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## Identification of YTH Domain-Containing Proteins as the Readers for *m*<sup>1</sup>A-Methyladenosine in RNA

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### Abstract

*m*<sup>1</sup>A-methyladenosine (*m*<sup>1</sup>A) is an important post-transcriptional modification in RNA; however, the exact biological role of *m*<sup>1</sup>A remains to be determined. By employing a quantitative proteomics method, we identified multiple putative protein readers of *m*<sup>1</sup>A in RNA, including several YTH domain family proteins. We showed that YTHDF1–3 and YTHDC1, but not YTHDC2, could bind directly to *m*<sup>1</sup>A in RNA. We also found that Trp<sup>432</sup> in YTHDF2, a conserved residue in the hydrophobic pocket of the YTH domain that is necessary for its binding to *N*<sup>6</sup>-methyladenosine (*m*<sup>6</sup>A), is required for its recognition of *m*<sup>1</sup>A. An analysis of previously published data revealed transcriptome-wide colocalization of YTH domain-containing proteins and *m*<sup>1</sup>A sites in HeLa cells, suggesting that YTH domain-containing proteins can bind to *m*<sup>1</sup>A in cells. Together, our results uncovered YTH domain-containing proteins as readers for *m*<sup>1</sup>A in RNA and provided new insight into the functions of *m*<sup>1</sup>A in RNA biology.

### Graphical abstract

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#### ASSOCIATED CONTENT

##### Supporting Information

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Detailed experimental procedures, list of PCR primers, LC–MS and MS/MS data, effects of knock-down of YTHDF1 and YTHDF2 on the lifetime and translation efficiency of *m*<sup>1</sup>A-containing RNA, and quantitative proteomic data (PDF)

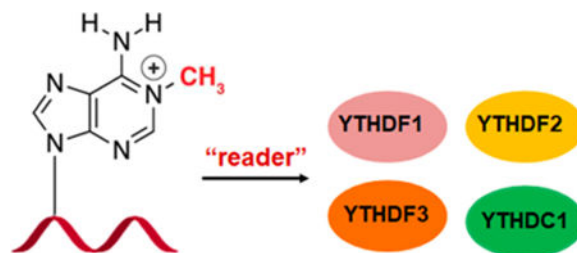
List of proteins quantified from the SILAC-based interaction screening for *m*<sup>1</sup>A-carrying RNA and the corresponding adenosine-carrying RNA using lysates from HEK293T cells and HeLa cells (XLSX)

List of proteins quantified from the SILAC-based interaction screening for *m*<sup>6</sup>A-carrying RNA and the corresponding adenosine-carrying RNA using lysates from HEK293T cells (XLSX)

*m*<sup>1</sup>A sites which overlap with YTHDF1–3 and YTHDC1 PAR-CLIP binding sites in HeLa cells (XLSX)

Comparison of RNA lifetime in cells with and without siRNA-mediated knockdown of YTHDF1 or YTHDF2 (XLSX)

Translation efficiency of mRNA in cells with and without siRNA-mediated knockdown of YTHDF1, YTHDF2, and YTHDF3 (XLSX)



RNA is known to contain more than 100 distinct types of post-transcriptional modifications, which can modulate its splicing, localization, stability, and translation.<sup>1</sup> It was found that methylation at the  $N^6$  position of adenosine ( $m^6A$ ) is involved in the epigenetic control of gene regulation.<sup>2-6</sup> Recent transcriptome-wide mapping also revealed the presence of  $m^1A$  in human mRNA,<sup>7-9</sup> though there is some controversy about the accuracy and specificity for some of the  $m^1A$  mapping methods.<sup>10</sup> These studies also suggested the potential role of  $m^1A$  in modulating mRNA splicing and translation.<sup>7-9</sup> TRMT6, TRMT61A, and TRMT10C are three known  $m^1A$  methyltransferases in human cells (writers),<sup>9</sup> and  $m^1A$  can be demethylated to adenosine by ALKBH1 and ALKBH3 (erasers).<sup>8,11</sup> However, it remains unknown which cellular proteins are involved in binding to  $m^1A$  in RNA (readers).

YTH domain-containing proteins can bind directly to  $m^6A$  in RNA. YTHDF1 promotes protein synthesis by interacting with the translation machinery;<sup>12</sup> YTHDF2 accelerates the degradation of  $m^6A$ -carrying mRNA and affects the translation of heat shock proteins;<sup>13,14</sup> YTHDF3 facilitates the translation and decay of  $m^6A$ -bearing RNA in synergy with YTHDF1 and YTHDF2;<sup>15,16</sup> YTHDC1 regulates mRNA splicing by modulating pre-mRNA splicing factors and mediates nuclear export of  $m^6A$ -harboring RNA;<sup>17,18</sup> YTHDC2 was found to be essential for meiosis in mammalian germline.<sup>19-21</sup> Together, the YTH domain-containing proteins modulate various biological processes through binding to  $m^6A$ -modified RNA.

In this study, we employed a mass spectrometry-based method to identify the  $m^1A$ -binding proteins in RNA. We revealed that several YTH domain-containing proteins directly interact with  $m^1A$  in RNA. In addition, functional analysis suggested important roles of YTH domain family proteins in the regulation of  $m^1A$ -carrying RNA. Our results provided a foundation for further understanding of the biological functions of this important RNA modification.

## RESULTS AND DISCUSSION

To understand better the biological functions of  $m^1A$  in RNA, we employed the stable isotope labeling by amino acid in cell culture (SILAC)<sup>22</sup>-based quantitative proteomics method to screen systematically for cellular proteins that bind to  $m^1A$ -bearing RNA (Figure 1a). To this end, we used a 5'-biotin-labeled  $m^1A$ -carrying RNA sequence derived from human SOX18 gene, which was previously shown to harbor an  $m^1A$  site near the start codon in both HEK293T and HeLa cells,<sup>7,8</sup> as the probe bait and the corresponding unmethylated sequence as the control bait. We cultured the cells separately in heavy and light culture media and extracted proteins from these cells. An equal amount of proteins from the heavy-

and light-labeled cells were incubated with biotin-conjugated m<sup>1</sup>A-bearing RNA and the corresponding unmethylated RNA, respectively, which was designated as the forward SILAC experiments. The opposite incubations in reverse SILAC experiments were conducted to remove experimental bias. After the incubation, the RNA-conjugated beads were extensively washed to remove nonspecific proteins, and the bound proteins were isolated from the beads, digested with trypsin, and analyzed using LC–MS/MS (Figure 1a).

The LC–MS/MS results revealed multiple proteins exhibiting preferential binding toward the m<sup>1</sup>A-bearing RNA over the corresponding rA-containing RNA (Figure 1b, Figure S1). In HEK293T cells, these proteins include YTH domain family members YTHDF3 and YTHDF2 and heterogeneous nuclear ribonucleoprotein hnRNPD, with the SILAC protein ratios (m<sup>1</sup>A/rA) being  $6.9 \pm 1.7$ ,  $4.1 \pm 2.7$ , and  $2.3 \pm 0.3$ , respectively (Figure 1b, Table S2). In HeLa cells, the proteins which bind more strongly to m<sup>1</sup>A-bearing RNA probe over rA-bearing RNA probe encompass TAR DNA-binding protein (TARDBP), DEAD-box helicase DDX56, YTHDF proteins, and others (Figure S1, Table S2). Among them, YTHDF1, YTHDF2, and YTHDF3 displayed SILAC ratios (m<sup>1</sup>A/rA) of  $2.3 \pm 0.7$ ,  $1.9 \pm 0.3$ , and  $2.1 \pm 0.4$ , respectively (Table S2). It is worth noting that our LC–MS/MS results also identified some proteins that bind more strongly to rA-containing RNA probe over m<sup>1</sup>A-bearing RNA probe, such as interleukin enhancer binding factors ILF2 and ILF3, which form a complex and negatively regulates microRNA processing,<sup>23</sup> KU70 and KU80, which forms a heterodimer and functions in DNA double strand break repair via the nonhomologous end-joining pathway,<sup>24</sup> etc. (Figure 1b and Table S2).

We also performed the SILAC experiments using the corresponding m<sup>6</sup>A and rA probes in HEK293T cells. As expected, the SILAC-based interaction screening resulted in the identification of YTHDF3, YTHDF2, and YTHDF1 as proteins displaying preferential binding toward the m<sup>6</sup>A-containing RNA over the rA-containing RNA substrate, with the SILAC protein ratios (m<sup>6</sup>A/rA) being  $14.6 \pm 7.9$ ,  $9.7 \pm 3.6$ , and  $9.0 \pm 2.7$ , respectively (Figure 1c, Table S3). This result is consistent with the previous finding,<sup>13</sup> validating that our SILAC-based method is capable of identifying specific RNA-binding proteins. Aside from YTH domain family proteins, our SILAC-based interaction screening also identified several other m<sup>6</sup>A-binding proteins, such as PCBP1 and CNBP, which exhibited the SILAC ratios (m<sup>6</sup>A/rA) of  $2.1 \pm 0.8$  and  $1.9 \pm 0.3$ , respectively (Figure 1c, Table S3). Moreover, our results showed that some proteins exhibit stronger binding affinities to the rA-containing RNA probe over the corresponding m<sup>6</sup>A-bearing RNA probe, such as heterogeneous nuclear ribonucleoprotein hnRNPA0, and the aforementioned ILF2, ILF3, KU70, and KU80, etc.

Representative LC–MS results for a tryptic peptide derived from YTHDF2, SINNYNPK are shown in Figure 2, which clearly revealed the stronger binding of YTHDF2 to the m<sup>1</sup>A-containing RNA than the corresponding unmethylated probe in both forward and reverse SILAC experiments (MS/MS shown in Figure S2). The selective binding of YTHDF3 toward m<sup>1</sup>A-containing RNA also found its support from a representative tryptic peptide derived from YTHDF3, i.e., IGGDLTAAVTK (Figure 2, MS/MS shown in Figure S3).

SILAC-based quantitative proteomic approach may also result in the discovery of those proteins that interact indirectly with m<sup>1</sup>A-bearing bait through protein–protein

interaction(s). Thus, we examined whether YTH domain-containing proteins can interact directly with m<sup>1</sup>A in RNA. Our results from electrophoretic mobility shift assay (EMSA) showed that YTHDF1–3 and the YTH domain of YTHDC1 (YTHDC1<sup>YTH</sup>), but not the YTH domain of YTHDC2 (YTHDC2<sup>YTH</sup>), bind more strongly to m<sup>1</sup>A-carrying RNA substrate than its unmethylated counterpart (Figure 3a–d, Figure S4). We also compared the binding difference of YTH domain-containing proteins toward m<sup>1</sup>A and m<sup>6</sup>A-bearing RNA baits. The EMSA results showed that YTHDF1–3 and YTHDC1<sup>YTH</sup> bind more strongly to m<sup>6</sup>A-carrying RNA than m<sup>1</sup>A-carrying RNA, with the  $K_d$  values being  $1.3 \pm 0.1/16.5 \pm 1.5$ ,  $1.3 \pm 0.1/5.8 \pm 1.7$ ,  $1.9 \pm 0.1/7.0 \pm 1.1$ , and  $0.7 \pm 0.1/23.3 \pm 2.1 \mu\text{M}$  (m<sup>6</sup>A/m<sup>1</sup>A), respectively (Figure 3a–d). Together, our results support that YTHDF1–3 and YTHDC1<sup>YTH</sup> can bind directly to m<sup>1</sup>A-containing RNA substrate, albeit at affinities that are lower than those toward the m<sup>6</sup>A-containing RNA probe, which are in keeping with the findings made from SILAC-based interaction screening.

The X-ray crystal structure of YTHDF2 revealed three aromatic amino acid residues in the hydrophobic pocket of the YTH domain of YTHDF2 that are crucial for its binding toward m<sup>6</sup>A.<sup>13,25</sup> To explore whether this hydrophobic pocket also assumes an important role in binding toward m<sup>1</sup>A, we conducted EMSA assay with a mutant form of YTHDF2 protein where the conserved Trp<sup>432</sup> was mutated to an alanine (W432A). The result indeed showed that the mutation led to a pronounced attenuation in binding affinity toward the m<sup>1</sup>A-containing probe (Figure 3b), suggesting that m<sup>1</sup>A may occupy the same hydrophobic pocket of YTHDF2 that is required for m<sup>6</sup>A recognition.

We further interrogated the publicly available transcriptome-wide m<sup>1</sup>A mapping data<sup>7</sup> and photoactivatable ribonucleoside cross-linking and immunoprecipitation (PAR-CLIP) data<sup>12,16,18,26</sup> of YTH domain-containing proteins in HeLa cells to assess whether YTH domain-containing proteins bind to m<sup>1</sup>A-bearing RNA in cells. In this aspect, we analyzed the overlap between the m<sup>1</sup>A sites (200 bp window centered on the m<sup>1</sup>A sites) and YTHDF1–3, and YTHDC1 PAR-CLIP binding sites in HeLa cells. The result showed that 24.4%, 23.1%, 4.5%, and 12.5% of the m<sup>1</sup>A sites overlapped with the YTHDF1, YTHDF2, YTHDF3, and YTHDC1 PAR-CLIP peaks, respectively (Figure 3e, Table S4). Therefore, these results suggest that YTH domain-containing proteins can recognize m<sup>1</sup>A in transcripts in cells. Analysis of the ribosome profiling data and RNA lifetime data<sup>12,16,26</sup> in HeLa cells revealed that, upon YTHDF2 knockdown, the YTHDF2-bound m<sup>1</sup>A-containing mRNA targets experienced a significant increase in lifetime compared with other targets ( $p = 1.0 \times 10^{-28}$ , Mann–Whitney U-test, Figure S5, Table S5), whereas the YTHDF1-binding m<sup>1</sup>A-containing targets exhibited a significant decrease in translation efficiency compared with other targets upon genetic depletion of YTHDF1 ( $p = 8.9 \times 10^{-12}$ , Mann–Whitney U-test, Figure S6, Table S6), suggesting that YTHDF1 may promote translation whereas YTHDF2 mainly accelerates the degradation of m<sup>1</sup>A-carrying transcripts.

## CONCLUSIONS

In summary, we identified by using an unbiased quantitative proteomic method, for the first time, YTH domain-containing proteins as the readers of m<sup>1</sup>A in RNA, which expands the repertoire of RNA modifications that can be regulated by this protein family. We

demonstrated that YTH domain-containing proteins directly interact with m<sup>1</sup>A-bearing RNA. Furthermore, we identified multiple other putative m<sup>1</sup>A-binding proteins by using SILAC-based quantitative proteomic screening. Further characterizations of these proteins in m<sup>1</sup>A recognition and RNA metabolism will improve our understanding about the biological functions of m<sup>1</sup>A in RNA.

## Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

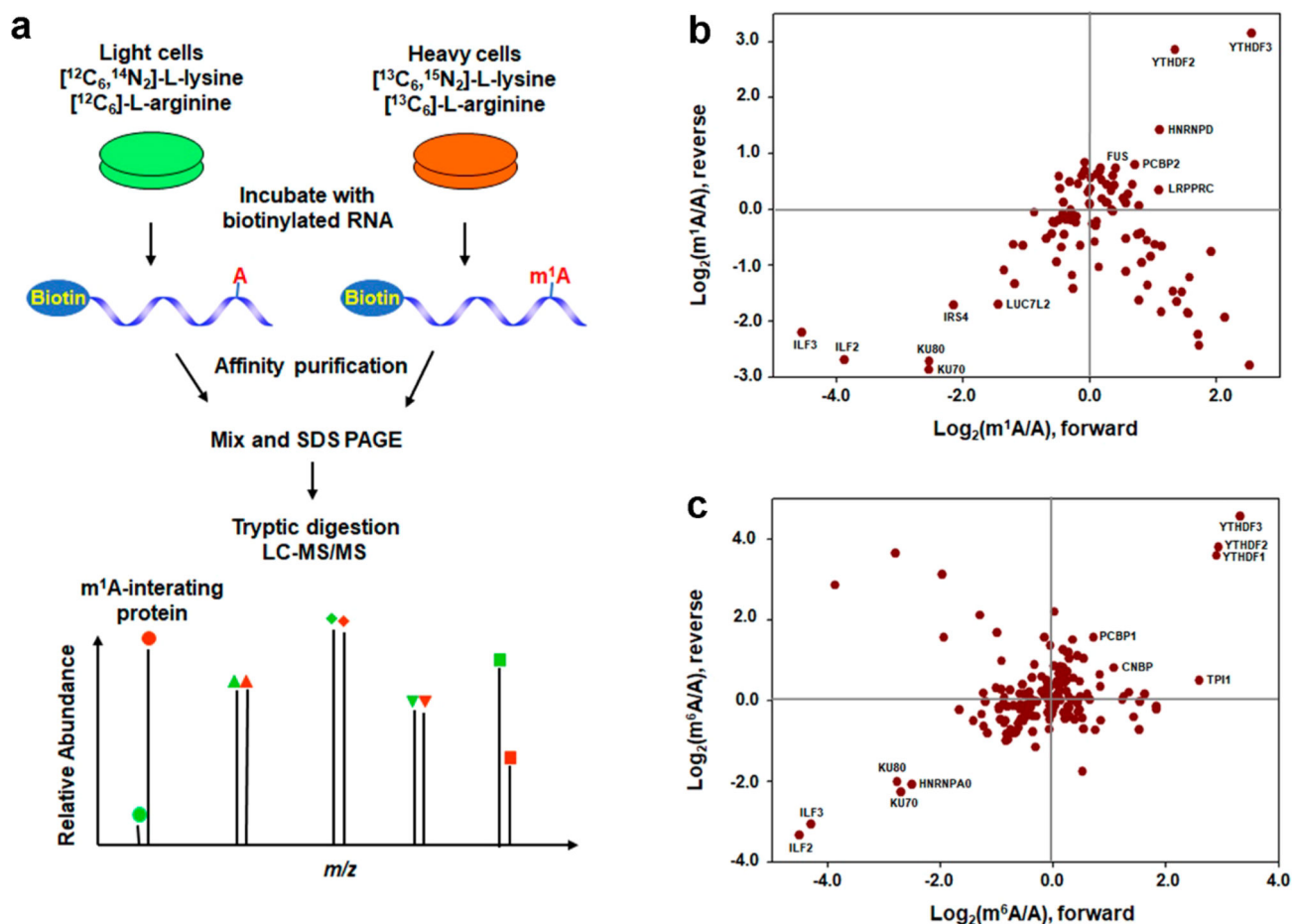
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## REFERENCES

- (1). Cantara WA; Crain PF; Rozenski J; McCloskey JA; Harris KA; Zhang X; Vendeix FA; Fabris D; Agris PF *Nucleic Acids Res.* 2011, 39, D195–201. [PubMed: 21071406]
- (2). Jia G; Fu Y; Zhao X; Dai Q; Zheng G; Yang Y; Yi C; Lindahl T; Pan T; Yang YG; He C *Nat. Chem. Biol.* 2011, 7, 885–887. [PubMed: 22002720]
- (3). Zheng G; Dahl JA; Niu Y; Fedorcsak P; Huang CM; Li CJ; Vagbo CB; Shi Y; Wang WL; Song SH; Lu Z; Bosmans RP; Dai Q; Hao YJ; Yang X; Zhao WM; Tong WM; Wang XJ; Bogdan F; Furu K; et al. *Mol. Cell* 2013, 49, 18–29. [PubMed: 23177736]
- (4). Jia G; Fu Y; He C *Trends Genet.* 2013, 29, 108–115. [PubMed: 23218460]
- (5). Schwartz S; Agarwala SD; Mumbach MR; Jovanovic M; Mertins P; Shishkin A; Tabach Y; Mikkelsen TS; Satija R; Ruvkun G; Carr SA; Lander ES; Fink GR; Regev A *Cell* 2013, 155, 1409–1421. [PubMed: 24269006]
- (6). Liu N; Dai Q; Zheng G; He C; Parisien M; Pan T *Nature* 2015, 518, 560–564. [PubMed: 25719671]
- (7). Dominissini D; Nachtergaele S; Moshitch-Moshkovitz S; Peer E; Kol N; Ben-Haim MS; Dai Q; Di Segni A; Salmon-Divon M; Clark WC; Zheng G; Pan T; Solomon O; Eyal E; Hershkovitz V; Han D; Dore LC; Amariglio N; Rechavi G; He C *Nature* 2016, 530, 441–446. [PubMed: 26863196]
- (8). Li X; Xiong X; Wang K; Wang L; Shu X; Ma S; Yi C *Nat. Chem. Biol.* 2016, 12, 311–316. [PubMed: 26863410]
- (9). Safra M; Sas-Chen A; Nir R; Winkler R; Nachshon A; Bar-Yaacov D; Erlacher M; Rossmann W; Stern-Ginossar N; Schwartz S *Nature* 2017, 551, 251–255. [PubMed: 29072297]
- (10). Grozhik AV; Jaffrey SR *Nat. Chem. Biol.* 2018, 14, 215–225. [PubMed: 29443978]
- (11). Liu F; Clark W; Luo G; Wang X; Fu Y; Wei J; Hao Z; Dai Q; Zheng G; Ma H; Han D; Evans M; Klungland A; Pan T; He C *Cell* 2016, 167, 1897.
- (12). Wang X; Zhao BS; Roundtree IA; Lu Z; Han D; Ma H; Weng X; Chen K; Shi H; He C *Cell* 2015, 161, 1388–1399. [PubMed: 26046440]
- (13). Zhu T; Roundtree IA; Wang P; Wang X; Wang L; Sun C; Tian Y; Li J; He C; Xu Y *Cell Res.* 2014, 24, 1493–1496. [PubMed: 25412661]
- (14). Zhou J; Wan J; Gao X; Zhang X; Jaffrey SR; Qian SB *Nature* 2015, 526, 591–594. [PubMed: 26458103]
- (15). Li A; Chen YS; Ping XL; Yang X; Xiao W; Yang Y; Sun HY; Zhu Q; Baidya P; Wang X; Bhattarai DP; Zhao YL; Sun BF; Yang YG *Cell Res.* 2017, 27, 444–447. [PubMed: 28106076]
- (16). Shi H; Wang X; Lu Z; Zhao BS; Ma H; Hsu PJ; Liu C; He C *Cell Res.* 2017, 27, 315–328. [PubMed: 28106072]

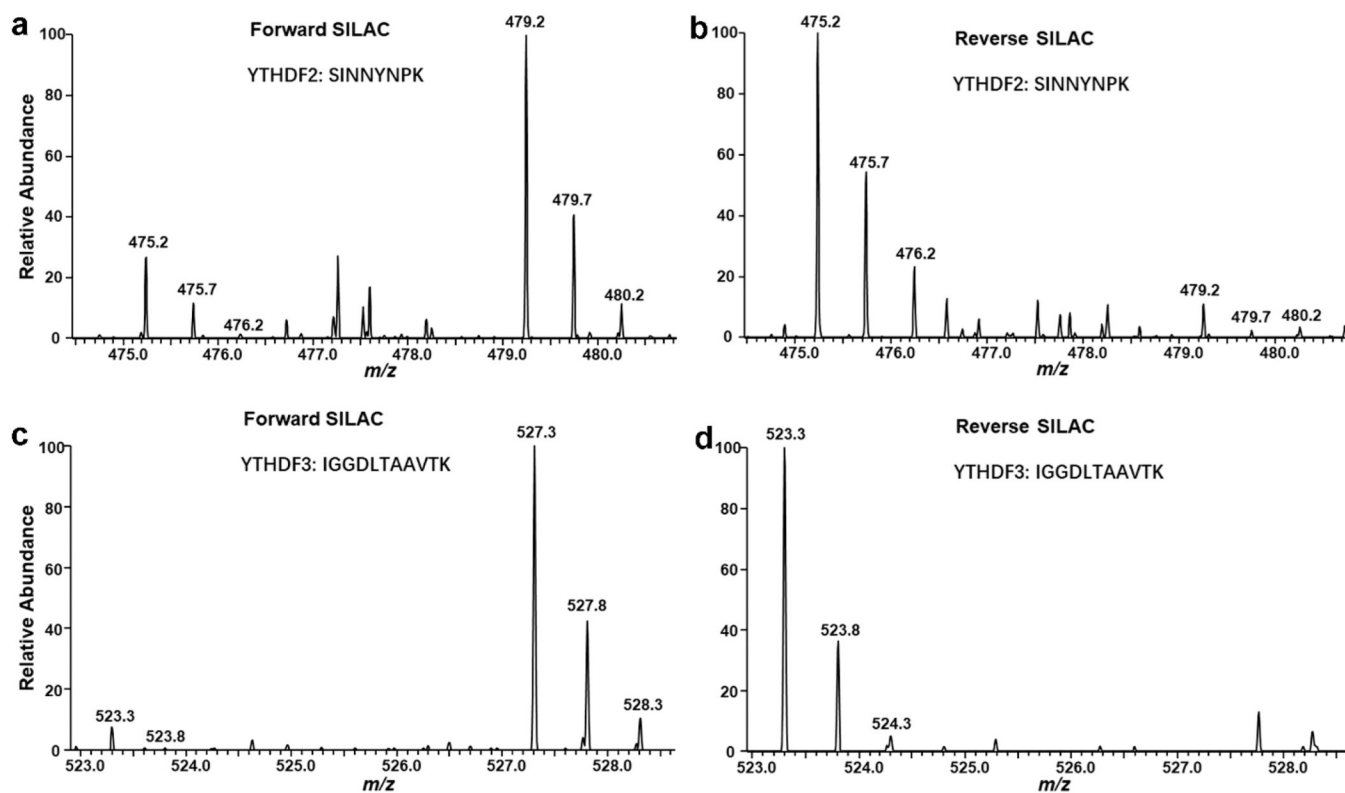
- (17). Xiao W; Adhikari S; Dahal U; Chen YS; Hao YJ; Sun BF; Sun HY; Li A; Ping XL; Lai WY; Wang X; Ma HL; Huang CM; Yang Y; Huang N; Jiang GB; Wang HL; Zhou Q; Wang XJ; Zhao YL; et al. *Mol. Cell* 2016, 61, 507–519. [PubMed: 26876937]
- (18). Roundtree IA; Luo GZ; Zhang Z; Wang X; Zhou T; Cui Y; Sha J; Huang X; Guerrero L; Xie P; He E; Shen B; He C *eLife* 2017, 6, No. e31311, DOI: 10.7554/eLife.31311. [PubMed: 28984244]
- (19). Hsu PJ; Zhu Y; Ma H; Guo Y; Shi X; Liu Y; Qi M; Lu Z; Shi H; Wang J; Cheng Y; Luo G; Dai Q; Liu M; Guo X; Sha J; Shen B; He C *Cell Res.* 2017, 27, 1115–1127. [PubMed: 28809393]
- (20). Wojtas MN; Pandey RR; Mendel M; Homolka D; Sachidanandam R; Pillai RS *Mol. Cell* 2017, 68, 374–387. [PubMed: 29033321]
- (21). Bailey AS; Batista PJ; Gold RS; Chen YG; de Rooij DG; Chang HY; Fuller MT *eLife* 2017, 6, No. e26116. [PubMed: 29087293]
- (22). Ong SE; Blagoev B; Kratchmarova I; Kristensen DB; Steen H; Pandey A; Mann M *Mol. Cell. Proteomics* 2002, 1, 376–386. [PubMed: 12118079]
- (23). Sakamoto S; Aoki K; Higuchi T; Todaka H; Morisawa K; Tamaki N; Hatano E; Fukushima A; Taniguchi T; Agata Y *Mol. Cell. Biol* 2009, 29, 3754–3769. [PubMed: 19398578]
- (24). Walker JR; Corpina RA; Goldberg J *Nature* 2001, 412, 607–614. [PubMed: 11493912]
- (25). Li F; Zhao D; Wu J; Shi Y *Cell Res.* 2014, 24, 1490–1492. [PubMed: 25412658]
- (26). Wang X; Lu Z; Gomez A; Hon GC; Yue Y; Han D; Fu Y; Parisien M; Dai Q; Jia G; Ren B; Pan T; He C *Nature* 2014, 505, 117–120. [PubMed: 24284625]



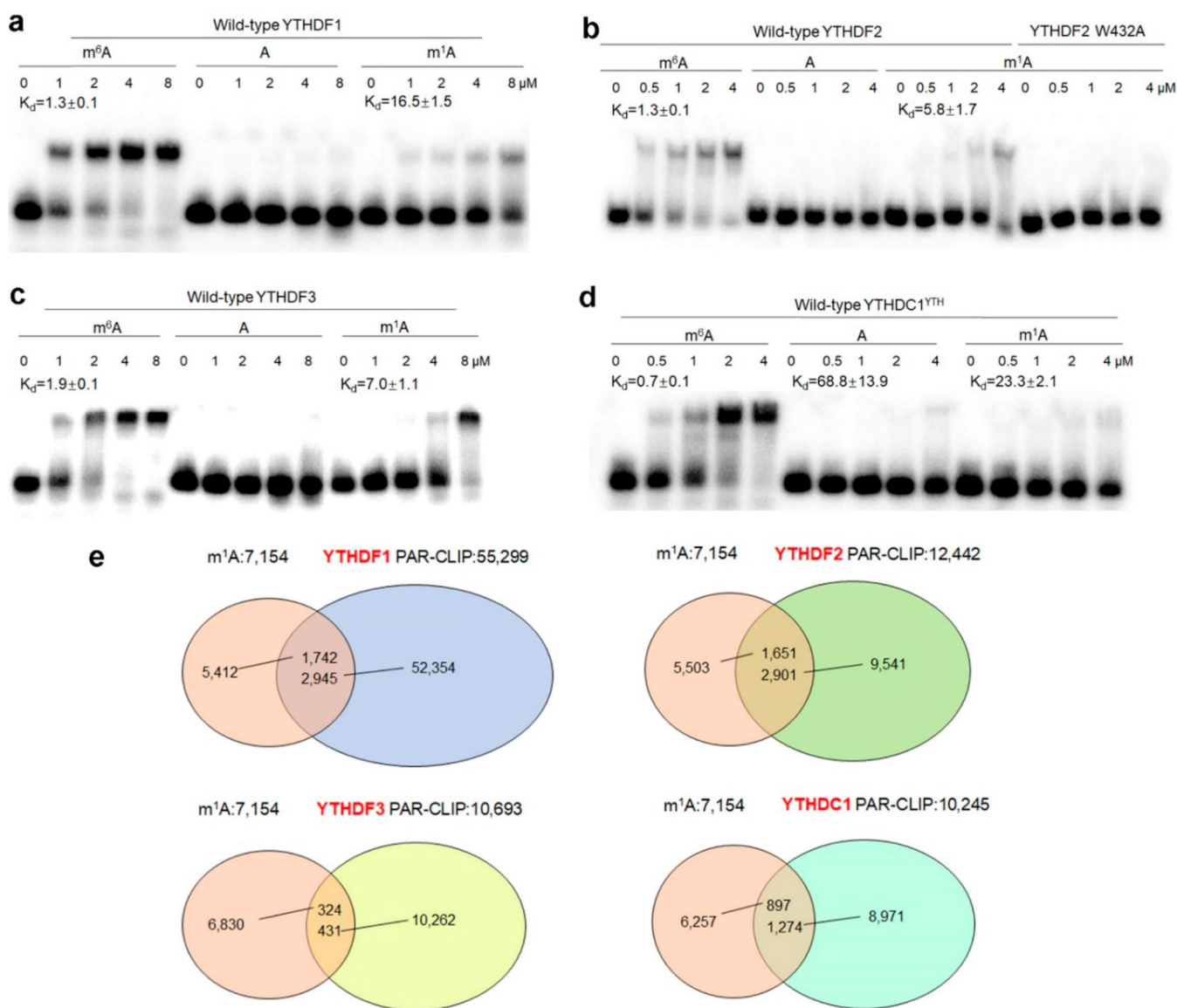
**Figure 1.**

Identification of m<sup>1</sup>A-binding proteins. (a) Schematic overview of the SILAC-based quantitative proteomics method for discovering m<sup>1</sup>A-binding proteins. Shown is the workflow for a forward SILAC labeling experiment. (b,c) Scatterplot of the proteins identified from pull-down assays using m<sup>1</sup>A- (b) or m<sup>6</sup>A- (c) carrying RNA over the corresponding unmodified RNA and the whole-cell protein lysate of HEK293T cells. The proteins present in top right quadrant are m<sup>1</sup>A- (b) or m<sup>6</sup>A (c)-binding proteins, whereas those in the bottom left quadrant are proteins which bind more weakly to the m<sup>1</sup>A- or m<sup>6</sup>A-bearing RNA than the control RNA probe. The proteins that fall in the bottom right quadrant represent those proteins displaying large light/heavy ratios in both forward and reverse SILAC labeling experiments, which are likely due to incomplete SILAC labeling (for those proteins with slow turnover rate). The data were based on results from two forward and one reverse SILAC experiments.





**Figure 2.** Representative ESI-MS for the  $[M + 2H]^{2+}$  ions of a tryptic peptide of YTHDF2, SINNYNPK (a, b) and a tryptic peptide of YTHDF3, IGGDLTAAVTK (c, d) revealing the preferential binding of YTHDF2 and YTHDF3 toward the  $m^1A$ -containing RNA probe in both forward (a, c) and reverse (b, d) SILAC experiments.



**Figure 3.** YTH domain-containing proteins bind directly to m<sup>1</sup>A-carrying RNA. (a–d) Electrophoretic mobility shift assay (EMSA) for measuring the binding affinity of YTHDF1 (a), YTHDF2 and W432A mutant proteins (b), YTHDF3 (c), and YTH domain of YTHDC1 (d) with m<sup>6</sup>A, A, and m<sup>1</sup>A-carrying RNA probes. The  $K_d$  values (in  $\mu\text{M}$ ) are listed in (a)–(d), and error bar represents the SEM ( $n = 3$ ). (e) Overlap between the m<sup>1</sup>A sites (200 bp window centered on the m<sup>1</sup>A sites) and YTHDF1–3, and YTHDC1 PAR-CLIP binding sites in HeLa cells.