formation of fewer MHC-positive cardiomyocytes when compared with P19 control cells (Figure 4A and B). Downregulation of cardiac muscle cell differentiation was confirmed by lower...
expression of MHC6 (Figure 4C, panel MHC6). The mRNA levels of Tbx5 and GATA-4 were downregulated since Day 4, whereas levels of Nkx2-5 were downregulated most significantly ($P < 0.05$) on Day 6 of P19[Gli/EnR] cellular differentiation (Figure 4C, panels Tbx5, Nkx2-5 and GATA-4). MEF2C mRNA was downregulated in P19[Gli/EnR] cells on Days 5 and 6 of differentiation (Figure 4C, panel MEF2C). The effect of MEF2C downregulation is not due to global gene repression by Gli/EnR fusion protein since Gli/EnR did not greatly affect expression of mesodermal markers such as BraT in DMSO-induced differentiation (47) (Table 4). P19[Gli/EnR] cells did not lose pluripotency or the ability to differentiate as they were shown to differentiate into glial and neuronal cells in the presence of retinoic acid (55). Therefore, Gli2, or genes bound by Gli2, is essential for maintaining normal expression of MEF2C during cardiomyogenesis in P19 EC cells (summarized in Table 4).

**Gli2 associates with Mef2c gene elements during cardiomyogenesis in P19 EC cells**

In order to determine if Gli2 binds Mef2c gene elements during cardiomyogenesis, we performed ChIP experiments. *In silico* analysis of the Mef2c gene (±100 kb) revealed nine conserved theoretical Gli binding sites (GBS) (Mef2c A–I, Figure 5A and B, summarized in Table 2), suggesting that MEF2C could be a target of a Gli transcription factor. Since the highest expression of Gli2 protein and of MEF2C mRNA was detected on Day 4 of P19[Gli2] cellular differentiation (Figure 3A and D, respectively), we performed ChIP analysis of Day 4 differentiating P19[Gli2] EC cells. After ChIP with a Gli2 antibody, a statistically significant enrichment was observed of chromatin fragments, corresponding to Mef2c sites B–I (Figure 5C). Mef2c site A appeared not to be associated with Gli2 protein (Figure 5C), therefore showing specificity of the ChIP assay. The Gili and Ptchl promoter regions were used as positive controls and Ascl1 gene element was used as a negative control (Figure 5C) based on previous observations (55,85,86). From this result, we observe for the first time that Gli2 is associated with Mef2c gene elements during cardiomyogenesis in P19 EC cells.

**MEF2C regulates expression of Gli2 during cardiomyogenesis in P19 EC cells**

To study the ability of MEF2C to regulate the expression of Gli2 during cardiomyogenesis *in vitro*, we examined P19 cells that stably overexpressed MEF2C-TAP [tandem affinity purification tag (25)] (Figure 6). Stable overexpression of MEF2C-TAP was confirmed by immunoblot analysis using CBP-specific antibodies that recognized the TAP-tag of the MEF2C-TAP protein (Figure 6A). Antigenic analysis using MHC-specific antibodies revealed an enhancement in cardiac muscle formation in P19[MEF2C-TAP] cells when compared with their respective control cells (Figure 6B and C). This is in agreement with previous findings, where MEF2C was shown to induce cardiomyogenesis in the absence of DMSO (45). We also observed significantly ($P < 0.01$)

![Figure 3](https://academic.oup.com/nar/article-abstract/40/8/3329/2411328/3337)
increased levels of Nkx2-5 and GATA-4, as well as a trend of increased MHC6 transcripts in Day 6 differentiated P19[MEF2C-TAP] cultures as compared with their control cell line (Figure 6D, panels Nkx2-5, GATA-4 and MHC6). Notably, Tbx5 mRNA was not affected by the expression of MEF2C-TAP (Figure 6D, panel Tbx5). Also, expression of MEF2C-TAP did not affect BraT transcript levels, suggesting that MEF2C-TAP affects cardiomyogenesis after mesoderm induction (Figure 6D, panel BraT). Furthermore, overexpression of MEF2C-TAP resulted in the elevation of Gli2 mRNA expression on Day 4 of P19 EC differentiation (Figure 3D, panel Gli2). We also observed an increase in Gli2 protein levels up to 2.4-fold on Days 5 and 6 of P19[MEF2C-TAP] differentiation as compared with Day 5 of differentiation in P19[TAP] cells (Figure 6A). Therefore, MEF2C upregulates Gli2 expression while enhancing cardiomyogenesis in P19 EC cells (summarized in Table 4).

To test if the expression of Gli2 can be regulated by a dominant-negative MEF2C, we examined P19 EC cells expressing MEF2C/EnR under the regulation of the Nkx2-5 enhancer (Figure 7A). As reported previously (21), P19[Nkx-MEF2C/EnR] cells failed to undergo cardiomyogenesis as evidenced by the absence of MHC-positive cardiac muscle cells (Figure 7B) and a severe downregulation of Nkx2-5, Tbx5, GATA-4 and MHC6 mRNA (Figure 7C, panels Nkx2-5, GATA-4, Tbx5 and MHC6). The phenotype of P19[Nkx-MEF2C/EnR] cells is in accordance with previous reports showing complete absence of cardiomyogenesis in vivo or in P19 EC cells (21). Expression of Nkx-MEF2C/EnR affected cardiomyogenesis only, as differentiating P19[Nkx-MEF2C/EnR] cells showed normal upregulation of
Pax3 and Myf5 transcripts (data not shown), therefore indicating normal skeletal myogenesis. Gli2 mRNA was downregulated in P19[Nkx-MEF2C/EnR] cells on Day 6 of differentiation (Figure 7C, panel Gli2). The loss of Gli2 appears to be specific for a subset of dominant-negative transcription factors, including Nkx-MEF2C/EnR (Figure 7), which caused inhibition of cardiomyogenesis,

Figure 5. Gli2 associates with Mef2c gene elements during cardiomyogenesis in vitro. (A) Custom tracks of murine Mef2c, Gli1, Ptc1 and Ascl1 genes using UCSC genome browser (http://genome.ucsc.edu). Letters designate conserved Gli binding sites (GBS), their genomic positions are listed in Table 2. Listed genes (±100 kb) from mouse and human genomes were searched for conserved theoretical GBS as described in ref. (61); (B) comparison of mouse and human sequences of Mef2c A-I sites from (A). The sequence of the Gli2 binding site is marked in bold. (C) ChIP analysis of Gli2-bound Mef2c, Gli1, Ptc1 and Ascl1 genes on Day 4 of P19[Gli2] cellular differentiation in the presence of 1% DMSO. Black bars designate genomic regions immunoprecipitated with Gli2-specific antibodies, and grey bars designate genomic regions precipitated with IgG-non-specific antibodies. Gli1 and Ptc1 promoters served as positive controls, Ascl1 gene element served as a negative control. Percent chromatin input was calculated using QPCR analysis and primers listed in Table 2. Error bars represent ±SEM from three biological replicas (n = 3). Statistical significance was determined as described in ‘Materials and Methods’ section. *P < 0.05, **P < 0.01.

Figure 6. MEF2C upregulates Gli2 expression while enhancing cardiomyogenesis in P19 EC cells. P19[MEF2C-TAP] and P19[TAP] cell lines were differentiated in the presence of 1% DMSO. (A) Immunoblot analyses of MEF2C-TAP and Gli2 protein expression in differentiating P19[MEF2C-TAP] and P19[TAP] cells on Days 5, 6 and 9 using Gli2-specific or CBP-specific antibodies. Arrow designates Gli2 protein band, and asterisk denotes non-specific binding of the Gli2 antibodies. α-Tubulin served as a loading control. Lanes were spliced out from the same autoradiogram as designated by vertical lines. Gli2 band densities were measured from one representative experiment using ImageJ program (56) and normalized to α-tubulin. Gli2 band density in P19[TAP] Day 5 sample was set at 100%; (B) Day 6 differentiated P19[MEF2C-TAP] and P19[TAP] cultures were reacted with MHC-specific antibodies to detect MHC expression. Nuclei were stained with Hoechst dye. Scale bar is 30 µM; (C) MHC-positive cardiomyocytes from (B) were counted in 10 random fields and expressed as percent of the total number of nuclei, n = 4. In total, 14,000 cells were counted; (D) QPCR analysis of the expression of the genes indicated at the times shown in P19[TAP] (black bars) and P19[MEF2C-TAP] (grey bars) cultures. Error bars represent ±SEM (continued)
and MyoD/EnR (76), Meox/EnR (47), and β-catenin/EnR (74), which caused inhibition of skeletal myogenesis (Table 4). In contrast, Gli2 levels were not downregulated in P19 cells expressing Pax/EnR (75) that resulted in the loss of skeletal myogenesis (Table 4). Thus, MEF2C, or genes bound by MEF2C, is essential for maintaining normal Gli2 expression during cardiomyogenesis in P19 EC cells (summarized in Table 4).

**MEF2C associates with Gli2 gene elements during cardiomyogenesis in stem cells**

To test if MEF2C can bind Gli2 gene elements, *in silico* analysis of the Gli2 gene (±100 kb) revealed six conserved theoretical MEF2 binding sites (MBS) (Gli2 A–F, Figure 8A and B, summarized in Table 2) suggesting that Gli2 could be a target of MEF2. ChIP analysis of Day 5 differentiating P19[MEF2C] cells revealed statistically significant enrichment, with a MEF2C antibody, of chromatin fragments corresponding to Gli2 A–D and F sites, but not to Gli2 E site (Figure 8C). The Hdac4 intron region, which is located on the same chromosome as the Gli2 gene (summarized in Table 2), was not enriched in the ChIP assay with MEF2C antibodies (Figure 8C), therefore serving as a negative control and demonstrating the specificity of ChIP analysis using MEF2C-specific antibodies. The Gata-4 and MyoG promoter regions were used as positive controls (Figure 8C). Therefore, MEF2C is associated with Gli2 gene elements during cardiomyogenesis in P19 EC cells.

**Gli2 and MEF2C factors form a protein complex that can function synergistically**

Based on the finding that Gli2 and MEF2C are co-expressed during later stages of cardiomyogenesis in mES and during P19 EC cardiomyogenesis (Figures 1 and 2, respectively) and they regulate each other's expression, as well as have similar abilities to regulate cardiomyogenesis in P19 EC cells (Figures 3 and 6), we hypothesized that these transcription factors may form a protein complex to synergistically regulate the expression of cardiomyogenesis-related genes.

To test this hypothesis, we first explored the ability of Gli2 and MEF2C to synergize in luciferase reporter assays. To this end, we took advantage of an endogenous Nkx2-5 promoter (−3059 to +223 nt relative to Nkx2-5 transcriptional start site), which was shown to drive gene expression in the cardiac crescent at E7.25, as well as outflow tract and right ventricle at E10.5, and require combinatorial function of multiple regulatory factors (59). This Nkx2-5 promoter contained 3XGli, 3XMEF2, 6XGATA-4, 3XNkx and 1XTbx5 binding sites and was positive controls (Figure 8C). Therefore, MEF2C is associated with Gli2 gene elements during cardiomyogenesis in P19 EC cells.

**Figure 6.** Continued from two biological replicas of two clonal populations (n = 4). Statistical significance using ANOVA test was determined between groups of P19[TAP] and P19[MEF2C-TAP] samples analysed on the same day. For Gli2 expression, ANOVA followed by post hoc Tukey HSD test was performed to determine statistically significant differences between P19[TAP] and P19[MEF2C-TAP] samples on all days analysed. *P < 0.05 and **P < 0.01. Primers are listed in Table 1.

**Figure 7.** Expression of Nkx-MEF2C/EnR downregulates Gli2 expression while inhibiting cardiomyogenesis in P19 EC cells. P19[Control] and P19[Nkx-MEF2C/EnR] cell lines were differentiated in the presence of 1% DMSO. (A) Northern blot analysis of the expression of MEF2C/EnR mRNA in P19 and P19[Nkx-MEF2C/EnR] cells on Days 0 and 2–6. Here, 18s served as a loading control. Lanes were spliced out from the same autoradiogram as designated by vertical lines. (B) Antigenic analysis of P19 and P19[Nkx-MEF2C/EnR] using MHC-specific antibodies on Day 6 of DMSO-induced differentiation. Nuclei were stained with Hoechst dye. Scale bar is 60 μM. (C) QPCR analysis of the expression of Nkx2-5, GATA-4, Tbx5, MHC6 and Gli2 mRNA in differentiated P19[Control] and P19[Nkx-MEF2C/EnR] cells on Days 0 and 6. Error bars represent ±SEM from two biological replicas of two clonal populations (n = 4). Statistical significance using ANOVA test was determined between groups of P19 and P19[Nkx-MEF2C/EnR] samples analysed on the same day. For Gli2 expression, ANOVA followed by post hoc Tukey HSD test was performed to determine statistically significant differences between P19 and P19[Nkx-MEF2C/EnR] samples on all days analysed. *P < 0.05, **P < 0.01. Primers are listed in Table 1.
Gli2 and MEF2C synergistically activate the Nkx2-5 promoter.

To test if Gli2 and MEF2C formed a protein complex, we co-transfected Gli2 and MEF2C-Flag or Flag-vector into HEK-293 cells. Gli2 co-immunoprecipitated with MEF2C-Flag but not with Flag-vector when transfected in HEK-293 cells. Transfection efficiency was monitored by assaying activity of co-transfected Renilla luciferase. Synergy and statistical significance was calculated as described in ‘Materials and Methods’ section. The average activation of Nkx2-5-luc by Gli2 was 2.3 ± 0.4 (data not shown) and was normalized to 1; (C) Gli2 was co-immunoprecipitated with MEF2C-Flag but not with Flag-vector when transfected in HEK-293 cells. Transfection efficiency was monitored by assaying autofluorescence of co-transfected GFP. Arrow designates Gli2 protein band, and asterisk denotes non-specific binding of the Gli2 antibodies. Lanes were spliced out from the same autoradiogram as designated by vertical lines. (D) Quantification of Gli2 band density from elution fractions in (C) using ImageJ program. Error bars represent ±SEM from three biological replicates (n = 3), *P < 0.05.
Gli2 anomalies were detected in the developing hearts of insight into the roles of Gli2 and MEF2C in participating in the Gli2–MEF2C protein complex are designated by ‘?’. Other protein(s) expression and bind each other’s gene elements (designated by thick line).

We propose a model in which Gli2 and MEF2C induce each other’s expression, as well as form a protein complex that synergizes on promoters containing both Gli and MEF2C transcription factors induce and maintain each other’s expression during cardiomyogenesis (Figure 10). It is likely that other transcription factors participate in the Gli2–MEF2C protein complex, designated as ‘?’. Our results obtained in vitro provide added mechanistic insight into the roles of Gli2 and MEF2C in cardiomyogenesis. Whereas only cardiac outflow tract anomalies were detected in the developing hearts of Gli2+/−Gli3−/− mice, MEF2C−/− mice showed heart looping defects along with gross abnormalities in the right ventricle, as well as in the outflow tract (19,20). This suggests that Gli2 and MEF2C may function cooperatively during cardiac outflow formation. Our studies are consistent with an overlapping pattern of defects that could be explained by Gli2 and MEF2C having both shared and distinct subsets of targets, likely dependent upon the unique protein complexes formed in each case.

The results from this study suggest that both Gli2 and MEF2C may be active at the cardiac muscle progenitor stage. This is supported by the embryonic expression patterns of MEF2C (starting from E7.5) (16,17) and Gli1 (E7.0–8.0) (3,10), which are both expressed after mesodermal markers BraT (E6.5) and MESP1 (E6.5), and concomitantly with the cardiac progenitor marker Nkx2-5 (E7.0) [reviewed in ref. (87)]. Although the expression of Gli2 resulted in downregulated expression of BraT, the decrease was minimal (15 ± 4%) in comparison with the upregulation of expression of other factors (Figure 3D). Furthermore, overexpression of MEF2C, Nkx-MEF2C/EnR and Gli/EnR did not affect the expression of BraT in P19 EC cellular differentiation [refs (21,45,47) and Figure 6D], while affecting the expression of cardiac muscle progenitor markers Nkx2-5 and GATA-4 [(21,41,45) and Figures 4C, 6D and 7C].

DISCUSSION

We have shown that Gli2 and MEF2C are associated with each other’s gene elements and are able to regulate each other’s expression during cardiomyogenesis in P19 EC cells. Further, Gli2 and MEF2C form a protein complex capable of synergistically activating gene promoters that participate in cardiomyogenesis. Thus, we propose a model similar to that for MEF2 and myogenic regulatory factors (MRFs) [reviewed in ref. (15)], where Gli2 and MEF2C transcription factors induce and maintain each other’s expression, as well as form a protein complex that synergizes on promoters containing both Gli and MEF2 binding elements during cardiomyogenesis (Figure 10). It is likely that other transcription factors participate in the Gli2–MEF2C protein complex, designated as ‘?’.

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cells in the absence of DMSO (41). Recently, it was shown that the activation of the Shh signalling pathway upregulated MEF2C mRNA transcript levels in the second heart field in vivo after chick embryos were treated with Shh signalling pathway activator, SAG agonist, at HH14 (98). Drosophila embryos expressing mutant loss-of-function MEF2F protein showed downregulated expression of Ci (Cubitus Interruptus, a single Drosophila homolog of vertebrate Gli transcription factors) (99). Our results support and extend previous results by showing that Gli2 and MEF2C can upregulate each other’s expression while enhancing DMSO-induced cardiomyogenesis in P19 EC cells, suggesting the possibility that they function in a regulatory loop.

Gli2 and MEF2C enhanced each other’s mRNA expression on Day 4 of differentiation, and not in undifferentiated cells (Day 0). It was previously shown that MyoD, a master-regulator of skeletal myogenesis capable of converting fibroblasts into skeletal myocytes (100), induced skeletal myogenesis in P19 EC cells only upon cellular aggregation (101,102). Indeed, all of the muscle transcription factors studied to date in P19 EC cells required cellular aggregation to be functional (21,40,45–47,50,52,74–76,78,101–103,114). It is therefore not surprising that the expression levels of Gli2 and MEF2C were not affected in undifferentiated (Day 0) P19 stable cell lines.

The expression of dominant-negative Gli2 resulted in downregulation of MEF2C expression and cardiomyogenesis. To our knowledge, this is the first indication that the expression of Gli/EnR results in impaired cardiomyogenesis in P19 EC cells. Previous reports showed that Shh signalling is not essential for cardiomyogenesis during embryogenesis (3,4) or in P19 EC cells (42), although a reduction or delay of cardiomyogenesis was observed. However, treatment of a cardiac lineage-restricted subline of P19 cells, P19CL6 cells, with the Shh signalling inhibitor cyclopamine blocked their differentiation into beating cardiac myocytes (43). The difference in the latter results may be due to a lack of compensatory signalling molecules, derived from other lineages, implicated in activating Gli due to a lack of compensatory signalling molecules, myocytes (43). The difference in the latter results may be blocked their differentiation into beating cardiac cells, with the Shh signalling inhibitor cyclopamine cardiomyogenesis was observed. However, treatment of a cardiac lineage-restricted subline of P19 cells, P19CL6 cells, with the Shh signalling inhibitor cyclopamine blocked their differentiation into beating cardiac myocytes (43). The difference in the latter results may be due to a lack of compensatory signalling molecules, derived from other lineages, implicated in activating Gli factors, such as TGFβ (104), Fgf (105) and Wnt (106). Finally, Zic factors have also been shown to bind genomic Gli binding sites and to modulate the transcriptional activity of Gli transcription factors (107). Our use of a dominant-negative approach would override compensatory activating signals (47,55), showing a role for Gli2 function during cardiomyogenesis.

While ubiquitously expressed Gli2 and MEF2C each enhanced cardiomyogenesis (Figures 3 and 6), and ubiquitously expressed Gli/EnR impaired cardiomyogenesis (Figure 4), surprisingly pgk-driven MEF/EnR enhanced cardiomyogenesis (46) and skeletal myogenesis (Karamboulas and Skerjanc, unpublished data). This is likely due to the dual nature of MEF2 factors, which have the ability to recruit HDAC4/5 to genes, inhibiting transcription (108), as well as recruiting histone acetyltransferases, activating transcription (108). A repressive role for Gli2 in regulating entry into various lineages was not observed, since the expression of Gli/EnR under the pgk promoter resulted in abolished skeletal myogenesis (47) and attenuated levels of neurogenesis and gliogenesis (55) at a specific stage of each respective differentiation program (47,55). Therefore, expression of Gli2/EnR under a general promoter such as pgk, impaired the formation of different stem cell differentiation pathways, including cardiomyogenesis, at specific stages.

While overexpression of MEF2C-TAP upregulated transcript levels of Gli2, Nkx2-5 and GATA-4, it did not affect Tbx5 mRNA expression. This correlates with previous studies, where knockdown of MEF2C using morpholinos disrupted zebrafish heart tube looping, but did not affect Tbx5 expression (109). It is possible that other MEF2 family proteins were able to rescue the expression of Tbx5 in zebrafish (109). Genetic redundancy of MEF2 family members is supported by our finding that the expression of Tbx5, Gli2 and cardiac muscle progenitor markers was drastically reduced in the presence of dominant-negative Nkx-MEF2C/EnR in vivo, and the loss of heart formation in vivo (21). It is likely that the Nkx2-5 enhancer driving the expression of MEF2C/EnR can be regulated by Gli2 and the previously described regulatory loop between MEF2C, GATA-4 and Nkx2-5 (21,40,45,78). Thus, as Nkx2-5 is expressed during formation of cardiac muscle progenitors, the overexpression of Nkx-MEF2C/EnR will inhibit the function of endogenous MEF2C and Gli2, as well as other cardiogenic factors, resulting in the loss of cardiomyogenesis. This is supported by the fact that both endogenous Gli2 and MEF2C were downregulated, but not abolished, by Day 6 in P19[Nkx-MEF2C/EnR] cells, whereas the expression of Nkx-MEF2C/EnR was reduced [ref. (21) and Figure 7].

We report for the first time that Gli2 and MEF2C associate with each other’s gene elements. Although existing literature supports identification of Gli2 as a target of MEF2C by demonstrating that the Drosophila MEF2 protein directly bound gene elements of Ci (a Drosophila homolog of vertebrate Gli transcription factors) (99), MEF2C was not previously reported to be a target of Gli3 in murine developing limb in a genome-wide ChIP-on-chip analysis (110). It is possible that MEF2C is a target of the Gli transcription factors at an earlier time-point than the time-point analysed in the study (E11.5) (110). Members of the Shh signalling pathway are expressed starting from E8.0 in murine paraxial mesoderm (81) and MEF2C is the first MEF2 family factor to be expressed in somites starting from E8.5–9.0 (16). Finally, it is possible that MEF2C is a target of Gli2 and not Gli3, during cardiomyogenesis.

Notably, the majority of the Gli2-bound sites are located in the intron regions of the Mef2c gene. It was previously shown that there are multiple promoter and enhancer regulatory elements within the Mef2c gene introns (17,111,112). Of those known regulatory elements in the Mef2c gene, the Mef2c D site, which was bound by Gli2 in this study, is located ~1500 bp relative to the 3' end of the reported cardiac promoter (17). Thus, it is possible that other Mef2c gene elements bound by Gli2 are located within novel regulatory regions, however, their identification is outside the scope of this study.
Much like the Mef2c gene sites bound by the Gli2 protein, Gli2 sites bound by the MEF2C protein are located in the intron regions of the Gli2 gene. Although regulatory elements in the Gli2 gene have not yet been reported, it is possible that its introns contain novel uncharacterized regulatory regions. For example, the Gli2 B site that was bound by MEF2C in this study, is part of the predicted genomic enhancer, which is well conserved between human, mouse, chicken, frog and fugu genomes [ref. (113), element hs1790]. Although this region was not found to have any enhancer activity at E11.5 when transfected into mouse embryos in a reporter cassette driving β-gal expression [ref. (113), element hs1790], it is possible that it can act as an enhancer at an earlier or later time point in development.

Both Gli2 and MEF2C are capable of enhancing the development of cardiac muscle through upregulation of the Nkx2-5, GATA-4 and BMP-4 regulatory loop [this study and refs (41,45)], skeletal muscle via upregulation of MRFs (47,48) and neurons through enhancement of Ascl1/Mash1 expression (55,114) in pluripotent P19 EC cells. The ability of Gli2 and MEF2C to regulate a similar subset of differentiation pathways in pluripotent P19 EC cells, led us to hypothesize that Gli2 and MEF2C could function together. We showed that Gli2 and MEF2C form a protein complex that functions synergistically on the Nkx2-5 promoter. MEF2 proteins are known to form synergistic protein complexes with cardiac muscle factors such as Hand1 (115) and GATA-4 (116). More recently, MEF2C, together with GATA-4 and Tbx5, was shown to convert mouse fibroblasts into cardiac myocytes (117). Furthermore, our findings are consistent with Pitx2, which is a homeodomain transcription factor involved in heart development, and MEF2 protein complex studies, where both MEF2 and Pitx2 DNA binding sites were necessary for the synergistic action of the protein complex (118).

In silico analysis of the mouse genome identified 957 genes containing both Gli and MEF2 conserved DNA binding clusters (Supplementary Table S1). Functional annotation analysis of these potential targets revealed a number of categories enriched among Gli and MEF2 clusters containing genes, such as cell differentiation, regulation of gene expression, chromatin organization, as well as development of nervous system, heart, the skeletal system and muscle organ (Table 3). While Nkx2-5 was not identified as a target gene for Gli2 and MEF2C, probably due to the presence of non-conserved Gli and MEF2 binding elements and/or due to stringent restrictions used in the screening process, MEF2C was identified as one of the potential target genes for both Gli and MEF2 transcription factors (Supplementary Table S1 and Table 3). The identification of numerous potential Gli and MEF2 targets regulating various lineages implies that the Gli2/MEF2C interactions identified here may be applicable to other lineages and require further study.

In summary, we propose a novel mechanistic model, in which Gli2 and MEF2C transcription factors induce and maintain each other’s expression, as well as form a protein complex capable of synergizing on gene promoters containing both Gli and MEF2 binding elements during cardiomyogenesis in vitro (Figure 10). This is the first evidence that the Shh signalling pathway is directly linked to the function of MEF2C protein. Our findings are similar to the MEF2 and MRF interaction model [reviewed in ref. (15)]. The Gli2/MEF2C protein complex may function during the induction of several lineages, including cardiac and skeletal muscle, as well as neurons. This is supported by the identification of numerous putative targets for both Gli2 and MEF2C, which are enriched in neurogenesis, as well as skeletal muscle and heart developmental processes (Table 3). Understanding the complex network of transcription factors that regulates lineage determination during stem cell differentiation is important for potential future cell therapies.

SUPPLEMENTARY DATA
Supplementary Data are available at NAR Online: Supplementary Table 1.

ACKNOWLEDGEMENTS
The authors would like to thank J. Fair and D. Ebadi for technical assistance, Dr Wallace for eye total protein extracts from Gli2+/− mice, Dr Brand and Dr Blais for helpful discussions and Dr Ryan, V. Mehta and T. LaRusso for eye total protein extracts from Gli2+/− mice, Dr Wallace, Dr Nemer for GATA-4 expressing plasmid, Dr Yutzey for Gli-responsive reporter construct, Dr Driscoll for Nkx2-5-luc, Dr Sasaki for Gli2 expression plasmid and Dr Hui for providing Gli2 specific antibodies, Dr Sasaki for Gli2 expression plasmid and Gli-responsive reporter construct, Dr Driscoll for Nkx2-5-luc, Dr Nemer for GATA-4 expressing plasmid, Dr Wallace for eye total protein extracts from Gli2+/− and Gli2−/− mice, Dr Wallace, Dr Brand and Dr Blais for helpful discussions and Dr Ryan, V. Mehta and D. Ebadi for critical reading of the manuscript. A.V. carried out mES, parental and stable P19 EC differentiation, IF, QPCR analysis, ChIP assay with Gli2 antibodies, IP from HEK-293 cells, bioinformatics analysis and manuscript writing. A.M. created and differentiated P19[TAP] and P19[MEF2C-TAP] stable cell lines and carried out ChIP assay with MEF2C antibodies. A.F. participated in ChIP assay with Gli2 antibodies. M.S. participated in differentiation and QPCR analysis of mES cells. C.K. differentiated P19[Nkx-MEF2C/EnR] cells, carried out northern blot and IF analysis of this cell line. I.S.S. conceived the study and participated in its design and coordination and helped to draft the manuscript. All authors read and approved the final manuscript.

FUNDING
Canadian Institutes of Health Research (MOP-53277 to I.S.S.); Heart and Stroke Foundation of Canada Doctoral Research Award (to A.V.); Ontario Graduate Scholarship (to A.V. and M.S.). Funding for open access charge: Canadian Institutes of Health Research (MOP-53277 to I.S.S.)

Conflict of interest statement. None declared.
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