Inhibition of GLI1 Expression by Targeting the CRD-BP–GLI1 mRNA Interaction Using a Specific Oligonucleotide

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Received November 6, 2015; accepted March 31, 2016

ABSTRACT
The stabilization of glioma-associated oncogene 1 (GLI1) mRNA by coding region determinant binding protein (CRD-BP) through the Wnt/β-catenin signaling pathway is implicated in the proliferation of colorectal cancer and basal cell carcinoma. Here, we set out to characterize the physical interaction between CRD-BP and GLI1 mRNA so as to find inhibitors for such interaction. Studies using CRD-BP variants with a point mutation in the GXXG motif at each KH domain showed that KH1 and KH2 domain are critical for the binding of GLI1 RNA. The smallest region of GLI1 RNA binding to CRD-BP was mapped to nucleotides (nts) 320–380. A 37-nt S1 RNA sense oligonucleotide, containing two distinct stem-loops present in nts 320–380 of GLI1 RNA, was found to be effective in blocking CRD-BP–GLI1 RNA interaction. Studies using various competitor RNAs with modifications to S1 RNA oligonucleotide further displayed that both the sequences and the structure of the two stem-loops are important for CRD-BP–GLI1 RNA binding. The role of the two-stem-loop motif in influencing CRD-BP–RNA interaction was further investigated in cells. The 2′-O-methyl derivative of the S1 RNA oligonucleotide significantly decreased GLI1, c-myc, and CD44 mRNA levels, in a panel of colon and breast cancer cells. The results from this study demonstrate the potential importance of the two-stem-loop motif as a target region for the inhibition of the CRD-BP–GLI1 RNA interaction and Hedgehog signaling pathway. Such results pave the way for the development of novel inhibitors that act by destabilizing the CRD-BP–GLI1 mRNA interaction.

Introduction
The Hedgehog (Hh) signaling pathway plays a crucial role in embryonic development and cell homeostasis (Hui and Angers, 2011). Its aberrant activation has been linked to tumorigenesis (Hui and Angers, 2011; Amakey et al., 2013). For instance, mutations in the key protein components leading to the constitutive activation of the Hh pathway have been identified in medulloblastoma and basal cell carcinoma (Scales and de Sauvage, 2009). In addition, exaggerated Hh signaling in the absence of known driver mutations has been linked to many types of cancers, including colon, breast, pancreatic, gastric, lung, and prostate cancers, as well as leukemia and multiple myeloma (Scales and de Sauvage, 2009). Key components of the mammalian Hh pathway include a positive regulatory transmembrane protein Smoothened (SMO) and the glioma-associated oncogene (GLI) zinc finger transcription factors (GLI1, GLI2, and GLI3), with GLI1 being predominantly a transcriptional activator (Hui and Angers, 2011). Although GLI1 mRNA level is widely accepted as an indicator of Hh pathway activity, GLI signaling can also be regulated by proteins involved in other signaling pathways. For instance, it has recently been shown that Wnt/β-catenin signaling can elevate GLI1 expression and transcriptional activity through the induction of expression of an RNA-binding protein called CRD-BP, which acts by binding to and stabilizing GLI1 mRNA (Noubissi et al., 2009). This mode of GLI1 regulation appears to be important for the survival and proliferation of colorectal cancer and basal cell carcinoma cells (Noubissi et al., 2009, 2014).

Coding region determinant–binding protein (CRD-BP) belongs to a highly conserved family of RNA binding proteins that have two N-terminal RNA recognition motifs followed by four C-terminal heterogeneous nuclear ribonucleoprotein-K-homology (KH) domains (Bell et al., 2013). In addition to GLI1, CRD-BP also has high affinity for several mRNAs whose protein products have been implicated in cancer. These include mRNAs for c-myc (Prokipcak et al., 1994), CD44 (Vikesa et al., 2006; King et al., 2014), K-Ras (Mongroo et al., 2011), βTrCP1 (Noubissi et al., 2006), microphthalmia-associated transcription factor (Goswami et al., 2015), mitogen-activated protein kinase 4 (Stohr et al., 2012), β-catenin (Gu et al., 2008), and multi-drug resistance 1 (Sparanese and Lee, 2007). The role of CRD-BP in cancer is further supported by the following findings: 1) Overexpression of CRD-BP in various human cancers, which include colon and breast cancers (Bell et al., 2013), and 2) mice genetically engineered to

This research was supported by the Natural Sciences and Engineering Research Council [Discovery Grant 227158].

dx.doi.org/10.1124/mol.115.102434

ABBREVIATIONS: ANOVA, analysis of variance; CRD-BP, coding region determinant binding protein; dsRNAi, double-stranded Dicer substrate RNA interference; EMSA, electrophoretic mobility shift assay; GLI, glioma-associated oncogene; Hh, Hedgehog; Kd, dissociation constant; KH domain, heterogeneous nuclear ribonucleoprotein-K-homology domain; nts, nucleotides; PCR, polymerase chain reaction; qPCR, quantitative real-time-polymerase chain reaction; SMO, Smoothened; TAMRA, 5-carboxytetramethylrhodamine.
overproduce CRD-BP in their mammary glands developed mammary adenocarcinoma (Tessier et al., 2004). Studies on various mRNA targets of CRD-BP have now confirmed that the oncogenic function of CRD-BP results from its physical association with the corresponding mRNAs, leading to their stabilization, increased expression, and ultimately the presentation of the cancerous phenotypes (Noubissi et al., 2006, 2009; Vikesaa et al., 2006; Gu et al., 2008; Mongroo et al., 2011). Hence, the ability to disrupt specific CRD-BP–RNA interaction could represent a novel therapeutic approach for the treatment of cancer.

We recently characterized CRD-BP–CD44 RNA interaction and showed that specific antisense oligonucleotides that disrupt CRD-BP–CD44 RNA interaction in vitro can indeed suppress CD44 mRNA expression in cells (King et al., 2014). In the present study, we characterized CRD-BP–GLI1 RNA interaction in vitro by first mapping the smallest GLI1 RNA that can still bind CRD-BP. On the basis of this information, we designed a 37-nt sense RNA oligonucleotide S1 as a potential competitor and hence an inhibitor of CRD-BP–GLI1 RNA interaction. We demonstrate that S1 RNA oligonucleotide can effectively inhibit CRD-BP–GLI1 RNA interaction in vitro and can suppress GLI1 mRNA expression in a panel of human colon and breast cancer cells.

Materials and Methods

Cell Culture. Colon cancer (HCT116 and HT29) and breast cancer (MCF-7, MDA-MB-231, and T47D) cell lines purchased from the American Type Culture Collection (ATCC; Rockville, Maryland) were maintained in minimal essential media (Life Technologies) supplemented with 10% fetal bovine serum in a humidified incubator at 37°C maintained in minimal essential media (Life Technologies) supplemented with 10% fetal bovine serum in a humidified incubator at 37°C supplemented with 5% CO2. Upon receipt of cells from ATCC, cells were harvested by 0.25% trypsin/0.02% EDTA treatment when they were in the logarithmic phase of growth for various experiments. and harvested by 0.25% trypsin/0.02% EDTA treatment when they were in the logarithmic phase of growth for various experiments.

Oligonucleotides and Primers. Table 1 shows sequences of oligonucleotides (S1 to S8 RNA oligo and D1 DNA oligo) used in this study. Oligonucleotides used for the electrophoretic mobility shift assays were standard phosphodiester derivatives (Integrated DNA Technologies Inc., Coralville, Iowa). For studies in cells, the RNA oligonucleotides were synthesized as 2′-O-methyl derivatives with a phosphodiester backbone (Integrated DNA Technologies Inc.). Table 2 shows the sequences of primers used to amplify the DNA template for use in synthesizing all the GLI1 RNA fragments described in Figs. 1 and 2.

Generation and Purification of Recombinant CRD-BP and Its Variants. The pET28b(+)–CRD-BP variant plasmids were constructed previously (Barnes et al., 2015). Recombinant CRD-BP was purified from Escherichia coli BL21 (DE3) using a column with 1-ml bed volume of nickel–nitrilotriacetic acid (NTA; QIAGEN, Valencia, CA) under denaturing conditions. Proteins eluted from the column at pH 5.4 were subjected to three steps of dialysis. The first step was for 24 hours in a pH 7.4 buffer containing 200 mM NaCl, 20 mM Tris-HCl, 1 mM reduced glutathione, 0.1 mM oxidized glutathione, 10% (v/v) glycerol, 2 M urea, and 0.01% (v/v) Triton X-100. The protein was then dialyzed twice, each for 2 hours in the same buffer as above but without urea and the glutathiones. Following dialysis, samples were centrifuged at 13,200 rpm for 30 minutes to remove any precipitated proteins. The purified protein solutions were quantified and analyzed for purity using Coomassie Brilliant Blue–stained 12% SDS-PAGE (Barnes et al., 2015).

Generation of DNA Templates and Radiolaabeled In Vitro Transcription. The human GLI1 cDNA (Accession #BC013000) purchased from Open Biosystems/GE Dharmacon (Lafayette, CO) was used as a template for the generation of PCR-amplified GLI1 DNA fragments. The different sets of forward and reverse primers used to amplify the different regions of GLI1 cdNA are shown in Table 2. PCR-amplified DNA templates were used directly for in vitro transcription by T7 RNA polymerase. One μg of DNA template was incubated for 1 hour at 37°C in a 20-μl reaction containing 1 × transcription buffer (Promega, Madison, Wisconsin), 10 mM dithiothreitol, 1 unit RNasin (Promega), 0.5 mM ATP, 0.5 mM CTP, 0.5 mM GTP, 12.5 μM UTP, 1.5 units T7 RNA polymerase (Promega), and 40 μCi [γ-32P]UTP (3000 Ci/ mmol). Following incubation, three units of RNase-free DNase I (Promega) were added and the reaction was further incubated for 10 minutes at 37°C. Upon addition of 10 μl of stopping dye (9 M urea, 0.01% Bromophenol blue, 0.01% xylene cyanole FF, 0.01% phenol), the entire sample was electrophoresed on an 8% polyacrylamide/7 M urea gel, and the band containing internally radiolabeled RNA was excised and eluted with elution buffer (10 mM Tris-HCl, pH 7.5, 0.1 M NaCl, 1 mM EDTA, 0.01% SDS) at 45°C for 6 hours. The purified, radiolabeled RNA was then phenol/chloroform extracted followed by ethanol precipitation. Specific activity of the RNA was then determined by scintillation counting.

Generation of Unlabeled RNA. For the synthesis of unlabeled GLI1 RNA nucleotides (nts) 230–420, 420–610, and 320–380 for use in competitive electrophoretic mobility shift assay, the following reactions and procedures were taken. Each in vitro transcription reaction contained 5 μg of gel-purified DNA template, 10 μl 10 × T7 buffer (400 mM Tris-HCl pH 7.6, 240 mM MgCl2, 20 mM spermidine, 0.1% Triton X-100), 5 μl each of 100 mM ATP, CTP, GTP, and UTP, 40 units RNasin, 5 units of T7 RNA polymerase to a total volume of 100 μl. The reaction was incubated at 37°C overnight and then treated with DNase I to remove the DNA template. Phenol-chloroform extraction was performed to remove protein impurities followed by passage through a G-50 spin column to remove unincorporated nucleotides. RNA was then ethanol-purified and resuspended in diethylpyrocarbonate (DEPC)-treated water. The purity of RNA was examined by visualization on 2.5% agarose gel, and concentrations were determined using a spectrophotometer.

Electrophoretic Mobility Shift Assay. The electrophoretic mobility shift assay (EMSA) was conducted in essentially the same manner as previously described (King et al., 2014; Barnes et al., 2015). The EMSA competition assays involved the preincubation between competitor molecules (oligonucleotides or RNA) and 300 nM CRD-BP for 10 minutes at 35°C. Following the preincubation, 13 nM [32P]-labeled GLI RNA nts 230–420 was added to the reaction. This was followed by the standard EMSA protocol. The molar excess concentrations of oligonucleotides or RNA over the [32P]-labeled GLI RNA are shown in Fig. 5.

Cell Transfection. Cells were plated in six-well plates at a density of 2.5 × 104 cells/ml. After 20 hours, cells were transfected with 20 nM double-stranded Dicer substrate RNA interference (dsRNAi), which was directed against CRD-BP mRNA (Integrated DNA Technologies) using Lipofectamine 2000 reagent (Life Technologies Inc.). The sense and antisense sequences were: r(GGAGGAGAACUUCUUUGGUCCCA) dAG and r(CUUGGAGAAAAGUGCACAGG). For the synthesis of unlabeled GLI1 mRNA, cells were transfected with the negative control duplex NC1 (Integrated DNA Technologies). For the synthesis of unlabeled GLI1 mRNA, cells were transfected with the negative control duplex NC1 (Integrated DNA Technologies). After a 48-hour incubation, cells from duplicate wells were subjected to RNA isolation as described below.

RNA Isolation and Quantitative Real-Time PCR. Total RNA was extracted from cells using the Ambion mirVana miRNA Isolation Kit (Thermo Fisher Scientific, Sunnyvale, CA) according to the manufacturer’s instructions. GLI1, CD44, c-myc, KRas, and β-actin mRNA levels were examined by quantitative real-time PCR (qPCR). After removing DNA using Ambion DNA Free kit (Thermo Fisher Scientific), the first strand of cDNA synthesis was performed using Scripto CDNA synthesis kit (Bio-Rad, Mississauga, Ontario) on 1 μg of total RNA. Following cDNA synthesis, qPCR reactions of GLI1, c-myc, KRas, and β-actin mRNAs were carried out using iQ SYBR Green Supermix (Bio-Rad) on an iQ5 Real-Time PCR System (Bio-Rad). The GLI1 PCR primers successfully used by
others (Noubissi et al., 2009) were: GLI1 forward primer, 5’-CCC AAT CAC AAG TCA GGT TCC T-3’, and GLI1 reverse primer, 5’-CTG ATG TGA AGC CCT ATT TGC C-3’. Specificity was confirmed by performing melt-curve analysis. The PCR primers used for measuring CD44, c-myc, and β-actin mRNA levels using the SYBR Green method have been previously described (King et al., 2014; Barnes et al., 2015). The cycling protocol consisted of 95°C for 3 minutes and 40 cycles of denaturation at 95°C for 10 seconds, and annealing at 52°C for 30 seconds. For measuring CRD-BP mRNA level, the iQ Supermix (BioRad) and FAM-TAMRA labeled probe included the following: CRD-BP forward primer, 5’-AGC TGG GAA AAG ACC TAC-3’; CRD-BP reverse primer, 5’-AAC CCT GAG AGG ACC ATC-3’; and CRD-BP TaqMan probe, 5’-/56-FAM/TGT TGC AGG GCC-b-actin forward primer, 5’-GGA TCC-3’; and β-actin reverse primer, 5’-AGG AAC CCT GAG AGG ACC ATC-3’. Specificity was confirmed by performing electrophoretic mobility shift assay in assessing these five samples relative to the control NC1-treated samples. One-way analysis of variance (ANOVA) was performed as statistical analysis to compare the target mRNA levels (GLI1, CD44, c-myc, and CRD-BP) of treated sample (siRNA- or S1-treated) with the negative control (NC1- or S4-treated) after normalization to reference gene (β-actin).

### Results

**Mapping the GLI1 RNA Region that Interacts with CRD-BP.** Using UV crosslinking coupled with immunoprecipitation of anti-FLAG CRD-BP, it was previously shown that CRD-BP has the strongest affinity for GLI1 mRNA within its coding region at nts 41–990 (Noubissi et al., 2009). To map the region within the first 990 nts of GLI1 mRNA that binds CRD-BP, we generated five RNA fragments of about 200-nt in size that span nts 36–990 of GLI1 mRNA. The five RNA fragments span GLI1 mRNA at the following nts: 36–231, 230–420, 420–610, 610–800, and 800–990 (Fig. 1A). The results of the electrophoretic mobility shift assay in assessing these five...
radiolabeled GLI1 RNA fragments in binding to the recombinant CRD-BP are shown in Fig. 2A. As shown in Fig. 2A, the $^{32}$P-labeled c-myc CRD RNA nts 1705–1886, which has a high affinity for CRD-BP (Barnes et al., 2015), resulted in a protein-RNA complex. Among the five GLI1 RNA fragments examined, only GLI1 RNA nts 230–420 showed affinity for CRD-BP (Fig. 2A). We found that some free RNAs, including GLI1 36–230 (Fig. 1A), exhibited more than one band on nondenaturing gel. We speculate that this could be the result of an alternative structure of the particular RNA or the formation of dimer RNA. We also noticed that some RNAs are more difficult to radiolabel, such as GLI1 610–800 and GLI1 800–990 (Fig. 1A). This could be the result of unusual secondary structure of RNA and/or the presence of low number of uridine residues. For instance, GLI1 800–990 has 29 uridine residues, whereas GLI1 230–420 and GLI1 420–610 have 45 and 41 uridine residues, respectively. We further mapped the region within nts 230–420 that is responsible for binding to CRD-BP. To do so, four GLI1 RNA fragments truncated from the 3’ end (Fig. 1B) and four RNA fragments truncated from the 5’ end were generated (Fig. 1C). Results of the EMSA in assessing their affinity for CRD-BP are shown in Fig. 2, B and C, respectively. Among the 3’-end-truncated fragments, only the RNA fragment corresponding to nts 230–380 displayed CRD-BP binding (Fig. 2B). On the other hand, we demonstrated that the three 5’-end-truncated fragments, nts 270–420, 320–420, and 350–420, had an affinity for CRD-BP (Fig. 2C). The combined experimental data from 3’-end- and 5’-end-truncated GLI1 RNA fragments suggests that the RNA region corresponding to nts 320–380 may be the smallest affinity site required for CRD-BP binding. To test this hypothesis, a GLI1 RNA fragment nts 320–380 was generated and then assessed using EMSA. As shown in Fig. 2D, CRD-BP does indeed have an affinity for this RNA fragment. The right panel in Fig. 1 summarizes the CRD-BP binding profiles of the GLI1 RNA fragments conducted in this study. 

Assessing Oligonucleotides for Ability to Inhibit the CRD-BP–GLI1 RNA Interaction. To determine if there is a distinct RNA structure present within the GLI1 RNA region that binds CRD-BP, the predicted RNA secondary structure of GLI1 nts 301–380 was generated using MFOLD (Zuker, 2003). Interestingly, the structure contains two distinct stem-loops within nts 346–382 (Fig. 3). This structure is remarkably similar to the two stem-loops generated for the secondary structure of CD44 RNA nts 2862–2930 (Fig. 3), which has previously been shown to have an affinity to CRD-BP (King et al., 2014). To test the hypothesis that the secondary structure, particularly the two stem-loops, as well as the sequence of GLI1 RNA are critical in binding CRD-BP, eight RNA and one DNA oligonucleotides of 37-nt were designed. The predicted secondary structures of these oligonucleotides were generated using MFOLD (Fig. 4). These sense oligonucleotides with identical lengths were designed to have different structures as well as slight variations in their nucleotide sequences as shown in Table 1. The sense S1 RNA oligonucleotide corresponds to GLI1 RNA nts 346–382, and the sense D1 RNA oligonucleotide is its DNA counterpart (Fig. 4). S2, S3, S4, S5, and S6 RNA oligonucleotides were designed with variations that included the removal of each loop (S2 and S3), different loop size (S4, S5, S6), or altered distance between loops (S4) so as to determine the importance of both stem-loops in CRD-BP binding. The S7 RNA oligonucleotide was designed to determine the importance of both stems in binding CRD-BP by altering the sequence at the stem, whereas S8 RNA oligonucleotide was designed to test the importance of the sequence within the loop. The ability of these oligonucleotides to compete with $^{32}$P-labeled GLI1 RNA nts 230–420 for binding to CRD-BP is shown in Fig. 5. As expected, the in vitro transcribed unlabeled GLI1 RNA nts 320–380 competed effectively (Fig. 5A). At 3 μM, the binding of $^{32}$P[GLI1 RNA nts 230–420 was inhibited by 74%, and at 22 μM, the binding was inhibited by 87%. As shown in Fig. 5, A and B, the S1 RNA oligo was able to inhibit $^{32}$P[GLI1 RNA binding by 89% at 147 μM. In contrast, the D1 oligo, a DNA oligonucleotide version of S1 RNA oligo, had no effect up to 147 μM, which indicates that the competition observed is RNA-specific and that the D1 oligonucleotide has no affinity for CRD-BP. The S2 RNA oligonucleotide, which is missing stem-loop 1 at the 5’ end (Fig. 4), effectively competed and inhibited CRD-BP binding by 40% at 147 μM, and the S3 RNA oligonucleotide missing stem-loop 2 at the 3’ end (Fig. 4) inhibited binding by 56% at similar concentration (Fig. 5B). Such reduced effectiveness suggests that the presence of both stem-loops is important for binding to CRD-BP. To determine whether the size of the loop region of both stem-loops is important, we designed and assessed S5 and S6 RNA oligonucleotides. The S5 RNA oligonucleotide with the smaller size loop at stem-loop 1 (Fig. 4) effectively inhibited CRD-BP binding by 76% at 147 μM (Fig. 5B). On the other hand, the S6 RNA oligonucleotide containing a smaller size loop at stem-loop 2 (Fig. 4) was markedly less effective, as it only inhibited CRD-BP binding by 16% at a similar concentration (Fig. 5B). These results suggest that the size of loops at both stem-loops is important and that stem-loop 2 likely plays a more important role in binding to CRD-BP. To determine if the distance between the two stem-loops is important, we designed S4 RNA
oligonucleotide. In contrast to S5 RNA oligonucleotide, which has three nts between stem-loops 1 and 2, S4 RNA oligonucleotide, which contains six nts between stem-loops 1 and 2 (Fig. 4), was completely ineffective as a competitor (Fig. 5B). This suggests that the close proximity of the two stem-loops is critical for CRD-BP–RNA binding. Finally, we assessed the significance of sequences at the stem regions and at the loop regions in binding CRD-BP. S7 RNA oligonucleotide has a secondary structure similar to S1 RNA oligonucleotide, differing only in the sequences at the two stem regions (Fig. 4). S8 RNA oligonucleotide also retained a similar secondary structure as S1 RNA oligonucleotide but differed in the sequences in the loop regions (Fig. 4). As shown in Fig. 5B, S7 RNA oligonucleotide was completely ineffective, and S8 RNA oligo only inhibited CRD-BP binding by 16% at 147 μM, suggesting that sequences at both the stem and loop regions are also important for CRD-BP binding.

Effect of Point Mutation in the GXXG Motif of KH Domains on the Ability of CRD-BP to Bind GLI1 RNA. Using site-directed mutagenesis to mutate the first glycine to an aspartate in the GXXG motif of heterogeneous nuclear ribonucleoprotein-K-homology domains (KH domains), we recently demonstrated that at least two KH domains of CRD-BP participate in binding c-myc and CD44.

Fig. 2. Mapping the CRD-BP binding site on GLI1 mRNA using electrophoretic mobility shift assay. (A) 5 different fragments (nts 36–230, 230–420, 420–610, 610–800, and 800–990) of [32P] GLI1 RNA (13 nM) covering nts 36–990 of GLI1 cDNA were incubated with various concentrations of recombinant CRD-BP as shown. (B) Four different 3’ truncated fragments (nts 230–380, 230–330, 230–300, and 230–275) of [32P] GLI1 RNA covering nts 230–420 of GLI1 cDNA were incubated with various concentrations of recombinant CRD-BP as shown. (C) Four different 5’ truncated fragments (nts 270–420, 320–420, 350–420, and 375–420) of [32P] GLI1 RNA covering nts 230–420 of GLI1 cDNA were incubated with various concentrations of recombinant CRD-BP. (D) [32P] GLI1 RNA nts 320–380 (13 nM) was incubated with various concentrations of CRD-BP. [32P] c-myc RNA nts 1705–1886 was used as the positive control as shown in (A) and (B). In each gel, free RNA and CRD-BP-bound RNA are indicated.
RNAs (Barnes et al., 2015). Here, we wished to further characterize CRD-BP–RNA interaction by determining whether CRD-BP has similar requirements for binding the GLI1 RNA. The previously described single-point-mutation KH variants (KH1, KH2, KH3, and KH4) and double-point-mutation KH variants (KH1-2, KH1-3, KH1-4, KH2-3, KH2-4, and KH3-4) were used in this study (Barnes et al., 2015). As additional controls, we used a Y5A variant, which has a point mutation in the RNA recognition motifs 1 domain and a D526E variant with a point mutation in the variable loop located between β2 and β3 in KH4 domain (Barnes et al., 2015). The representative results of electrophoretic mobility shift assay in assessing these variants for binding to [32P]-labeled GLI1 RNA nts 230–420 are shown in Fig. 6A, and the data summarized from three biologic replicates is shown in Fig. 6B. Both control variants, Y5A and D526E, exhibited a dissociation constant (Kd) that was comparable to that of the wild-type CRD-BP. Surprisingly, the KH2 variant could not bind to GLI1 RNA and the KH1 variant demonstrated significantly weak binding and its Kd could not be determined (Fig. 6B). The KH3 variant displayed a binding profile similar to that of wild-type CRD-BP, whereas the KH4 variant appeared to have a higher affinity for GLI1 RNA (Fig. 6B). All the double-point-mutation KH variants, including the KH3-4 variant, displayed inability to bind to GLI1 RNA.

Effect of Knocking Down CRD-BP on GLI1 and Other mRNA Expression in Breast and Colon Cancer Cells.

To further characterize CRD-BP–GLI1 RNA interaction and the downstream implications of this interaction, the ability of S1 RNA oligonucleotide to downregulate GLI1 mRNA expression was assessed. First, we needed to find cell line models in which the CRD-BP–mRNA interaction exists. To do this, we used a dsRNAi directed against CRD-BP on the basis of siRNA sequences previously shown to successfully knock down CRD-BP (Kato et al., 2007). The dsRNAi directed against CRD-BP had no effect on the level of β-actin mRNA in all cell lines tested, and hence it was used as a reference gene for normalization. Cells transfected with the negative control duplex NC1 were used as controls and the mRNA levels of all genes were expressed relative to the level in NC1-treated cells. As shown in Fig. 7, the dsRNAi directed against CRD-BP was effective in knocking down CRD-BP mRNA in two colon cancer cell lines, HCT116 and HT29, and three breast cancer cell lines, MCF-7, MDA-MB-231, and T47D. The knockdown appears to be most effective in HT29, MDA-MB-231, and T47D cells when the level of CRD-BP mRNA was reduced to 20% or less. The levels of CRD-BP mRNA were reduced in HCT116 and MCF-7 cells to 45% and 38% respectively. We found that the CT values of CRD-BP mRNA obtained in all NC1-treated cell lines were not significantly different. In the colon cancer cell lines (HCT116 and HT29) CRD-BP–knocked down cells, the GLI1 mRNA levels were significantly suppressed: A reduction to 35–38% was observed (Fig. 7). GLI1 mRNA was also suppressed to levels similar to the three breast cancer cell lines (Fig. 7). These results suggest that the CRD-BP–GLI1 mRNA interaction clearly exists in colon and breast cancer cells. C-myc and CD44 mRNAs, which are also targets of CRD-BP (Vikesaa et al., 2006; Barnes et al., 2015), were also measured to confirm the CRD-BP–RNA interaction in the CRD-BP–knocked down cells. Our results showed that a significant downregulation of both c-myc and CD44 mRNA is observed in all five cell lines when CRD-BP was knocked down (Fig. 7).
Fig. 4. Predicted secondary structures of oligonucleotides used against CRD-BP–GLI1 RNA interaction. The predicted secondary structures of DNA oligo D1 and RNA oligonucleotides S1 to S8 were generated using the MFOLD program (Zuker 2003).
Effect of S1 RNA Oligonucleotide on GLI1 and Other mRNA Expression in Breast and Colon Cancer Cells. Having shown that the physical interaction between CRD-BP and its target mRNAs exists in the cell lines studied above and on the basis of the hypothesis that CRD-BP binds to and protects GLI1 mRNA from degradation, we next tested the ability of S1 RNA oligonucleotide, an effective competitor of CRD-BP–GLI1 RNA interaction in vitro (Fig. 5), to influence GLI1 mRNA levels. The 2′-O-methyl-modified version of S1 RNA oligonucleotides was used. As a negative control, 2′-O-methyl-modified S4 RNA oligonucleotide was used, because the S4 RNA oligonucleotide is not an inhibitor of the CRD-BP–GLI1 RNA interaction (Fig. 5B). Our results showed that S4 RNA oligonucleotide had no effect on β-actin, GLI1, c-myc, and CD44 mRNA levels compared with NC1-treated cells (data not shown). Hence, we normalized the S1-treated mRNA level to that of S4-treated cells. The results in Fig. 8 show that S1 RNA oligonucleotide was indeed effective in suppressing GLI1 mRNA expression in all five cell lines. S1 RNA oligonucleotide was most effective in the two breast cancer cell lines MDA-MB-213 and T47D, where suppression to 18% and 25% was observed (Fig. 8). The suppression of GLI1 mRNA levels down to 30% and 35% was observed in HT29 and MCF-7 cells, respectively (Fig. 8). Though still statistically significant, the RNA oligonucleotide appears to be less effective in suppressing GLI1 mRNA in HCT116 cells, with an inhibition of only

Fig. 5. Inhibition of the CRD-BP–GLI1 RNA interaction by specific oligonucleotides as determined by EMSA. (A) Purified recombinant CRD-BP (300 nM) was incubated with [32P] GLI1 RNA nts 230–420 in the presence of different concentrations (μM) of GLI1 RNA nts 320–380, D1 DNA oligo, or S1 RNA oligonucleotide. (B) As in (A) different concentrations of competitor RNA oligonucleotides S1 to S8 were added to the EMSA reaction containing CRD-BP and [32P] GLI1 RNA nts 230–420. Lane 1 on each gel has no CRD-BP added. Lane 2 on each gel has 300 nM CRD-BP added but had no competitor added.
To investigate whether the S1 RNA oligonucleotide is specific for suppressing GLI1 mRNA or whether it can inhibit the general CRD-BP–RNA interaction leading to suppression of other CRD-BP mRNA targets, we also measured the levels of c-myc and CD44 mRNAs in S1-treated cells. Our results show significant downregulation of c-myc mRNA, although to a lesser extent than GLI1 mRNA, in HCT116, MCF-7, MDA-MB-231, and T47D cells transfected with the S1 RNA oligonucleotide (Fig. 8). CD44 mRNA level was also significantly downregulated in all cell lines compared with the control S4-treated cells (Fig. 8). But it is clear that S1 RNA oligonucleotide is less effective in downregulating CD44 mRNA, because suppression down to 60–70% is observed in all the cell lines. We found that there was no effect on KRas mRNA level in cells transfected with S1 RNA oligonucleotide (data not shown). These results imply that the S1 RNA oligonucleotide which targeted CRD-BP–GLI1 RNA interaction has some level of specificity in targeting the GLI1 mRNA expression in cells but also supports the hypothesis regarding the potential importance of the distinct two-stem-loop motif as a region for targeted therapy.

**Discussion**

There is now significant cumulative evidence to implicate CRD-BP in tumor progression. Although the mechanism by which CRD-BP exerts its oncogenic function may still be unknown, it has become increasingly clear that its ability to physically associate with its target mRNAs is one important criterion. To date, only a few studies have described the use of molecules capable of blocking the CRD-BP–RNA interaction (Coulis et al., 2000; Mao et al., 2006; King et al., 2014; Mahapatra et al., 2014), but none have specifically described potential inhibitors of the CRD-BP–GLI1 mRNA interaction. In this study, we further characterize the CRD-BP–GLI1 RNA interaction and report that a sense RNA oligonucleotide specifically designed on the basis of the two-stem-loop motif region mapped in GLI1 mRNA nts 320–380 is not only capable
of blocking CRD-BP–GLI1 RNA interaction in vitro but also able to inhibit GLI1 mRNA expression in a panel of colon and breast cancer cells.

On the basis of a previously reported study that CRD-BP has a high affinity for nts 41–990 in the coding region of GLI1 mRNA (Noubissi et al., 2009), we proceeded to map the smallest nucleotide sequence within this region to which CRD-BP still binds. Using electrophoretic mobility shift assay, we found the 61-nt GLI1 RNA fragment spanning the coding region nts 320–380 of GLI1 mRNA to be the smallest region with which CRD-BP could still physically associate (Fig. 2). This information is invaluable for the development of inhibitors of CRD-BP–GLI1 RNA interaction for: 1) effective design of oligonucleotides that could block the CRD-BP–GLI1 RNA interaction; 2) comparison of GLI1 RNA sequence and structures with those of other RNA binders, leading to a better understanding of the RNA sequences and structural requirements for CRD-BP–RNA binding; and 3) use of small fluorescently labeled GLI1 RNA in fluorescent polarization assays to screen for small molecule inhibitors of CRD-BP–GLI1 RNA interaction. With respect to the second point, the comparison of the MFOLD-predicted structure of GLI1 and CD44 RNAs led us to hypothesize that the two distinct common stem-loops (Fig. 3) are important for RNA interactions with CRD-BP. We therefore designed an S1 sense RNA oligonucleotide spanning the two-stem-loop regions of GLI1 RNA at nts 346–382 (Fig. 4), which turned out to be an effective inhibitor of the CRD-BP–GLI1 RNA interaction (Fig. 5). Our results using a number of variations of S1 RNA oligonucleotide showed that changes in the sequences and structure of the two stem-loops led to a significant decrease in its effectiveness as an inhibitor (Fig. 5), further corroborating the significant role played by the two stem-loops of GLI1 RNA in binding CRD-BP.

In addition to understanding the GLI1 RNA sequence and structural requirements for binding CRD-BP, we also investigated the role of each of the KH domains on CRD-BP in binding to GLI1 RNA. This was examined using KH variants of CRD-BP with a point mutation in the GXXG motif (Barnes et al., 2015). We found that the KH1 variant had significant retardation, and the KH2 variant was completely unable to bind to GLI1 RNA (Fig. 6). These results are startling and in contrast to our previous findings, where we demonstrated that the two mutated variants were effective binders of c-myc and CD44 RNAs (Barnes et al., 2015). Our results in this study, therefore, suggest that KH1 and KH2 domain of CRD-BP plays a greater role in the binding of GLI1 RNA compared with c-myc and CD44 RNAs. Surprisingly, we found that all the KH double-mutation variants, including KH3-4, had complete abrogation in ability to bind GLI1 RNA (Fig. 6). Again, this is in contrast to our previous findings that KH3-4 variant was an effective binder to c-myc and CD44 RNAs (Barnes et al., 2015). On the basis of these results, we propose that the other regions of the KH3 and KH4 domains play a less critical role in binding GLI1 RNA compared with c-myc and CD44 RNAs, and that the GXXG motif in KH3-4 di-domain play similar role as

Fig. 7. Effect of knocking down CRD-BP on GLI1 and other mRNA expression in breast and colon cancer cell lines. Colon cancer cells (HCT116 and HT29) and breast cancer cells (MCF-7, MDA-MB-231, and T47D) were plated and transfected with 20 nM of dsiRNA-CRD-BP or the negative control duplex NC1 as described in Materials and Methods. Total RNA extracted was subjected to quantitative real-time PCR for measurements of GLI1, c-myc, CD44, and β-actin mRNAs. The steady-state mRNA levels of GLI1, c-myc, and CD44 were normalized to the reference gene β-actin. Data shown are expressed as relative to the negative control NC1, which is taken as 1.0, and were averaged from five biologic replicates (n = 5) with S.E. as error bars. One-way ANOVA was performed as statistical analysis. Asterisk indicates results are significantly different from the negative control, NC1.
other di-domains in the binding of GLI1 RNA. Overall, our results further corroborate the notion that CRD-BP physically interacts with different RNAs in a distinct manner, and therefore an understanding of each unique CRD-BP–RNA interaction will be important for the design of inhibitors specific to the unique CRD-BP–RNA interaction.

A previous study showed that the overexpression of CRD-BP in 293T human embryonic kidney cells led to the stabilization of GLI1 mRNA and ultimately increased GLI1 protein level, whereas CRD-BP knockdown by shRNA prevented β-catenin/Tcf–dependent GLI1 upregulation and Wnt/β-catenin–dependent signaling induction of GLI1 transcriptional activity in 293T cells and NIH3T3 cells (Noubissi et al., 2009). Such results demonstrate the existence of the CRD-BP–GLI1 mRNA interaction in those cells. Furthermore, there was a positive correlation between the activation of β-catenin signaling and expression of CRD-BP and GLI1 in a panel of colorectal cancer cell lines (Noubissi et al., 2009). In this study, we knocked down CRD-BP using siRNA in two colorectal cancer cell lines, HCT116 and HT29, and three breast cancer lines, MCF-7, MDA-MB-231, and T47D, and showed that the selected mRNA targets of CRD-BP, including GLI1 mRNA, were suppressed (Fig. 7). These results suggest that the physical relationship between CRD-BP and its target mRNA in these cells exists. This is consistent with the model that CRD-BP binds to and protects its target mRNAs from decay. Having established that CRD-BP–GLI1 mRNA interaction exists in the five cell lines, the effect of the 2′-O-methyl derivative of the S1 RNA oligonucleotide on mRNA expression was evaluated. Our results show significant decrease in GLI1 mRNA level in all cell lines upon treatment with the S1 RNA oligonucleotide (Fig. 8). Although to a lesser extent, the S1 RNA oligonucleotide also significantly decreases c-myc and CD44 mRNA levels in all five cell lines (Fig. 8). These results suggested that the sense S1 RNA oligonucleotide designed on the basis of specific GLI1 RNA sequences and structure to target the CRD-BP–GLI1RNA interaction has some level of specificity in inhibiting GLI1 expression. While elucidating the exact mechanism of action of the S1 RNA oligonucleotide in cells is beyond the scope of this study, it is possible to speculate on its mode of action on the basis of in vitro studies and the chemical nature of the molecule. We reason that destabilization of GLI1 mRNA is not attributable to the RNase H pathway, because the S1 RNA oligonucleotide used in this study is a sense RNA and the 2′-O-methyl derivative is resistant to RNase H-mediated degradation (Schneider et al., 2011). We propose that the 37-nt sense S1 RNA oligonucleotide binds to the same site on CRD-BP as the GLI1 mRNA transcripts, and thereby occludes CRD-BP from binding to GLI1 mRNA, which leads to GLI1 transcript degradation. We hypothesize that this physical association also prevents c-myc and CD44 mRNAs from binding to CRD-BP.

Given that the aberrant Hh signaling pathway is implicated in the development of a number of human cancers, efforts have focused on finding effective inhibitors for this pathway.

Fig. 8. Effect of S1 RNA oligonucleotide on GLI1 and other mRNA expression in breast and colon cancer cell lines. Colon cancer cells (HCT116 and HT29) and breast cancer cells (MCF-7, MDA-MB-231, and T47D) were plated and transfected with 100 nM 2′-O-methyl derivative of S1 RNA oligonucleotide as described in Materials and Methods. Total RNA extracted was subjected to quantitative real-time PCR for measurements of GLI1, c-myc, CD44, and β-actin mRNAs. The steady-state mRNA levels of GLI1, c-myc, and CD44 were normalized to the reference gene β-actin. Data shown are expressed as relative to the negative control S4 RNA oligonucleotide which is taken as 1.0, and were averaged from five biologic replicates (n = 5) with S.E. as error bars. One-way ANOVA was performed as statistical analysis. Asterisk indicates results are significantly different from the negative control, oligo S4.
(Amakye et al., 2013). Unfortunately, most inhibitors which target the upstream effector protein SMO have turned out to be disappointing owing to the side effects produced by the SMO inhibitors and the presence of so-called “non-classical” Hh activation that is SMO-independent (Amakye et al., 2013; Shevde and Samant, 2014). To this end, there has been an increase in the development of inhibitors against GLI1 function (Amakye et al., 2013) and GLI1-DNA interaction (Infante et al., 2015). One of the nonclassic Hh activation pathways includes the Wnt/β-catenin signaling, which involves the stabilization of GLI1 mRNA by CRD-BP (Noubissi et al., 2009). Here, our characterization of the CRD-BP–GLI1 mRNA interaction has led to the design and discovery of a 37-nts sense S1 RNA oligonucleotide that can block CRD-BP–GLI1 RNA interaction in vitro and significantly suppress GLI1 mRNA expression in cells. The generated knowledge of the smallest GLI1 RNA binder of CRD-BP will now allow the development of a fluorescent polarization method to rapidly screen for small-molecule inhibitors of the CRD-BP–GLI1 RNA interaction. These findings, together with the new information on the important KH domains for GLI1 RNA interaction, can pave the way for the development of new inhibitors of the Hh signaling pathway that will act by destabilizing GLI1 mRNA via the inhibition of the CRD-BP–GLI1 mRNA interaction.

Acknowledgments

The authors thank Dr. Wai Ming Li for discussion and input on the manuscript.

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