Genomic profiling identifies genes and pathways dysregulated by HEY1–NCOA2 fusion and shines a light on mesenchymal chondrosarcoma tumorigenesis

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Abstract
Mesenchymal chondrosarcoma is a rare, high-grade, primitive mesenchymal tumor. It accounts for around 2–10% of all chondrosarcomas and mainly affects adolescents and young adults. We previously described the HEY1–NCOA2 as a recurrent gene fusion in mesenchymal chondrosarcoma, an important breakthrough for characterizing this disease; however, little study had been done to characterize the fusion protein functionally, in large part due to a lack of suitable models for evaluating the impact of HEY1–NCOA2 expression in the appropriate cellular context. We used iPSC-derived mesenchymal stem cells (iPSC–MSCs), which can differentiate into chondrocytes, and generated stable transduced iPSC–MSCs with inducible expression of HEY1–NCOA2 fusion protein, wildtype HEY1 or wildtype NCOA2. We next comprehensively analyzed both the DNA binding properties and transcriptional impact of HEY1–NCOA2 expression by integrating genome–wide chromatin immunoprecipitation sequencing (ChiP-seq) and expression profiling (RNA-seq). We demonstrated that HEY1–NCOA2 fusion protein preferentially binds to promoter regions of canonical HEY1 targets, resulting in transactivation of HEY1 targets, and significantly enhances cell proliferation. Intriguingly, we identified that both PDGFβ and PDGFRα were directly targeted and upregulated by HEY1–NCOA2; and the fusion protein, but not wildtype HEY1 or NCOA2, dramatically increased the level of phospho-AKT (Ser473). Our findings provide a rationale for exploring PDGF/PI3K/AKT inhibition in treating mesenchymal chondrosarcoma.

Keywords: mesenchymal chondrosarcoma; HEY1–NCOA2 fusion; RNA-seq; ChIP-seq

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Introduction
Gene fusions derived from chromosome rearrangements have been recognized as driver mutations in neoplasia. Especially in bone and soft-tissue tumors, more than 200 different fusions have been reported [1–4]. Most of these fusions can serve as ideal molecular diagnostic markers because they are strongly associated with particular histological subtypes. Furthermore, detailed studies of their functions have provided pivotal knowledge about pathogenetic mechanisms for targeted cancer therapy.

The HEY1–NCOA2 fusion was identified as a recurrent fusion in mesenchymal chondrosarcoma by Wang et al in 2012 [5]. This fusion has been used successfully as a molecular marker for the diagnosis of mesenchymal chondrosarcoma [6–22]. HEY1 (hairy/enhancer-of-split related with YRPW motif 1) is a member of the basic helix-loop-helix-orange (bHLH-O) family of transcriptional repressors and a downstream effector of Notch signaling [23]. The protein consists of a basic helix-loop-helix (bHLH) domain, the Orange domain and two conserved C-terminal motifs of which the function remains to be elucidated (Figure 1A). HEY1 protein, via its basic domain, directly contacts DNA and preferentially binds E-box sequences (CACGTG, CACGCG) in the promoter regions of target genes as a dimer, recruiting corepressors to repress the target genes of Notch signaling [24].

NCOA2 (Nuclear Receptor Coactivator 2) is a transcriptional coactivator for nuclear hormone receptors,
including steroid, thyroid, retinoid, and vitamin D receptors. NCOA2 has three distinct regions, i.e. an N-terminal bHLH-PAS domain consisting of a bHLH sequence followed by a tandem PAS domain that mediates DNA binding and protein dimerization, a central area of nuclear receptor interaction domain (NID) with three LXLL motifs that drive NCOA2 interaction with nuclear hormone receptors, and a C-terminal region that contains two transcriptional activation domains: AD1/CID (transcriptional activation domain 1/CBP/p300 interaction domain) and AD2 (transcriptional activation domain 2), which respectively recruit CBP/p300 and histone methyltransferases to facilitate transcription [25]. NCOA2 has been reported to be involved in translocations that results in fusions with multiple genes in various cancers [26–39]. Of note, in cancer-associated NCOA2 fusions, only the C-terminal AD1/CID and AD2 domains are consistently preserved in the fusion proteins. The involvement of the same functional domains of NCOA2 in various cancer gene fusions support a model in which the C-terminal transcriptional activation domains (AD1/CID and AD2)
of NCOA2 are aberrantly directed and utilized by the DNA-binding domain contributed by the N-terminal fusion partner.

The HEY1–NCOA2 fusion transcript identified in mesenchymal chondrosarcoma represents an in-frame fusion of HEY1 5’-exons (NM_012258, exons 1–4) to NCOA2 3’-exons (NM_006540, exons 13–23) [5]. As such, it is predicted that the HEY1-NCOA2 fusion protein replaces the C-terminal portion of HEY1 with the NCOA2 transcriptional activation domains, AD1/CID and AD2, while retaining the HEY1 bHLH domain (Figure 1A). It was hypothesized that the HEY1-NCOA2 fusion protein may act as a transcriptional activator by recruiting coactivators through its NCOA2 component to HEY1 target genes. In this study, we present a comprehensive analysis of both the DNA binding properties and transcriptional impact of HEY1-NCOA2 expression. We demonstrate that the fusion protein preferentially binds to regions that are occupied by the wild-type HEY1 transcription factor and provide a correlation with its effect on gene expression.

Materials and methods

Cells and reagents

The human cell lines HEK293T (CRL-3216) and iPSC-derived Mesenchymal Stem Cells (iPSC-MSCs, ACS-7010) were purchased from the ATCC (Manassas, VA, USA) and maintained in the recommended medium and fetal bovine serum with 100 units/ml penicillin/streptomycin at 37 °C in a humidified atmosphere containing 5% CO2. The polybrene infection/transfection reagent (TR-1003), doxycycline hydrochloride, 3xFlag peptide and anti-Flag M2 affinity gel (A2220) were purchased from Sigma-Aldrich (St. Louis, MO, USA). Anti-Flag M2 antibody (1:1,000, mouse monoclonal, F1804/M2) was from Sigma-Aldrich. Anti-CCND1 (1:1,000, rabbit monoclonal, 55506/E3P5S), anti-HES1 (1:500, rabbit monoclonal, 11988/D6P2U), anti-BCL2 (1:500, mouse monoclonal, 15071/124), anti-phospho-Akt (Ser473) (1:500, rabbit monoclonal, 4060/D6P2U), anti-phospho-Akt (Ser473) (1:500, rabbit monoclonal, 4060/D6P2U), anti-GAPDH (1:1,000, rabbit monoclonal, 2118/14C10) were from Cell Signaling Technology (Danvers, MA, USA).

Construction of cell lines for ChIP-seq, RNA-seq and cell proliferation studies

Human HEY1 (CAT#: RC200257) and NCOA2 (CAT#: RC212235) cDNA clones, from OriGene Technologies (Rockville, MD, USA), were used to construct N-terminal Flag-tag full-length coding sequences, i.e. FLAG-HEY1, FLAG-NCOA2, and FLAG-HEY1-NCOA2. In brief, the amplified coding sequence with Flag-tag were cloned into the expression vector pINDUCER21 (#46948, Addgene, Watertown, MA, USA) using Gateway LR clonase II enzyme system (Invitrogen, Waltham, MA, USA). The selected clones were transfected into the HEK293T cells by Sanger sequencing. For the HEY1-NCOA2 fusion construct, the sequence represents the fusion point as shown in supplementary material, Figure S1. For the generation of lentivirus, one million HEK293T cells were plated in each well of a 6-well plate for 24 h, then transiently cotransfected with lentiviral packaging helper plasmids pHDM-G, pCAGG-HIVgpc, pCAG4-RTR2 and the plasmid of interest at a 1:1 ratio using FuGene HD (Promega, Madison, WI, USA) according to the manufacturer’s instructions. Cells were incubated for 48 h and the supernatant containing the virus was collected for cell transduction. The iPSC-MSC cells were infected by the prepared lentivirus with 6.6 μg of polybrene for 72 h and GFP-positive cells were selected by fluorescence-activated cell sorting (FACS).

Intracellular localization of the HEY1–NCOA2 fusion protein

The coding sequences of HEY1, NCOA2, and HEY1–NCOA2 were cloned into the vector CL20-mEGFP to generate fusions of EGFP with HEY1, NCOA2, or HEY1-NCOA2, respectively. The iPSC-MSC cells were cultured in μ-Slide 4 Well (ibidi GmbH, Gräfelfing, Germany) and transiently transfected with the EGFP-constructs for 24 h. Hoechst was directly added into the cultures and the subcellular localization of HEY1, NCOA2, and HEY1-NCOA2 was observed using fluorescence microscopy. Images were capture using a Marianas 2 confocal microscope (Zeiss, White Plains, NY, USA).

ChIP-seq

Stably transduced iPSC-MSCs were treated with or without doxycycline (50 ng/ml) for 24 h, then subjected to the ChIP-seq process following the protocol reported previously [40]. Immunoprecipitation was performed using anti-Flag M2 affinity gel (Sigma, A2220) according to the manufacturer’s protocol. Eluted samples were incubated with RNase A (50 μg/ml final concentration) at 37 °C for 30 min, overnight with proteinase K to a final concentration of 200 μg/ml at 65 °C, and cleaned up using a Monarch PCR & DNA Cleanup Kit (TI030; New England BioLabs, Ipswich, MA, USA). Two biological replicates were prepared and sequenced independently. In brief, DNA was quantified using the QuantiT PicoGreen ds DNA assay (ThermoFisher, Waltham, MA, USA). Libraries were prepared with the HyperPrep Library Preparation Kit (PN 07962363001, Roche, Indianapolis, IN, USA). Libraries were quantified using an Quant-iT PicoGreen ds DNA assay (ThermoFisher) or by low-pass sequencing with a MiSeq nano kit (Illumina, San Diego, CA, USA). Single or paired end 50 cycle sequencing was performed on a NovaSeq 6000 platform (Illumina) at the Hartwell Center for Biotechnology at the St. Jude Children’s Research Hospital. Details for ChIP-seq data analysis are presented in Supplementary materials and methods.
RNA-seq and gene expression analysis

Stably transduced iPSC-MSCs were treated with or without doxycycline (50 ng/ml) for 24 h, then subjected to RNA extraction using the Quick-RNA Mini Prep kit (Zymo Research, Irvine, CA, USA). RNA-sequencing was performed by the St. Jude Hartwell Center of Biotechnology. In brief, RNA was quantified using the Quant-IT RiboGreen RNA assay (ThermoFisher) and quality checked using a 2100 Bioanalyzer RNA 6000 Nano assay (Agilent, Santa Clara, CA, USA) or 4200 TapeStation High Sensitivity RNA ScreenTape assay (Agilent) prior to library generation. Libraries were prepared from total RNA with a TruSeq Stranded mRNA Prep Kit according to the manufacturer’s instructions (PN 20020595, Illumina). Paired end 100 cycle sequencing was performed using a NovaSeq 6000 (Illumina). Details for RNA-seq data analysis as well as pathway and gene set enrichment analyses (GSEA) are described in Supplementary materials and methods.

Immunoblotting

Cell lysates were prepared with Laemmli buffer and separated on a 4–15% Mini-Protein precast gels (Bio-Rad, Hercules, CA, USA). The proteins were then transferred onto a nitrocellulose membrane and nonspecific binding was blocked by incubating with 5% nonfat milk in TBST buffer (0.01 M Tris–Cl, 0.15 M NaCl, 0.05% Tween-20, pH 8.0) at room temperature for 1 h. The membrane was subjected to the indicated primary antibody with gentle agitation overnight at 4 °C and then horseradish peroxidase (HRP)-conjugated secondary antibody at room temperature for 1.5 h. The proteins were detected using a ChemiDoc touch imaging system (Bio-Rad) after the membrane was incubated in Clarity western ECL substrate (Bio-Rad) for 5 min.

Real-time quantitative polymerase chain reaction (RT-qPCR)

Stably transduced iPSC-MSCs were treated with or without doxycycline (50 ng/ml) for 24 h, then subjected to RNA extraction using the Quick-RNA Mini Prep kit (Zymo Research). One μg of RNA was reverse-transcribed into cDNA using a High-Capacity RNA-to-cDNA kit (ThermoFisher, Cat#: 4387406), and quantitative polymerase chain reaction (qPCR) was performed in triplicate using iTaq Universal SYBR Green Supermix (Bio-Rad, Cat#: 172-5120) on the Applied Biosystems QuantStudio 7 Flex Real-Time PCR platform. Expression of mRNA was normalized to GAPDH as a reference transcript, and the fold-change of expression was calculated using the $2^{-\Delta \Delta Ct}$ method. The sequences of PCR Primers used are listed in supplementary material, Table S1.

Cell proliferation assays

Stably transduced iPSC-MSCs were seeded in a 6-well plate (1 x 10^4 per well) for 24 h and then subjected to a cell proliferation study. Medium with or without doxycycline (50 ng/ml) respectively was used and replenished every 3 days. The cell number was counted using a Countess II FL Automated Cell Counter (Life Technologies, Waltham, MA, USA) at seeding and at day 6 and day 12 after the initial doxycycline treatment. Additionally, Click-iT EdU flow cytometry cell proliferation assays (Invitrogen, C10634) were performed according to the manufacturer’s protocol. In brief, cells were cultured with or without doxycycline (50 ng/ml) for 1 week, labeled with 10 μM EdU for 3 h, and then harvested, fixed, and stained using the Click-iT Plus reaction cocktail with fluorescent picolyl azide and analyzed by flow cytometry using FlowJo software (v10.7.1, FlowJo, Ashland, OR, USA).

Three-dimensional (3D) cell culture assay

3D cell cultures were performed in 96-well, round black/clear bottom, ULA (Ultra-Low Attachment surface) Corning spheroid microplates (Sigma, CLS4520). In each well, 1,500 stably transduced iPSC-MSCs were cultured with or without doxycycline (50 ng/ml). Spheroid formation and growth were visualized microscopically at 24 h (day 1), 48 h (day 2), 72 h (day 3), 96 h (day 4), and day 7 after cell seeding. The sizes (radius) of spheroids were measured using TCapture imaging software (v5.1.1, Tuscen Photonics Co., Fuzhou, Fujian, PR China).

Results

Intracellular localization of HEY1–NCOA2 fusion protein

In order to evaluate the intracellular localization of HEY1–NCOA2, HEY1, and NCOA2, we generated EGFP-tagged constructs and monitored protein localization by fluorescence microscopy in transfected iPSC-MSCs. We confirmed that HEY1–NCOA2, like wildtype HEY1, is localized to the nucleus. In contrast, wildtype NCOA2 is located in both the nucleus and cytoplasm (Figure 1B). In comparing the GFP signal patterns between EGFP-HEY1-NCOA2 and EGFP-HEY1, our data showed similar intranuclear distribution of HEY1–NCOA2 and wildtype HEY1.

Identification of HEY1–NCOA2 DNA-binding sites

The induced expression of FLAG-tagged HEY1–NCOA2, HEY1, and NCOA2 proteins in stably transduced iPSC-MSCs were verified by immunoblotting at 8, 16, 24, and 48 h after doxycycline induction (Figure 1C), and the genome-wide binding profiles of HEY1–NCOA2, HEY1, and NCOA2 were investigated through anti-FLAG ChIP-seq in each iPSC-MSC cell model at 24 h after doxycycline (50 ng/ml) induction. By genome-wide ChIP-seq analyses, a total of 30,142 DNA-binding peaks were identified for HEY1 and HEY1–NCOA2, respectively. No specific DNA-binding peaks were identified with the induced expression of NCOA2 in comparison to controls, which
is consistent with prior knowledge about NCOA2. NCOA2 is one of the members of the p160/steroid receptor coactivator (SRC) family. SRCs have not been shown to directly bind DNA, despite possessing the bHLH domain [41]. In reviewing the location of HEY1 and HEY1–NCOA2 binding sites in the genome, we observed remarkable overall similarity between those two (supplementary material, Figure S2A). Further comparison revealed that 92% (n = 3,445) of HEY1-NCOA2 DNA-binding sites were shared with HEY1 (Figure 2A). Interestingly, out of all those shared DNA-binding targets, 88.5% are located within promoter regions (Figure 2A); in contrast, only 60.3% of HEY1-only DNA-binding peaks were located in promoter

Figure 2. HEY1–NCOA2 fusion protein DNA-binding pattern in the genome. (A) Genomic context of HEY1 and HEY1–NCOA2 binding peaks, with a breakdown into three groups: (1) Peaks that are overlapping between HEY1 and HEY1–NCOA2 fusion protein, (2) HEY1-only binding peaks, and (3) HEY1–NCOA2-only binding peaks. (B) Signal enrichment heatmap for the peaks specific for HEY1 only (blue bar), HEY1–NCOA2 only (red bar), and overlapping peaks (purple bar). (C) Normalized enrichment of HEY1-binding targets, comparing HEY1-only peaks and HEY1-binding peaks overlapping with HEY1–NCOA2-binding peaks. (D) E-box sequences bound by the HEY1 and HEY1–NCOA2 fusion protein. Distribution of the distance between binding peaks and the closest E-box motif(s); the distance equals 0 for the direct overlap between peak and motif, which fraction is visualized with pie plots.
regions and HEY1–NCOA2-only binding peaks ($n = 295$) exhibited a dramatically different binding profile across the genome (Figure 2A). The enrichment analysis of the ChIP-seq data further demonstrated the remarkable similarity in the genome-wide DNA binding feature between HEY1–NCOA2 fusion protein and wildtype HEY1 (Figure 2B), and suggested that the DNA regions bound by both HEY1 and HEY1–NCOA2 are enriched in high-affinity binding targets of HEY1 (Figure 2C).

The E-box (CAGCGT, CACGCC) DNA elements are HEY1-preferred DNA binding motifs [24]. Therefore, we searched next for the E-box sequences in our ChIP-seq data. Indeed, as shown in Figure 2D, the E-box sequences were identified in 12,157 out of 30,142 HEY1 binding peaks (40%) and 2,502 out of 3,740 HEY1–NCOA2 binding peaks (66%). Strikingly, about 98.8% (2,472 out of 2,502) of HEY1–NCOA2 binding peaks that contain E-box sequences overlap with HEY1 binding peaks (supplementary material, Figure S2B).

Gene expression profile associated with HEY1–NCOA2

Our ChIP-data demonstrated that HEY1–NCOA2 fusion protein preferentially binds to promoter regions across the genome that are also high-affinity binding targets of HEY1 with canonical E-box binding motif, which confirmed our hypothesis based on the predicted structure of HEY1–NCOA2 fusion protein. We speculated further that the expression of those target genes, which are normally repressed by HEY1, may be transactivated in the presence of HEY1–NCOA2 fusion protein, given that the NCOA2 transcriptional activation domains replace the C-terminal portion of HEY1. To verify this hypothesis, we performed RNA-seq in stably transduced iPSC-MSCs expressing exogenous HEY1, NCOA2, and HEY1–NCOA2 after 24 h doxycycline induction, which we will refer to as ‘MSC-HEY1(+),’ ‘MSC-NCOA2(+),’ and ‘MSC-HEY1–NCOA2(+),’ respectively, in this article. Transduced iPSC-MSCs that lacked doxycycline treatment, which we will refer to as ‘MSC-HEY1crtl,’ ‘MSC-NCOA2ctrl,’ and ‘MSC-HEY1–NCOA2ctrl,’ respectively, were similarly profiled as controls. Overall, MSC-HEY1-NCOA2(+), MSC-HEY1(+), and MSC–NCOA2(+) cells can be robustly separated by gene expression profiling (supplementary material, Figure S3A,B). Differential gene expression (DGE) analysis was performed to compare gene expression values between MSC-HEY1-NCOA2(+) and MSC-HEY1(+) cells and identified a total of 2,440 differentially expressed genes (fold change >2, false discovery rate [FDR] < 0.05), about two-thirds of which (1,563; 64%) showed increased expression in MSC-HEY1-NCOA2(+) cells (Figure 3A, supplementary material, Table S2). In addition, DGE analysis was performed to compare gene expression values between MSC-HEY1-NCOA2(+) and MSC-HEY1-NCOA2ctrl, MSC-HEY1(+) and MSC-HEY1crtl, as well as MSC-NCOA2(+) and MSC-NCOA2ctrl, respectively. Differentially expressed genes identified in each matched pair were subjected to Gene Set Enrichment Analysis (GSEA). As shown in Figure 3B, genes downregulated by HEY1 [MSC-HEY1(+)] versus MSC-HEY1crtl] were significantly enriched among HEY1-NCOA2-upregulated genes ($NES = 1.8, p \text{ value} \approx 0, \text{FDR} \approx 0$). GSEA analysis showed no significant correlation in the expression between NCOA2-regulated and HEY1–NCOA2-regulated genes (data not shown). Further on, as shown in Figure 3C, the additional GSEA showed that genes that were directly targeted by both HEY1 and HEY1-NCOA2 (binding peaks identified in their promoters by ChIP-seq, $n = 3,194$), were significantly enriched in the gene set that were upregulated by HEY1–NCOA2 in comparison to HEY1 [MSC-HEY1-NCOA2(+) versus MSC-HEY1(+)]. The integrated genome-wide ChIP-seq and RNA-seq expression profiling demonstrated that HEY1-NCOA2 fusion protein activated the expression of canonical HEY1 target genes that were transcriptionally repressed by wildtype HEY1.

Correlation between HEY1 and HEY1–NCOA2 binding profiles and gene expression

Cross-comparison between ChIP-seq and RNA-seq expression data revealed that 674 out of 1,563 genes (43%) upregulated by HEY1–NCOA2 [MSC-HEY1–NCOA2(+) versus MSC-HEY1(+)] had binding peaks for HEY1–NCOA2 fusion protein as well as HEY1 in their promoter regions (the list of 674 genes, which we will refer to as ‘HEY1–NCOA2 key-direct targets’ in this article, is available in supplementary material, Table S3). This finding indicates that almost half of the genes upregulated by HEY1–NCOA2 may result from direct transcriptional activation by the fusion protein. The 674 HEY1–NCOA2 key-direct targets include previously identified targets of HEY1, such as HES1, KLF10, BMP2, and FOXC1 [42], as well as a variety of novel findings, including PDGFRB, PDGFRAB, SOX4, etc. HEY1–NCOA2 and HEY1 binding peaks in the promoter regions of HES1, PDGFRB, PDGFRAB, and SOX4, demonstrated by ChIP-seq analysis, are presented in supplementary material, Figure S4.

Validation of the data derived from the in vitro iPSC-MSC models on human mesenchymal chondrosarcoma tumor samples

To validate if the transcriptional dysregulation profile that were identified in the in vitro iPSC-MSC-HEY1-NCOA2 model recapitulate the molecular phenotype of human mesenchymal chondrosarcoma, we performed single-sample GSEA (ssGSEA) analysis to assess the expression of the 674 HEY1–NCOA2 key-direct targets in a set of RNA-seq data derived from 107 sarcoma patients’ samples (RNA-seq data downloaded from St. Jude Cloud, https://platform.stjude.cloud/), including one mesenchymal chondrosarcoma sample as well as 106 samples from eight other sarcoma entities, i.e. Gastrointestinal Stromal Tumor (GIST), Rhabdomyosarcoma (RHB), Infantile fibrosarcoma.
IFS, Alveolar Soft Part Sarcoma (ASPS), Synovial Sarcoma (SS), Desmoplastic Small Round Cell Tumor (DSRCT), Ewing’s Sarcoma (EWS), and Osteosarcoma (OS). The same ssGSEA strategy was also applied to the analysis of Exon-Array data of 57 sarcoma samples published by Wang et al [5], in which the HEY1–NCOA2 fusion was identified as a novel and recurrent genetic alteration in mesenchymal chondrosarcoma (Exon-Array data downloaded from http://cbio.mskcc.org/Public/sarcoma_array_data/). The detailed

Figure 3. Gene expression profile associated with HEY1–NCOA2. (A) Heatmap showing row-normalized expression level of genes differentially regulated by HEY1–NCOA2 in comparison to wildtype HEY1 [MSC-HEY1–NCOA2(+)] versus MSC-HEY1[+]. (B) GSEA Enrichment plot of the expression of genes that were identified as downregulated by HEY1 in MSC-HEY1(+) versus MSC-HEY1ctrl. The gene rank was based on the log2(fold-change) of the expression of genes in MSC-HEY1–NCOA2(+) versus MSC-HEY1–NCOA2ctrl. (C) GSEA enrichment plot for the expression of genes of which their promoters had binding peaks of both HEY1 and HEY1–NCOA2. The gene rank was based on the log2(fold-change) of the gene expression in MSC-HEY1–NCOA2(+) versus MSC-HEY1(+). (D) Boxplots visualizing the enrichment of the 674 HEY1–NCOA2 direct-target-and-transactivating genes identified using the iPSC-MSCs cell models in sarcoma tumor samples. The enrichment was visualized across patients’ samples from a total of eight sarcoma entities. GSEA, Gene Set Enrichment Analysis; NES, Normalized Enrichment Score; FDR, False Discovery Rate; MCS, Mesenchymal Chondrosarcoma; GIST, Gastrointestinal Stromal tumor; RHB, Rhabdomyosarcoma; IFS, Infantile Fibrosarcoma; ASPS, Alveolar Soft Part Sarcoma; SS, Synovial Sarcoma; DSRCT, Desmoplastic Small Round Cell Tumors; EWS, Ewing’s Sarcoma.
information about all samples accrued for the ssGSEA analysis are presented in supplementary material, Table S4. Our ssGSEA analysis, both in the RNA-seq dataset and in the Exon-array dataset, demonstrated that the 674 HEY1–NCOA2 key-direct targets are highly enriched in the expression profile of mesenchymal chondrosarcoma tumor samples. As shown in Figure 3D, the Normalized Enrichment Scores (NESs) of the 674 genes

Figure 4. Functional pathways enriched in HEY1–NCOA2 upregulated genes. (A) Bar plot showing the $-\log_{10}(FDR)$ of the top 10 enriched KEGG pathways, computed for 1,563 HEY1–NCOA2 genes upregulated by HEY1–NCOA2 as compared with HEY1 [MSC-HEY1-NCOA2(+) versus MSC-HEY1(+)]. (B) Heatmap summarizing the GSEA (Gene Set Enrichment Analysis) of selected gene sets. Each row represents one gene set. Row #4–#8, gene signatures were selected based on the KEGG pathway enrichment analysis. The values in the heatmap indicate the $-\log_{10}$ of p value, multiplied by $-1$ if the NES score was indicating the enrichment biased toward the negative phenotype. (C) GSEA enrichment plot of MSC-HEY1-NCOA2(+) versus MSC-HEY1-NCOA2(−) and MSC-HEY1-NCOA2(+) versus MSC-HEY1(+). The gene set “Chondrocyte Differentiation” is from Gene Ontology (GO). The false discovery rate (FDR), nominal p value and normalized enrichment score were calculated by GSEA.
in mesenchymal chondrosarcoma tumor samples are around 1.5; in contrast, the mean NESs in other sarcoma cases is between −0.5 and 0.5. Of note, samples of tumor types that are available in both the St. Jude cloud RNA-seq dataset and the Exon-array dataset are presented in Figure 3D. The ssGSEA NESs of all 164 samples are available in supplementary material, Table S5, and the corresponding box plots of NESs are presented in supplementary material, Figure S5. These results, although limited by the data availability, suggest that the key HEY1–NCOA2-upregulated genes identified in our in vitro iPSC-MSC cell models indeed represent the genes highly expressed in mesenchymal chondrosarcoma patients’ tumors.

Functional classification of genes upregulated by the HEY–NCOA2 fusion

Functional classification of the 1,563 HEY1–NCOA2 upregulated genes was performed by pathway enrichment analysis. The top 10 significantly enriched pathways in KEGG are presented in Figure 4A, including cell cycle pathway, Hedgehog and WNT signaling pathways, as well as the PI3K–Akt signaling pathway, all of which were linked with promoting cell proliferation in cancer (refer to supplementary material, Table S6 for all KEGG pathways significantly enriched). The enrichment of these pathways was further supported by GSEA from the comparisons: (1) MSC-HEY1-NCOA2(+) versus MSC-HEY1-NCOA2(-)

Functional characterization of HEY1–NCOA2 upregulated genes, the fusion protein also rewires the transcriptional network during the differentiation, which may result in the development of mesenchymal chondrosarcoma.

Verification of the expression of selected targets upregulated by HEY1–NCOA2

The expression of 13 selected HEY1–NCOA2-upregulated genes was verified by RT-qPCR in an independent set of RNAs. As shown in Figure 5A, the significantly increased expression of these genes in HEY1–NCOA2-expressing cells [MSC-HEY1-NCOA2(+)] was verified by RT-qPCR.

All selected genes except CCND1 are among the 674 HEY1–NCOA2 key-direct targets (supplementary material, Table S3). According to RNA-seq, 10 out of the 12 genes barely had a detectable expression [Transcripts Per Million (TPM) << 1] in control cells (MSC-HEY1(+), MSC-NCOA2(+), and MSC-HEY1-NCOA2(+)). As for the remaining two genes, SOX4 and PDGFRα, RNA-seq revealed ~4 and ~8 transcripts per million RNA molecules in control cells, respectively. In contrast, RNA-seq revealed relatively high expression

Figure 5. HEY1–NCOA2 target gene expression validation. (A) RT-qPCR validation of genes regulation by inducible HEY1, NCOA2, or HEY1–NCOA2 expression in iPSC-MSC. The graph represents the fold change between the induced and uninduced condition. (B) Immunoblotting was further confirm the expression of BCL2, CCND1, and HES1 after HEY1–NCOA2 induction in stably transduced iPSC-MSCs. GAPDH was used as a loading control.
of CCND1 in control cells (~245 TPM). Moreover, the differential gene expression analysis demonstrated that HEY1 significantly downregulates CCND1 expression (>2-fold), while HEY1–NCOA2 dramatically upregulates its expression (>4-fold) in our iPSC-MSC cell models (supplementary material, Table S2). In addition to verifying the expression of CCND1 by RT-qPCR, immunoblotting was performed that demonstrated decent CCND1 expression in controls cells, decreased CCND1 expression in MSC-HEY1 (+), and dramatically increased CCND1 expression in MSC-HEY1–NCOA2 (+) (Figure 5B). In addition, as shown in Figure 5B, there was no detectable expression of HES1 or BCL2 by immunoblotting in cells without HEY1–NCOA2 expression, which is compatible with the transcription expression data (TPM) obtained from RNA-seq. Overall, the analysis of the expression of selected genes by RT-qPCR and immunoblotting, including 12 HEY1–NCOA2 direct targets and the indirect-dysregulated gene CCND1, confirmed the RNA-seq results.

HEY1–NCOA2 fusion enhances cell proliferation in iPSC MSC

RNA-seq followed by pathway enrichment analysis revealed that the cell cycle pathway was significantly upregulated by the HEY1–NCOA2 fusion and downregulated by wildtype HEY1. We verified the effects of the HEY1–NCOA2 fusion, as well as wildtype HEY1 and NCOA2, on cell growth in stably transduced iPSC-MSCs. As shown in Figure 6A, HEY1–NCOA2 expressing iPSC-MSCs displayed significantly accelerated proliferation when compared with matched control cells. In contrast, cell proliferation was slightly inhibited with the induced expression of wildtype HEY1, while the induced expression of exogenous NCOA2 had no significant impact on cell growth. We next assessed cell cycle transit by an EdU incorporation assay. Compared to control cells, the proportion of HEY1–NCOA2-expressing cells in S-phase was increased by 8.2% ($p = 0.0328$) (Figure 6B), demonstrating a significant increase in cell cycle transit as a result of HEY1–NCOA2 expression.

Human iPSC-MSCs require the addition of rhFGF basic (recombinant human fibroblast growth factor, basic) and rhIGF-1 (recombinant human insulin-like growth factor-1) growth factors to the growth medium as per the ATCC cell culture instruction. Since our RNA-seq data revealed several FGF and IGF family genes were upregulated by HEY1–NCOA2 (supplementary material, Table S2), we tested whether HEY1–NCOA2 expression can induce cytokine-independent growth. Indeed, as shown in Figure 6C,
only HEY1–NCOA2 expressing iPSC-MSCs displayed significantly accelerated proliferation in the absence of rhFGF basic and rhIGF-1 growth factors in the culture medium \( (p = 0.0078) \).

**HEY1–NCOA2 fusion increases cell growth in a 3D model**

As 3D spheroid cell culture more closely resembles in vivo tissue in terms of cell–cell and cell–matrix interactions [45], we used a scaffold-free technique to evaluate whether the fusion protein can also affect cell growth in a 3D model. FACS sorted iPSC-MSCs stably transduced for inducible expression of HEY1, NCOA2 or HEY1–NCOA2 were seeded (1,500 cells per well, 96-well plates) in wells with a round bottom and ultralow attachment surface. Doxycycline (50 ng/ml) was added into selected wells to induce the expression of transduced constructs, and no doxycycline in control cells’ wells. After 24 h, cells of each well formed a single spheroid with a clear boundary. In addition, in the MSC-HEY1-NCOA2(+) wells, cell satellites around the main spheroid were visible (supplementary material, Figure S7B). As the culture continued, the spheroids shrunk in all wells except those in MSC-HEY1-NCOA2(+) wells. The size (radius) of each spheroid was measured at 24, 48, 72, and 96 h and 7 days. The graph in supplementary material, Figure S7A presents the mean value of radius ± SD (\( n = 10 \)). Representative 3D cell growth images taken at 24 h and 72 h after seeding are shown in supplementary material, Figure S7B. The 3D culture experiment demonstrated the dramatic difference in size and morphology of spheroids between HEY1–NCOA2-expressing iPSC-MSCs [MSC-HEY1-NCOA2(+)] and wildtype protein expression cells [MSC-HEY1(+)] and MSC-NCOA2(+)].

**Discussion**

Mesenchymal chondrosarcoma is a rare malignant tumor. It arises most commonly in the second and third decades of life, and the outcome for these patients appears to be poor [46]. In a recent study, the overall 5-year and 10-year survival rate for mesenchymal chondrosarcoma patients was 51% and 43%, respectively [47]. The discovery of the HEY1–NCOA2 fusion as a molecular marker for mesenchymal chondrosarcoma was an important breakthrough for characterizing this disease. The identification of the pathways modulated by this fusion could help elucidate the pathogenesis of this disease and potentially develop new treatment options for mesenchymal chondrosarcoma.

We hypothesized that the HEY1–NCOA2 fusion protein would drive the expression of canonical HEY1 targets through the bHLH DNA-binding and dimerization domains of HEY1, as well as the transactivation domains of NCOA2. This was confirmed by our ChIP-seq data, which revealed extreme similarity in genome-wide DNA-binding patterns between HEY1–NCOA2 and wildtype HEY1. The intracellular localization and ChIP-seq data indicate that the oncogenic activities of HEY1–NCOA2 likely reflect its impact on HEY1-dependent pathways. Hence, in the subsequent differential gene expression analysis of RNA-seq data, we focused on the relationship between HEY1–NCOA2 and HEY1 in searching for HEY1–NCOA2 key target genes.

The most intriguing key targets of HEY1–NCOA2 discovered in this study are PDGFB and PDGFA. We demonstrated that both PDGFB and PDGFA were directly targeted and transactivated by the HEY1–NCOA2 fusion protein. PDGFB is a signaling ligand, a member of the protein family comprised of both platelet-derived growth factors (PDGF) and vascular endothelial growth factors (VEGF). PDGFs act via two receptors, PDGFRα and PDGFRβ, and PDGF-PDGFR interactions activate PDGF receptor tyrosine kinases. By RNA-seq and RT-qPCR, we demonstrated that PDGFB gene expression was extremely upregulated by HEY1–NCOA2 (~105-fold by RNA-seq and ~120-fold by RT-qPCR). In comparison to PDGFB, the upregulation of PDGFA expression was moderate (~3.75-fold).

PDGF–PDGFR signaling mediates a variety of downstream signaling effector pathways, including PI3K–AKT, among others [48]. Additionally, functional classification of HEY1-NCOA2 upregulated genes highlighted PI3K–AKT signaling as one of top enriched pathways in the current study. We next evaluated the activation of PI3K–AKT. Our preliminary data demonstrated that with the expression of the HEY1–NCOA2 fusion protein, but not wildtype HEY1 or NCOA2, the level of phosphor-AKT (Ser473) dramatically increased (supplementary material, Figure S8). Of note, using approaches based on immunohistochemistry (IHC), some previous studies of mesenchymal chondrosarcoma demonstrated increased expression of PKC-α, PDGFR-α, and phosphorylated-mTOR in the malignant mesenchymal chondroblasts of mesenchymal chondrosarcoma tumor samples, indicating that these pathways’ potential roles in the development or maintenance of mesenchymal chondrosarcoma [49,50]. Hence, our data, to some extent, elucidated the biologic mechanism underlying the histopathologic features that had been identified in mesenchymal chondrosarcoma patients’ tumor samples. Clearly, much more work is needed to better define the role(s) of PDGF signaling in mesenchymal chondrosarcoma tumorigenesis, but our study suggests that small-molecule inhibitors targeting PDGF signaling should be evaluated in preclinical models of mesenchymal chondrosarcoma.

Another interesting finding in this study that may provide a fundamental rationale for exploring a new treatment approach for mesenchymal chondrosarcoma is the direct transactivation of BCL2 by the HEY1–NCOA2 fusion protein. BCL2 has been reported to exhibit stronger expression in malignant mesenchymal chondroblasts in comparison to more mature-appearing chondrocytic counterparts in
mesenchymal chondrosarcoma tumor samples, by IHC staining [49]. In addition, a recent study reported that the inhibition of Bcl-2 family members restores the apoptotic machinery in mesenchymal chondrosarcoma, rendering the cells sensitive to conventional chemotherapy [51]. In the present report, we integrated genomewide ChIP-seq and expression profiling (RNA-seq) identified BCL2 as one of the 674 key-direct targets of the HEY1–NCOA2 fusion protein. Further on, our RT-qPCR and immunoblotting studies verified the RNA-seq findings of BCL2 transactivation by the HEY1–NCOA2 fusion protein (Figure 5). Overall, our findings provide a biological rationale for exploring BCL2 inhibitors in treating mesenchymal chondrosarcoma.

Other targets of interest, such as HES1 and SOX4, have a well-credentialled role in the transcriptional control of chondrocyte specification and differentiation [52]. Chondrocytes arise in development from mesenchymal stem cells/multipotent skeletal progenitor-stem cells (SSCs). HES1 was reported to delay the differentiation of SSCs into chondrocytes [53], and SOX4, along with its closely related family members SOX11 and SOX12, is expressed strongly in SSCs and prechondrocytes, and weakly in chondrocytes [54]. In a previous study, Bhattaram et al demonstrated that Sox4/Sox11/Sox12 act redundantly to keep mesenchymal progenitor cells alive in a mouse model [55]. In our study, the promoters of both HES1 and SOX4 were bound directly by HEY1–NCOA2, and both genes were significantly upregulated by HEY1–NCOA2. Further study is needed to elucidate the interplay between HES1, SOX4, and HEY1–NCOA2 in the tumorigenesis of mesenchymal chondrosarcoma.

Our study represents the first comprehensive, genome-wide analysis of HEY1–NCOA2 DNA binding sites and correlates these to HEY1–NCOA2-associated gene expression profiles. Our results point to a pivotal role for the HEY1–NCOA2 fusion in mesenchymal chondrosarcoma tumorigenesis. While we do not yet know whether HEY1–NCOA2 alone is sufficient to induce malignant transformation in vivo, in our model system using iPSC-MSC, HEY1–NCOA2 appeared to act on the key aspects of increasing cell cycle transition, a recognized facet of malignant transformation, and likewise induces genes that are implicated in chondrocyte differentiation. The findings of our present study, which were generated by using a forced gene expression system based on the iPSC-MSC cell model, remains to be verified in isogenic cell lines with the HEY1–NCOA2 fusion. Additionally, the intriguing HEY1–NCOA2 downstream targets identified in this study will be evaluated in a larger cohort of mesenchymal chondrosarcoma tumor samples to establish more precisely the specificity of these dysregulated genes and pathways in mesenchymal chondrosarcoma development. Finally, further studies on the potential downstream target genes and pathways of HEY1–NCOA2 may help to develop new therapeutic approaches for this often-lethal malignancy.

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Author contributions statement

LW, WQ and WR conceived and designed the study. WQ, WR, ZY and HJ acquired data. WR, WQ, BX and LW analyzed and interpreted data. WR, BX, SW, YF and GW were responsible for bioinformatics. All the authors critically revised the article and read and approved the final version.

Data availability statement

The ChIP-seq and RNA-seq data in this study are available at https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE196003.

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