Oncogenic and Tumor-Suppressive Functions of the RNA Demethylase FTO

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ABSTRACT

The epitranscriptome represents the more than 140 types of chemically varying and reversible RNA modifications affecting RNA fate. Among these, the most relevant for this review are the mRNA modifications N6-methyladenosine and N6,2′-O-dimethyladenosine. Epitranscriptomic mRNA biology involves RNA methyltransferases (so-called “writers”), RNA demethylases (“erasers”), and RNA-binding proteins (“readers”) that interact with methylation sites to determine the functional outcome of the modification. In this review, we discuss the role of a specific RNA demethylase encoded by the fat mass and obesity–associated gene (FTO) in cancer. FTO initially became known as the strongest genetic link for human obesity. Only in 2010, 16 years after its discovery, was its enzymatic function as a demethylase clarified, and only recently has its role in the development of cancer been revealed. FTO functions are challenging to study and interpret because of its genome-wide effects on transcript turnover and translation. We review the discovery of FTO and its enzymatic function, the tumor-promoting and suppressive roles of FTO in selected cancer types, and its potential as a therapeutic target.

Discovery of the Fto Gene

In an unsuccessful attempt to generate a transgenic mouse model overexpressing the human Harvey rat sarcoma viral oncogene homolog (HRAS), in 1994 a mutant was identified with syndactyly in both forelimbs that was named Fused toes (Ft; ref. 1). Homozygosity of the Ft mutation led to embryonic lethality around day 10 with embryos showing severely altered head morphology including missing large parts of the telencephalon and mesencephalon (1, 2). The mutation was later found to also cause a defect in establishing the left-right axis during embryonic development (3). Full mapping revealed a 1.6 Mb deletion on the long (q) arm of chromosome 8 covering genes currently known as Fused toes 1 (Ft1) or Akt-interacting protein (Aktip), retinins pigmentosa GTPase regulator interacting protein 1 like (Rgrip1L), Fat mass and obesity–associated (Fto) and three members of the Iroquois homeobox protein family Irx3, Irx5, and Irx6 (4–6). Inactivation of mouse Fto/Aktip results in telomeric defects associated with a progeroid phenotype (7, 8). Phenotypes of mice with a homo-morphic mutation in Rgrip1L closely resemble those of Ft1-mutant mice (9). Rgrip1L has a critical function in the assembly of primary cilia and mutations in human RPGRIP1L cause the life-threatening diseases Joubert syndrome and Meckel syndrome (9). Rpgrip1L has a critical function in the assembly of primary cilia and mutations in human RPGRIP1L cause the life-threatening diseases Joubert syndrome and Meckel syndrome (9). Rpgrip1L has a critical function in the assembly of primary cilia and mutations in human RPGRIP1L cause the life-threatening diseases Joubert syndrome and Meckel syndrome (9). Rpgrip1L has a critical function in the assembly of primary cilia and mutations in human RPGRIP1L cause the life-threatening diseases Joubert syndrome and Meckel syndrome (9).

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The Genetic Link of FTO to Metabolism and Obesity

The distal region of mouse chromosome 8q covering the deletion in the Ft1 mutant is homologous to the long arm of human chromosome 16 (16q12-14; ref. 6). In 2007, four independent studies linked SNPs in FTO to obesity in humans (15–18). One of these studies focused on type 2 diabetes mellitus (T2DM) and identified several SNPs in the FTO gene to be significantly associated with T2DM in nearly 40,000 European adults (15). Obesity is a known risk factor for T2DM and the association between the SNPs and T2DM disappeared when correcting for body mass index (BMI), indicating the association of the SNPs with T2DM is mediated through BMI (15). Thereupon, “Fatso” was renamed to “Fat mass and obesity–associated,” maintaining the FTO gene symbol. Although over 100 different loci have been found to robustly associate with BMI to date, the association with FTO remains the strongest of all genes in multiple ancestries (19). The first experimental link between FTO and obesity came with the discovery that whole-body knockout of Fto in mice leads to reduced bodyweight in both males and females, and significantly reduced weight gain in both sexes upon high-fat diet feeding (20). Offspring from heterozygous parents are born to Mendelian ratio and homozygous Fto−/− mice show normal development and weight at birth. However, postnatal survival of Fto−/− mice is decreased with 60% (20), which can be improved by reduction in litter size or postponed weaning, suggesting that the altered metabolism causes reduced survival after weaning. A knockin mouse model introducing the Fto1674/1674 mutant with reduced catalytic activity showed similar phenotypes as the Fto−/− model, albeit only in males and not in females (21). Ubiquitous overexpression of FTO in mice on the other hand leads to an increase in body weight and fat mass in both males and females, on a normal diet as well as on a high-fat diet (22). In humans a recessive loss-of-function mutation in FTO was discovered in an Arab family that leads to postnatal growth retardation, microcephaly, facial dimorphism, and functional brain deficits (23). All patients homozygous for the mutation died between 1 and 30 months of age, indicating additional functions of human
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FTO compared with murine Fto (23). In another human study, SNPs in FTO were associated with reduced brain volume in healthy elderly (24). Combined, these findings establish FTO as an important regulator of fat mass in murine models, with a strong genetic link to human obesity.

**FTO Is an RNA-N⁶-methyl Adenine Demethylase**

At the time SNPs in the FTO gene were linked to obesity, the molecular function of FTO was unknown. Computational analysis revealed sequence similarity between FTO’s N-terminus and members of the family of non-heme Fe(II)- and 2-oxoglutarate (also known as α-ketoglutarate)-dependent dioxygenases (2OGX; ref. 25). 2OGX proteins are found in species from mammals to green algae and are characterized by similar overall protein fold and the positioning of highly conserved residues involved in substrate binding and catalysis, despite a large overall divergence in primary amino acid sequence (26, 27). In silico predictions of FTO’s protein structure closely resembled structures for the E. coli alpha-ketoglutarate–dependent dioxygenase AlkB, a member of the 2OGX family, despite a mere 17% sequence identity between the two (25). Studies using structural analysis and comparative analysis of enzymatic activities between FTO and other 2OGX proteins (25, 27–31) and enhanced UV cross-linking immunoprecipitation analysis altogether revealed that FTO enzymatic function is to demethylate N⁶-methyadenosine (m⁶A) residues in mRNA, small nuclear RNA (snRNA), small nuclear RNAs (snRNA) as well as of m1A in transfer RNA (Fig. 1A; refs. 32–34). m⁶A modification of mRNAs are associated with various functions, often context specific, through regulation of mRNA fate and translation. A more elaborate description relevant for this review is found in the next section as well as we would like to refer other published reviews (35, 36).

m⁶A modifications seem to accumulate near stop codons and in the 3’ UTR (untranslated region), are less frequent in the coding sequence, and are mostly depleted from the 5’ UTR (37–39). By genome-wide methylated RNA immunoprecipitation sequencing (MeRIP-seq) an m⁶A-methylation consensus site was identified as RRACH (H = A,C,U and R = A,G; ref. 38). Yet, some studies have not found enrichment of RRACH sequence motifs for FTO in CLIP analyses (40, 41), or only found them upon exogenous FTO overexpression (42). Furthermore, FTO has a relatively low substrate affinity for m⁶A-mRNA and the conversion of m⁶A into A is relatively slow (32, 34, 43). Noteworthy here is that the catalysis involves two steps of conversion of m⁶A into hydroxymethyladenosine (hm⁶A) and release of A and formaldehyde. FTO generates hm⁶A as the major product, followed by a release of A and formaldehyde over longer times, which is in contrast to the ALKBH5 2OGX-protein that catalyzes conversion into A with concomitant release of formaldehyde (44, 45). Transcriptome-wide MeRIP-seq studies using an antibody recognizing methylated m⁶A and cap-specific N⁶,2’-O-dimethyladenosine (m⁶Am) revealed m⁶Am as FTO responsive sites in addition to m⁶A (Fig. 1B; refs. 39, 43, 46, 47). Subsequent biochemical experiments revealed the catalytic efficiency (k₅₆A/K₅₆m) for FTO toward m⁶Am to be around 100-fold higher than for m⁶A (43). Depletion of FTO using siRNA from HEK293 cells resulted in increased methylation of cap-specific m⁶Am but not m⁶A as was analyzed by thin layer chromatography (43). The proposition that FTO does not affect m⁶A in mRNA to a biologically meaningful extent has however been questioned by others in the field, and clashes with a substantial amount of literature. For example, increased mRNA-m⁶A levels were found in brain tissue from Fto knockout mice (46), liver-specific FTO overexpression reduced mRNA-m⁶A levels in mice (48) and knockdown of FTO increased mRNA m⁶A levels in different acute myeloid leukemia (AML) cell lines (49, 50). Hep-G2 cells (51) and HeLa, HEK293T and JT3-L1 cells (34). The ability of FTO to demethylate cap-specific m⁶Am but also m⁶A was additionally confirmed by mass spectrometry analysis of purified mRNA (34, 52). Furthermore, the metabolite NADP was shown to directly bind FTO and stimulates its m⁶A demethylation activity to regulate adipogenesis (53). In addition, when FTO was overexpressed in rice and potato plants, m⁶A levels of mRNA and nonribosomal RNA and gene expression were significantly affected, resulting in grain yield and increased biomass (54). Remarkably, both rice and potato do not contain an FTO or an m⁶Am-methyltransferase homolog and have no m⁶Am at their mRNA 5’-cap (54). For interpretation of the data, several issues have to be considered. Some of the aforementioned studies rely on semiquantiative mRNA dot blotting or m⁶A mass spectrometry measurements to determine mRNA-m⁶A levels, both of which are highly sensitive to the purity of input mRNA and can be affected by for example ribosomal RNA if sample preparation is suboptimal (55).

Furthermore, studies that rely on MeRIP-seq might not be as reliable as initially thought. Reanalysis of MeRIP-based studies revealed considerable irreproducible variation and poor reproducibility across studies due to noise and difficulties in separating underlying expression changes from changes in m⁶A level (56). Additional more advanced techniques to quantify m⁶A modification at specific sites transcriptome-wide have recently been developed and rely on mazF endonuclease that cleaves unmethylated ACA motifs, but leaves “m⁶A-CA” motifs intact (MAZTER-seq/m⁶A-REF-seq; refs. 57, 58). When mRNA isolated from HEK293T cells was transcribed in vitro with FTO, m⁶A-REF-seq identified a significant decrease in m⁶A levels (58). However, when FTO was either knocked out or overexpressed in human embryonic stem cells or HEK293T cells, no impact on m⁶A levels was found in any condition with MAZTER-seq, in contrast to overexpression of ALKBH5 or knockout of METTL3 (57). It should be noted techniques based on mazF nuclease can by their nature only identify around 16% of mammalian m⁶A sites, due to the requirement for the ACA motif and ACA-site spacing constraints (57), and therefore the technique would greatly benefit from the development of other/less context specific m⁶A-sensitive nucleases.

Notably, in vitro studies identify cap-specific m⁶Am as the most efficiently converted substrate in biochemical assays with purified components (34, 43, 52). Nonetheless, in mRNAs from different mammalian cell lines and human and mouse tissues including cortex, liver, heart, skin, adipose, neurons and digestive tissues, m⁶A is on average 10- to 20-fold more abundant, with some AML cell lines having 30-fold more m⁶A over cap-specific m⁶Am (34, 50, 52, 59–62). This makes m⁶A in mRNAs the numerically more turned-over modification in vitro and might explain why many studies show an increase in transcriptome wide m⁶A levels upon FTO depletion, despite its preference for cap-specific m⁶Am demethylation over m⁶A in experiments with synthetic single-stranded RNA (ssRNA; refs. 34, 52). In addition, subcellular localization of FTO has been argued to differ between cell lines (34) or during the cell cycle (63) and thereby determine what substrates are most readily available (34).

Besides mRNAs, cap-associated m⁶Am modification of Sm-class snRNAs are increased by a substantial 10- to 20-fold upon FTO depletion in mouse liver and HEK293 cells and their demethylation by FTO was demonstrated in biochemical assays and cells (64). This was further supported by development of a novel method to quantify m⁶A modifications transcriptome-wide at specific sites (m⁶ACE-seq). m⁶ACE-seq makes use of cross-linking an m⁶A-antibody (also...
recognizing m^6Am) to m^6A, which blocks subsequent XRN1 exo-
nuclease digestion and thereby marks sites of m^6A modification (65).

Also in this study FTO depletion in HEK293T cells led to accumu-
lation of m^6Am at the first nucleotide of several small RNAs (sRNA),
snoRNAs, and Sm-class snRNAs, which was confirmed in biochem-
ical assays, without significantly affecting mRNA m^6A levels (65).

Hence m^6Am in snRNA was proposed to be the major target for

Figure 1.
Molecular structure of m^6A and m^6Am. A, Adenosine residues in RNA (left) can be methylated on the primary amine in position 6 to m^6A (right). The methyl group depicted in blue is added by a multiprotein methyltransferase writer complex consisting of the catalytic METTL3-METTL14 heterodimer and additional proteins such as Wilms' tumor 1-associating protein (WTAP), Vir like m^6A methyltransferase associated (VIRMA), RNA-binding motif protein 15 (RBM5), zinc finger CCCH-type containing 13 (ZC3H13), and Cbl proto-oncogene like 1 (CBLL1), also known as HAKAI). Demethylation of m^6A is catalyzed by two m^6A eraser proteins: FTO and AlkB homolog 5 (ALKBH5). B, Structure of a mammalian mRNA 7-methylguanosine cap linked to the first transcribed nucleotide via a 5'-5' triphosphate bridge. Nucleotides at position +1 relative to the cap are often methylated on the 2'-hydroxyl of the ribose ring (red); for adenosine this results in 2'-O-methyladenosine (Am). Am at the +1 position can be further methylated on the primary amine (blue) to N^6,2'-O-dimethylenosine (m^6Am).

Functional Consequences of m^6A Demethylation
To better understand the functional consequences of FTO's
demethylation activities one has to consider the effects of mRNA
(de)methylation. The (patho)physiologic effects of m^6A are mediated
through m^6A-binding proteins. The most well studied of these so-
called "m^6A-readers" are proteins that contain a YTS21-B homology
The study of YTH domain-containing proteins was intensified upon the discoveries that they bind to m^6^A (38) and YTHDF2, similar to the yeast homolog of YTHDC1, promotes degradation of its target mRNA in mammalian cells (69). Some proteins lacking a YTH domain have also been proposed to bind m^6^A directly, such as fragile X-mental retardation protein (FMRP), IGF 2 mRNA-binding proteins (IGF2BP) and subunits of eukaryotic initiation factor 3 (70–73). However, these proteins might interact with known m^6^A-binding proteins, as has been shown for FMRP and YTHDF2, and further studies are required to confirm direct m^6^A binding (74).

The most well-known downstream effect of m^6^A modification is regulation of mRNA stability (reviewed in ref. 75). In particular under stress conditions, YTHDF proteins and their bound m^6^A-modified mRNAs have been found to relocate to different phase separated compartments including P bodies (69, 76, 77), which are subcellular compartments where mRNAs are degraded through decapping and subsequent 5' to 3' exonucleolytic cleavage by XR1 (78). In addition, direct interaction of YTHDF2 with the CCR4-NOT (carbon catabolite repression—negative on TATA-less) deadenylase complex (independent of P body factors) has been observed (79), which is notable given mRNA deadenylation is known to precede P body formation. A plausible model could be that YTHDF2 promotes recruitment of the multi complex CCR4-NOT to initiate transcript degradation by deadenylase with subsequent translocation to P bodies for decapping and further XR1-mediated decay. Recently, an additional degradation pathway was proposed where YTHDF2 indirectly binds RNaseP/MRP to promote endonucleolytic cleavage of mRNA (80).

Nuclear export of mRNA is another process affected by m^6^A modification. Nuclear YTHDC1 was shown to interact with serine and arginine rich splicing factor (SSRF) 3 and thereby promote export of m^6^A-modified mRNAs via nuclear RNA export factor (NXF1) (81). NXF1 mediates the bulk of mRNA export, which requires interactions with the transcription export complex (TREX; reviewed in ref. 82). Components of the m^6^A methyltransferase complex were shown to interact with subunits of the TREX complex and thereby stabilize YTHDC1 binding to mRNAs, which in turn stimulated mRNA export (83). Separate from NXF1, specific m^6^A-modified mRNAs can be exported from the nucleus by exportin 1 (XPO1, also known as CRM1), the major mediator of protein nuclear export (72, 73, 84).

An m^6^A-binding motif is not always required for interaction of a protein with m^6^A-modified RNA. RNA structure mapping in RNAs from lymphoblastoid cells revealed that adjacent to RRACH m^6^A consensus sites with increased paired conformation the RNA adopts an increased ssRNA conformation (85). These properties allow m^6^A to execute a type of structural switch where a region of structured RNA is either stabilized or destabilized depending on the position of the modified residue. For example, in the MALAT1 IncRNA, m^6^A modification destabilizes a hairpin structure and thereby exposes the single-stranded binding domain for binding by the RNA-binding protein (RBP) heterogeneous nuclear ribonucleoprotein C (HNRNPC; refs. 86, 87). Global reduction of m^6^A on mRNAs reduces HNRNPC binding at m^6^A consensus sites transcriptome wide because of preferred duplex over ssRNA conformation (88). It will be important to assess the actual contribution of m^6^A over a potential structural rearrangement following mRNA (de)methylation to fully understand RNA–protein interactions. Finally, it should be noted m^6^A can also repel certain RBPs from their binding sites, for example, G3BP1 and LIN28A, although the physiologic relevance of this process remains unexplored (89, 90).

**Functional Consequences of m^6^Am Demethylation**

Although the m^6^Am modification has been known for quite some time, studying of how it affects mRNA fate was hampered due to a lack of identified writers, readers or erasers. The limited number of studies published so far delivered no consensus on how m^6^Am affects mRNA on a transcriptional or translational level. For example, m^6^Am was proposed to promote transcript stability by preventing decapping mRNA 2 (DCP2)-mediated mRNA decapping in HEK cells (43), where another study only observed increased transcript stability when in addition to m^6^Am, m^6^A was present (34).

With the identification of phosphorylated CTD interacting factor 1 (PCIF1) as a cap-specific methyltransferase (CAPAM), experimental assessment of this modification has become more accessible (91). Concerning mRNA stability, some studies show that PCIF1 knockout has no effect (91, 92) or only for mRNAs in the lower half of expression levels (93). Concerning mRNA translation, studies report an increase (91), no difference (93, 94) or a decrease (92) of translation efficiency upon m^6^Am modification. Potential mechanisms are still obscure, and m^6^Am modification did not affect binding of the eukaryotic initiation factor 4E (eIF4E) to the cap as the initiating step in mRNA translation (91, 95). Comparing transcript regulation in testes, brain, and spleen upon Pcf1 knockout in mice, the sets of upregulated and downregulated genes showed hardly any overlap between the tissues (94), questioning a substantial global effect of m^6^Am on transcript stability across tissues. Also, transcripts starting with an adenosine showed lower expression in Pcf1 knockout mice compared with wild-type mice only in the testes and not in the brain or spleen (94). In a study using colorectal cancer stem cells (CSC) cytoplasmic mRNA-m^6^Am was identified as FTO’s main target (96). Upon FTO knockdown in these cells, in vivo tumorigenicity was increased along with m^6^Am levels despite marginal effects on both transcriptome and translatome. Both of these phenotypes were completely reversed upon knockdown of PCIF1/CAPAM (96). A systematic study on the effect of the identity and methylation status of the first transcribed nucleotide on protein expression found large differences between the HeLa, MEF, and JAWS II (murine immature dendritic cell line) cells (95). m^6^Am compared with A as first nucleotide was shown to have no (MEF) a small (HeLa) or a large (JAWS II) effect on the expression of a reporter protein (95). Mapping of m^6^Am and m^6^A in different human and mouse tissues and human cell lines additionally showed extensive tissue specificity, and although not in all tissues, m^6^Am was in general negatively correlated with protein levels from the Human Proteome Map (97). Apart from mRNA regulation, m^6^Am modifications in mRNAs are deposited by METTL4 (98, 99) and mice with Mettl4 knockout display anatomic defects including craniofacial dysmorphism with incomplete penetrance (100).

**FTO in Cancer**

In the past decade, FTO emerged as a relevant factor in cancer development. Not surprising in light of the global effects of FTO m^6^A (m)-mRNA demethylation a picture emerges of pleiotropic and context-depending functions of FTO ranging from stabilization of
Oncogenic mRNAs, cancer-specific changes in cellular metabolism and tumor immune evasion. FTO expression and function has been linked to several cancer types (Supplementary Table S1). Compromising FTO function experimentally in cancer cells in most cases reduces oncogenic potential measured by cell proliferation and migration. Nonetheless, tumor-suppressive functions of FTO also have been reported. Often, the mechanisms by which FTO promotes or inhibits tumorigenesis are not clarified. Yet, in several cases, the involved oncogenic mechanisms have been revealed, which are discussed here. However, the data that suggest an oncogenic role of FTO are in contradiction with genome-wide screening approaches that have been used in recent years to determine gene essentiality and their effect on cancer growth, including over 1,000 cancer cell lines in the Cancer Dependency Map (DepMap; depmap.org). For example, in AML cell lines where growth effects were observed as discussed below (49, 50), no effects are seen in genome-wide screens (101, 102). These findings extend to the rest of the DepMap that overall indicate no requirement for FTO in cancer growth. Although of great value the DepMap dataset and related analysis have also important limitations. The DepMap addresses the genetic requirements for cancer cell proliferation and survival in cell culture, which greatly differs from in vivo tumorigenesis in patients or in animal models. To mention a few; use of (different) nutrient-enriched and growth factor–enriched growth media, a two-dimensional rather than a three-dimensional environment, and the presence of a physiologic microenvironment and associated functional immune system. Therefore, how FTO as a factor that does not appear to be essential for cancer cell growth in large-scale drug and genetic perturbation platforms is identified as an oncogene in so many other studies (Supplementary Table S1), is a matter of concern, yet needs to be scrutinized by further studies.

**Oncogenic Activities of FTO**

m^6^A modification is associated with a lower mRNA stability. FTO expression has been found elevated in several cancers and FTO-mediated m^6^A-mRNA demethylation is shown to stabilize mRNAs encoding oncogenic factors in several cases.

**AML**

A couple of publications describe various oncogenic roles of FTO in the development of AML. Analysis of microarray and transcriptome data revealed that FTO is highly expressed in AML with t(15;17)/PML-RARA, FLT3-ITD, and/or NPM1 mutations (49). Fusion proteins resulting from MLL rearrangements cause enhanced demethylation of CpG islands in the FTO locus, resulting in upregulation of FTO transcription. In an MLL-AF9 fusion model of leukemia induction [AF9 is a component of the super elongation complex (SEC; ref. 103)], the enhanced Fto expression accelerated leukemogenesis while Fto deficiency delays it. Taken together, the data from *in vitro* and *in vivo* experiments suggest that certain cancer cells become reliant on increased FTO expression. The high expression of FTO cause m^6^A demethylation and degradation of a subset of transcripts, among them are ankyrin repeat and SOCS box containing 2 (ASB2) and retinoic receptor alpha (RARA) as key regulators of AML. Which m^6^A reader proteins might be involved remained unclear as YTHDF1/2 knockdown did not affect the transcript stability (49). Subsequent studies discovered that for R-2-hydroxylglutarate (R-2HG)-sensitive subtypes of AML, R-2HG acts tumor suppressive by inhibiting FTO catalytic activity (50). R-2HG is produced by mutant isocitrate dehydrogenase 1 and 2 (IDH1/2), which are frequent somatic lesions found in glioma (~80%) and AML (10%–20%; refs. 104–106). Inhibition of FTO by R-2HG results in accumulation of m^6^A on MYC transcripts, decreasing its stability and downregulation of oncogenic MYC functions. In addition, R-2HG caused downregulation of FTO by suppression of CGAAT/enhancer binding protein alpha (C/EBPα) through accumulation of m^6^A on CEBPA-mRNAs and its destabilization. C/EBPα is one of the transcriptional activators of FTO. Here, data indicate that YTHDF2-dependent mechanisms are involved in the degradation of the mRNAs. Remarkably, cap-specific m^6^A levels were found to be very low in AML, challenging a pathophysiologic role for this modification in hematopoiesis and leukemia. In another study, metabolomics analysis showed that FTO inhibition by R-2HG attenuates glycolysis flux in AML cells through m^6^A accumulation on lactate dehydrogenase B (LDHB) and phosphofructokinase platelet (PFKP) transcripts and degradation through binding to YTHDF2 (107). Combined, these data suggest AML cells with high FTO expression rely on FTO-mediated m^6^A demethylation to degrade transcripts of tumor suppressors and promote stability of oncogenes like MYC (50) and metabolic regulators such as PFKP and LDHB (107), contributing to cell proliferation and a block in differentiation. AML subtypes with either low endogenous FTO or IDH1 mutation (R-2HG–sensitive subtypes) on the other hand have found other ways to activate protumorigenic pathways and remain refractory to FTO inhibition.

**Glioma**

In the same study addressing FTO inhibition in R-2HG–sensitive AML, R-2HG also was found to act tumor suppressive by inhibiting FTO in glioma cells through accumulation of m^6^A on MYC transcripts and downregulation of MYC oncogenic functions. Glioblastomas are tumors arising from glial cells in the brain or spinal cord, among which, glioblastoma is the most well-known and deadly subtype (108). These heterogeneous tumors contain glioblastoma stem cells (GSC) that display increased radiotherapy and chemotherapy resistance (108). In primary GSC cultures, differentiation of GSCs to neurons and astrocytes was found accompanied by a substantial increase in accumulation of mRNA-m^6^A (109). Reduction of m^6^A levels by knockdown of methyltransferase (METTTL)3 or METTTL14 led to increased maintenance of stem cell properties, while overexpression of METTTL3 had an opposite effect (109). Increasing m^6^A levels in primary GSCs by treatment with the FTO inhibitor MA2 (ethyl ester form of meclofenamic acid) furthermore reduced tumor growth in xenograft experiments (109). Among other m^6^As, ADAM19 mRNA encoding a transmembrane metalloprotease that is overexpressed in glioblastoma and promotes cell growth and invasion was found to be destabilized by accumulation of m^6^A upon FTO inhibition (109). In an unrelated study, a negative feedback loop of MYC and MAX interactor 1 (MXI1) mediated by miR-155/23a cluster was identified that affected proliferation of glioma cells (110). MXI1 inhibits the transcriptional activity of MYC by competitive binding to MAX, which is the heterodimerization partner of MYC in an active transcription-promoting complex (111). The study shows that FTO-dependent upregulation of MYC results in miR-155/23a–mediated suppression of MXI1 (110). In addition, treatment with FTO-inhibitor MA2 decreased MYC in glioma cells and led to reduced proliferation (110). Patients with glioma are often treated with the chemotherapeutic temozolomide, which has been shown to reduce MYC levels through increased activity TAp63, a repressor of MYC transcription (112). Treatment of glioma cell lines with both MA2 and temozolomide had a stronger effect on MYC and cell proliferation than either single treatment, although effects were additive at best (110).
Lung cancer

Around 80% of lung cancers can be classified as non–small cell lung cancer (NSCLC), of which, lung adenocarcinoma (LUAD) and lung squamous cell carcinoma (LUSC) are two major subtypes (113). The role of FTO in NSCLC remains poorly understood and while some find poorer patient survival in high FTO-expressing LUSC but not in LUAD (113), others found no differences in LUSC and rather an increase in survival in high FTO LUAD (114). Generally, however, FTO appears to promote growth of multiple NSCLC cell lines from both subtypes. Knockdown of FTO impaired proliferation and migration in various LUSC cell lines while overexpression of FTO but not a catalytically inactive FTO mutant led to opposite phenotypes (113). Mechanistically, FTO was shown to demethylate m6A and thereby stabilize myeloid zinc finger 1 (MZF1)-mRNA (113), a factor shown to induce MYC transcription and thereby promote tumorigenesis in LUAD (115). Although this link was not investigated explicitly in the aforementioned study, ectopic MZF1 expression was sufficient to restore cell growth defects upon FTO knockdown (113). The positive effect of FTO on NSCLC growth was also found in mouse xenograft experiments using A549 human NSCLC cells with FTO knockdown, which exhibited reduced tumor growth compared with control cells (116). The study shows that ubiquitin-specific peptidase 7 (USP7)-mRNA becomes hypermethylated upon FTO knockdown with a concomitant decrease in its transcript levels (116). In addition, ectopic USP7 expression could rescue the growth defect induced by FTO knockdown by yet unclear downstream molecular mechanisms (116). It should be noted that in above-mentioned studies targeted either on profiling of FTO targets in AML or on previously described relevance to lung cancer, precluding the unbiased identification of additional and/or lung cancer specific targets of FTO.

Breast cancer

Expression of the estrogen receptor [ER− (negative)/ER+ (positive)], the progesterone receptor (PR−/+) and amplification of the HER2 (HER2+) serve as markers for the different subtypes of breast cancer, including triple-negative breast cancer (TNBC; ER−, PR−, HER2−), luminal A (ER+, PR−/+, HER2−), luminal B (ER+, PR+, ER−/+, HER2−, HER2+, ER+, PR+, HER2+; ref. 117). In luminal A and TNBC cell lines, FTO knockdown both in vitro and in vivo increased apoptosis, reduced migratory capacity and reduced cell growth (118). The study revealed FTO demethylates the mRNA encoding BCL2 interacting protein (BNIP)3, a proapoptotic BCL2-family member, and BNIP3 protein levels are increased upon FTO knockdown through an unidentified, YTHDF2-independent mechanism (118). In a different study, FTO was shown to promote migration and invasion but not proliferation in HER2+ cells (119). Focusing on miRNA regulation, increased miR181b-3p levels were observed upon FTO knockdown or treatment with FTO inhibitor MA2 and the ADP ribosylation factor like GTPase (ARL)5B was identified as a major target of miRNA miR-181b-3p (119). ARL5B is an ADP ribosylation factor-like (ARL) family member that regulates lysosome transport and has been implicated in the regulation of pancreatic cancer cell migration (120). Indeed, ARL5B modulation mimics or restores the FTO-induced migration phenotype in HER2+ breast cancer cell lines (119). How miR-181b-3p levels are regulated by FTO remains unclear; although a substantial number of miRNAs was shown to contain m6A, miR-181b-3p was not among them and FTO deficiency did not affect its transcript level in HEK cells (121).

FTO Regulation of Immune Surveillance

Tumor neantigens generated by cancer cells provoke spontaneous immune response as well as they offer targets for immunotherapies. However, a sufficient and lasting anticancer immune response leading to complete tumor elimination is rare. Cancer cells may escape immune surveillance by mounting mechanisms of immune evasion. Evidence is mounting that the epitranscriptome is involved in immune evasion. It was shown that transcripts encoding lysosomal proteases accumulate m6A and that subsequent binding of YTHDF1 enhances their translation in dendritic cells (122). The enhanced lysosomal degradation of engulfed neoantigens suppresses the cross-presentation of neoantigens by the dendritic cells to CD8⁺ T cells as a mechanism of immune evasion. Consequently, Ythdf1 deficiency in mice results in an elevated antigen-specific CD8⁺ T-cell antitumor response (122). It should however be noted here the effect of YTHDF1–3 proteins on translation has been disputed and a redundant role in promoting degradation of m6A-modified transcripts has been proposed (123, 124).

In a different study, bioinformatic analysis revealed that FTO exhibit the most significant negative association with cytotoxic T lymphocyte score in melanomas and other cancer types. This led to the suggestion that FTO acts as an immune inhibitory factor for tumor-infiltrating function of CD8⁺ T cells (125). The study concludes that FTO promotes tumor immune evasion as an epitranscriptomic regulator of glucose metabolism through m6A demethylation of transcripts encoding the transcription factors c-Jun, JunB, and C/EBPβ. The resulting enhanced expression of these factors transcriptionally activates downstream glycolytic genes. High consumption of glucose by tumor cells is known to restrict T cells metabolically. Fto knockdown in melanoma (B16-OVA) and lung cancer cell lines inoculated in immune proficient mice resulted in smaller tumor sizes while in immunodeficient mice inoculation showed no difference between wild-type and Fto-knockdown cells. In addition, a 2-fold increase in infiltration of tetramer CD8⁺ T cells was observed in Fto-knockdown versus wild-type tumors. Transcriptome analysis showed enhanced expression of cytokines and cytotoxic molecules upon Fto knockdown in T cells cocultured with Fto-knockdown tumor cells show a higher cytotoxic state. Moreover, similar to Fto knockdown, treatment with the FTO-inhibitor Dac51 resulted in accumulation of m6A and reduction in expression of c-Jun, JunB, and C/EBPβ as well as an inhibition of glycolysis in patient-derived organoids. Further in vivo studies in mice showed that treatment with Dac51 increases T-cell infiltration of B16-OVA melanoma cells similarly to Fto-knockdown tumor cells. Moreover, Dac51 treatment was shown to synergize with anti-PD-L1 immune checkpoint blockade.

Tumor-Suppressive Activities of FTO

Notwithstanding the well-described oncogenic roles of FTO as discussed above several studies present strong cases of FTO in a tumor-suppressive role.

Ovarian cancer

In ovarian tumors and CSCs, FTO expression was found suppressed and associated with increased global m6A RNA levels (126). Moreover, FTO overexpression inhibited CSC proliferation and tumor initiation capacity in cell culture and xenograft mouse models. Furthermore, expression of a mutant FTO lacking demethylase function was ineffective in altering stemness phenotypes. Transcriptome-wide RNA sequencing and m6A mapping identified over 700 significantly deregulated transcripts as a consequence of m6A modifications with pathways
related to mRNA transcription and splicing, stem cell signaling and DNA repair. As the most prominent FTO targets two phosphodiesterase genes (PDE4B and PDE1C) were selected, which regulate cAMP signaling as a key pathway in maintaining stemness. The authors claim to provide the first solid evidence that FTO function acts tumor-suppressive in certain cancer types or stages (126).

Colorectal cancer
In a recent study involving CSCs, the cap-specific m^6^A modification was identified as an epitranscriptomic mark for maintaining stem-cell phenotype of colon cancer cells (96). Knockdown of FTO in various experimental settings promotes CSC phenotypes, including initiation of tumor formation and resistance to chemotherapeutic drugs in mouse xenograft models. Analyses of tumor arrays revealed that although global FTO expression does not change in different colorectal stages, the subcellular localization of FTO changes from strictly nuclear to cytoplasmic in metastatic submucosal invasion. This may be related to altered function of FTO. However, the mechanisms involved in suppressing FTO functions is CSC remain obscure also because no large differences were found in transcriptome analysis and a translation assay by polyosome mapping (96).

Hepatocellular carcinoma
Studies using hepatocellular carcinoma (HCC) cell lines initially suggested FTO as a protumorigenic factor. In mouse xenograft experiments using HepG2 cells, FTO deficiency led to a significantly lower tumor volume, possibly through demethylation and stabilization of pyruvate kinase M2 (PKM2) mRNA (127). In addition, knockdown of FTO in HepG2 cells reduced the expression of lipogenesis genes, resulting in deficiency of lipid accumulation and induction of cellular apoptosis, suggesting a role for FTO in cancer cell survival and hepatocellular metabolism (128). However, in a recent study using a mouse model with hepatic Fto deficiency (FTOL-KO), only minor systemic metabolic changes were found (129). Long-term treatment with diethylthiosamine (DEN), which provokes HCC, produced fewer and smaller tumors in the Fto-proficient compared with Fto-deficient mice (129). The study proposes the involvement of Cullin 4A (Cul4a) as a member of E3 ubiquitin ligase complexes that regulate cell-cycle progression and DNA replication. In wild-type livers, short-term high-dose DEN treatment resulted in an increase in FTO expression and a decrease in Cul4a protein levels, whereas in Fto-deficient livers Cul4a expression is maintained (129). Although in Fto-deficient livers Cul4a-mRNA showed an increase in m^6^A modification compared with wild-type livers upon DEN treatment, transcript levels remained unaffected. The authors therefore proposed that sustained translation of hypermethylated Cul4a induced the expression of its target genes such as cyclin E1 to promote HCC development (129).

Epithelial–Mesenchymal Transition–Mediated Tumor Progression
By analyzing multiple datasets on cancer, Jeschke and colleagues noted that FTO is downregulated in various epithelial tumors (132). Experiments in cell culture showed that FTO depletion in several breast and prostate cancer cell lines increased clonogenicity, invasiveness and migratory potential. Moreover in vivo xenotransplant experiments in mice showed increased tumor progression after FTO depletion as well as by inhibition of FTO with meclofenamic acid. Gene expression analysis revealed an activated epithelial–mesenchymal transition (EMT) program in FTO knockdown cancer cells and xenograft tumors, while in patient samples lower expression of FTO was significantly correlated with a high EMT signature. In particular, many mRNAs encoding factors in the Wingless (Wnt)-β-catenin signaling pathway showed increased m^6^A deposition. In breast cancer cells, knockdown of FTO resulted in stabilization of the Wnt-ligand Wnt family member 34 (WNT3A) while at the same time destabilization of transcripts encoding casein kinase 1 subunits and 2 involved in β-catenin destruction, illustrating the diverse effects of m^6^A modifications. Because FTO-low type epithelial tumors rely on Wnt signaling to activate EMT, treatment with the Wnt inhibitor icrT3 reduced tumor clonogenicity and invasiveness of tumor cells specifically after FTO knockdown or tumor progression in xenotransplant mouse models. Note that the conclusions from the study described in this paragraph (132) contradict with the studies described above that infer an oncogenic role of FTO in breast cancer (118, 119).

Therapeutic Implications
Several small-molecule inhibitors for FTO with varying potency and specificity have been developed since its crystal structure was solved in 2010 (31). The first identified inhibitor, rhein, has a broad inhibitory effect on AlkB family proteins by competitive binding at the catalytic active site preventing binding of m^6^A-mRNA (133). Despite the weak substrate specificity, allograft experiments in mice using 4T1 breast cancer cells demonstrated rhein treatment can reduce tumor growth albeit less efficiently than Fto knockdown (118).

As a 2OGX/AlkB family member, FTO requires bound Fe(II) and 2-oxoglutarate as cofactors for oxidative demethylation of m^6^A (134). Many inhibitors for 2OGX proteins chelate the Fe(II) moiety, thereby preventing 2-OG and substrate binding, while others chemically mimic 2-OG (135). In silico docking and molecular modeling studies have been used to define a molecular blueprint for potential inhibitors so to further optimize low-affinity inhibitors. For example, compound 12 mimics 2-OG and protrudes into the nucleotide-binding site and was the first reported inhibitor with specificity for FTO over other AlkB family proteins (around 30-fold with an IC_{50}=1 μmol/L, ref. 136). The cell permeable ethyl ester of compound 12 was partially taken up and active in HeLa cells, resulting in a dose-dependent increase of m^6^A in mRNA (136). Meclofenamic acid (MA) is an FDA-approved NSAID that was also found to inhibit FTO, but not ALKBH5, with an IC_{50}=0.7 μmol/L (137). Although this affinity is quite low, at high (80–120 μmol/L) concentrations the ethyl ester of MA (MA2) can increase mRNA m^6^A levels in HeLa cells through FTO inhibition (137). MA2 treatment of glioblastoma stem cell lines at similar or slightly lower concentrations increased m^6^A levels and inhibited cell growth both in vitro and in vivo xenograft experiments (109). Although MA2 had to be delivered intratumorally, this provided another example of how small-molecule targeting of FTO can reduce cancer growth. On the basis of the crystal structure of the FTO/MA complex, MA was optimized in silico for improved biological activity and cellular uptake while retaining target specificity, resulting in FB23 and FB23-2 (138). FB23-2...
showed an IC$_{50}$ of 2.6 µmol/L toward FTO while resulting in moderate uptake in different AML cell lines, increased mRNA m$^6$A levels, growth inhibition, and a relatively high specificity exemplified by a large overlap in transcriptomes of cells treated with FB23-2 compared with genetic FTO depletion (138). FB23-2 was well tolerated in vivo and could effectively reduce growth of both patient-derived AML cells and AML cell lines in mouse xenograft experiments upon intraperitoneal injection (138). Next, FB23 and FB23-2 were further optimized, resulting in Dac51 used to study the role of FTO in immune surveillance as discussed in the paragraph above (125). Because MA-derived inhibitors showed specificity for FTO over ALKBH5, a molecular docking approach was used to predict the structure of compounds binding in the same pocket with preferable membrane permeability, hydrophilicity, and molecular weight (139). Using this approach FTO-02 and FTO-04 (IC$_{50}$ = 2.2 and 3.4 µmol/L, respectively) were identified as novel FTO inhibitors (139). Although these compounds are yet to be tested in vivo, FTO-04 showed high selectivity for FTO over ALKBH5 and prevented formation of neurostem cells from patient-derived GSCs without affecting healthy controls, concomitant with an increase in mRNA m$^6$A levels in treated cells (139).

In a similar experimental approach, The NCI developmental therapeutics program library of around 260,000 compounds were virtually docked to FTO’s catalytic center (140). Two compounds, CS1 (bisantrane) and CS2 (brequinar), were identified that inhibited AML cell growth with IC$_{50}$ concentrations around 100 nmol/L in multiple cell lines; considerably lower than the concentration of FB23-2 needed in the same cell lines (140). Both compounds substantially prolonged survival by inhibiting AML progression in patient-derived xenograft (PDX) models of AML, and at the same concentration CS1/2 were more effective than FB23-2 (140). As both compounds were also tolerated in mice at four times the concentration used for treatment in the PDX model (140), they are good candidates for bioavailability and efficacy studies for different cancer types that would benefit from FTO inhibition.

Yet another promising starting point for FTO inhibition might be entacapone, an FDA-approved inhibitor of catechol-O-methyltransferase canonically used as adjunctive treatment for Parkinson disease. In an effort to obtain FTO inhibitors for treatment of metabolic disease, FDA-approved drugs were screened for virtual molecular docking to FTO (51). On the basis of this screening approach, entacapone was identified and biochemically verified to bind FTO and inhibit its m$^6$A demethylase activity in vitro (IC$_{50}$ = 3.5 µmol/L) and in mouse liver upon addition of entacapone to the diet (51). Although the chemical structure is distinct from MA-based inhibitors, entacapone displayed good specificity for FTO over other dioxygenases and did not affect DNA methylation or histone methylation patterns in Hep-G2 cells (51).

Taken together, structure-based design and virtual screening approaches have led to the development of a series of small-molecule FTO inhibitors. With multiple candidates showing biological activity against and specificity for FTO, future efforts towards improving bioavailability and pharmacokinetics might yield valuable FTO inhibitors for treatment of multiple diseases including cancer.

**Discussion and Perspectives**

It has become clear that m$^6$A and cap-specific m$^6$Am modifications on mRNAs and mRNA demethylation by FTO play important roles in posttranscriptional regulation of gene expression. The presence of m$^6$A can affect processes such as mRNA splicing, nuclear export, translation and stability by facilitating or preventing interaction with various RBPs. m$^6$A-induced binding by RBPs can either involve dedicated m$^6$A-recognizing proteins or be induced by changes in mRNA folding facilitating binding of other RBPs. Cellular mRNA is in constant and complex interaction with various RBPs throughout every stage of its life cycle, and it is therefore not surprising that perturbation of the demethylase function of FTO affects many steps in the mRNA processing and translation. The contribution of FTO to mRNA m$^6$A demethylation is however not uncontroversial and pinpointing downstream targets and reliably quantifying their m$^6$A level has often been difficult. The advent of transcriptome-wide quantitative methods to detect m$^6$A such as MAZTER-seq (57) and m$^6$ACE-seq (65) will improve the assessment of m$^6$A RNA deposition in response to various cellular perturbations, and proposed alterations in m$^6$A level should be confirmed by these transcript-specific quantitative methods (141, 142). In contrast to m$^6$A, the function of cap-specific m$^6$Am in regulating mRNA fate and cellular (patho)physiology is still poorly understood. The recent discoveries of PCIF1/CAPAM as a m$^6$Am writer (91) and FTO as the factor adjusting m$^6$Am levels required for inducing a CSC phenotype in colorectal cancer (96) is likely the first of further discoveries of m$^6$Am functionality. Similarly, the identification of snRNA m$^6$Am demethylation by FTO (64) provides a new avenue of research to be explored, which is facilitated by discovery of the snRNA m$^6$Am methyltransferase METTL4 (98, 99). Because the methyltransferases for m$^6$A in mRNA and m$^6$Am in mRNA as well as in snRNA have now been identified, this allows validation of m$^6$A or m$^6$Am as FTO substrates by investigating the phenotypes of cells lacking the corresponding methyltransferase.

Subcellular compartmentalization of FTO may be different between cell types (34) and might be dynamically regulated in the cell cycle (63) or by external stimuli (143), which has to be taken into account as it directly influences substrate availability. In patients with colorectal cancer, for example, FTO was mainly nuclear in precursor lesions and healthy adjacent tissue but also found in the cytoplasm of more advanced stage colorectal cancer (96). Yet, care should be taken relying solely on antibody staining because some have been shown to cause artifactual staining and therefore knockout strategies should be used for validation (65). How FTO recognizes its targets remains another open question in the field because RNA-interaction studies identified no clear consensus binding site. In addition, our knowledge of FTO binding partners and how they might affect demethylation activity or substrate specificity is limited (144). In general, increased expression of FTO leads to reduced mRNA m$^6$A content, which in turn stabilizes transcripts by preventing degradation through canonical RNA degradation pathways facilitated by m$^6$A-binding proteins such as YTHDF2. However, one cannot rely on these general pathways to hold up in all tissue types or disease states. This is exemplified in AML where FTO-mediated demethylation of m$^6$A in 5’ terminal and internal exons led to stabilization of MYC-mRNA, while METTL14 silencing reduced m$^6$A in the MYC 3’ terminal exon, leading to transcript destabilization (50, 145). In addition, both oncogenic and tumor suppressive roles for FTO in the same tissue have been described in, for example, liver, kidney, and colorectal cancer. As the control of mRNA fate is largely dictated by interaction with different RBPs, identifying the landscape of RNA-protein interactions upon FTO modulation will undoubtedly shed light on the different roles of FTO in different cancer (sub) types. Expression patterns of RBPs differ between tissues and might
explain why some of FTO’s effects are common among different cell types while other are restricted to a specific (potentially malignant) state of a particular tissue. Protumorigenic roles of FTO are now apparent in different cancer subtypes and targeted pharmacologic inhibition of FTO led to reduced tumor growth in preclinical models of AML and glioma in vivo (49, 50, 107). In combination, a better understanding of the molecular processes downstream of FTO-mediated RNA demethylation and continued development of targeted therapies aimed at inhibiting FTO might prove a powerful strategy to combat certain cancers.

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