Recognition of RNA N⁶-methyladenosine by IGF2BP proteins enhances mRNA stability and translation

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N⁶-methyladenosine (m⁶A) is the most prevalent modification in eukaryotic messenger RNAs (mRNAs) and is interpreted by its readers, such as YTH domain-containing proteins, to regulate mRNA fate. Here, we report the insulin-like growth factor 2 mRNA-binding proteins (IGF2BPs; including IGF2BP1/2/3) as a distinct family of m⁶A readers that target thousands of mRNA transcripts through recognizing the consensus GG(m⁶A)C sequence. In contrast to the mRNA-decay-promoting function of YTH domain-containing family protein 2, IGF2BPs promote the stability and storage of their target mRNAs (for example, MYC) in an m⁶A-dependent manner under normal and stress conditions and therefore affect gene expression output. Moreover, the K homology domains of IGF2BPs are required for their recognition of m⁶A and are critical for their oncogenic functions. Thus, our work reveals a different facet of the m⁶A-reading process that promotes mRNA stability and translation, and highlights the functional importance of IGF2BPs as m⁶A readers in post-transcriptional gene regulation and cancer biology.

As the most abundant messenger RNA (mRNA) modification, N⁶-methyladenosine (m⁶A) modification is reversible and plays critical roles in multiple fundamental biological processes (for example, cell differentiation, tissue development and tumorigenesis)⁴–⁵⁹. High-throughput sequencing revealed that m⁶A is especially enriched in the 3′ untranslated regions (UTRs) and near the stop codons of mRNAs with a consensus sequence of RRACH (R corresponds to G or A; H corresponds to A, C or U)⁴–⁵. The biological importance of m⁶A modification relies on m⁶A-binding proteins (that is, readers). Thus, it is crucial to identify and characterize m⁶A readers that directly guide distinct bioprocesses. A group of YT521-B homology (YTH) domain-containing proteins (YTHDFs) have been identified as m⁶A readers that control mRNA fate by regulating pre-mRNA splicing, facilitating translation or promoting mRNA decay⁶–⁹. In the YTHDF2-mediated decay pathway⁹, mRNA levels are expected to increase when m⁶A abundance is reduced. However, our recent data showed that a large portion of mRNAs with reduced m⁶A abundance tended to be downregulated due to decreased RNA stability⁹, suggesting the presence of alternative mechanisms to stabilize m⁶A-modified mRNAs. Here, we report the insulin-like growth factor 2 (IGF2) mRNA-binding proteins 1, 2 and 3 (IGF2BP1/2/3) as a new family of m⁶A readers that guard m⁶A-modified mRNAs from decay. IGF2BPs, a conserved family of single-stranded RNA-binding proteins (RBPs)⁴⁰, are composed of six canonical RNA-binding domains, including two RNA recognition motif (RRM) domains and four K homology (KH) domains⁴⁰. Besides IGF2, a few other well-known mRNAs (for example, MYC, ACTIN and LIN28B) have been reported as targets of IGF2BPs⁴⁰–⁴⁹. However, the exact molecular mechanisms by which IGF2BPs recognize and regulate the expression of their targets remain elusive. Here, we provide compelling evidence showing that IGF2BPs preferentially recognize m⁶A-modified mRNAs and promote the stability (and probably also translation) of thousands of potential mRNA targets (including MYC) in an m⁶A-dependent manner, thereby globally affecting gene expression output. Furthermore, as m⁶A readers, IGF2BPs play oncogenic roles in cancer cells, probably by stabilizing methylated mRNAs of oncogenic targets (for example, MYC).

Results

Identification of IGF2BPs as m⁶A-binding proteins. To identify m⁶A-binding proteins, we applied two independent methods:
Selective binding of IGF2BPs to m^6^A-modified RNAs. a, Identification of m^6^A-specific binding proteins by RNA affinity chromatography using single-stranded RNA probes with methylated (red) or unmethylated (green) adenosine. Consensus sequence shown in bold. Silver staining (left) and western blotting (right) showed selective pulldown of ~68 kDa IGF2BP proteins from the HEK293T nuclear extract. Western blot images are representative of three independent experiments. b, Enrichment of the m^6^A consensus sequence ‘GGAC’ in the binding sites of RBPs. The three IGF2BP paralogues are shown in red, whereas the YTHDFs were shown in orange. Enrichment is shown as a percentage of IGF2BP high-confidence target genes that contain m^6^A peaks. The m^6^A-seq data were reported in ref.3. c, Overlap of IGF2BP target genes identified by RIP–seq and published PAR-CLIP in HEK293T cells. RIP–seq was performed once. d, Enrichment (bottom) of IGF2BP-binding peaks within different gene regions. Enrichment was determined by the proportion of IGF2BP-binding peaks within each gene region. Enrichment was determined by the proportion of IGF2BP-binding peaks within each gene region. e, Pie charts showing the numbers and percentages of IGF2BP high-confidence target genes that contain m^6^A peaks. The m^6^A-seq data were reported in ref.3.
(1) using methylated single-stranded RNA bait (ss-m^6A), with the consensus sequence GG(m^6A)CU or unmethylated control RNA (ss-A) for RNA pull-down (Fig. 1a and Supplementary Fig. 1a), followed by mass spectrometry analysis; and (2) developing a computational pipeline for screening potential m^6A-binding proteins by using published RBP crosslinking and immunoprecipitation followed by high-throughput sequencing (CLIP–seq) data sets and known m^6A modification sites. All three IGF2BP proteins were identified by mass spectrum and were confirmed to selectively bind to the methylated bait (ss-m^6A) with a 3–4-fold higher affinity than the unmethylated control (ss-A) (Fig. 1a and Supplementary Fig. 1b), and the binding seems to be independent of RNA secondary structure (Supplementary Fig. 1c). Similar to endogenous proteins, recombinant IGF2BP proteins purified from human cells also preferentially bound to methylated RNA probe over the unmethylated one (Supplementary Fig. 1d,e). In addition to ss-m^6A, IGF2BP2 also preferentially bound to methylated hairpin RNA (hp-m^6A) probes over control (hp-A) probes (Supplementary Fig. 1f,g). Meanwhile, our computational pipeline revealed that all three IGF2BP were among the top 15 of 112 RBPs in terms of both the significance (Fig. 1b) and the frequency (Supplementary Fig. 1h) of m^6A motifs enriched in their RNA-binding sites. Thus, IGF2BP proteins are potential m^6A-binding proteins.

We then overexpressed FLAG-tagged IGF2BP2 in HEK293T cells and immunoprecipitated ribonucleoprotein complexes to evaluate the in cellulo binding. A significant enrichment of m^6A modifications in FLAG-IGF2BP2-bound RNA was observed (Fig. 1c and Supplementary Fig. 1i), similar to that in RNA immunoprecipitates (RIPs) of endogenous IGF2BP2 (Supplementary Fig. 1j). Sequencing purified RNA from FLAG-RIP samples identified >5,000 genes from each RIP sample; among them, >50% overlapped with published photoactivatable ribonucleoside-enhanced (PAR)-CLIP–seq targets (P<5×10^-14, Fisher's exact test; Fig. 1d). The 3,747, 3,211 and 3,914 transcripts identified by both RIP and PAR-CLIP methods can be considered as high-confidence targets of IGF2BP1, IGF2BP2 and IGF2BP3, respectively (Fig. 1d and Supplementary Table 1). The three IGF2BP proteins shared 2,149 (55–70%) high-confidence RNA targets (Fig. 1e). All three IGF2BP preferably bind to the ‘UGGAC’ consensus sequence containing the ‘GGAC’ m^6A core motif (Fig. 1f), and >80% of the high-confidence targets contain at least one m^6A peak as detected by m^6A-seq (Fig. 1g). Moreover, most of the IGF2BP-binding sites (92%) are located in protein-coding transcripts (that is, mRNAs) and are highly enriched near stop codons and in 3'UTRs, coinciding with the m^6A distribution (Fig. 1b–j). In addition, we analysed ENCODE (Encyclopedia of DNA Elements) enhanced CLIP (eCLIP)–seq data in HepG2 cells and human embryonic stem cells (hESCs), and found that the ‘UGGAC’ motif was also enriched in the targets of IGF2BP in both cell types (Supplementary Fig. 1k,l).

Methyltransferase-like protein 3 (METTL3) and METTL14 are two critical components of the methyltransferase complex, which catalyses methylation at N^6-adenosine. We performed CLIP of FLAG-tagged IGF2BP2 and IGF2BP3 in HEK293T cells with or without METTL14 knockdown. Four representative high-confidence targets, including MYC, FSCN1, TK1 and MARCKSL1, exhibit strong binding with IGF2BP2 around their m^6A motifs in control cells (Fig. 1k). Such binding was largely impaired upon METTL14 knockdown (Fig. 1k), suggesting that there is a requirement of cellular m^6A modification for the binding. Taken together, these data demonstrated the role of IGF2BP2 as m^6A-binding proteins in vitro and in cellulo.

Silencing of IGF2BP2 globally downregulates target gene expression. We next conducted RNA sequencing (RNA-seq) in individual IGF2BP2–knockdown and control HepG2 cells (Supplementary Fig. 2a). The global transcripts were grouped into non-targets, CLIP targets and CLIP+RIP targets according to their binding by IGF2BP2 in HEK293T cells (see Fig. 1d), considering the availability of CLIP data for all three IGF2BP2 in this cell line. Knockdown of individual IGF2BP2 globally and preferentially inhibited the expression of CLIP targets and especially of CLIP+RIP targets, with much more CLIP+RIP targets being downregulated than upregulated (Fig. 2a,b). Gene-set enrichment analysis (GSEA) also showed that genes highly expressed in the control groups were enriched with the IGF2BP2 CLIP+RIP targets (false discovery rate (FDR)<0.05; Supplementary Fig. 2b). Functional annotation indicated that target genes with reduced expression were enriched in DNA replication, cell cycle, proliferation and cancer-related biological processes and pathways (Supplementary Fig. 2c). In addition, an enrichment of cell cycle genes and MYC target genes was observed in controls versus short hairpin IGF2BP2 (shIGF2BP2; Supplementary Fig. 2d). Downregulation of representative targets was confirmed by quantitative PCR (qPCR; Fig. 2e).

To determine whether the expression of IGF2BP targets is also affected by the level of cellular m^6A, we performed m^6A-seq and RNA-seq in control or METTL14-knockdown HepG2 cells. Upon METTL14 knockdown, 1,516 genes showed reduced m^6A modifications (that is, m^6A-Hypo genes; fold change (FC)<0.667). Among these genes, 418 had reduced (that is, m^6A-Hypo-down genes; FC<0.8) mRNA levels, whereas 335 had increased levels (that is, m^6A-Hypu-up genes; FC>1.2) (Fig. 2d). As expected, IGF2BP high-confidence targets showed a global and significant reduction in mRNA level upon METTL14 knockdown (Fig. 2e). The expression of individual targets, including FSCN1, TK1, MARCKSL1 and MYC, was confirmed by qPCR to be significantly downregulated upon METTL3 or METTL14 knockdown (Fig. 2f). Similar changes were observed for FSCN1 and MYC, two shared targets of IGF2BP2 and YTHDF2, in IGF2BP2/YTHDF2 double-knockdown cells (Supplementary Fig. 2e). Conversely, the m^6A-Hypo-down genes were globally downregulated upon knockdown of IGF2BP2, as compared to their non-target counterparts (Supplementary Fig. 2f). The correlated regulation of gene expression by IGF2BP2 and METTL14 indicates that IGF2BP2 are responsible for the expression output of m^6A-regulated genes.

Regulation of mRNA stability by IGF2BP2s. mRNA stability profiling in HepG2 cells revealed that high-confidence targets of IGF2BP1–3 tend to have longer half-lives (P<0.001) than their non-target counterparts (Fig. 3a), whereas YTHDF2 targets tend to have shorter half-lives (Supplementary Fig. 3a). Similar results were observed in HeLa cells (Supplementary Fig. 3a) and human cord blood CD34^+ cells (Supplementary Fig. 3e). As IGF2BP3 silencing affected target gene expression most potently (Fig. 2a,b), we performed mRNA stability profiling in IGF2BP3-knockdown cells. The median half-life of IGF2BP3 high-confidence targets as well as CLIP targets was significantly reduced in IGF2BP3-depleted cells to approximately 50% of that in control cells (Fig. 3b,c and Supplementary Fig. 3b,c). Accelerated mRNA decay of MYC, FSCN1, TK1 and MARCKSL1 upon knockdown of each IGF2BP was confirmed in HepG2 cells (Fig. 3d and Supplementary Fig. 3d) and human cord blood CD34^+ cells (Supplementary Fig. 3e). Furthermore, the stability of MYC, FSCN1, TK1 and MARCKSL1 mRNAs was also reduced when m^6A writers were silenced (Fig. 3e and Supplementary Fig. 3f).

To identify co-factors of IGF2BP2 that may enhance stability of m^6A targets, we pulled down the IGF2BP2 complexes and conducted mass spectrometry analysis. Notably, ELAV-like RNA-binding protein 1 (ELAVL1; also known as HuR), matriosome 3 (MATR3) and poly(A)-binding protein cytoplasmic 1 (PABPC1), three known mRNA stabilizers were identified. Western blotting confirmed their binding to ectopically expressed IGF2BP2 in HEK293T cells (Fig. 3f), consistent with previous reports. Moreover, co-localization of HuR and IGF2BP2 was observed in cytoplasmic granules...
Fig. 2 | IGF2BPs regulate transcriptome-wide mRNA levels. a, Volcano plots displaying enrichment of dysregulated target genes in IGF2BP-knockdown (shIGF2BP) versus control (shNS) HepG2 cells. The numbers of significantly downregulated (log₂ FC < -1, P < 0.05, two-tailed Student’s t-test) or upregulated (log₂ FC > 1, P < 0.05, two-tailed Student’s t-test) genes in the CLIP target group and CLIP + RIP target group are shown. Vertical dashed lines indicate cut-off of log₂FC (1 or -1), whereas the horizontal dashed lines indicate cut-off of P value (0.05). FC, fold change. b, Cumulative frequency of mRNA log₂ FC in non-target, CLIP target and CLIP + RIP target genes upon IGF2BP silencing. P values were calculated using two-sided Wilcoxon and Mann–Whitney test. c, Relative changes in FSCN1, TK1, MARCKSL1 and MYC mRNA levels upon IGF2BP silencing. Results from two shRNAs for each IGF2BP are shown. Values are the mean ± s.d. of n = 3 independent experiments. Two-tailed Student’s t-tests were used (**P < 0.01; ***P < 0.001). d, Distribution of genes with a significant change in both the m6A level and the gene expression level in METTL14-knockdown HepG2 cells compared with control cells. e, Cumulative frequency of mRNA log₂ FC showing global reduction of IGF2BP high-confidence target genes in shMETTL14 versus shNS cells. Values are the mean ± s.d. of n = 3 independent experiments. Two-tailed Student’s t-tests were used (**P < 0.01; ***P < 0.001). Source data for c and f are in Supplementary Table 3.
Polysome profiling showed that FLAG-IGF2BP1 and FLAG-IGF2BP2 were present in most of the sucrose gradient fractions, whereas FLAG-IGF2BP3 was accumulated in 60S and 40S (Supplementary Fig. 4a). A similar distribution of endogenous IGF2BP1/2/3 proteins was observed in HepG2 cells (Supplementary Fig. 4b). Interestingly, HuR was detected along with each FLAG-tagged IGF2BP (Supplementary Fig. 4a). Moreover, knockdown of IGF2BP1 in HEK293T cells significantly reduced MYC mRNA in the translating pool (Supplementary Fig. 4c, fractions 13–18), suggesting a role of IGF2BPs in the active translation of their target genes. In agreement with the localization of IGF2BPs to stress granules during heat shock, IGF2BP2 shifted to non-ribosome fractions, and was found to gradually return to ribosome fractions during recovery from heat shock (Supplementary Fig. 4d).
**shMETTL14**

**shNS**

**Fig. 4 | IGF2BPs regulate MYC expression through binding to methylated CRD.** a. Distribution of m^6^A peaks across MYC mRNA transcript. The CRD region is highlighted in yellow. m^6^A-seq was repeated twice, whereas RIP-seq was performed once. IP, immunoprecipitation. b. RIP-qPCR showing the association of MYC CRD with FLAG-tagged IGF2BP2 in HEK293T cells. c. Enrichment of m^6^A modification in MYC CRD as detected by a gene-specific m^6^A qPCR assay. d. RIP-qPCR showing the binding of METTL3 and METTL14 to the MYC CRD. e. RNA pulldown of endogenous IGF2BP proteins from HEK293T nuclear extract using synthetic CRD RNA fragments, CRD1 and CRD2, with (m^6^A) or without (A) m^6^A modifications. Images are representative of three independent experiments. f. Relative luciferase activity (that is, protein level) and Fluc mRNA level of wild-type (CRD-WT) or mutated (CRD-mut) CRD reporters in HEK293T cells with ectopically expressed IGF2BP1, IGF2BP2 or IGF2BP3. g. RIP-qPCR detecting the in vivo binding of FLAG-IGF2BPs to the transcripts of CRD-WT or CRD-mut luciferase reporter in HEK293T cells. h. Relative luciferase activity of CRD-WT or CRD-mut in HeLa cells with or without stable knockdown of IGF2BPs. i. Relative luciferase activity of CRD-WT or CRD-mut in HeLa cells with ectopic expression of IGF2BP2. For all luciferase assays, the Fluc/Rluc ratio (representing luciferase activity) of CRD-WT with empty vector or shNS was used for normalization. Values are the mean ± s.d. of n = 3 independent experiments, and two-tailed Student’s t-tests were used in b–d and f–j. (**P < 0.01; ***P < 0.001). Unprocessed scans of western blot analysis are available in Supplementary Fig. 8. Source data for b–d and f–j are in Supplementary Table 3.
IGF2BPs regulate MYC expression in an m6A-dependent manner.

To determine whether IGF2BP-mediated gene regulation is m6A dependent, we chose MYC, a well-known target of IGF2BP1 (refs [92][93]), for a systematic study. A approximately 250 nucleotide (nt) cis-acting element called coding region instability determinant (CRD) resides in the 3′ terminus of the MYC coding region and has been proven to be critical for IGF2BP1 binding [94]. As shown in Fig. 4a, m6A modifications are accumulated across MYC transcript, and the m6A peaks coincide well with IGF2BP-binding sites. Notably, the CRD-containing region has a high abundance of m6A modifications that decreases remarkably upon METTL14 knockdown (Fig. 4a). By conducting RIP and gene-specific m6A assays, we confirmed in cellulo IGF2BP binding (Fig. 4b) and Supplementary Fig. 5a) and m6A modification (Fig. 4c), as well as METTL3 and METTL14 binding (Fig. 4d), in CRD. Moreover, the m6A modifications in the consensus sites of the synthetic CRD RNA oligos greatly facilitated their binding by endogenous IGF2BPs (Fig. 4e).

We next inserted the 249-nt wild-type or mutant CRD sequence into a firefly luciferase (Fluc) reporter (Supplementary Fig. 5b). Mutations in the m6A sites of CRD (CRD1 and CRD2 RNA oligos) dramatically abrogated the association with IGF2BP proteins in vitro (Supplementary Fig. 5c). As expected, ectopic IGF2BPs induced a significant increase in Fluc activity of the wild-type reporter in a dose-dependent manner (Supplementary Fig. 5d). Such increases were largely impaired by mutations in the m6A consensus sites (Fig. 4f, left, and Supplementary Fig. 5b). Consistently, the relative Fluc mRNA level of reporters with wild-type CRD, but not those with mutant CRD, was increased by IGF2BP overexpression (Fig. 4f, right). RIP-qPCR demonstrated a strong binding of IGF2BPs with wild-type CRD reporters and a much less or no binding with mutant CRD in cellulo (Fig. 4g). Conversely, knockdown of individual IGF2BPs, similar to knockdown of METTL14, caused inhibited Fluc activity, which also relies on the presence of wild-type m6A motifs within CRD (Fig. 4h,i). Noticeably, IGF2BP-mediated increase of luciferase activity could be partially or completely blocked by METTL14 knockdown (Fig. 4j). Taken together, our data demonstrate that m6A modifications in CRD are required for the binding of IGF2BPs to MYC and for IGF2BP-mediated regulation of MYC expression.

Recognition of m6A by the KH domains of IGF2BPs. RRM and KH, the RNA-binding domains of IGF2BPs, are different from the YTH domain, the known m6A-binding domain [2][9][13][14]. We constructed IGF2BP mutants with truncation of the two RRM domains, or with mutations of GxxG to GEGG in the KH domains as reported [10]. We showed that mutations in the KH domains (KH1–4), but not RRM truncation, completely abolished the interaction between IGF2BPs and ss-m6A probes (Fig. 5b and Supplementary Fig. 5e). Interaction of IGF2BPs with ss-m6A probes was only partially reduced by KH1–2 di-domain mutation, but was almost completely abolished by KH3–4 mutation (Fig. 5b), indicating that the KH3–4 di-domain is dispensable for m6A recognition and binding, whereas KH1–2 might play an accessory role. KH domain mutations, including KH3–4 and KH1–4, but less likely KH1–2

Fig. 5 | The KH domains of IGF2BPs are critical for m6A recognition and binding. a, Schematic structures showing RNA-binding domains within IGF2BP proteins and a summary of IGF2BP variants used in this study. Blue boxes are RRM domains, red boxes are wild-type KH domains with GxxG core and grey boxes are inactive KH domains with GxxG to GEGG conversions. b, RNA pulldown followed by western blotting showed in vitro binding of single-stranded RNA (ssRNA) baits with wild-type or KH domain-mutated IGF2BPs, representative of three independent experiments. c, In vitro binding of CRD RNA probes with wild-type or KH3–4-mutated IGF2BPs, representative of three independent experiments. d, The association of wild-type and KH3–4-mutated IGF2BPs with MYC CRD in HEK293T cells, as assessed by RIP-qPCR. e, Relative luciferase activity of CRD reporters in HEK293T cells with forced expression of wild-type or mutated IGF2BP2 variants. f, Changes in MYC mRNA levels in HeLa cells with empty vector or forced expression of wild-type or KH3–4-mutated IGF2BPs 1 h post-heat shock (HS). Ctrl, control. Values are the mean ± s.d. of n = 3 independent experiments, and two-tailed Student’s t-tests were used in d–f (**P < 0.01; ***P < 0.001). Unprocessed scans of western blot analysis are available in Supplementary Fig. 8. Source data for d–f are in Supplementary Table 3.
Indeed, knocking down of each individual IGF2BPs in HeLa (cervical cancer) and HepG2 (liver cancer) cells significantly repressed MYC expression (Fig. 6a) and inhibited cancer cell proliferation, colony formation ability and cell migration/invasion (Fig. 6b–d and Supplementary Fig. 6c–e), which mimics the effect of MYC silencing (Supplementary Fig. 6f–j).

By utilizing the clustered regularly interspaced short palindromic repeats (CRISPR)–CRISPR-associated protein 9 (Cas9) system, we generated IGF2BP-knockout (KO) cells (Fig. 7a) and performed rescue experiments. The decreased proliferation and colony formation ability of KO cells could be reversed by forced expression of wild-type IGF2BP, but not the KH3–4 mutants (Fig. 7b,c), suggesting that the oncogenic function of IGF2BPs relies on their role as m6A readers. Ectopic expression of MYC also restored the proliferative ability of IGF2BP-KO cells (Fig. 7d), which further supports MYC as a critical target of IGF2BPs.

**Discussion**

The characterization of the YTHDFs as direct m6A readers has provided profound insights into our understanding of the effects of m6A modification on genetic information flow. Our findings add a distinct RBP family, IGF2BPs, into the catalogue of m6A readers, and reveal their roles in mRNA stabilization and translation (Fig. 7e). Notably, >3,000 mRNA transcripts were identified as targets of each individual IGF2BP protein, whereas >5,000 mRNAs were being targeted by at least one protein and >2,000 mRNAs were being co-targeted by all three IGF2BPs. The binding sites of IGF2BPs are enriched with m6A motif ‘GGAC’. Given that only around 7,000 mRNA transcripts are m6A modified in individual mammalian cells and that >80% of IGF2BP targets have at least one m6A peak (Fig. 1g), IGF2BPs probably have a broad impact on m6A-associated gene regulation. Accordingly, reduction of cellular...
**Fig. 7 | IGF2BPs are oncogenic m^6^A readers.** a, CRISPR–Cas9-mediated KO of IGF2BPs and the subsequent depletion of MYC in HepG2 cells, as detected by western blotting. Images are representative of three independent experiments. b, Effect of wild-type or KH3–4-mutated IGF2BPs on restoring cell proliferation in IGF2BP-KO cells. Data shown represent the mean value of viable cell numbers of two independent experiments. c, Colony formation assay using wild-type or IGF2BP-KO (sgIGF2BP) HepG2 cells. Representative images of crystal violet staining of cells are shown beside the histograms of colony numbers. Colonies were counted from three replicate wells, and two independent experiments were performed. The colony number of each experiment represents the average count of three replicate wells. d, MTT assays displaying the effect of MYC on restoring cell proliferation in IGF2BP-KO cells. Values are the mean ± s.d. of n = 3 independent experiments. Two-tailed student t-test were used (**P < 0.01; ***P < 0.001). e, Working model of IGF2BP-mediated regulation of m^6^A-modified mRNAs. mRNAs were methylated de novo by the methyltransferase complex, which is composed of METTL3, METTL14 and a regulatory subunit WTAP. The naive mRNA with m^6^A modifications were preferentially recognized by IGF2BP proteins. By recruiting mRNA stabilizers, such as HuR and MATR3, IGF2BPs protect target mRNAs from degradation in the P-body while facilitating translation after being exported to the cytoplasm. Under stress conditions such as heat shock, IGF2BP-containing messenger ribonucleoproteins are translocated to stress granules for the storage of their mRNA targets. ALKBH5, alkB homolog 5 RNA demethylase; FTO, alpha-ketoglutarate dependent dioxygenase FTO. Dashed arrows indicate translocation to or from stress granules. Unprocessed scans of western blot analysis are available in Supplementary Fig. 8. Source data for b–d are in Supplementary Table 3.
m’A levels upon METTL14 knockdown impairs the in cellulo binding of FLAG-IGF2BP2/3 to their RNA targets. Furthermore, we demonstrated that IGF2BPs bind directly to MYC CRD and promote MYC expression in an m’A-dependent manner.

The direct binding of IGF2BPs to m’A RNAs through their KH domains was demonstrated both in vitro and in cellulo. The finding from all significant YTHDF2-binding sites (Supplementary Fig. 7c,d), suggesting that a significantly lower guanine–cytosine (GC) content than tidies alone showed poor selectivity for m’A RNA compared with full-length proteins (data not shown), which is consistent with previous reports that post-translational modifications in the KH-3–4 flanking domains may be important for IGF2BP selectivity15,26. It is also possible that KH domains are different from the well-defined YTH domains and may not possess discrete pocket for m’A recognition, and that other mechanisms (e.g., reducing solvation penalty, as previously proposed14) may be involved in the recognition of m’A by KH domains. Future structural studies are warranted to understand how specific KH domains bind to m’A-modified RNAs. It will also be interesting to investigate other KH domain proteins as potential m’A-binding readers.

Our RNA stability profiling revealed that IGF2BPs stabilize target RNAs. The opposite role of IGF2BPs versus YTHDF2 imposes a significantly lower guanine–cytosine (GC) content than 3’UTRs and near stop codons. In fact, YTHDF2-binding sites show a lower density in the 3’UTRs than in the coding regions2, which is distinct from IGF2BP-binding sites (Fig. 1h). Analysis of the ENCODE PAR-CLIP data revealed only a very small proportion (0.85–1.20%) of IGF2BP-binding sites (Fig. 1h). Further analysis found from all significant YTHDF2-binding sites (Supplementary Fig. 7c,d), suggesting that the local nucleotide composition may also contribute to the binding preference of different readers. Together, these data indicate that IGF2BPs and YTHDF2 have a distinct pattern in target recognition and regulation.

The mRNA stabilizing function of IGF2BPs was also supported by its co-factors, HuR and MATR3. In particular, HuR was previously identified as an indirect m’A-binding protein, which increased the stability of bound RNA and blocked microRNA targeting22,25,27. Interestingly, we showed here that HuR was co-localized with IGF2BPs in P-bodies, which are locations for mRNA fate decision34–40. Our findings suggest that HuR could be recruited by IGF2BPs to protect m’A-containing mRNAs from degradation and facilitate their translation (Fig. 7e). We found that IGF2BPs co-localize with stress granules and shuttle between ribosome and non-ribosome fractions during heat shock and recovery, suggesting a role of IGF2BPs in mRNA translation in stress response. Collectively, IGF2BPs can promote stability by inhibiting mRNA degradation or enhancing mRNA storage under stress, and facilitate their translation (Fig. 7e). We recently reported that METTL14 promotes the stability and translation of MYC mRNA and plays an essential oncogenic role in leukaemia44, which is probably also attributed to IGF2BP-mediated, m’A-dependent regulation of MYC expression.

Dysregulation of IGF2BPs could result in abnormal accumulation of oncogenic products such as MYC, and therefore support the malignant state of cancer cells. Consistent with the frequent amplification of IGF2BP genes in various types of cancers, our finding that IGF2BPs exhibit oncogenic roles as m’A readers demonstrates the functional importance of IGF2BPs and their associated m’A reading processes in tumorigenesis, and highlights the therapeutic potential of targeting IGF2BPs in cancers.

Methods

Methods, including statements of data availability and any associated accession codes and references, are available at https://doi.org/10.1038/s41556-018-0045-z.

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**Author contributions**

H.H., H.Weng and J.C. conceived and designed the entire project. H.H., H.Weng, C.He, J.Y. and J.C. designed and supervised the research. H.H., H.Weng, X.Q., H.S., H.Wu, B.S.Z., A.M., C.Liu, C.L.Y., J.R.S., X.D., M.S., C.Li, S.N., C.Hu, K.F. and J.C. performed the experiments and/or data analyses. H.H., H.Weng, WS., L.D. and J.Y. performed the genome-wide or transcriptome-wide data analyses. Y.-C.H., S.H., K.D.G., X.I., M.W., L.Q., J.-L.G., C.He, J.Y. and J.C. contributed reagents/analytic tools and/or grant support. H.H., H.Weng, WS., H.S., B.S.Z., A.M., S.N., C.He, J.Y. and J.C. wrote and revised the paper. All authors discussed the results and commented on the manuscript.

**Competing interests**

C.He is a scientific founder of Accent Therapeutics, Inc.

**Additional information**

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Methods
Plasmids and short interfering RNAs. pcDNA3-based vectors encoding wild-type and KH domain mutant FLAG-tagged chicken ZBP1 (refer to as IGF2BP2), human IGF2BP2 and IGF2BP3 were kindly provided by Dr. Hüttelmaier (Martin Luther University, Halle-Wittenberg, Germany). The human IGF2BP2 (IGF2BP2-A) and IGF2BP3 (IGF2BP3-A) were produced by PCR using the Q5 Site Directed Mutagenesis Kit (NEB) with forward primer 5′-GAAAGGTGACCTCCCT-3′ and reverse primer 5′-GAATTCTGGTCGTGCTGCC-3′. The plasmid encoding human MYC (pcDNA3-HA-HA-humanCMYC) was obtained from Addgene. The RNAi Consortium (TRC) lentiviral vectors encoding shRNAs against IGF2BP1 (TRCN0000075149 and TRCN0000075152), IGF2BP2 (TRCN0000149002 and TRCN0000148565), IGF2BP3 (TRCN0000074677 and TRCN0000074673), METTL3 (TRCN0000034715), METTL14 (TRCN0000159393) and their non-specified construct (shRS, RH56848) were purchased from GE Dharmacron, whereas the packing vectors, pMD2.G, pMDLg/pRRE and pRSV-Rev, were obtained from Addgene. The Myc short interfering RNA (siRNA) was purchased from Santa Cruz Biotechnology, whereas the HuR siRNA was from GE Dharmacron.

Cell culture and transfection. The human hepatocellular carcinoma cell line HepG2 (ATCC HB-8085) was maintained in EMEM medium (American Type Culture Collection (ATCC)) supplemented with 10% FBS (Invitrogen), 2 mM glutamine, 100 U/ml penicillin–streptomycin. All cell lines were purchased from the ATCC and were not authenticated by ourselves. All cell lines were routinely tested for mycoplasma contamination. The cell lines were not a part of a database of commonly misidentified cell lines maintained by the International Cell Line Authentication Committee (ICLAC). For heat-shock treatment, HeLa cells were incubated at 42 °C for 1 h. Plasmids and siRNAs were transfected into cells with Lipofectamine 2000 (Invitrogen) according to the manufacturer’s instructions.

Isolation and lentiviral infection of CD34+ cells. Cord blood of healthy donors was obtained from Cincinnati Children’s Hospital Medical Center (CCHMC, Ohio, USA) and subjected to isolation of mononuclear cells using Ficoll–Paque PLUS (GE Healthcare Life Sciences). Human CD34+ haematopoietic stem/progenitor cells were then purified from mononuclear cells by using human CD34 MicroBead Kit (Miltenyi Biotec). The CD34+ cells were cultured in StemSpan SFEM medium (StemCell Technologies) supplemented with 1% Lipid Mixture 1 (L0288, Sigma–Aldrich), 2 mmol per litre l-glutamine, 1% penicillin–streptomycin, 100 ng/ml per ml stem cell factor (SCF), and 2 ng/ml IL-3. Cells were infected with concentrated lentiviral particles through two rounds of ‘spinoculation’.

RNA affinity chromatography and nLC-EISI-MS/MS. Biotin-labelled RNA oligonucleotides containing adenosine or m^6^A were synthesized by GE Dharmacron. The hpRNA baits were denatured at 99 °C for 10 min and slowly cooled down to room temperature to allow the formation of stem-loop structure. The hpRNA baits were denatured at 99 °C for 10 min and slowly cooled down to room temperature. For each reaction, 1 μl RNA probes (4 nM final concentration) and 1 μl protein (10% of the concentration gradient indicated in Supplementary Fig. 1d) were incubated in 8 μl binding buffer (10 mM Tris–HCl pH 7.5, 50 mM KCl, 1 mM EDTA, 0.05% Triton X-100, 5% glycerol, 1 mM dithiothreitol and 40 μM per ml RNasin) on ice for 30 min. The RNA–protein mixtures were separated in 5% native polyacrylamide gels (in 0.5xTris-borate-EDTA buffer) at 4 °C for 60 min at 13 V per cm and was visualized by Odyssey Imaging System (LI-COR Biosciences) and quantified by ImageMaster TotalLab (GE Healthcare). The dissociation constant (K_d) was calculated with non-linear curve fitting (function one-site-specific binding) using GraphPad Prism with y = B_max⋅x/(K_d+x), where y is the ratio of [RNA–protein]/([free RNA] + [RNA–protein]), x is the input protein concentration, and B_max is set to 1.

m^6^A dot blot. The m^6^A dot blot assay was conducted as previously described. Briefly, the indicated amount of total cellular RNA or synthesized RNA oligonucleotide was denatured in 3-fold volume of RNA incubation buffer (65.7% formamide, 7.77% formaldehyde and 1.33xMOPS) at 65 °C for 5 min, followed by chilling on ice and mixing with 1-fold volume of 3xSSC. RNA samples were applied to Amersham Hybond-N+ membrane (GE Healthcare) with a Bio-Dot Apparatus (BioRad). After UV crosslinking, the membrane was stained with 0.02% methylene blue in 0.3M sodium acetate. The membrane was then washed with 1x PBST buffer, blocked with 5% fat-free milk in PBST, and incubated with anti-m^6^A antibody (ab206033, 1:1000, Synaptic Systems) overnight at 4 °C. After washing with horseradish peroxidase-conjugated anti-rabbit IgG secondary antibody (Santa Cruz Biotechnology), the membrane was visualized using Amersham ECL Prime Western Blotting Detection Reagent (GE Healthcare).

Western blotting. Cells were lysed using 1x SDS buffer and sonicated. Equal amounts of proteins were loaded and separated by 10% SDS–PAGE, transferred to polyvinylidene fluoride membranes, and detected by immunoblotting with the Pierce ECL Western Blotting Substrate (Thermo Fisher Scientific) or Amersham ECL Prime Western Blotting Detection Reagent. Antibodies used for western blotting were as follows unless otherwise specified: IGF2BP1 (IMPI, clone D33A2, no. 8482), IGF2BP2 (IMP2, clone D2R8, no. 14672), MYC (clone D3H9, no. 13987), HuEl (ELAVL1, clone D9W7E, no. 12582) were from Cell Signaling Technology (CST), IGF2BP3 (IMP3, A303–426A) and MATR3 (A300–591A-T) were from Bethyl Laboratories, and FLAG (M2, F3165) was from Sigma–Aldrich. GAPDH (sc-47724, Santa Cruz) was used as a loading control.

Prediction of m^6^A-binding proteins. We developed a computational pipeline to discover the potential m^6^A-binding proteins from ENCODE and published RBP–CLIP data sets generated by various CLIP methods, including high-throughput sequencing of RNA isolated by CLIP (HITS–CLIP), PAR–CLIP, eCLIP, individual nucleotide resolution CLIP (iCLIP) and infrared–CLIP (irCLIP). The CLIP–seq peaks of each RBP were intersected with known m^6^A sites” to calculate the ratio between the number of peaks containing m^6^A and the total number of peaks (m^6^A-containing peak number/total peak number). These peaks were imported into HOMER software for de novo motif identification. The potential m^6^A-binding proteins should meet the following requirements: (1) the ratio (m^6^A-containing peak number/total peak number) should be >10%; and (2) for the identified motifs of each RBP, the P value should be <1x10^-10.

LC-MS/MS. The recombinant proteins were purified using an Eksigent nanoLC ultra 2Dflow system attached to a TripleTOF 5600 plus (Scienx) for nano-liquid chromatography–electrospray ionization mass spectrometry (nLC-EISI-MS/MS) analyses.

Expression data and purification. FLAG-tagged IGF2BP1, IGF2BP2 and IGF2BP3 were expressed in HEK293T cells. For each protein, 1.5 cm^2 of cells were prepared and lysed in 6 ml lysis buffer (50 mM Tris–HCl pH 7.5, 300 mM KCl, 0.5% NP-40, 5% glycerol, 2 μg/ml aprotinin, 1 μg/ml dithiothreitol) at 4 °C for 1 h and sonicated (5 s on, 25 s off, for 24 cycles). The lysate was then cleared by centrifugation at 4 °C. The proteins were affinity purified using 40 μl anti-FLAG M2 resin (Sigma–Aldrich) at 4 °C for 4 h. After extensive wash buffer with 300 mM NaCl, 0.5% NP-40, 5% glycerol, 1 mM dithiothreitol, proteins were eluted in 500 μl 1x FLAG elution solution (0.5 mg per ml FLAG peptide in wash buffer) at 4 °C for 1 h. Protein purity was verified with SDS–PAGE followed by coomassie staining.

Electrophoretic mobility shift assay/gel shift assay. The Cy5.5-labelled RNA oligonucleotides (ss: Cy5.5-CCUGACUGACUGACUGACGCU; ss-mA6: Cy5.5-CCUGACUGACG(mA6)CGUGCU) with the same sequences of biotin-labelled ss-A and ss-m^6^A were synthesized by GE Dharmacron. The gel shift assay was performed as previously described. Briefly, RNA probes were denatured by heating at 65 °C for 5 min and slowly cooling down to room temperature. For each reaction, 1 μl RNA probes (4 nM final concentration) and 1 μl protein (10% of the concentration gradient indicated in Supplementary Fig. 1d) were incubated in 8 μl binding buffer (10 mM Tris–HCl pH 7.5, 50 mM KCl, 1 mM EDTA, 0.05% Triton X-100, 5% glycerol, 1 mM dithiothreitol and 40 μM per ml RNasin) on ice for 30 min. The RNA–protein mixtures were separated in 5% native polyacrylamide gels (in 0.5xTris-borate-EDTA buffer) at 4 °C for 60 min at 13 V per cm and visualized by Odyssey Imaging System (LI-COR Biosciences). The signal intensities of bands were quantified by ImageJ software. The experiments were repeated three times, and the results were normalized to the values of the control. The band intensities were compared by t-test and corrected by Bonferroni multiple comparisons.

To identify m^6^A-binding proteins, we further focused on the proteins that have top motifs containing ‘GGAC’ , which represents the most common m^6^A consensus sequence.
with RiboMinus Eukaryote Kit v2 (Ambion) followed by depleting transfer RNA (tRNA) with RNA Clean and Concentrator-5 (Zymo Research). 50 ng purified mRNA of each sample was subjected to LC-MS/MS quantification of m6A levels as reported previously27.

RIP. RIP was performed as previously described28 with some modifications. Briefly, cells seeded in a 10-cm dish at 70–80% confluency were crosslinked by UV and harvested by trypsinization. Nuclear extraction was isolated and sonicated. 1 × 106 cells seeded in a 10-cm dish at 70–80% confluency were treated with 200 µl 0.6 M hydroxylamine. After 24 h incubation, cells were washed with ice-cold PBS, crosslinked with 150 mJ per cm2 of 365 nm UV light at 4 °C, followed by washing three times and incubation with pre-cleared nuclear immunoprecipitation buffer supplemented with RNase inhibitors at 4 °C overnight. After washing with RIP buffer for three times, a second round of RNA T1 digestion was conducted under 30 U per µl at 22 °C for 15 min. Input and co-immunoprecipitated RNAs were recovered and analysed by qPCR or RNA-seq.

CLIP. CLIP was performed following previously reported protocol29 with some modifications. HEK293T cells seeded in four 15-cm plates at approximately 80% confluence were treated with 200 µg/ml 4-thiouridine. After 16 h, cells were washed with ice-cold PBS, crosslinked with 150 µl per cm² of 365 nm UV light and harvested by trypsinization. Nuclear extraction was isolated and sonicated. 2–3 µg of nuclear lysates were serially digested by 0.05 U per µl DNase RQ1 at 37 °C for 5 min, and 0.2 µl per µl RNase T1 at 22 °C for 15 min. 10 µg of FLAG (F3165, Sigma-Aldrich) or IFGBP2 (14672, CST) antibody was conjugated to protein A/G magnetic beads (Thermo Fisher Scientific) by incubation for 4 h at 4 °C overnight. After washing with RIP buffer for three times, a second round of RNase T1 digestion was conducted under 30 U per µl at 22 °C for 15 min. Input and co-immunoprecipitated RNAs were recovered and analysed by qPCR.

RNA-seq. Total RNA was isolated from IFGBP knockdown or control HepG2 cells using mirNeasy Kit (Qiagen). Poly(A) RNA was subsequently purified from 50–100 ng total RNA using NEBNext Poly(A) mRNA Magnetic Isolation Module. NEBNext Ultra Directional RNA Library Prep Kit (New England Biolabs) was used for library preparation. Each group was sequenced in triplicate.

RNA stability assay and sequencing for mRNA lifetime. HepG2 cells with stably expressed shRNAs against IFGBP or shNS were seeded into 6-well plates to get 50% confluency after 24 h. Cells were treated with 5 µg per ml actinomycin D and collected at indicated time points. The total RNA was extracted by mirNeasy Kit (Qiagen) and analysed by RT–PCR and RNA-seq. For RNA-seq, an equal amount of external RNA control consortium (ERCC) RNA spike-in control (Thermo Fisher Scientific) was added to the total RNA samples as internal controls before library construction. Sequencing libraries were prepared using NEBNext Ultra Directional RNA Library Prep Kit. RNA stability profiling was generated from two biological replicates. The turnover rate and half-life of mRNA was estimated according to a previously published paper30. As actinomycin D treatment results in transcription stalling, the change of mRNA concentration at a given time (dC/dt) is proportional to the constant of mRNA decay (Kdec) and the mRNA concentration (C), leading to the following equation:

\[
\frac{dC}{dt} = -K_{\text{decay}}C
\]

Thus, the mRNA decay rate Kdecay was estimated by:

\[
\ln\left(\frac{C_f}{C_0}\right) = -K_{\text{decay}}t
\]

To calculate the mRNA half-life (t1/2), when 50% of the mRNA is decayed (that is, C/t0 = 1/2), the equation was:

\[
\ln\left(\frac{1}{2}\right) = -K_{\text{decay}}t_1/2
\]

From where:

\[
t_1/2 = \ln2 / K_{\text{decay}}
\]

m6A-seq. Total RNA was extracted by homogenizing cells in TRIzol reagent and purifying with Direct-zol RNA MiniPrep Kit (Zymo). mRNA was further purified using Dynabeads mRNA Purification Kit (Thermo Fisher). RNA fragmentation was performed by sonication at 10 ng per µl in 100 µl RNase-free water using Bioruptor Pico (Diagenode) with 30 s on and 30 s off for 30 cycles. m6A immunoprecipitation and library preparation were performed according to published protocol31. Sequencing was carried out on Illumina HiSeq 2000 according to the manufacturer's instructions.

Sequencing data analysis. For RIP–seq data: samples were sequenced by Illumina HiSeq 1000 with a single-end 51-bp pair (bp) read length. The RIP–seq reads were mapped to human genome version hg19 by TopHat2 version 2.0.13 with default settings41. Differential gene expression was calculated by Cuffdiff version v2.2.1 (ref.32). The RIP targets were defined as genes with reads per kilobase, per million reads (RPKM) ≥ 1, immunoprecipitation/input ≥ 2, and P < 0.05. For PAR-CLIP data: the IFGBP PAR-CLIP data were obtained from the public database Gene Expression Omnibus (GEO; accession No. GSE21918)33. Ten replicates were trimmed to cut-off version 1.9.1 (ref.32). The processed reads were mapped to human genome version hg19 by bowtie version 1.1.2 with parameters: -v 3 -m 5 -b -strata34. The mapped results were analysed by PARalyzer v1.5 with default settings35. The results were further filtered by ModeScore ≥ 0.6. For peaks that were larger than 50 nt, we extracted 50 nt centered on the ModeLocation site. GENCODE v24 was used to annotate the filtered peaks, and finally, the PAR-CLIP target genes of IFGBP1 (7511), IFGBP2 (7974) and IFGBP3 (9228) were identified. We then chose peaks that overlapped with RIP targets for de novo motif analysis using HOMER software with default RNA analysis parameters.

For eCLIP data: to identify IFGBP footprints, all mapped reads to consecutive genomic peaks were first assembled. Significant peaks were calculated by determining the read-number cutoffs using the Poisson distribution as previously described36. The Poisson distribution assumes all intervals are independent and have equal probability of an occurrence happening. Peaks with significantly high read-number values (P < 10 × 10−5 and the minimum peak height ≥ 10) and occurring in both basal and stimulated duplicates were defined as IFGBP footprints. We performed de novo motif identifications on eCLIP peaks using the HOMER software37 with default RNA analysis parameters.

For RNA-seq data: all RNA-seq samples were sequenced by Illumina HiSeq 1000 with single-end 51-bp read length. All reads were mapped to human genome version hg19 by hisat2 v2.0.4 with default settings38. Read counts were calculated using HTSeq39, and was converted to RPKM by our custom Perl script. The average gene expression values of three independent studies were used for the following analysis.

For mRNA lifetime profiling: all RNA-seq samples for mRNA lifetime profiling were sequenced by Illumina HiSeq1000 with single-end 51-bp read length. All reads were mapped to human genome version hg19 by hisat2 v2.0.4 with default settings40. Read counts were calculated using HTSeq40, and was converted to RPKM by our custom Perl script. RPKM was converted to atomole by linear fitting of the RNA spike-in as previously described41. The degradation rate of RNA and the mRNA half-life were calculated according to the aforementioned formula. The final half-life was calculated by using the average value of 1 h, 3 h and 6 h. For m6A-seq samples were sequenced by Illumina HiSeq 2000 with single-end 51-bp read length. All reads were mapped to human genome version hg19 by TopHat v2.0.13 with default settings42. The m6A level changes for shMETTL14/shNS were calculated by using exomePeak43. Gene expression level changes for input and treatment were analysed using Cuffdiff.

Integrative data analysis and statistics. Three biological replicates of RNA-seq data were conducted for shNS, shIFGBP2, shIFGBP2 and shIFGBP3. The gene expression values used were the average of the three replicates. The differential expression patterns of non-targets, PAR-CLIP targets and CLIP + RIP targets were compared for each IFGBP isoform by log, transformed FC (that is, log(shIGF2BP2/shNS), log(shIGF2BP2/shNS) and log(shIGF2BP3/shNS)). Non-parametric Mann–Whitney U-test or Wilcoxon rank-sum test, two-sided significance level = 0.05) was applied for calculating the P value44. The same analysis was applied for mRNA half-life.

Gene-specific m6A qPCR. m6A modifications on individual genes were determined using Magna MeRIP m6A Kit (Millipore) following the manufacturer’s instructions. Briefly, 100 µg of total RNA was sheared to about 100 nt in length by metal-ion-induced fragmentation, then purified and incubated with anti-m6A antibody (200/03, Synaptic Systems)-conjugated dynabeads (CS200621, Millipore)-conjugated beads in 500 µl immunoprecipitation buffer supplemented with RNase inhibitors at 4 °C overnight. Methylated RNA was immunoprecipitated with beads, eluted by competition with free m6A, and recovered with RNasey kit (Qiagen). One-tenth of the fragmented RNA was saved as input control, and further analysed by qPCR along with MeRIPed RNA. The related enrichment of m6A in each sample was calculated by normalizing to total input.

Immunofluorescence and microscopy. HEK cells were grown on cover slides, fixed and stained with indicated antibodies as previously described45. Antibodies used for immunofluorescence were as follows: FLAG (F3165, 1:200, Sigma–Aldrich), HuR (1:2582, 1:200, CST), DCPI1 (A303–590A T–1:200, Bethyl Laboratories), Alexa Fluor 488 anti-rabbit IgG (4412, 1:500, CST), Alexa Fluor 594 anti-mouse IgG (8890, 1:500, CST). Nuclei were stained by 4,6-diamidino-2-phenylindole (DAPI). Image acquisition was performed on a Zeiss LSM-710 confocal microscope under a x63 oil objective (Zeiss).

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Polysome profiling. We followed the procedure reported previously40 with the following modifications. We started with one 15-cm dish of confluent HEK293T cells transiently overexpressing FLAG-tagged IGF2BP2. Before collection, cycloheximide was added to the media at 100 μg per ml for 7 min. The lysis buffer was formulated as 20 mM HEPEs, pH 7.6, 100 mM KCl, 5 mM MgCl₂, 100 μg per ml cycloheximide, 1% Triton X-100, freshly added 1:100 protease inhibitor (Roche), 40 μg per ml SUPEResin (Ambion). The sample was then frozen into 30 fractions, 0.5 ml per fraction, and analyzed by Gradient Station (BioCamp) equipped with ECONOV monitor (BioRad) and Gison FC203B fraction collector (Mandell Scientific). Sample from each fraction was subjected to western blot analysis for FLAG (A5892, Sigma-Aldrich), eIF3A (3411, CST), eIF3B (sc-16377, Santa Cruz) and HuR (A-21277, Molecular Probes), or to qPCR analysis of the MYC transcript.

For detection of endogenous IGF2BP2 proteins in ribosomal fractions, three 15-cm dishes at approximately the same number of cells were counted under a microscope as described above. Sample from each fraction was subjected to western blot analysis for IGF2BP1, IGF2BP2 and IGF2BP3.

Dual-luciferase reporter assay. To generate the CRD firefly luciferase reporter construct, DNA fragments of wild-type and mutant CRD were synthesized by Integrated DNA Technologies and cloned into the Xhol site of the pMIR-REPORT vector (Ambion). HEK293T or HeLa cells were seeded in triplicate in 24-well plates to allow 70% confluence in the next day. 100 ng reporter plasmids with wild-type or mutant CRD (pMIR-CRD-WT and pMIR-CRD-mut, respectively) and 20 ng IGF2BPs (pRL-TK) were co-transfected with or without IGF2BP2 expression vectors using Lipofectamine 2000 (Invitrogen). Fluc and Rluc activities were measured 24 h later with the Dual-Luciferase Reporter Assay System (Promega) according to the supplier's instructions. The relative luciferase activity was calculated by dividing Fluc by Rluc and normalized to individual control for each assay. For measuring Fluc mRNA changes, lysates left from luciferase activity measurement were lyzed with Qiazol reagent and total RNAs were extracted for qPCR analysis of Fluc and Rluc mRNA abundance. The relative luciferase mRNA was calculated as described above.

Protein co-immunoprecipitation and nLC-ESI-MS/MS. Cells grown in 10-cm dishes at 70–80% confluency were lysed with 500 μl NP40 buffer (150 mM NaCl, 1.5 mM MgCl₂, 0.5% NP40, 50 mM Tris-HCl at pH 8.0) and sonicated. Proteins were immunoprecipitated from 500 μg of cell lysates with FLAG or IGF2BP1 antibody and the corresponding IgG as (described above). After applying a magnet, proteins associated with Protein A/G Magnetic Beads were washed three times and analyzed by western blotting. For mass spectrometry analysis, the protein complexes were eluted from beads by incubation with 0.2 mol per litre glycine buffer (pH 2.6), followed by neutralization with equal volume of 1 mol per litre Tris pH 8.0, and identified using nLC-MS/MS by the Proteomics Laboratory (University of Cincinnati, Ohio, USA). Briefly, samples from immune-enriched with IgG and FLAG antibodies were solubilized in Laemmli gel buffer and loaded onto separated lanes of a 4–12% Tris-HCl gel at pH 8.0 and sonicated. Proteins were immunoprecipitated from 500 μg of cell lysates with FLAG or IGF2BP1 antibody and the corresponding IgG as (described above). After applying a magnet, proteins associated with Protein A/G Magnetic Beads were washed three times and analyzed by western blotting.

Cell proliferation, migration and invasion assays. HeLa and HepG2 cells were seeded in 96-well plates at 1,000 cells per well. Cell proliferation was evaluated by MTT Assay (Promega). For the colony formation assay, 2,000 HeLa cells or 10,000 HepG2 cells were seeded in 6-well plates and stained with crystal violet 7–10 days later. Colonies were counted in three random fields under a ×20 microscope.

For the cell migration and invasion transwell assays, 25,000 HepG2 cells or 50,000 HeLa cells in 500 μl starvation media were plated on the top chambers of Transwell Clear Polyester Membrane Inserts (for the migration assay, Corning Costar) and BioCoat Matrigel Invasion Chambers (for the invasion assay, Corning Costar), while culture media with 20% FBS was applied on the bottom. After 48–72 h, migrated or invaded cells were stained with crystal violet and counted under a ×20 microscope.

For the wound-healing assay, HeLa and HepG2 cells were seeded in 24-well plates and allowed to grow to an confluent monolayer in 24 h. Cells were then scratched and allowed to continue to culture in complete culture media for migration for 24–72 h. Images were taken at the indicated time under microscope. The scratched areas were quantified using Adobe Photoshop and were used to calculate migration rate.

RNA isolation and quantitative RT–PCR. Total RNA was isolated from cultured cells using miNeasy Kit or TRizol reagent. First-strand complementary DNA (cDNA) was synthesized by reverse transcription of 500 ng RNA using QuantiTect Reverse Transcription Kit (Qagen). qPCR was carried out using QuantiTect SYBR Green PCR Kit (Qagen) and mRNA expression was normalized to reference genes, GAPDH and TATA-binding protein (TBP). The primers used in all qPCR assays are listed in Supplementary Table 2.

CRISPR–Cas9 KO. Human HepG2 cells were transiently transfected with a modified pSpCas9(BB)-2A-GFP plasmids (48138, Addgene) containing IGF2BP1, IGF2BP2 or IGF2BP3 sgRNAs with optimized scaffold29 and a high-fidelity eSpCas9(1.1)44 using Lipofectamine 2000 (Invitrogen) following the manufacturer's instructions. Single cells were isolated 48 h after transfection by FACS (BD FACs Aria III) into 96-well plates. Only cells with high levels of green fluorescent protein (GFP; top 4%) were selected. Independent clones were allowed to grow for 3 weeks. KO efficiency was tested by western blotting with specific antibodies. The sgRNA sequences used are listed below: IGF2BP1, TATCCACCCACCGCTCGGAT; IGF2BP2, GAGAAGTGCCCCAGGGGCGC; IGF2BP3, TGGCCACCGACTGATAGAGCT.

Statistics and reproducibility. Comparisons were performed by using t-tests (two-tailed) or Wilcoxon and Mann–Whitney test as indicated in the figure legends. Data are presented as mean ± s.d. P < 0.05 was considered significant. The number of biological (non-technical) replicates for each experiment is indicated in the figure legends. Three independent sets of RNA samples were used for RNA-seq. Two independent sets of RNA samples were used for mRNA stability profiling and two sets of RNA samples were used for mRNA-seq. All western blot, dot blot and immunofluorescence images are representative of three independent experiments.

No statistical method was used to predetermine sample size.

Life Sciences Reporting Summary. Further information on experimental design is available in the Life Sciences Reporting Summary.

Code availability. The custom Perl and R scripts used in this study are available on request to the corresponding authors.

Data availability. All sequencing data that support the findings of this study have been deposited in NCBI's GEO under accession numbers GSE90639 (for RIP-seq), GSE90642 (for m6A-seq), and GSE90684 (for RNA-seq). Data for LC/MS/MS of IGF2BP2 co-immunoprecipitation have been deposited to Figshare (https://doi.org/10.6084/m9.figshare.5693410). Previously published ENCODE PAR-CLIP and eCLIP data that were re-analyzed here are available under accession codes GSE21918 (for PAR-CLIP of IGF2BP3). GSE57886 (for IGF2BP1 eCLIP in HepG2), GSE92220 (for IGF2BP3 eCLIP in HepG2) and GSE85909 (for IGF2BP2 eCLIP in HeLa). The human cancer data were derived from TCGA Research Network (http://cancergenome.nih.gov/). The data set derived from this resource that supports the findings of this study is available in the cbioPortal for Cancer Genomics (http://www.cbioportal.org/), using TCGA pan-cancer studies.

Source data for Figs. 1c,k, 2c, 4c, 5a, 6b–f, 7b–d and 8b–d and Supplementary Figs. 2e–f, 3c, 4c, 5d and 6d–f are provided in Supplementary Table 3. All other data supporting the findings of this study are available from the corresponding authors on reasonable request.

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Experimental design

1. Sample size
Describe how sample size was determined.

No sample size was pre-determined. Sample size and number of independent experiments are always clearly stated in the figure legend or in the Methods section. Three to more independent results were used to perform statistical analyses. If less, no statistics were performed from these samples. All raw data required for statistical tests are indicated in supplementary Table 3 (Statistics data source).

2. Data exclusions
Describe any data exclusions.

No data were excluded from analysis.

3. Replication
Describe whether the experimental findings were reliably reproduced.

Experiments in the article were reliably reproduced, replication were described in the figure legends.

4. Randomization
Describe how samples/organisms/participants were allocated into experimental groups.

No formal randomization techniques was used. No animals and/or human research participants were involved.

5. Blinding
Describe whether the investigators were blinded to group allocation during data collection and/or analysis.

Investigators were not blinded to group allocation during data collection and/or analysis.

Note: all studies involving animals and/or human research participants must disclose whether blinding and randomization were used.

6. Statistical parameters
For all figures and tables that use statistical methods, confirm that the following items are present in relevant figure legends (or in the Methods section if additional space is needed).

n/a

Confirmed

☐ The exact sample size (n) for each experimental group/condition, given as a discrete number and unit of measurement (animals, litters, cultures, etc.)

☐ A description of how samples were collected, noting whether measurements were taken from distinct samples or whether the same sample was measured repeatedly

☐ A statement indicating how many times each experiment was replicated

☐ The statistical test(s) used and whether they are one- or two-sided (note: only common tests should be described solely by name; more complex techniques should be described in the Methods section)

☐ A description of any assumptions or corrections, such as an adjustment for multiple comparisons

☐ The test results (e.g. P values) given as exact values whenever possible and with confidence intervals noted

☐ A clear description of statistics including central tendency (e.g. median, mean) and variation (e.g. standard deviation, interquartile range)

☐ Clearly defined error bars

See the web collection on statistics for biologists for further resources and guidance.
Software

Describe the software used to analyze the data in this study.

Microsoft excel 2016 was used to calculate mean, standard deviation and P value. TotalLab 2.0 was used to quantify signal of gel shift assay. The custom Perl and R scripts used in this study are available on request to the corresponding authors.

For manuscripts utilizing custom algorithms or software that are central to the paper but not yet described in the published literature, software must be made available to editors and reviewers upon request. We strongly encourage code deposition in a community repository (e.g. GitHub). Nature Methods guidance for providing algorithms and software for publication provides further information on this topic.

Materials and reagents

Indicate whether there are restrictions on availability of unique materials or if these materials are only available for distribution through for-profit company.

No unique materials used in the study.

Describe the antibodies used and how they were validated for use in the system under study (i.e. assay and species).

Mouse anti-m6A antibody, Supplier: Synaptic Systems, Cat.: #202003, RRID: AB_2279214.
Mouse anti-FLAG M2 antibody, Supplier: Sigma-Aldrich, Cat.: F3165, Clone: M2, Lot: SLBN8915V, RRID: AB_259529.
Rabbit anti-IGF2BP1 antibody, Supplier: Cell Signaling Technology, Cat.: #8482, Clone: clone D33A2, Lot: 1.
Rabbit anti-IGF2BP2 antibody, Supplier: Cell Signaling Technology, Cat.: #14672, Clone: clone D4R2F, Lot: 1.
Rabbit anti-MYC antibody, Supplier: Cell Signaling Technology, Cat.: #13987, Clone: clone D3N8F, Lot: 1.
Rabbit anti-HuR antibody, Supplier: Cell Signaling Technology, Cat.: #12582, Clone: clone D9W7E, Lot: 1.
Rabbit anti-IGF2BP3 antibody, Supplier: Bethyl Laboratories, Cat.: A303-426A, RRID: AB_10951696.
Rabbit anti-MATRIN3 antibody, Supplier: Bethyl Laboratories, Cat.: A300-591A-T, RRID: AB_495514.
Rabbit anti-DCP1A antibody, Supplier: Bethyl Laboratories, Cat.: A303.590A-T, RRID: AB_11125540.
Mouse anti-GAPDH antibody, Supplier: Santa Cruz Biotechnology, Cat.: sc-47724, Clone: 411, RRID: AB_627678.
rabbit IgG, Supplier: Millipore, Cat.: #NI01, Lot: D00168753.
HRP-conjugated anti-rabbit IgG secondary antibody, Supplier: Santa Cruz Biotechnology, Cat.: sc-2357, Lot: A1817.
HRP-conjugated anti-mouse IgG secondary antibody, Supplier: Santa Cruz Biotechnology, Cat.: sc-2055, Lot: E1116.
Alexa Fluor 488 anti-rabbit IgG, Supplier: Cell Signaling Technology, Cat.: #4412, Lot: 11.
Alexa Fluor 594 anti-mouse IgG, Supplier: Cell Signaling Technology, Cat.: #8890, Lot: 2.

Antibodies were validated by the use of negative control and/or positive control (such as knockdown or overexpression) for IGF2BP1, IGF2BP2, IGF2BP3, MYC, HuR and FLAG antibodies. Antibodies were used at 1:1000 dilution for western blot or at 1:200 for immuno staining, while 1 microgram or 10 microgram antibody was used for each co-ip/RIP or CLIP assay.
10. Eukaryotic cell lines
   a. State the source of each eukaryotic cell line used. All cell lines were purchased from the American Type Culture Collection (ATCC).
   b. Describe the method of cell line authentication used. Cell lines were not authenticated by ourselves.
   c. Report whether the cell lines were tested for mycoplasma contamination. All cell lines were tested to be mycoplasma negative.
   d. If any of the cell lines used are listed in the database of commonly misidentified cell lines maintained by ICLAC, provide a scientific rationale for their use. No cell lines used in this study were found in the database of commonly misidentified cell lines that is maintained by ICLAC and NCBI Biosample.

Animals and human research participants

Policy information about studies involving animals; when reporting animal research, follow the ARRIVE guidelines

11. Description of research animals
    Provide details on animals and/or animal-derived materials used in the study. No animals were used.

Policy information about studies involving human research participants

12. Description of human research participants
    Describe the covariate-relevant population characteristics of the human research participants. The study did not involve human research participants.