m^6^A mRNA demethylase FTO regulates melanoma tumorigenicity and response to anti-PD-1 blockade

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Melanoma is one of the most deadly and therapy-resistant cancers. Here we show that N^6^-methyladenosine (m^6^A) mRNA demethylation by fat mass and obesity-associated protein (FTO) increases melanoma growth and decreases response to anti-PD-1 blockade immunotherapy. FTO level is increased in human melanoma and enhances melanoma tumorigenesis in mice. FTO is induced by metabolic starvation stress through the autophagy and NF-κB pathway. Knockdown of FTO increases m^6^A methylation in the critical protumorigenic melanoma cell-intrinsic genes including PD-1 (PDCD1), CXCR4, and SOX10, leading to increased RNA decay through the m^6^A reader YTHDF2. Knockdown of FTO sensitizes melanoma cells to interferon gamma (IFNγ) and sensitizes melanoma to anti-PD-1 treatment in mice, depending on adaptive immunity. Our findings demonstrate a crucial role of FTO as an m^6^A demethylase in promoting melanoma tumorigenesis and anti-PD-1 resistance, and suggest that the combination of FTO inhibition with anti-PD-1 blockade may reduce the resistance to immunotherapy in melanoma.
Melanoma is one of the most deadly and therapy-resistant human cancers in the United States. The incidence of melanoma continues to rise at an alarming rate each year, faster than any of the other common cancers. In the past decades, tremendous progress has been made in elucidating the mechanism of melanoma development at the molecular, cellular and organismal levels. Both genetic, including mutations in oncogenes and tumor suppressor genes, and epigenetic mechanisms, including DNA methylation, microRNAs, and other non-coding RNAs, have been demonstrated to play critical roles in melanoma pathogenesis. In particular, the discoveries of BRAF mutations and their functions in most melanomas have led to the development of molecular therapy targeting the mutant BRAF. However, the majority of the patients treated with an inhibitor of mutant BRAF eventually suffer continuous disease progression. Recent breakthroughs in immunotherapy, including anti-PD-1 checkpoint blockade therapy, have benefitted a growing number of melanoma patients. Still, more than half of these patients do not show a durable response to immunotherapy. Multiple mechanisms, such as driver mutations, epigenetic mechanisms, tumor plasticity, and immunosuppression, may cooperate to maintain the melanoma phenotype and mediate resistance to therapies, including immunotherapy, which makes targeting a single pathway less effective. Our understanding of the molecular mechanisms for melanoma development and therapeutic response is still limited.

An emerging molecular mechanism regulating gene expression at the post-transcription level is N6-methyladenosine (m6A) RNA methylation. m6A RNA methylation is the most prominent chemical modification found in messenger RNA (mRNA) and non-coding RNA in eukaryotic cells. Recently, the first m6A demethylase FTO was discovered, suggesting that m6A RNA methylation is reversible and dynamic and may have crucial physiological and pathological functions. Subsequently, two studies independently mapped the transcriptome-wide m6A distribution. These studies also suggest that, similar to the reversible epigenetic modifications to DNA and histone, post-transcriptional methylation of adenosines in RNA provides an epitranscriptomic layer of regulation that may control mRNA fate and gene expression.

In the past several years, compelling evidence has demonstrated that m6A methylation plays critical roles in controlling RNA metabolism and function in physiological processes and stress response. m6A modification regulates development, stem cell homeostasis, response to stresses such as heat-shock and genotoxic damage, and control of the circadian clock. m6A modification regulates RNA fate and such functions as mRNA stability, nuclear processing, transport, localization, translation, primary microRNA processing, and RNA-protein interactions. m6A modulators have been shown to play important roles in several cancers by regulating cancer-type-specific target genes and functions, suggesting the unique regulatory and functional significance of the intrinsic m6A mRNA modifications and their regulators in different cancers. Recent work has also suggested that the genetic variations of FTO are associated with increased melanoma risk in humans. The association of these FTO variations in exon 8 with melanoma risk seems to be independent of body mass index (BMI). As the first m6A demethylase identified, FTO has been shown to act as an oncogenic factor in leukemia and glioblastoma. However, the role of FTO as an m6A eraser in melanoma pathogenesis and response to anti-melanoma therapies remains poorly understood.

Here we report that the m6A demethylase FTO regulates melanoma growth and mediates melanoma resistance to anti-PD-1 antibody in vitro and in vivo. Our results identify a unique FTO-mediated and m6A-mediated mechanism in promoting melanoma tumorigenesis and resistance to anti-PD-1 blockade, and suggest that the combination of FTO inhibition with anti-PD-1 blockade may reduce resistance and improve the anti-melanoma response.

Results

FTO as a protumorigenic factor in melanoma. To determine the role of FTO in human melanoma, we first analyzed FTO protein levels specifically in the melanocytes of normal human skin and malignant melanoma samples using immunofluorescence. We used MART1 immunofluorescence co-staining to specifically identify melanocytes in normal skin and melanoma cells (Fig. 1a). FTO (Red) was low in epidermal melanocytes in normal skin (n = 16), while it was significantly increased in all diagnoses of melanoma, including metastasis (n = 64), primary melanoma (n = 36), and melanoma at stages 1–IV (n = 65) (Fig. 1a, b). There was no significant difference in FTO levels between different diagnosis groups of melanomas except between Stage I and IV. Next, we determined the FTO protein levels in normal human epidermal melanocytes (NHEM) and a panel of melanoma cell lines. FTO remained unchanged in WM35 (BRAF mutation, Human) cells (Fig. 1c). However, it was up-regulated in six of the tested cell lines, particularly in Mel624 (BRAF mutation, Human) and B16F10 (BRAF wild-type, Mouse) cells, two aggressive model melanoma cell lines, as well as WM115 (BRAF mutation, Human), WM793 (BRAF mutation, Human), WM3670 (BRAF and NRAS mutation, Human), and CHL-1 (BRAF and NRAS wild-type, Human) (Fig. 1c). These results demonstrate that FTO is upregulated in human melanoma samples and multiple melanoma cell lines, suggesting a protumorigenic role of FTO in melanoma development.

To determine the biological function of FTO in melanoma cells, we performed both gain- and loss-of-function studies in melanoma cell lines (Fig. 2a). Knockdown of FTO in FTO-high Mel624, CHL-1, and B16F10 cells decreased cell growth/proliferation, migration, invasion, and cell viability in cells placed in suspension (Fig. 2b–e, and Supplementary Fig. 1a–n). Furthermore, forced overexpression of FTO in WM35 with an FTO protein level similar to NHEM (Fig. 1c). To elucidate the role of FTO in melanoma pathogenesis in vivo, we performed xenograft and syngeneic melanoma tumorigenesis assays. Knockdown of FTO markedly reduced tumor volume and tumor growth in both Mel624 and CHL-1 cells in immunocompromised nude mice, and in B16F10 cells in immunocompetent C57BL/6 mice (Fig. 2f, g, and Supplementary Fig. 2a–j). Furthermore, forced overexpression of FTO in WM35 induced tumor growth in immunocompromised nude mice (Fig. 2f, g, and Supplementary Fig. 2d). These findings suggest that FTO is required for the malignant traits of melanoma cells in vitro and plays a protumorigenic role in melanoma tumor growth in vivo.

Role of m6A in FTO function in melanoma cells. To examine the role of FTO in m6A regulation, we investigated the effect of FTO knockdown on m6A levels in mRNA. Knockdown of FTO increased m6A levels in both total RNA and purified mRNA in both Mel624 and B16F10 cells (Fig. 3a–d). To determine the role of m6A in melanoma cell function, we performed gain-of-function tests of the m6A writers METTL3 (Methyltransferase Like 3) and METTL14 (Methyltransferase Like 14). Forced overexpression of METTL3 and METTL14 increased the
total m6A level and decreased cell growth/proliferation (Fig. 3e–g), migration, and invasion (Fig. 3h, i, and Supplementary Fig. 3a, b). Furthermore, knockdown of METTL3 and METTL14 prevented the effect of FTO knockdown on cell growth/proliferation, migration, invasion, and cell viability in melanoma cells in suspension (Fig. 3j–m, and Supplementary Figs. 3e–j, 4a–f). The opposite was also true. Overexpression of METTL3 and METTL14 prevented the effect of FTO overexpression on these malignant traits (Supplementary Fig. 4g–l).

These findings indicate that m6A modifications in RNA reduce melanoma cell proliferation and cell viability, suggesting a tumor-suppressive role of m6A in melanoma.

Identification of potential target genes of FTO in melanoma.
We next used both candidate and unbiased screening approaches to determine the potential mRNA targets of FTO in its melanoma-promoting function (Fig. 4a). First, we started with a candidate approach, focusing on the genes that are known to be associated with melanoma growth and in particular response to immunotherapy, including known melanoma-promoting genes PD-1 (PDCD1)35, PD-L1 (CD274)11,36, and CD4737–39. All three genes were expressed in Mel624 cells. We found that serum starvation induced the expression of FTO, PD-1 (PDCD1), and CD47, but not PD-L1 (CD274) (Supplementary Fig. 5a). FTO knockdown significantly decreased PD-1 (PDCD1) expression, but had no effect on PD-L1 (CD274) or CD47 (Fig. 4b). These data suggested that PD-1 is a potential FTO target gene in melanoma. Second, we performed microarray analysis of gene expression in MEL624 cells with or without FTO knockdown to determine transcriptome-wide changes upon FTO knockdown. We identified 106 genes that were significantly altered by FTO knockdown (Fig. 4a). Third, we used m6A-seq in combination with RNA-seq to identify more than 1000 potential transcriptome-wide m6A-modified gene targets for FTO (Fig. 4a), including the melanoma-promoting genes CXCR4 and SOX10. m6A peaks detected in poly(A)+-enriched RNAs showed reproducible patterns of methylation (Supplementary Fig. 5b). FTO knockdown increased m6A enrichment in 5'UTR and 3'UTR, while it had little effect on the CDS region (Fig. 4c), and altered total peak distribution and unique peak distribution (Fig. 4d).
FTO knockdown cells also showed an increased number of common m6A genes (Fig. 4c) and fewer m6A peaks (Supplementary Fig. 5c), likely due to the degradation of m6A-modified mRNAs. Sequence analysis of m6A peaks showed the previously identified m6A target sites (GGACU)17 (Fig. 4f). However, PD-1 (PDCD1) was undetectable in either microarray or RNA-seq analysis (Supplementary Fig. 5g–i), possibly due to the limitations of these assays. m6A IP seq analysis showed that FTO knockdown increased the m6A levels in the transcripts of CCR4 and SOX10 (Supplementary Fig. 5e, f). Next, we validated a set of downregulated genes (PD-1 (PDCD1), CXCR4, SOX10, ANGPTL2, CTSV, FCMR, NOP16, and RAB40), and upregulated genes (CDKN1A) upon FTO knockdown (Fig. 4g), and Supplementary Fig. 5k. Among the genes identified above, PD-1 has been shown to be

Identification of functionally critical target genes of FTO.

Among the genes identified above, PD-1 has been shown to be
expressed in some melanoma cells and act as an intrinsic pro-tumorigenic factor to promote melanoma tumor growth. It activates the downstream of mTOR signaling, demonstrating a critical intrinsic role of PD-1 in melanoma. In addition, PD-1 has been successfully targeted in immune cells for cancer immunotherapy. Therefore understanding the regulation of PD-1 expression of melanoma cells in FTO knockdown cells increased phosphorylation of p70s6K, a substrate of mTOR activation, whereas demethylase-inactive mutants had no effect (Supplementary Fig. 6e–h). These findings indicate that FTO is responsible for promoting the expression of melanoma-intrinsic PD-1, CXCR4, and SOX10.

Next we investigated whether these melanoma-promoting genes are responsible for FTO’s function. Forced overexpression of PD-1 (PDCD1) in FTO knockdown cells increased phosphorylation of p70s6K (Fig. 5c), accompanied by significantly increased cell growth/proliferation (Fig. 5d) and migration (Fig. 5e, and Supplementary Fig. 7a). Similarly, forced overexpression of CXCR4 and SOX10 in FTO knockdown cells significantly increased cell growth/proliferation and migration (Fig. 5f–i, and Supplementary Fig. 7b–e). These results demonstrate that PD-1, CXCR4, and SOX10 are critical downstream
targets of FTO responsible for its function in melanoma. Indeed, all melanoma cell lines showed increased PD-1 protein levels as compared with normal melanocytes (Supplementary Fig. 7f). PD-1 (PDCD1) mRNA expression is positively associated with FTO expression in melanoma (Supplementary Fig. 7g, h). In addition, we found a positive association between FTO and PD-1 protein levels in Mel624 cells (Supplementary Fig. 7i, j) and human melanoma tissue (Supplementary Fig. 7k).

To determine whether regulation of these melanoma-promoting genes by FTO is m6A-dependent, we assessed the role of m6A by knocking down the m6A methyltransferases METTL3 and METTL14 (Supplementary Fig. 8a–c). In FTO knockdown cells, subsequent knockdown of both METTL3 and METTL14 significantly increased the mRNA levels of PD-1 (PDCD1), CXCR4, SOX10, CTSV, and NOP16 (Fig. 5j–n, and Supplementary Figs. 8a–d, 9a, b), and the protein levels of PD-1, CXCR4, and SOX10 (Fig. 5o), respectively, to a level similar to that in the control cells (Fig. 5j–o). These findings demonstrate that FTO regulates the expression of its target genes via m6A RNA modification and its demethylase activity.

**Role of YTHDF2 in FTO’s function in melanoma.** The function of m6A in regulating gene expression is executed mostly
through the readers, including YTH domain-containing family proteins in mammalian cells\(^1\). FTO knockdown did not affect the expression of critical m\(^6\)A regulators (Fig. 6a). To determine how m\(^6\)A RNA methylation regulates gene expression, we analyzed the role of YTHDF1-3. Knockdown of YTHDF1 or YTHDF3 had no effect on the mRNA levels of PD-1 (PDCD1), CXCR4, or SOX10 in control and FTO-knockdown cells. However, knockdown of YTHDF2 significantly increased the mRNA levels of all three genes in both control and FTO-knockdown cells (Fig. 6b–d, and Supplementary Fig. 10a–c). Moreover, knockdown of FTO decreased the mRNA stability of PD-1 (PDCD1), CXCR4, and SOX10 (Fig. 6e–g). Knockdown of YTHDF2 increased the mRNA stability of PD-1 (PDCD1), CXCR4, and SOX10 in shFTO cells (Fig. 6h–i). Consistently, knockdown of YTHDF2 increased, while forced overexpression of YTHDF2 decreased, cell proliferation and migration in melanoma cells in vitro (Fig. 6k, l, and Supplementary Fig. 10d–f) as well as tumor growth in vivo (Fig. 6m, and Supplementary Fig. 10g, h), supporting a tumor suppressor role for YTHDF2. These findings indicate that YTHDF2-mediated RNA decay controls the expression of the FTO target genes in melanoma cells.
FTO expression is induced by metabolic stress. To investigate potential pathways by which FTO is upregulated in melanoma cells, we analyzed the role of metabolic stress, since cancer cells acquire adaptive capabilities to survive and grow under metabolic stress. We found that metabolic stress conditions, such as low serum (0.2% FBS) or model starvation medium ( Hank’s Balanced Salt Solution, HBSS, lacking serum, amino acids and glucose) increased the expression of FTO and PD-1 (PDCD1) at the mRNA (Fig. 7a, and Supplementary Fig. 5a) and protein levels (Fig. 7b, and Supplementary Fig. 11a, b). Consistently, HBSS decreased the m6A level (Fig. 7c). Since these conditions are also known to induce autophagy (Fig. 7b, and Supplementary Fig. 11a, b), we reasoned that FTO induction by metabolic stress is mediated through the autophagy pathway. Indeed, knockdown of the autophagy essential genes ATG5 or ATG7 reduced starvation-induced FTO and PD-1 (PDCD1) expression, as well as NF-κB activity (Fig. 7d, e). Knockdown of FTO decreased PD-1 (PDCD1) expression (Fig. 7f). Knockdown of the NF-κB subunit p65 (RELA) markedly reduced the induction of FTO and PD-1 (PDCD1) by metabolic stress (Fig. 7g). These results demonstrate that FTO and PD-1 (PDCD1) are induced by metabolic stress through the autophagy and NF-κB pathways.

Role of FTO on anti-PD-1 blockade and IFNγ. Although immunotherapy has been effective in producing long-lasting therapeutic effects on several aggressive cancers in a subset of patients7–9, understanding the mechanism of resistance is critical to broaden the clinical application of immunotherapy for patients with advanced cancers11. Since FTO promotes melanoma tumorigenicity and regulates the expression of tumor-promoting melanoma cell-intrinsic PD-1 as well as CXCR4 and SOX10, we reasoned that FTO may also regulate the response of melanoma to immunotherapy. To determine whether FTO inhibition affects...
immunotherapy response, we treated C57BL/6 mice bearing B10F10 melanomas with anti-PD-1 antibody or isotype control IgG. Anti-PD-1 antibody had no effect on tumor growth of control B16F10 melanomas with anti-PD-1 antibody or isotype control (Fig. 8a), consistent with previous studies35,43, mimicking resistance to PD-1 blockade in melanoma treated as in Fig. 8a). The PD-1 blockade increased CD4+ T cell infiltration into tumors after PD-1 blockade (Supplementary Fig. 12a). The PD-1 inhibition sensitizes melanoma to anti-PD-1 blockade, we were assessed whether the immune system is required for the effect of FTO knockdown on anti-PD-1 blockade. In immunodeficient NSG (severely combined immunodeficient) mice, anti-PD-1 antibody had no effect on tumor growth in either control or B16F10 cells with FTO knockdown (Fig. 8b). These results indicate that the effect of FTO knockdown on melanoma response to anti-PD-1 immunotherapy is dependent on the immune system.

To understand the mechanism by which FTO inhibition sensitizes melanoma to anti-PD-1 blockade, we first analyzed whether FTO knockdown affects T cell infiltration into tumors after PD-1 blockade (Supplementary Fig. 12a). The PD-1 knockdown increased CD4+ tumor-infiltrating lymphocyte (TIL) numbers in B16F10 melanoma with FTO knockdown, but not in control cells (Fig. 8c). Notably, no significant difference was seen in CD8+ TIL numbers in either control or FTO-knockdown cells (Fig. 8c). We also assessed the IFNγ production of CD4+ and CD8+ TILs on day 14 after inoculation. However, FTO knockdown in melanoma cells did not affect the number of IFNγ-producing CD4+ or CD8+ TILs (Fig. 8d). Although further investigation is necessary to establish the effect of FTO knockdown and anti-PD-1 treatment on other immune cells and other immune mediators, these data suggest that FTO knockdown did not affect the infiltration of IFNγ-producing cytotoxic TILs.

Fig. 7 Metabolic stress induces FTO expression through NF-kB and autophagy. a qPCR analysis of FTO and PD-1 (PDCD1) mRNA levels in Mel624 cells cultured with control medium (10% FBS DMEM), 0.2% FBS DMEM, serum-free DMEM, Hank's balanced salt solution containing calcium and magnesium (HBSS), or a combination of DMEM and HBSS. b Immunoblot analysis of FTO, PD-1, p62, LC3-I/II (short and long exposure), and GAPDH in Mel624 cells treated as in a. c m6A dot blot assays using total RNA of Mel624 cultured with control medium or HBSS. Methylene blue staining was used as a loading control. d qPCR analysis of FTO and PD-1 (PDCD1) mRNA levels in Mel624 cells treated with or without metabolic stress. e Luciferase reporter analysis of NF-kB response element in Mel624 cells as in d. f qPCR analysis of FTO and PD-1 (PDCD1) mRNA levels in Mel624 cells with or without FTO knockdown treated with or without metabolic stress. g qPCR analysis of FTO and PD-1 (PDCD1) mRNA levels in Mel624 cells with or without siRNA knockdown of RELA treated as in e. Data are shown as mean ± S.E. (a, d, f, g), or mean ± S.D. (e) (n > 3). *P < 0.05; **P < 0.01; ***P < 0.001; Student’s t-test.
Next we analyzed the role of FTO in the response of melanoma cells to IFNγ. First, we assessed the effect of IFNγ on m^6A regulators. IFNγ downregulated FTO in a dose-dependent manner, but had no effect on the protein levels of other regulators such as ALKBH5, METTL3, or METTL14 (Fig. 8e). Consistently, IFNγ also increased m^6A levels (Fig. 8f), while it decreased PD-1 protein levels (Supplementary Fig. 12b). Next, we investigated whether FTO knockdown sensitizes melanoma cells to cell death induced by IFNγ. FTO knockdown did indeed sensitize melanoma cells to IFNγ-induced growth inhibition and cell killing (Fig. 8g, and Supplementary Fig. 12c). The converse was also true: Overexpression of WT FTO suppressed IFNγ-induced melanoma cell death, while overexpression of FTO mutants had no effect (Fig. 8h). These data suggest a critical role of FTO in the resistance of melanoma cells to IFNγ. Furthermore, we found that overexpression of PD-1 (PDCD1), CXCR4, or SOX10 inhibited IFNγ-induced cell death in FTO-knockdown melanoma cells (Fig. 8i–k). Consistently, blocking IFNγ by anti-IFNγ antibody increased tumor growth of both control and B16F10 melanoma cells with FTO knockdown in C57BL/6 mice (Fig. 8l). These results demonstrate that FTO inhibition enables a response to anti-PD-1 blockade in melanoma in vivo, and to...
IFNγ-induced killing in melanoma cells in vitro through regulating melanoma cell-intrinsic PD-1, CXCR4, and SOX10. Taken together, these results demonstrate a critical role of melanoma cell-intrinsic FTO in promoting melanoma resistance to anti-PD-1 blockade, suggesting that FTO inhibition can reduce resistance to anti-PD-1 therapy (Fig. 8m).

Discussion

Here we found that the obesity-associated protein FTO, also an m6A eraser, plays an important role in melanoma (Fig. 8m). FTO is increased in melanoma, suggesting a protumorigenic role of FTO in melanoma. FTO increases proliferation, migration, and invasion in melanoma cells in vitro and melanoma tumor growth in vivo. FTO can be upregulated by metabolic stress and starvation, a metabolic challenge that tumor cells are constantly facing in vivo, suggesting that the induction of FTO serves as an adaptive mechanism to metabolic stress in melanoma cells to promote proliferation, invasion, and migration. At the molecular level, FTO knockdown increases m6A enrichment in the 5′UTR and 3′UTR regions across the whole transcriptome. FTO knockdown increases m6A enrichment in the critical melanoma-promoting genes including PD-1 (PDCD1), CXCR4, and SOX10, and decreases their mRNA stability. The RNA decay of these genes in FTO knockdown cells is mediated by the m6A reader YTHDF2. YTHDF2 knockdown increases melanoma growth, whereas YTHDF2 overexpression decreases it, supporting a tumor suppressor role for YTHDF2 in melanoma. The regulation of these three genes by FTO is mediated through m6A demethylation and FTO’s demethylase activity. Lastly, FTO plays an important role in therapeutic resistance to anti-PD-1 immunotherapy and cell killing by IFNγ, a critical mechanism in immunotherapy. Overall, our findings demonstrate a crucial role of the m6A demethylase FTO in melanoma tumorigenesis and response to immunotherapy (Fig. 8m).

Our findings demonstrate that melanoma genes are regulated by FTO through m6A RNA methylation. FTO has been shown to be an RNA demethylase for m6A16 and also works on near cap m6A44, with opposite effects on RNA stability. FTO-mediated m6A demethylation was reported to promote mRNA decay44 and thus FTO could decrease transcript levels with m6Am demethylation. However, recent studies indicated that the effect of FTO on mRNA stability is mostly through m6A but not m6A demethylation. Importantly, the cap m6A methyltransferase PCIF1 was recently identified, and PCIF1 deletion eliminated cap m6A, but had minimal effects on target mRNA stability46, inconsistent with the previous report44, confirming the minimal effect of cap m6A on mRNA stability. We found that the regulation of PD-1 (PDCD1), CXCR4, and SOX10 all appear to be partially mediated through the latter m6A/YTHDF2 mechanism, because: (1) FTO knockdown decreased the mRNA levels and stability of these melanoma genes, in parallel with increased m6A enrichments; (2) knockdown of the m6A reader YTHDF2 increased the mRNA levels and stability of these genes, reversing the effect of FTO knockdown; (3) knockdown of METTL3 and METTL14 reversed the effect of FTO knockdown; (4) forced overexpression of YTHDF2 phenocopies the effect of FTO knockdown in cell proliferation, migration, and tumor growth. These results suggest that m6A enrichment lead to RNA decay of these genes through a YTHDF2-dependent mechanism in melanoma. Nevertheless, it is still possible that FTO-regulated demethylation could have other additional effects that remain to be uncovered.

We found that FTO plays a crucial role in metabolic pathways in melanoma cells. First, FTO regulates PD-1 (PDCD1) expression, thus FTO-mediated demethylation promotes mTOR signaling and cell growth/proliferation. Previously, FTO has been shown to regulate nutrient sensing and mTOR47. Here we identified an epitranscriptomic mechanism by which FTO regulates mTOR signaling through m6A-mediated tuning of the PD-1 (PDCD1) gene. Second, we found that metabolic stress and starvation induces FTO expression in melanoma cells. This metabolic stress-induced FTO expression requires autophagy and NF-xb. Due to uncontrolled proliferation of tumor cells in vivo, tumor cells likely face constant metabolic stress and lack of sufficient nutrient supply. It is possible that FTO induced by starvation promotes melanoma cell proliferation and/or survival. Our findings suggest FTO as a critical adaptation mechanism in melanoma cells under metabolic stress.

Anti-PD-1 checkpoint blockade therapy has demonstrated an unprecedented anti-tumor response rate in advanced cancers, including melanoma48, and reduced immune-related adverse side effects compared to ipilimumab48. However, more than half of such patients do not show a durable response to immunotherapy10. Recent studies have shown that melanoma intrinsic PD-1 is one of the critical intrinsic tumor-promoting regulators in melanoma, even in mice lacking adaptive immunity35. However, it has been unclear how PD-1 expression is regulated in melanoma cells. We found that PD-1 expression is positively regulated by FTO and m6A mRNA methylation. m6A modification of the melanoma cell-intrinsic PD-1 (PDCD1) gene leads to increased mRNA decay via YTHDF2. PD-1 (PDCD1) expression is induced via the autophagy/NF-xb/FTO axis by metabolic stress and starvation, suggesting the FTO/PD-1 axis as a critical component of metabolic stress adaptation. Furthermore, FTO knockdown sensitized melanoma to anti-PD-1 blockade (Fig. 8a). Therefore, inhibiting the FTO pathway in melanoma may provide a new opportunity to reduce resistance to anti-PD-1 immunotherapy.

Functional intrinsic IFNγ signaling machinery is not only critical for promoting anti-tumor immunity49, but also required for the therapeutic response to immune checkpoint blockade11. Consistently we found that IFNγ downregulates FTO in melanoma cells, and FTO mediates resistance to melanoma cell killing by IFNγ through its m6A demethylase activity and downstream targets PD-1 (PDCD1), CXCR4, and SOX10. FTO overexpression inhibits, while FTO knockdown promotes, IFNγ-induced cell death. It is possible that, at least in part, IFNγ exerts its cytotoxic effect by inhibiting FTO in melanoma cells. Indeed, anti-IFNγ antibody increased melanoma tumor growth in both control and FTO-knockdown cells in mice. Although FTO knockdown inhibits melanoma tumor growth in both immunocompromised and immunocompetent hosts (Figs. 2f, 8a, b), the response of FTO knockdown cells to anti-PD-1 requires host adaptive immunity (Fig. 8a, b). It appears that the melanoma intrinsic FTO pathway is not only critical for melanoma malignant traits and tumorigenicity in immunocompromised hosts, but also crucial for promoting resistance to IFNγ-induced killing in vitro and anti-PD-1 blockade in immunocompetent hosts in vivo. Although FTO knockdown in B16F10 increased the number of CD4+ TILs, it did not affect the number of IFNγ-producing CD4+ or CD8+ TILs (Fig. 8c, d). It is possible that the intrinsic function of the FTO pathway in IFNγ resistance mediated, at least in part, the effect of FTO knockdown in PD-1 blockade. Further studies are warranted to assess the full mechanism of immune regulation by PD-1 blockade with FTO inhibition in melanoma.

In summary, we have demonstrated that FTO inhibition suppresses melanoma tumorigenicity and the expression of melanoma cell-intrinsic genes PD-1 (PDCD1), CXCR4, and SOX10, at least partially through YTHDF2-mediated mRNA decay. We found that FTO inhibition enabled an anti-melanoma response to anti-PD-1 immunotherapy in mice. Our findings may provide new opportunities for developing improved melanoma therapies.
by inhibiting the FTO pathway in combination with anti-PD-1 immunotherapy.

**Methods**

**Human skin tumor samples.** All human specimens were studied after approval by the University of Chicago Institutional Review Board. The human melanoma array was obtained from US Biomax.

**Cell culture.** WM35, WM115, WM793, and WM3670 were kindly provided by Dr. DeMeer Harlyn (Wistar Institute, Philadelphia, PA). Mel624, CHL-1, SK-mel30, and B16F10 cells were purchased from ATCC or provided by the Comprehensive Cancer Center Core Facilities at the University of Chicago. Melanoma cells were maintained in DMEM (Dulbecco’s modified Eagle’s medium) (Invitrogen) supplemented with fetal bovine serum (FBS, 10% HyClone), penicillin (100 U/ml), and streptomycin (100 μg/ml, Invitrogen, Carlsbad, CA). NHEM cells (Normal Human Epidermal Melanocytes) were purchased from Lonza and cultured in 95% air and 5% CO2 at 37°C in NHEM defined medium obtained from Lonza according to the manufacturer’s instructions.

**Lentiviral generation and infection.** Lentiviral particles of shNC (Negative control) and shFTO for humans and mice were purchased from Santa Cruz Biotechnology. plKO.1 plasmids of shNC, shFTO, shATG5, and shYTHDF2 were obtained from Sigma. pLenti plasmids of overexpression control (mGFP) and overexpression of PDCD1 (mGFP-PDCD1), CXCR4, and SOX10 for humans were obtained from Origene (OriGene, Rockville, MD). Lentivirus was produced by co-transfection into HEK-293T cells with lentiviral vectors in combination with the pCMVdela8a2 packaging vector and pVSV-G envelope vector using GenJet Plus DNA In Vitro Transfection Reagent (Signagen). Virus-containing supernatants were collected 24–48 h after transfection and used to infect recipients. Target cells were infected in the presence of Polybrene (8 μg/ml) (Sigma-Aldrich) and selected with puromycin (Santa Cruz Biotechnology, 1 μg/ml) for 6 days.

**siRNA transfection.** Using GenMute siRNA Transfection Reagent (Signagen), cells were transfected with siRNA (Dharmacon), targeting negative control (NC) METTL3, METTL14, YTHDF1, YTHDF2, and YTHDF3, according to the manufacturer’s instructions. Luciferase reporter assays (Promega) were carried out according to the manufacturer’s instructions.

**Quantitative real-time PCR (qPCR).** qPCR assays were carried out using a CFX Connect real-time system (Bio-Rad, Hercules, CA) with Bio-Rad IQ SYBR Green Supermix (Bio-Rad). The threshold cycle number (CQ) was analyzed in triplicate for each sample. The CQ values for ANGPTL2, RAB40B, CTSV, RBBP9, NOP16, FCMR, PDCD1, CD274, CD47, CDKN1A, METTL3, METTL14, FTO, ALKBH5, YTHDF1, YTHDF2, YTHDF3, ATG5, ATG7, GAPDH, β-actin, and HPRT1 were normalized against HPRT1, GAPDH, or 18S, according to the manufacturer’s instructions.

**Flow cytometric analysis of apoptosis.** Apoptosis was analyzed using the annexin V–FITC apoptosis detection kit (eBioscience), following the manufacturer’s instructions. Cell were then by a BD FACSLessibur flow cytometer (BD Biosciences). For cells viability assay in cells in suspension, poly-hydroxyethyl methacrylate (poly-HEMA) plates were used to culture cells in suspension. A solution of poly-HEMA (Sigma–Aldrich, St. Louis, MO) mixed in ethanol was poured onto polystyrene bacteriological dishes. After the poly-HEMA had dried, the same procedure was repeated once followed by extensive washing with phosphate buffered saline (PBS).

**Migration and invasion assay.** Migration and invasion analysis, cell (5 × 104) suspension (150 μl of serum-free medium) were seeded onto 8-mm Pore Transwell Inserts (Corning) coated with Matrigel for invasion assay, or without Matrigel for migration assay. Lower chambers were filled with complete medium (900 μl). Cells on the Transwell Inserts were then fixed with paraformaldehyde/PBS (4%) for 30 min. Next, fixed cells were stained with hematoxylin solution (Sigma-Aldrich) for 1 h. Then microphotographs of the cells migrated onto the lower side of the filter were imaged using a microscope. From the microphotographs, cells that migrated or invaded onto the lower side of the filter were manually counted. Cell numbers were quantified from ten randomly selected fields with the same area (500 μm × 500 μm) per Transwell insert.

**Immunofluorescence and immunohistochemistry.** For immunofluorescence analysis, cells were first fixed with paraformaldehyde/PBS (4%) for 30 min, followed by permeabilized in Triton X-100/PBS (0.5%) for 20 min. Next cells were washed with PBS, blocked with PBS supplemented with 2% normal goat serum (Invitrogen) for 30 min, and stained for actin filaments with rhodamine-labeled phallolidin (Invitrogen, Carlsbad, CA). Then cells were washed three times with Triton X-100/TBS (0.1%) for 10 min and fixed in Prolong Gold Antifade with DAPI (Invitrogen), followed by observation under a fluorescence microscope (Olympus IX71).

For Human Melanoma Tissue Microarray staining, melanoma tissue array slides were obtained from US Biomax (Rockville, MD). After removing the blocking solution (3% albumin from chicken egg white (Sigma-Aldrich) in PBS), tissue array slides were incubated at 4°C with primary rabbit anti-FTO, goat anti-CD1, or mouse anti-MART1 for 18 h. After removing the primary antibodies, slides were washed with PBS solution with 0.025% TritonX-100. Tissue array slides were then incubated at room temperature with Alexa Fluor 594-conjugated secondary rabbit IgG (Jackson ImmunoResearch), Alexa Fluor 488-conjugated secondary mouse IgG (Jackson ImmunoResearch), and Alexa Fluor 405-conjugated secondary goat IgG (Jackson ImmunoResearch), then washed with PBS, and mounted with Prolong Gold Antifade with DAPI (Invitrogen) for cell nucleus counterstaining for FTO/MART1 staining, or with Fluoromount Mounting Medium (Sigma-Aldrich) for FTO/PD-1/MART1 staining. Two investigators independently scored the immunofluorescence intensity and transferred the results to a “scoring grid,” as 3 (strong), 2 (medium), 1 (weak), and 0 (negative). ImageJ (NIH) was also used for analyzing melanoma cells and tissue arrays. Areas with the same size were selected. The intensity of a selected area was measured followed by background subtraction to calculate mean pixel density. Stained samples were analyzed using a fluorescence microscope (Olympus IX71).

**immunohistochemistry.** Antibodies used are as follows: Anti-FTO (Abcam, ab26605, 1:100); Anti-PD-1 (Novus, NBP 2-15197, 1:100); Rhodamin-phallolidin (Life technologies, R415, 1:100); anti-PD-1 (R&D Systems, Cat# AF1086 Minneapolis, 1:100).

**Cell proliferation assay.** Cell proliferation was assessed using the Cell Counting Kit-8 (CCK-8) (Sigma–Aldrich, St. Louis, MO) following the manufacturer’s protocol.

**Immunoblotting.** Cell lysates were obtained using RIPA buffer (Fierce) containing inhibitors for proteases and phosphatases. Protein abundance were then analyzed by SDS-PAGE, and transferring onto nitrocellulose membranes followed by immunoblotting. Antibodies used are as follows:

- **Anti-ALKBHS5 (Millipore Co., ABEE 1013, 1:2000);** anti-Beta-actin (Santa Cruz, Sc-8432, 1:5000); anti–CXCR4 (Santa Cruz, Sc-3533A, 1:2000); anti–CXCR4 (Novusbio, NBPI-770675S, 1:5000); FTO (Santa Cruz, Sc-271713, 1:200); GAPDH (Santa Cruz, Sc-47724, 1:5000); GF (Cell Signaling Technology, 23555, 1:10000); LC3B (Cell Signaling Technology, 3868S, 1:10000); METTL14 (Millipore Co., ABEE 1388, 1:10000); METTL3 (Proteintech, 15073-1-AP, 1:10000); p62 (Progen Biotechnik GmbH, GP62-2, 1:10,000); p70S6K (Cell Signaling Technology, 2708S, 1:2000); PD-1 (Proteintech, 66220-1-Ig, 1:5000); p-p70S6K (Cell Signaling Technology, 92345, 1:10000); and SOX10 (Santa Cruz, Sc-365692, 1:2000). The unprocessed blots are provided in Source Data.

**Mouse tumorigenesis and treatment.** All of the animal procedures used were approved by the institutional animal care and use committee at the University of Chicago. Nude mice were purchased from Harlan Sprague-Dawley. C57BL/6 mice were obtained from Envigo. NSG (severely combined immunodeficient (NOD/SCID) interleukin-2 receptor (IL-2R) gamma chain null) mice were obtained from Jackson Laboratory. For xenograft experiments, one million cells were injected subcutaneously into the right flanks of 6-week-old female nude or C57BL/6 mice. Tumor growth was monitored and measured weekly by a caliper, and tumor volume was calculated using the formula, Tumor volume (mm³) = d² × D/2, where d and D are the shortest and the longest diameters, respectively. For treatment with anti-PD-1 antibody (BioXcell, clone RMP1-14) or isotype control IgG antibody (BioXcell, clone 2A3), B16F10 melanoma cells (5 × 10⁵) were inoculated subcutaneously into C57BL/6 or NSG mice. When the tumors reached a volume of 80–100 mm³, mice were treated with anti-PD-1 or isotype control antibody (200 μg/mouse) by i.p. injection, every other day for three times. For IFNγ blockade treatment, C57BL/6 mice were treated with anti-IFNγ antibody (BioXcell, Clone XMG1.2) or isotype control IgG (BioXcell, Clone HRPN) (250 μg/mouse) every other day after tumor cell inoculation.61,63
Analysis of tumor infiltrating lymphocytes (TILs). Tumor tissue from B16F10 tumor-bearing mice (Day 14 after tumor cell inoculation) was dissociated by digestion with 2 mg/ml collagenase type IV (Warthington Biochemical, LS004188) and 100 µg/ml DNase (Sigma-Aldrich, DN25) in RPMI 1640 with 5% FBS for 45 min at 37 °C. After digestion, tumor tissue was passed through 70-µm filters and mononuclear cells collected on the interface fraction between 40 and 80% Percoll. Live cells (Zombie NER negative) were gated using Zombie-violet (Catalog: 421035) staining. Next, cells were sorted by FSC-SSC and FSC–H to exclude doublets. Lymphocytes were gated on SSC-A and FSC-A. CD4+ and CD8+ TILs were gated on CD45+ cells. Gating strategies are shown in Supplementary Fig. 12A. The following mAbs recognizing the indicated antigens were used: FITC-anti-CD8 (Clone: 17A2, Catalog: 100204, 1:100), BV605-anti-CD4 (Clone: GK1.5, Catalog: 100451, 1:200), PE-CD7-anti-CD8 (Clone: 53-6.7, Catalog: 100722, 1:200), PerCP-Cy5.5-anti-CD45 (Clone: 30-F11, Catalog: 103129, 1:400), Zombie-violet (Catalog: 423105), and APC-anti-IFNγ (Clone: XMG1.2, Catalog: 308510, 1:100) (BioLegend). For assessment of IFNγ, cells were stimulated with 50 ng/ml phorbol 12-myristate 13-acetate (Sigma-Aldrich, P8139) and 1 µg/ml ionomycin (Fisher Scientific, BP25271) in the presence of Brefeldin A (BioLegend, 420601) for 4 h. After incubation, cells were then fixed. After surface staining, cell werepermeabilized using the BioLegend Kit (Catalog: 421002) and. Data were analyzed using FlowJo (version 10.5.3, FlowJo LLC).

m6A dot blot assay. Total RNA was extracted using an RNeasy plus Mini Kit (QIAGEN, Hilden, Germany), following the manufacturer’s protocol. For mRNA isolation, first total RNA was extracted using an RNeasy mini kit with DNase 1- column, followed by purifying RNA extraction using a Dynabeads mRNA Purification Kit (Life technology, Carlsbad, CA). Then mRNA was concentrated with an RNA Clean & Concentrator-5 kit (Zymo Research, Irvine, CA). Briefly, RNA samples were loaded onto Amersham Hybond-N + membrane (GE Healthcare, Chicago, IL) and crosslinked to the membrane with UV radiation. Then the membrane was blocked with 1% BSA in TBST (100 mM Tris, pH 7.4, 0.15% NaCl, 0.05% Tween20) for 1 h, incubated with a specific anti-m6A antibody (Synaptic Systems, 202003, 1:2000) overnight at 4 °C followed by HRP-conjugated anti-rabbit IgG (Cell Signaling Technology) for 1 h at room temperature, and then developed with Thermo ECL SuperSignal Western Blotting Detection Reagent (Thermo Fisher Scientific, Waltham, MA).

mRNA stability assay. A transcriptional inhibitor, actinomycin D (2 µM), inhibits mRNA transcription. Each sample was harvested at 0, 3, and 6 h after treatment with actinomycin D. Total RNA was isolated with an RNeasy mini kit (QIAGEN). The HPRT1 housekeeping gene was used as a loading control. HPRT1 mRNA does not contain m6A modifications, is not bound by YTHDF2, and is rarely affected by actinomycin D treatment2,25.

m6A IP. 100–150 µg total RNA was extracted from cells using TRIzol following the manufacturer’s protocol. mRNA was purified using a Dynabeads mRNA DIRECT Kit following the manufacturer’s protocols. One microgram mRNA was sonicated to ~200 bp. 2% of fragmented mRNA was saved as input, and m6A containing mRNA fragments were enriched with an EpImark N6-Methyladenosine Enrichment Kit following the manufacturer’s protocols. Finally, together with the input, IP RNA was extracted using RNA Clean and Concentrator (Zymo Research), followed by library preparation using the TruSeq stranded mRNA sample preparation kit (Illumina).

Sequencing data analysis. General pre-processing of reads: sequencing were performed using Illumina Hisoe4000 with single end 80 bp read length. The adapters were removed by using cutadapt for m6A-seq, reads were aligned to the reference genome (hg38) in Tophat v2.0.14 using the parameter -g 1 to mimic the exclusion of potential reads from the non-annotated regions. To mimimize bias from potential inaccuracy in gene structure annotation, mRNA fragments were enriched with an EpiMark N6-Methyladenosine Enrichment Kit following the manufacturer’s protocol. For mRNA isolation, first total RNA was extracted using an RNeasy mini kit with DNase 1-column, followed by purifying RNA extraction using a Dynabeads mRNA Purification Kit (Life technology, Carlsbad, CA). Then mRNA was concentrated with an RNA Clean & Concentrator-5 kit (Zymo Research, Irvine, CA). The same amount of the concentrated IP RNA or input RNA from each sample was used for the cDNA library. The mRNA expression was determined by the number of amplification cycles (Cq). The relative m6A levels in genes were calculated by the m6A levels (m6A IP) normalized using the expression of each gene (Input).

Microarray analysis. RNA was extracted from Mel624 cells with or without FTO knockdown with the RNeasy Mini Kit (Qiagen, Hilden, Germany), according to the manufacturer’s instructions. Total RNA concentration and purity were determined with NanoDrop (Thermo Scientific, Waltham, MA), and total RNA integrity was confirmed with the Agilent Bioanalyzer (Agilent Technologies, Santa Clara, Calif.). The RNA samples were processed with the HumanHT-12 v4.0 Gene Expression BeadChip microarray (Illumina) at the Functional Genomics Core Facility of the University of Chicago. The data have been deposited in the GEO repository with the accession numbers GSE128961.

Statistical analyses. Statistical analyses were carried out using Prism 6 and 7 (GraphPad). Data were obtained from at least three independent experiments and for statistical significance were analyzed using Student’s t-test or Mann–Whitney U-test. P < 0.05 was considered statistically significant.

Reporting summary. Further information on experimental design is available in the Nature Research Reporting Summary linked to this article.

Data availability
m6A IP sequencing and RNA sequencing data are accessible at the GEO repository, under accession number GSE112902. Microarray data are accessible at the GEO repository, under accession number GSE128961. Other data from this study are available from the corresponding author upon request. The source data of immunoblot is provided as a Source Data file.

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Author contributions

S.Y. and Y.Y.H. conceived the project. S.Y., J.W., Y.C., G.P., S.H., C.H., and Y.Y.H. designed the research and supervised the experiments conducted in the laboratories. S.Y., J.W., Y.C., G.P., and P.S. carried out the experiments and/or performed data analyses; Y.D., A.A., and Z.L. contributed the reagents/analytic tools/data analysis; S.Y. and Y.Y.H. wrote the manuscript. All authors reviewed the results and the manuscript.

Additional information

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Competing interests: C.H. is a scientific founder and a member of the scientific advisory board of Accent Therapeutics, Inc. The remaining authors declare no competing interests.

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