In Vivo Genome-Wide CRISPR Activation Screening Identifies Functionally Important Long Noncoding RNAs in Hepatocellular Carcinoma

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SUMMARY

Our integrated transcriptomic data and functional CRISPR-based screening identified functional long noncoding RNAs that are frequently overexpressed in hepatocellular carcinoma. Cancer Susceptibility 11 (CASC11), the top candidate in the library screening, is shown to regulate the expression of MYC proto-oncogene in a cis-regulatory manner, which consequently dysregulates a subset of cell-cycle regulators and drives hepatocellular carcinoma tumor growth.

BACKGROUND & AIMS: Long noncoding RNAs (lncRNAs) are found to have profound impacts on diverse cellular processes. Although high-throughput sequencing studies have shown the differential lncRNA expression profiles between hepatocellular carcinoma (HCC) and nontumor livers, the functional impacts of lncRNAs on HCC development await further investigation. Herein, we sought to address the functional roles of lncRNAs in HCC pathogenesis by in vivo functional screening.

METHODS: We performed genome-wide clustered regularly interspaced short palindromic repeats (CRISPR)/dead CRISPR-associated protein 9 (dCas9) lncRNA activation screening in HCC xenografts. We characterized the clinical relevance of positively selected lncRNAs using transcriptomic data sets. We used CRISPR-based gene activation and knockdown approaches to show the functional roles of positively selected lncRNAs including Cancer Susceptibility 11 (CASC11) in HCC. RNA sequencing and chromatin isolation by RNA purification sequencing were used to investigate the molecular mechanisms of CASC11 in HCC progression.

RESULTS: The in vivo functional screening identified 1603 positively selected lncRNAs, 538 of which were overexpressed in HCC patients. Systematic transcriptomic data analysis and clinical investigation showed that patients with high expression of these lncRNA candidates correlated with aggressive tumor behaviors. Overexpression of these lncRNAs aggravated HCC cell growth. Detailed characterization of a lncRNA candidate, CASC11, showed its pivotal role in cell proliferation and tumor growth. Mechanistically, chromatin isolation by RNA purification sequencing showed that CASC11 was bound to the CASC11/MYC proto-oncogene shared promoter region on chromosome 8q24. CASC11 modulated the transcriptional activity of MYC in a cis-regulatory manner, which affected the expression of MYC downstream target genes, consequently promoting G1/S progression.

CONCLUSIONS: Our study showed the power of in vivo CRISPR screening, which comprehensively investigated the functionality of lncRNAs in HCC progression, providing a rationale for targeting these lncRNAs clinically.
 Hepatocellular carcinoma (HCC) is a highly aggressive primary liver malignancy that frequently occurs in patients with liver cirrhosis. HCC causes more than 800,000 deaths annually as a result of delayed diagnosis and lack of effective therapeutic interventions. Therefore, a better understanding of the underlying mechanisms of HCC development may provide valuable insights into the development of new predictive biomarkers and treatment strategies for HCC patients.

HCC development traditionally has been associated with the mutations of oncogenes and tumor-suppressor genes. However, beyond the protein-coding genome, the importance of the noncoding genome in HCC development has not been investigated sufficiently. The power of high-throughput sequencing technology has made a serendipitous discovery of a vast landscape of regulatory elements in the human genome. More than 80% of the human genome is actively transcribed into a myriad of noncoding RNAs. Among these, long noncoding RNAs (lncRNAs) represent the largest class of noncoding RNA species in the human transcriptome. lncRNAs, once considered as transcriptional noise, have shown their tremendous biological importance in chromatin remodeling, gene transcription, and RNA processing. Aberrant expression of lncRNAs has been shown to have profound effects on cancer hallmarks, including sustaining proliferative signaling; inhibiting cell death and growth suppressors; inducing immortality, angiogenesis, invasion, and metastasis; rewriting the metabolic program; and evading the immune system.

Although the differential expression of lncRNAs in human cancers has been accomplished by RNA-sequencing (RNA-seq) or microarray studies, such approaches do not provide evidence suggesting which lncRNAs are functional, let alone what roles they play in cancer development. Recent genome-wide functional screening approaches such as clustered regularly interspaced short palindromic repeats (CRISPR)/CRISPR associated protein 9 (Cas9) knockout screening have become a popular tool to identify protein-coding genes that are crucial for cancer cell growth and resistance to chemotherapeutic drugs in a high-throughput manner. However, the knockout approach attained by generating frameshift mutations within exons is not applicable to lncRNAs lacking open reading frames. For this reason, recent studies have exploited the power of CRISPR/dead Cas9 (dCas9)-based systems to manipulate the transcription of lncRNAs in in vitro functional screening. Previous studies used a cell culture system to identify functional lncRNAs promoting drug resistance. However, the expression of lncRNAs is influenced strongly by various factors where cancer cells reside, including the culture condition, the components of extracellular matrix, as well as the crosstalk between tumor cells and other cells in a tumor. To our knowledge, there currently is no systematic study showing the functions of lncRNAs at a large scale in HCC. Here, to precisely infer the functional roles of lncRNAs in HCC development, we took the advantage of genome-wide CRISPR activation screening in a xenograft mouse model. We identified a group of lncRNA candidates whose up-regulation significantly promoted HCC cell growth. Further gain- and loss-of-function experiments confirmed the oncogenic roles of Cancer Susceptibility 11 (CASC11) in HCC progression by positively regulating the transcription of MYC proto-oncogene in a cis-regulatory manner. CASC11 depletion evoked global expression changes of genes involved in MYC signaling and cell-cycle progression. Overall, our study provides a systematic framework that integrates the clinical transcriptomic data with functional CRISPR activation screening to uncover functional lncRNAs that influence HCC tumor growth.

### Results

**In Vivo Genome-Wide CRISPR Activation Screening Identifies a Subset of HCC Promoting lncRNAs**

To identify the functional lncRNAs that promote HCC development, we performed a genome-wide CRISPR/dCas9 lncRNA activation screening in xenograft mouse model (Figure 1A). MHCC97H cells expressing dCas9/VP64 and MS2 RNA-binding domain fused to the transcription activators p65 and heat shock factor 1 (MS2-p65-HSF1) were transduced with the human lncRNA activation library, which contains 96,458 single-guide RNAs (sgRNAs) targeting 10,504 lncRNAs, with 10 sgRNAs tiling the 800-bp upstream region of each lncRNA transcriptional start site. The successful infected clones were injected into both flanks of each mouse at a density of $2 \times 10^6$ cells subcutaneously. Considering the tumor viability, tumors from 20 mice were pooled for amplicon sequencing to achieve the 800× library representation. sgRNA abundance in tumors and

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**Abbreviations used in this paper:**

CASC11, cancer susceptibility 11; CDK, cyclin-dependent kinase; cDNA, complementary DNA; ChIP, chromatin immunoprecipitation; ChIRP, chromatin isolation by RNA purification; CRISPR, clustered regularly interspaced short palindromic repeats; CRISPRa, CRISPR activation; dCas9, dead clustered regularly interspaced short palindromic repeats–associated protein 9; FDR, false discovery rate; GAPDH, glyceraldehyde-3-phosphate dehydrogenase; GBA1, glucosylceramidase beta pseudogene 1; HCC, hepatocellular carcinoma; HKU, The University of Hong Kong; LNA, locked nucleic acid; lncRNA, long noncoding RNA; MS2, MS2 RNA-binding domain; MYC, MYC proto-oncogene; pCR, quantitative polymerase chain reaction; RNA-seq, RNA sequencing; SAM, synergistic activation mediator; SDS, sodium dodecyl sulfate; sgRNA, single-guide RNA; shRNA, short hairpin RNA; siRNA, small interfering RNA; TCGA, The Cancer Genome Atlas; TSS, transcriptional start site; VP64, transactivation component composed of four tandem copies of Herpes Simplex Viral Protein 16.

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**Figure 1. In vivo genome-wide CRISPR activation screening.** (A) Schematic representation of genome-wide CRISPR activation screening. (B) Scatterplot showing the enrichment of sgRNAs targeting the positively selected lncRNAs (FDR, <0.05) compared with nontargeting sgRNAs in tumors vs pretransplantation cells. (C) Pipeline for the identification of functional lncRNA candidates. FPKM, fragments per kilobase per million; MOI, multiplicity of infection.
pretransplantation cells was evaluated by high-throughput sequencing.

More than 90% of sgRNA library constructs were retained in all samples, collectively suggesting that our library sequencing had sufficient library coverage and read depth. We also evaluated the screen performance based on the distribution of nontargeting sgRNA controls, which were clustered mostly at the center of the scatterplot, suggesting that they were neither enriched nor depleted throughout the library screening (Figure 1B). We used the Model-based Analysis of Genome-wide CRISPR/Cas9 Knockout (MAGECK) algorithm with additional filtering steps to identify positively selected lncRNAs in our library screening (Figure 1C). We hypothesized that if a lncRNA is beneficial for HCC growth in a harsh tumor microenvironment, its corresponding sgRNAs would be enriched in tumors as compared with the pretransplantation cells. A total of 7211 sgRNAs (log₂ fold change > 1, false discovery rate [FDR] < 0.05) were significantly enriched in tumors, while only 3 sgRNAs were significantly depleted in tumors. These 7211 enriched sgRNAs targeted the promoter regions of 5283 lncRNAs, among which 1603 lncRNAs were targeted by at least 2 enriched sgRNAs (FDR < 0.05) (Supplementary Table 1).

To identify high-confidence positively selected lncRNAs, we evaluated the expression levels of lncRNA candidates by using RNA-seq data of HCC/non-tumor (NT) liver samples. Because some lncRNAs were not available in The Cancer Genome Atlas (TCGA)–HCC RNA-seq data, we also evaluated the expression profiles of positively selected lncRNAs in our RNA-seq from the University of Hong Kong (HKU) cohort. After filtering out lncRNAs with overall negligible fragments per kilobase per million (FPKM) values, we observed that most positively selected lncRNAs showed distinct differential expression patterns between tumors and nontumor tissues in both the TCGA–HCC and HKU cohort (Figure 2A). More than 60% of positively selected lncRNAs were overexpressed in HCC, whereas less than 10% of positively selected lncRNAs were down-regulated in HCC clinical samples (Figure 2B). For further analysis, we ranked the positively selected lncRNA candidates by calculating the average log₂ fold change of normalized sgRNA read counts between xenograft samples and pretransplantation cells. As a result, 538 lncRNAs of 1603 positively selected lncRNAs were overexpressed significantly in human HCC (Figure 2C). Interestingly, a subset of positively selected lncRNAs were reported previously to be oncogenic drivers in HCC, inclusive of LINC01138 and RAET1K, which served as positive controls of our in vivo library screening. In addition, we identified a subset of lncRNA candidates whose regulatory functions in HCC require more rigorous investigation, such as CASC11, Glucosylceramidase Beta Pseudogene 1 (GBAP1), LOC730101, LOC100499489, and LINC01133 (Figure 2D).

Intriguingly, there was no correlation between the level of sgRNA enrichment in xenograft tumors versus pretransplantation cells and the genomic locations of sgRNAs, suggesting that sgRNAs targeting closer to transcriptional start sites (TSSs) did not necessarily achieve greater sgRNA enrichment (data not shown).

**Clinical Relevance of Top lncRNA Candidates in HCC Patients**

We compiled a list of positively selected lncRNAs in the library screening for further characterization (Supplementary Table 1). Before the extensive experimental validation of our library screening, we first evaluated the clinical relevance of positively selected lncRNAs. We selected 7 lncRNAs among the top candidates who showed higher expression in tumors relative to nontumor tissues for further validation (Figure 3A). Importantly, Kaplan–Meier analysis showed that higher expression of lncRNAs, namely CASC11, GBAP1, LOC730101, LOC100499489, LINC01138, and LINC01133, was associated with poor overall survival in HCC patients (Figure 3B). To evaluate the hazard ratios associated with the expression of each positively selected lncRNA in 2 independent patient cohorts, we constructed a multivariate Cox regression model. Interestingly, CASC11, LINC01138, LOC730101, LOC100499489, and TCONS_00011697 appeared to have high a hazard ratio in relation to other clinical covariates, suggesting that they could be independent prognostic biomarkers for the risk assessment of HCC patients (Figure 3B). To predict the biological pathways associated with the up-regulation of the selected lncRNA candidates, we stratified HCC patients from TCGA-HCC and HKU-HCC cohorts into high and low expression groups based on the median expression level of selected lncRNA candidates in tumors, and performed gene set enrichment analysis using hallmark annotations retrieved from Molecular Signature Database. We identified positive enrichment of classical tumor pathways associated with more aggressive tumor behaviors (Figure 3C). Notably, patients with higher expression of CASC11, GBAP1, LINC01138, LOC730101, and TCONS_00011697 were associated with biological pathways that modulated cancer cell proliferation and cell-cycle progression, including E2F transcription factors (E2F) targets and Growth 2 phase/mitotic phase (G2/M) checkpoints (P < .05). Moreover, patients with higher expression of LINC01133 and LINC01138 were more likely to be associated with epithelial mesenchymal transition (P < .05). Patients with increased expression of CASC11, LOC730101, and LINC01133 showed higher enrichment of genes that modulated inflammatory response, cytokine production, and immune signature, suggesting that these lncRNAs may act as critical players in regulating the dynamics and plasticity of inflammatory networks and tumor immunity. To further explore the functional importance of our positively selected lncRNA candidates, we individually expressed the top enriched sgRNAs targeting the representative lncRNA candidates, showing that most enriched sgRNAs significantly induced the expression of their corresponding lncRNAs. In addition, transcriptional activation of selected lncRNA candidates significantly augmented HCC cell proliferation (Figure 4). Collectively, our clinical investigations and experimental validation reify our library screening as a powerful platform to identify clinically relevant lncRNAs that foster multiple cancer hallmark functions in HCC.
Figure 2. Expression levels of positively selected lncRNAs in HCC tumors. (A) Differential expression of positively selected lncRNAs between tumor and nontumor tissues. (B) Proportion of overexpressed lncRNAs as shown in the TCGA–HCC and HKU–HCC cohort. (C) Expression level of positively selected lncRNAs in tumors vs nontumors. Seven positively selected lncRNAs are indicated in the plot. (D) Rank of positively selected lncRNA candidates according to the average log2 fold change (FC) of their corresponding enriched sgRNAs (FDR, <0.05). NT, non-tumor; T, tumor.
CASC11 with 4 significantly enriched sgRNAs hits (log₂ fold change > 1, FDR < 0.05) was identified as a top-ranked positively selected lncRNA in the library screening. However, little is known about the functional roles of CASC11 in liver carcinogenesis. We confirmed the noncoding nature of CASC11 by using the ORf finder software from the National Centre for Biotechnology Information (Bethesda, MD). PhyloCSF codon substitution frequency analysis (Cambridge, MA), and our ribosome-sequencing data13,14 (data not shown). We observed higher expression of CASC11 in human HCC than nontumor livers, which was detected in 54% and 42% of HCC patients with paired HCC and NT liver samples in the TCGA-HCC (n = 50) and HKU-HCC (n = 63) cohorts, respectively (Figure 5A). Notably, hepatitis B virus and hepatitis C virus carriers were shown to have higher expression of CASC11 compared with patients with no viral infection background, suggesting that viral infection may influence the transcription of CASC11 (Figure 5B). The up-regulation of CASC11 was associated with advanced neoplasm histologic grades and the presence of vascular invasion in HCC patients (Figure 5C and D), indicating that CASC11 may serve as a prognostic indicator for the risk assessment of HCC patients.

CASC11 Drives HCC Cell Proliferation and Cell Progression In Vitro and In Vivo

To further corroborate the functional roles of CASC11 in HCC progression, we knocked down CASC11 expression in MHCC-97H and HepG2 cells by specific small hairpin RNAs (shRNAs) and locked nucleic acid (LNA) (Figure 6A). Knockdown of CASC11 significantly impaired HCC cell proliferative and colony-forming abilities (Figure 6B and C). We further examined the effects of CASC11 on tumor growth by introducing CASC11 overexpressing cells and CASC11 knockdown cells into nude mice subcutaneously. Consistent with our in vitro findings, CASC11 depletion impeded HCC growth in nude mice (Figure 6D). On the other hand, overexpression of CASC11 further aggravated HCC tumor growth in the CASC11-overexpressing group compared with the nontargeting control group (Figure 6E). Together, our in vitro and in vivo data collectively suggest that CASC11 is a prominent driver for HCC progression.

ChIRP-Seq Shows Significant Enrichment of CASC11 Binding on CASC11/MYC Shared Promoter

The oncogenic nature of CASC11 prompted us to investigate the mechanistic details of CASC11 in HCC further. We performed chromatin isolation of RNA purification sequencing (ChIRP-seq) to investigate the genomic binding region of CASC11. Two nonoverlapping ChIRP probe sets (ie, A and B pools) targeting the full length of CASC11 transcript were used to pull down CASC11 interacting DNA. ChIRP-seq signals that were enriched concordantly in both samples were considered as high-confidence CASC11 binding sites. Of the 2789 high-confidence peaks across the genome, more than 50% of peaks were mapped to genomic regions, with 12.6% of peaks annotated to gene promoters (Figure 7A). Kyoto Encyclopedia of Genes and Genomes pathway enrichment analysis showed that genes whose promoters were bound by CASC11 were associated with multiple cancer-related pathways, leading to the transcriptional misregulation in cancer (Figure 7B). Importantly, we found that CASC11 was enriched dramatically at the CASC11/MYC shared promoter region on chromosome 8q24 (Figure 7C and D, and Supplementary Table 2). To evaluate the binding efficacy and specificity of CASC11 on its target chromatin regions, we quantified CASC11 binding at MYC promoter by ChIRP–quantitative polymerase chain reaction (qPCR), using glyceraldehyde-3-phosphate dehydrogenase (GAPDH) as the promoter control. CASC11 probes successfully retrieved more than 70% of CASC11 transcripts, whereas probes tiling β-galactosidase (LacZ) retrieved neither CASC11 nor GAPDH, further confirming the pull-down efficacy of CASC11 probes (Figure 7E). Subsequent qPCR quantification showed that CASC11 binding was detected at the CASC11/MYC shared promoter (-1500 to -2000 and +100 to +400 relative to MYC TSS), whereas the GAPDH promoter did not show any CASC11 binding signals (Figure 7F). Collectively, our data showed that CASC11 is bound preferentially to the CASC11/MYC shared promoter region, indicating a potential role of CASC11 on cis-regulation of MYC transcription.

CASC11 Modulates MYC Transcription

CASC11 is localized mainly in the nucleus (Figure 8A). CASC11 lies next to MYC in a head-to-head manner, with its promoter 1467-bp apart from MYC (Figure 9A). To predict the functional connection between CASC11 and MYC, we looked into the RNA-seq profile from the TCGA–HCC cohort, which showed a strong correlation between CASC11 and MYC expression (Figure 8B). Notably, patients with MYC amplification had higher expression of CASC11 (P < .001) (Figure 8C). HCC samples with MYC gain/MYC amplification appeared to have significant expression correlation with CASC11, whereas MYC wild-type HCC samples did not show the expression correlation in MYC and CASC11 (Figure 8D).

Considering the strong enrichment of CASC11 at the CASC11/MYC shared promoter, we next asked whether CASC11 exerts its influence on HCC growth through trans
identified 683 differentially expressed genes upon CASC11 knockdown, of which 292 genes were down-regulated and 391 genes were up-regulated ($P < .05$) (Figure 10A and Supplementary Table 3). Gene set enrichment analysis showed that gene sets, namely G2M checkpoint and MYC targets from the Molecular Signatures Database Hallmark collection, were dysregulated significantly in CASC11 knockdown cells (Figure 10B and Supplementary Table 4). Consistently, we observed in the TCGA-HCC cohort that the expression of MYC target genes was enriched significantly in the CASC11-high-HCC group compared with the CASC11-low-HCC group (Figure 10C). MYC target genes and cell-cycle-associated genes were enriched positively in patients with high expression of CASC11, in concordance with the findings from the RNA-seq of CASC11 knockdown cells (Figure 10D).

To further analyze the functional relevance of CASC11 in MYC-driven HCC, we knocked down CASC11 in HCC cell lines with or without MYC overexpression (Figure 11A). We observed that MYC and CASC11 were overexpressed in HepG2 and MHCC97H, but had a low expression in Huh7 cells (Figure 11B). Knockdown CASC11 induced a pronounced reduction in MYC expression at both the transcriptional and translational levels in all tested cell lines (Figure 11C and D). We also investigated the status of several cell-cycle master regulators, such as cyclin-dependent kinase (CDK4), CDK6, DNA binding protein inhibitor 1 (ID1), DNA binding protein inhibitor 2 (ID2), and E2F transcription factor 1 (E2F1) in CASC11-depleted MHCC97H, HepG2, and Huh7 cells. These genes were down-regulated significantly at the transcriptional level upon CASC11 depletion. However, we found the reduced protein expression of CDK4 and CDK6 only in MHCC97H and HepG2 cells, but not in Huh7. Conversely, CRISPRA-induced overexpression of CASC11 profoundly up-regulated the expression of a subset of cell-cycle–related genes (Figure 11E). These findings collectively suggest CASC11 is the main driver of activated MYC signaling with MYC amplification. Previous studies have suggested that CDK4 and CDK6 were direct transcriptional targets of MYC. Supporting this, we observed significant MYC peaks on the promoter regions of CDK4 and CDK6 in the chromatin immunoprecipitation sequencing (ChIP-seq) profiles of HepG2 retrieved from the Encyclopedia of DNA Elements (Figure 12A). In addition, we observed that MYC and CDK4 were highly correlated with the expression of CASC11 (Figure 12B). We reasoned that CASC11 would modulate CDK4 and CDK6 expressions via MYC. To explore the functional connection between CASC11 and MYC, we silenced MYC and CASC11 expression, which consistently decreased the expression of MYC-regulating cell-cycle genes (Figure 12C). Pathway enrichment analysis showed that

CASC11 Drives HCC Progression Through a Subset of Cell-Cycle–Related Genes

Our in vitro and in vivo data, as well as the mechanistic studies, shows the important role of CASC11 in promoting HCC progression. To decipher CASC11 downstream pathways, we performed transcriptome sequencing analysis in MHCC-97H cells treated with LNA targeting CASC11. We observed that MYC and CASC11 were overexpressed in CASC11 knockdown cells (Figure 10C). MYC target genes and cell-cycle-associated genes were enriched positively in patients with high expression of CASC11, in concordance with the findings from the RNA-seq of CASC11 knockdown cells (Figure 10D).

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**Figure 5. Frequent up-regulation of CASC11 in HCC patients.** (A) Relative expression of CASC11 in HCC and nontumor liver tissues in TCGA cohort and HKU cohort. Data were presented as FPKM. (B) Expression of CASC11 in control (non–hepatitis B virus [HBV] or hepatitis C virus [HCV] carriers), HBV carriers, and HCV carriers (1-way analysis of variance with the Dunnett test). (C) Expression of CASC11 was associated positively with more advanced neoplasm histologic grade (1-way analysis of variance with the Dunnett test) and (D) vascular invasion in HCC patients (Student t test). All T, all tumor; NT, non-tumor; T, tumor; FPKM: fragments per kilobase per million.
Knockdown (KD) of CASC11 significantly inhibits HCC cell proliferation in vitro and in vivo. (A) Knockdown efficiency of CASC11 in MHCC-97H and HepG2 transfected with shRNAs or LNAs targeting CASC11 was measured by RT-qPCR. Relative expression was normalized to U6. (B) Relative cell viability of MHCC-97H and HepG2 after CASC11 KD. (C) The number of colonies was assessed after 2 weeks postseeding of stable CASC11 KD MHCC-97H and HepG2 cells. (D) Tumor growth of xenografts injected with stable CASC11 KD MHCC-97H cells or nontargeting cells (n = 10). Tumor volume and tumor weight were measured in the nontargeting control (NTC) and CASC11-KD group. (E) Percentage of total CASC11 RNA retrieved by biotinylated CASC11 antisense probes. The LacZ probe serves as a negative control. (F) ChIRP qPCR analysis measuring CASC11 enrichment across different regions of the CASC11/MYC shared promoter. GAPDH serves as a negative control. Results represent mean values of 3 independent experiments and the statistical significance was evaluated by 1-way analysis of variance with the Dunnett multiple comparison test. Data are plotted as means ± SD, *P < .05, **P < .01, ***P < .005, and ****P < .001. GABA, gamma-amino-butyric acid; LacZ, β-galactosidase.

Figure 7. ChIRP-seq shows genome-wide CASC11 binding sites. (A) Distribution of CASC11 binding sites at different genomic regions. Promoter regions are defined as ±3000 bp of the nearest gene’s TSS. (B) Kyoto Encyclopedia of Genes and Genomes pathway enrichment analysis of promoter-binding ChIRP-seq peaks. (C) Circos plot showing genome-wide CASC11 binding sites. (D) ChIRP-seq signals (fold enrichment over input) of CASC11-bound peaks at promoter regions. (E) Percentage of total CASC11 RNA retrieved by biotinylated CASC11 antisense probes. The LacZ probe serves as a negative control. (F) ChIRP qPCR analysis measuring CASC11 enrichment across different regions of the CASC11/MYC shared promoter. GAPDH serves as a negative control. Results represent mean values of 3 independent experiments and the statistical significance was evaluated by 1-way analysis of variance with the Dunnett multiple comparison test. Data are plotted as means ± SD, *P < .05.
Myc-knockout and CASC11-silencing cells shared many common downstream target genes that were associated with cell-cycle G1/S phase transition, DNA-dependent DNA replication, and nucleic acid metabolic processes (Figure 12D).

We examined the functional impact of CASC11 on cell-cycle progression. In concordance with differentially expressed gene analysis, CASC11 depletion induced the accumulation of cells at G1-phase and S-phase defects, whereas overexpression of CASC11 accelerated G1/S progression (Figure 11F and G). Intriguingly, the delay of G1/S progression as a result of MYC silencing was reversed by the ectopic expression of CASC11, indicating that CASC11 may play its role in cell-cycle control through MYC (Figure 11H). Taken together, CASC11 may promote cell-cycle progression through the modulation of MYC.

Discussion

In view of the high mortality rate of HCC, we are now endeavoring to understand the underlying mechanism of HCC pathogenesis. The exquisite complexity of the human genome portends a daunting task for HCC treatment. It becomes apparent that dysregulation of the noncoding genome has profound effects on cancer phenotypes. Despite extensive annotations of IncRNAs, the functional mechanisms of IncRNAs in HCC remain largely elusive, accentuating the necessity to devise an integrated pipeline that helps uncover functional IncRNAs in a high-throughput manner. One of the greatest challenges in studying human IncRNAs is the genomic complexity of IncRNAs and a lack of effective tools that can define their physiological functions in a high-throughput manner. CRISPR/Cas9 knockout screening has been used widely to identify essential protein-
coding genes that promote tumor growth and metastasis in xenograft mouse models, but an in vivo CRISPR activation screening targeting lncRNAs has not yet been reported.7,17–19

A variety of local and environmental factors including hypoxic conditions, proliferating tumor cells, infiltrating inflammatory cells, vascular system, stromal cells, and extracellular matrix create a unique environment that strongly influences molecular and cellular events, thereby promoting tumor growth.12 Considering the unique expression and tissue-specificity of lncRNAs, an in vivo screening has a greater potential to discover HCC-promoting lncRNAs in mouse models than in vitro cell cultures. Here, we conducted a CRISPR/dCas9 library activation screening in the xenograft mouse model to address the functional roles of lncRNAs in HCC progression. In contrast to CRISPR
knockout screening, in which the consistent enrichment of multiple sgRNAs indicates high-confidence screening hits, not every sgRNA can induce the transcriptional activation of target lncRNA loci. CRISPRa acts effectively within a small window around the targeted lncRNA TSS, in which the efficacy of targeted sgRNA is influenced strongly by several genomic features including chromatin accessibility, transcription factor binding sites, and the position of sgRNAs relative to the TSS. sgRNAs with low efficacy would greatly impede the overall statistical power in the gene-level analysis as designed for CRISPR knockout screening. Hence, we performed sgRNA-level analysis by comparing the relative abundance of individual lncRNA between tumors and pre-transplantation cells. Considering the potential of false-negative results from the enriched singleton sgRNA, only the lncRNA loci that were targeted by at least 2 corresponding sgRNAs were considered positively selected in tumors vs pretransplantation cells. As a result, we identified 1603 positively selected lncRNAs in our library screening, and the majority of them showed distinct expression patterns in tumors as indicated in transcriptomic data sets in the TCGA–HCC and HKU–HCC cohorts. In addition to the well-characterized lncRNA oncogenic drivers in HCC progression, we also identified a wealth of novel lncRNAs whose expression levels were comparatively higher in tumors, among which LINC01133, LOC100499489, LOC5730101, GBAP1, TCONS_00011697, and CASC11 promoted HCC cell proliferation upon CRISPRa overexpression, warranting efforts for more detailed functional characterization. Hence, the functional CRISPR screening facilitates the discovery of a wealth of novel predictive biomarkers. Importantly, overexpression of these positively selected lncRNAs promoted HCC growth, further confirming the reliability of our library screening and highlighting the importance of using an appropriate experimental system with the relevant physiological condition to define the functionality of lncRNAs.

For a better understanding of the modes of actions of lncRNAs in promoting HCC pathogenesis, we furthered our investigation on the functional importance of CASC11. Strikingly, our in vitro and in vivo studies showed the cell proliferation and tumorigenesis-promoting effects of CASC11. CASC11 is located at the 8q24.21 genomic region, where many cancer-associated single-nucleotide polymorphisms alter the transcription of lncRNAs, consequently affecting the transcriptional activities of MYC. MYC, regulating as many as 15% of genes in the human genome, acts as an important hub for orchestrating a broad spectrum of biological functions. Although scientists have made valiant efforts on studying the roles of MYC in various human cancer diseases for decades, it still is challenging to inhibit cancer growth by direct MYC inhibition owing to its potential detrimental consequences for normal cell development. Hence, there is a voracious appetite for strategies that indirectly interfere with MYC transcription. Indeed, several cis-acting lncRNAs have been reported to regulate MYC transcription, through facilitating the activity of MYC promoter-binding transcription factors and the formation of higher-order chromatin architecture. For example, MYC-modulating lncRNA (MYMLR) was found to regulate the MYC promoter activity by binding to MYC enhancer. Although MYMLR shows some overlap with CASC11, they are transcribed from different promoters and they showed different regulatory cascades, indicating the genomic complexity of this locus.

Realizing that CASC11 is transcribed divergently near the MYC promoter, we pursued our investigation on the complex interplay between MYC and CASC11. Notwithstanding multiple lines of evidence showing the oncogenic role of CASC11 in different cancers, to our knowledge, there is a paucity of studies on the functional mechanism of the CASC11/MYC axis in HCC progression. Considering an increasing body of evidence showing that lncRNAs regulate adjacent protein-coding genes, we speculated that CASC11 functions through MYC, consequently dysregulating its downstream targets. ChiRP-seq analysis showed that CASC11 preferentially is bound on the CASC11/MYC promoter region. In addition, our functional assays unequivocally suggested that the perturbation of CASC11 expression drastically affected MYC expression through cis-regulatory function. The explanation could be that CASC11 exerts its local function—either in the act of its transcription or in its own transcript—to modulate the chromatin-associated processes and influence the transcription of its neighboring gene MYC. These findings also highlight the importance of using the CRISPR activation system to activate the target lncRNA transcription at the endogenous level, thereby capturing the local functions of lncRNAs and recapitulating all lncRNA isoforms at the targeted loci, whereas this is not the case in traditional cDNA overexpression.

Although we showed the cis-regulatory role of CASC11 on MYC transcription, several questions regarding the complex regulatory circuitry of CASC11 and MYC await further investigation. Increasing evidence has shown that antisense lncRNA transcript mediates R-loop formation in the local chromatin that favors the binding of the transcriptional machinery, thereby inducing the transcriptional activation of its neighboring gene. Toward this, we observed an enriched DNA:RNA immunoprecipitation sequencing signal in the CASC11/MYC shared promoter. The involvement of CASC11 in MYC transcriptional activity may be pertinent to a burgeoning concept of liquid–liquid...
phase separation, in which IncRNAs facilitate the assembly of RNA binding proteins to form a spatial compartment in the nucleus for gene transcription.\(^3\)

Cell-cycle dysregulation is a central hallmark of cancer progression.\(^3\) We showed that CASC11 promoted HCC cell growth by dysregulating cell-cycle pathways and MYC-associated pathways. However, because our ChIRP-seq data did not show significant CASC11 enrichment on the CDK4 and CDK6 promoters, we speculated that the down-regulation of CDK4 and CDK6 expression in CASC11-depleted cells presumably was caused by the down-regulation of MYC. Interestingly, CASC11 knockdown and MYC knockout cells showed consistent enrichment in many common downstream targets, which were particularly associated with G1/S-phase progression and DNA replication. Importantly, MYC silencing decelerated the rate of G1/S progression driven by CASC11 overexpression, lending support to the idea that CASC11 exerts its functional role through MYC.

Conclusions

In summary, our study has unveiled the power of genome-wide CRISPR activation screening in tumor xenografts to uncover oncogenic IncRNAs, providing a rationale to target these IncRNAs clinically. We integrated computational analysis of clinical transcriptomic data sets with functional CRISPR activation screening, which systematically showed the physiological relevance of IncRNAs in promoting HCC progression. As proof-of-principle, we showed that CASC11 is a prominent driver in HCC progression through regulating the expression of MYC and its downstream targets. Further investigations are required to translate the experimental findings into clinical application by developing functional IncRNA candidates as potential biomarkers and therapeutic targets for HCC patients.

Materials and Methods

Clinical Specimens

HCC and their corresponding nontumorous liver tissues were obtained from 16 patients with surgical resection at Queen Mary Hospital between 1997 and 2007. The use of clinical specimens was approved by the Institutional Review Board of the University of Hong Kong and the Hong Kong Hospital Authority.

Cell Culture

The HCC cell line MHCC97H was obtained from Dr Z. Y. Tang (Fudan University, Shanghai, China). Huh-7 was a gift from Dr H. Nakabayashi (Hokkaido University, Hokkaido, Japan). 293FT was obtained from American Type Culture Collection (Manassas, VA). All of these cell lines were cultured in Dulbecco’s modified Eagle medium (Gibco, Grand Island, NY) supplemented with 10% fetal bovine serum (Life Technologies) and 100 U/mL penicillin/streptomycin (Invitrogen). MHCC97H cells were supplemented with 1 mmol/L sodium pyruvate.

Cell Line Authentication

Human HCC cell lines were authenticated by the Authentication PCR Amplification Kit (Life Technologies). To evaluate whether the HCC cell lines used in this study resembled the primary HCC tumors, a comprehensive pan-cancer analysis was performed by comparing the transcriptomic profiles of our cell lines with more than 9753 primary tumors from TCGA and 1378 cell lines from the Cancer Cell Line Encyclopedia encompassing 32 cancer types. Reference-Based Single-Cell RNA-seq Annotation (Single R, San Francisco, CA) was used to infer the cell of origin of our tested cell lines by using the aforementioned reference transcriptomic data sets.

Pooled Library Amplification and Transduction

Human CRISPR 3–plasmid IncRNA activation pooled library (1000000106; Addgene, Watertown, MA) was a gift from Professor Feng Zhang. Briefly, the Refseq noncoding RNAs (release 69) catalog and the Broad Institute IncRNA catalog were combined to filter for all major classifications of IncRNAs no less than 200-bp long and their TSSs ≥50-bp away from the neighboring gene\(^3\)\(^5\)\(^3\). This results in a library composed of 95,958 sgRNAs targeting 10,504 IncRNAs. Each IncRNA TSS was targeted by 10 sgRNAs. Each sgRNA was individually cloned into BsmB1 sites of lentivirus (MS2) backbone (61427; Addgene). Five hundred nontargeting sgRNAs were included as nontargeting controls. sgRNA library cloning was described in a protocol by Joung et al.\(^1\) The sgRNA library was amplified at 50–100 ng/μL using Endura Electrocompetent cells (60242; Lucigen, Middleton, WI) according to the manufacturer’s instructions.

Figure 11. (See previous page). CASC11 regulates cell-cycle progression through MYC. (A) Western blot showing the protein expression of MYC in a panel of HCC cell lines and 2 immortalized normal liver cell lines. α-tubulin served as housekeeping normalization. (B) Transcript level of MYC and CASC11 in a panel of HCC cell lines and 1 pair of HCC clinical samples as determined by RT-qPCR. Data were normalized with U6 and compared with the nontumor (NT) liver sample (67 NT). (C) RT-qPCR validation of cell-cycle–related genes upon LNA-mediated CASC11 knockdown for 48 hours. Relative expression was normalized to U6. (D) Western blot showing protein expression of CDK4, CDK6, MYC, and α-tubulin upon CASC11 silencing in MHCC-97H, HepG2, and Huh7 cells. (E) RT-qPCR measurement showing the up-regulation of CASC11-regulated genes as a result of CASC11 overexpression. (F) Cell-cycle profiles of MHCC-97H and HepG2 cells by flow cytometry using propidium iodide (PI) staining upon CASC11 knockdown for 72 hours. (G) Cell-cycle profiles of MHCC-97H cells by flow cytometry using PI staining after activation of CASC11. (H) Cell-cycle profiles of CASC11-overexpressing cells infected with MYC targeting siRNAs. Result is presented with respect to the control. (A–H) Results represent mean values of 3 independent experiments. Statistical significance was evaluated by 1-way analysis of variance with the Dunnett multiple comparison test. Data are presented as means ± SD, *P < .05, **P < .01, ***P < .005, and ****P < .001. RT-qPCR, Quantitative reverse transcription PCR.
For transduction, plasmids were packaged into lentivirus with the appropriate lentiviral packaging plasmid system (pPACKH1-GAG, pPACKH1-REV, and pVSV-G; System Biosciences, Palo Alto, CA) in HEK293FT cells using Lipofectamine 3000 (Life Technologies) according to the manufacturer’s instructions. CRISPR SAM is a 3-vector system: dCas9–VP64–blast (61425; Addgene), MS2–P65–HSF1 activator complex with a 2A Hygromycin resistance marker (MPhV2, 89308; Addgene), and lenti-sgRNA (MS2) zeo (61427; Addgene). MHCC97H cells were transduced with dCas9–VP64 and MPhV2. The successfully transduced HCC cells were selected by blasticidin (10 \( \mu \)g/mL) (Life Technologies) and hygromycin (300 \( \mu \)g/mL) (Life Technologies) for 5 days.

Before the sgRNA library transduction, the concentration of the selection reagents (ie, zeocin) was determined by a kill curve (Life Technologies). Lentiviral titer was calculated by adding different doses of lentiviral supernatant to a fixed number of cells and determining the cell viability after complete selection for 7 days. The multiplicity of infection (MOI) was calculated as the number of transduced cells under antibiotic selection divided by the number of cells without antibiotic selection. MHCC-97H cells expressing SAM were transduced with lentiviral sgRNA library at a low MOI (MOI, <0.3) to make sure each cell contained only 1 sgRNA. Transduced cells were selected in 300 \( \mu \)g/mL zeocin (Life Technologies) for 7 days. Cells were maintained at more than 500 cells per sgRNA during passing to ensure sufficient library coverage. After a week of zeocin selection, successful clones were divided into pretransplantation cells and cells for in vivo screening. For sgRNA validation, sgRNAs were individually cloned into the BsmB1 site of
lenti-sgRNA (MS2) optimized puromycin backbone. The cloned sgRNAs were delivered in SAM complex expressing MHCC97H cells, followed by 2 µg/mL puromycin selection for 5 days.

**In Vivo Library Screening**

All animal works were performed under the guidelines of the Animal Experimentation Ethics Committee of the University of Hong Kong. A total of $8 \times 10^7$ sgRNA library transduced MHCC97H cells were used to ensure library representation. Cells were resuspended in phosphate-buffered saline and Matrigel (BD Biosciences, Franklin Lakes, NJ) (1:1 ratio) and injected into both flanks of 4- to 6-week-old male BALB/c nude mice at a density of $2 \times 10^6$ cells per flank. At 4 weeks after transplantation, mice were killed.

**Genomic DNA Extraction**

To maintain the full library representation, 330 µg genomic DNA was used. Genomic DNA was extracted from tumors and pretransplantation cells by salt precipitation as genomic DNA was used. Genomic DNA was extracted from analyzing by the MAGeCK algorithm (0.5.7). Results from 2 independent deep-sequencing results were ended sequencing analysis was performed by Novogene and ethanol precipitation. Massive parallel amplion pair-precipitation (Qiagen) and proteinase K (Life Technologies) at 55 mmol/L Tris, 50 mmol/L EDTA, 1% sodium dodecyl sulfate (SDS), pH 8) and proteinase K (Life Technologies) at 55°C overnight. The lysate mixture was incubated with RNaseA (Life Technologies), 7.5 mol/L ammonium acetate, isopropanol, and ethanol for genomic DNA precipitation.

**PCR Amplification of sgRNA Regions and sgRNA Readout by Deep Sequencing**

Before deep-sequencing analysis, sgRNA regions were amplified from genomic DNA using NEBnext High Fidelity 2× Master Mix (New England Biolabs, Ipswich, MA) with 24 cycles of PCR reactions. U6 primers for sgRNA amplification are listed in Supplementary Table 5. A total of 330 µg genomic DNA was used per sample to maintain the library representation. PCR products were purified by gel electrophoresis at 100 V for 50 minutes, followed by visualization under UV illumination in Alphalmager 2200 (Alpha Innotech, San Leandro, CA), followed by gel purification (Qiagen) and ethanol precipitation. Massive parallel amplion pair-ended sequencing analysis was performed by Novogene Technology to evaluate sgRNA abundance (Beijing, China). Results from 2 independent deep-sequencing results were analyzed by the MAGeCK algorithm (0.5.7).

**Computational Analysis of CRISPR Activation Screening**

We used the MAGeCK algorithm (0.5.7) to identify positively selected lncRNAs in the library screening. Briefly, reads from Illumina sequencing (San Diego, CA) were demultiplexed by Trim-Galore (version 0.6.5) (RRID:SCR_011847). The MAGeCK algorithm count was used to annotate the sequencing reads and adjust the read count distribution and sequencing depth of all samples using median normalization. sgRNA sequences were filtered further based on the following criteria: duplicate entries of the same sgRNA sequence; and sgRNA sequence representing more than 1 lncRNA loci. The average fold change of normalized sgRNA read counts, negative binomial P value, and FDR of each sgRNA across independent experimental replicates were calculated using the MAGeCK test function. lncRNAs were considered to be positively selected if they were targeted by at least 2 corresponding sgRNAs with a FDR of less than 5%.

**cDNA Overexpression**

A lentiviral expression Cytomegalovirus vector (CMV) with multiple cloning sites (MCS) carrying Flag and HA tags (pCDH–CMV–MCS–3X–Flag–3X–HA–puro vector) was used to express the full length of CASC11 cDNA. Briefly, pCDH–CMV–MCS–3X–Flag–3X–HA–puro vector was digested with NotI and XbaI to remove HA and Flag sequence. The full length of CASC11 cDNA was cloned into the digested pCDH vector. The confirmed clone was verified by sequencing. MHCC97H cells and the Alexander hepatoma (PLC/PRF/5) cells were transfected with either pCDH–CASC11 vector or pCDH empty vector using the Lipofectamine 3000 transfection kit (Life Technologies) at a MOI < 0.5 and subsequently selected with puromycin for 3 days. The transcript levels were measured by qPCR. The sequence of cDNA overexpression is listed in Supplementary Table 5.

**Nuclear and Cytoplasmic Fractionation**

Nuclear and cytoplasmic fractionation of MHCC97H cells were prepared using the RNA subcellular Isolation Kit (Active Motif, Carlsbad, CA) as described by the manufacturer’s protocol. GAPDH glyceraldehyde-3-phosphate dehydrogenase (GAPDH) was used as a cytoplasmic endogenous control, while U6 small nuclear 1 (U6) was used as the nuclear endogenous control. The efficiency of fractionation was determined by qPCR, followed by 2% agarose gel electrophoresis in 1× Tris–borate–EDTA buffer.

**Cell Proliferation Assay and Colony Formation Assay**

For the cell proliferation assay, cells were seeded at a density of 10,000 cells/well in a 24-well plate in triplicate and incubated in a 37°C humidified CO₂ incubator. The number of cells was quantified for 6 days by a Z1 Coulter Counter Cell and Particle Counter (Beckman Coulter, Brea, CA). For colony formation assays, 1000 cells were seeded in a 6-well plate until colonies formed in complete medium (approximately 14 days). Colonies were fixed in methanol and stained with 0.1% crystal violet (Sigma-Aldrich, Burlington, MA) for visualization. ImageJ (National Institutes of Health, Bethesda, MD) was used to count the number of colonies.

**Cell-Cycle Analysis**

Cells were fixed with 75% ethanol at 4°C overnight. The next day, cells were treated with 10 µL of 2 mg/mL
propidium iodide (Sigma-Aldrich) and 10 mg/mL RNase-A (Life Technologies) at 37°C for 30 minutes. Analysis was performed on a FACSCanto II Flow Cytometer (BD Biosciences).

**Reverse-Transcription and Real-Time PCR**
Total RNA was extracted by TRIzol reagent (Invitrogen). First-strand cDNA was synthesized from 1 μg total RNA using Fast PrimeScript Reverse Transcription (RT) Master Mix (TaKaRa, San Jose, CA). qPCR analysis was performed using SYBR green master mix (Life Technologies) on an Applied Biosystem StepOnePlus Real-time PCR system (Life Technologies). Sequences for qRT-PCR primers are listed in Supplementary Table 5. Hypoxanthine-guanine phosphoribosyltransferase (HPRT) or U6 served as an internal control. Relative RNA levels were calculated using comparative cycle threshold (CT) (2^ΔΔCT).

**Western Blot**
Cells were lysed in RIPA buffer (50 mmol/L Tris-HCl, 150 mmol/L NaCl, 0.1% SDS, 5 mmol/L EDTA, and 0.5% sodium deoxycholate) containing proteinase and phosphatase inhibitors (Roche, Basel, Switzerland). Protein samples were separated on SDS–polyacrylamide gel electrophoresis, transferred to a hydrophobic polyvinyl iodine difluoride membrane (GE Healthcare, Chicago, IL), and probed with primary antibodies at 4°C overnight and secondary antibodies at room temperature for 1 hour. Antibodies used for Western blot were obtained from Cell Signalling Technology and Santa Cruz: anti-MYC antibody (sc-23896; Cell Signalling Technology), anti-CDK4 antibody (sc-23896; Santa Cruz, Santa Cruz, CA), and anti–α-tubulin antibody (2144S; Cell Signalling Technology).

**RNA Interference**
CASC11 targeting shRNA sequences were cloned into the pLKO.1 puro plasmid. siRNAs targeting MYC were designed by Integrated DNA Technology (Coralville, IA). GapmeR antisense (Exiqon, Skelstedet, Vedbaek) oligonucleotides (LNA) were designed by Qiagen. siRNAs and LNAs were transfected at 500 nmol/L and 50 nmol/L using RNAiMAX transfection reagent (Life Technologies) or Lipofectamine 3000 (Life Technologies) according to the manufacturer’s protocol. Sequences of shRNAs, siRNAs, and LNAs are listed in Supplementary Table 5.

**ChIRP-seq**
ChIRP-qPCR assay was performed as described previously.35 A total of 11 antisense biotinylated probes against CASC11 was designed by LGC Biosearch Technologies (Petaluma, CA). The probes were split into 2 pools (probe set A and B). Two independent ChIRP-seq runs with 2 pools were performed separately. Briefly, 1 × 10^7 cells were cross-linked with 1% glutaraldehyde for 10 minutes (Sigma-Aldrich) and quenched with 0.125 mol/L glycine (Sigma-Aldrich) for 5 minutes. Cells were lysed in complete lysis buffer (50 mmol/L Tris-HCl, pH 7.0, 10 mmol/L EDTA, 1% SDS, Protease inhibitor Cocktail III (Calbiochem, San Diego, CA) and RNase inhibitor (Life Technologies) and sonicated with a Bioruptor sonicator (Bioruptor; Diagenode, Liège, Belgium) to an average length of 100–500 bp. Chromatin was hybridized with 100 pmol of CASC11 antisense probes and complete hybridization buffer (Protease inhibitor Cocktail and RNase inhibitor) and incubated at 37°C overnight. The RNA-chromatin hybrid was captured by Streptavidin C1 magnetic beads (Life Technologies) and washed with wash buffer (2 × saline sodium citrate, 0.5% SDS, and Protease inhibitor Cocktail). Ten percent of bead samples were purified for RNA isolation and 90% of bead samples for DNA isolation. For RNA isolation, bead samples were subjected to proteinase K and RNA elution buffer (Tris, pH 7.0, 1% SDS) and incubated at 95°C for 15 minutes, followed by TRIzol-chloroform extraction and the miRNeasy Mini column purification (Qiagen). One-step, real-time, reverse-transcription qPCR was performed to quantify the enriched transcripts in eluted RNA. For DNA isolation, bead samples were resuspended in complete DNA elution buffer (50 mmol/L) and incubated at 37°C for 30 minutes with gentle rotation. Then, samples were subjected to RNaseA, RNaseH, and proteinase K treatment and incubated at 50°C for 45 minutes, followed by phenol:chloroform:isoamyl:alcohol extraction (Sigma). Eluted DNA was quantified by qPCR or sequencing library construction. For ChIRP-qPCR, bound chromatin was assessed using primers targeting the MYC promoter region. ChIRP pull-down of GAPDH served as a negative control. For ChIRP-seq, the library preparation and deep sequencing were conducted by the Centre for PanorOmic Sciences, HKU. ChIRP-seq raw reads were uniquely mapped to the human genome (GRCh38) using Bowtie2 (version 2.4.1) (RRID:SCR_016368), followed by a peak calling algorithm using MACS2 (version 2.2.7) (RRID:SCR_013291) with a q-value cut-off of 0.01. Fold enrichment of chromatin association of CASC11 was calculated by normalizing the common peak signals of 2 pools to input. Peak signals of probe set A and probe set B samples were normalized to input. Probes and primer sequences are listed in Supplementary Table 5.

**Clinical Investigation of Positively Selected IncRNAs in HCC Clinical Samples**
Positively selected IncRNAs were dichotomized into high- and low-expression groups based on either mean or median expression cut-off level, whichever gave the greatest degree of discrepancy. The overall survival rate in HCC patients above and below the cut-off expression level was calculated using the Kaplan–Meier method. To investigate the effect of positively selected IncRNAs on survival, IncRNA expression in relation to clinical parameters including tumor stage (tumor stages 1 and 2 vs tumor stages 3 and 4), histologic grade (histologic grades 1 and 2 vs histologic grades 3 and 4), gender (male vs female), and age (< median age vs > median age) were assessed in a multivariate Cox regression model. Hazard ratios were calculated to evaluate the prognostic effect of IncRNAs.
RNA-Seq and Bioinformatics Analysis

Transcriptome sequencing (RNA-seq) was performed in 16 pairs of HCC samples and the corresponding nontumor samples (Bioproject Accession ID: 294031). Library preparation and RNA-seq of HCC11 knockdown cells and MYC knockout cells were performed as described previously.\textsuperscript{5} Differential expression analysis was performed using EdgeR (RRID:SCR_012802) with default parameters. Gene set enrichment analysis was performed using fast gene set enrichment analysis R package (version 3.10) (RRID:SCR_001905) with default parameters. Pathway annotations of Hallmark gene sets were retrieved from the Molecular Signatures Database and Gene Ontology.

ChIP-Seq Analysis

ChIP-seq of MYC was obtained from the Encyclopedia of DNA Elements. Read alignment was performed by Bowtie2 and peak calling by MACS2.

Pathway Enrichment Analysis

Gene set enrichment analysis was performed to evaluate genes that are over-represented in the experimental condition vs the control by fast gene set enrichment analysis R package (version 3.10) with default parameters. Annotations of different gene sets were retrieved from the Molecular Signatures Database.

TCGA Data

Clinical information of HCC patients including RNA-seq expression profiles of HCC patients (FPKM value), disease-free and overall survival, as well as histologic grades were downloaded from TCGA via the Broad Institute (http://gdac.broadinstitute.org). Gene set enrichment analysis on TCGA data was performed using gene set enrichment analysis version 4.0.3 (https://www.gsea-msigdb.org/gsea/downloads.jsp).

Prediction of Protein-Coding Capacity of IncRNA

The protein-coding potential of IncRNAs was predicted using CPAT (http://lilab.research.bcm.edu/cpat) and ORF finder (https://www.ncbi.nlm.nih.gov/orffinder). Homo sapiens HOX transcript antisense RNA (HOTAIR) served as the noncoding RNA control while GAPDH served as coding control.

Statistical Analysis

All statistical analysis was performed using RStudio (version 3.6.1) and GraphPad PRISM software (version 9.2; GraphPad Software, San Diego, CA). Data are presented as means ± SEM. The Student t test or 1-way analysis of variance was used to determine the difference between 2 groups or more than 2 groups, followed by the Dunnett multiple comparison test. A Kaplan–Meier plot was used to analyze patient survival data. In all comparisons, \( P < .05 \) was considered statistically significant. At least 3 biological replicates were performed. Graphical abstract was created with Biorender.com.

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Ethical Approval and Consent to Participate
The use of clinical specimens was approved by the institutional review board of the University of Hong Kong and the Hong Kong Hospital Authority. Animal experiments were performed with the approval of the Committee on the Use of Live Animals in Teaching and Research of the University of Hong Kong. All experimental procedures strictly followed the animals (Control of Experiments) ordinance of Hong Kong.

Data Availability Statement
The authors confirmed that the data supporting the findings of the study are available in the article and Supplementary materials. Supplementary tables are provided as Microsoft Excel tables (Redmond, WA). The genome-wide transcriptomic sequencing and chromatin isolation by RNA purification sequencing are available in the Sequence Read Archive (SRA) database: PRJNA786081. This article does not generate original code. Any additional information required to re-analyze the data reported in this article is available from the lead contact upon request.

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