**Fusobacterium nucleatum** stimulates cell proliferation and promotes PD-L1 expression via IFIT1-related signal in colorectal cancer

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Abstract

*Fusobacterium nucleatum* (*F. nucleatum*) is enriched in colorectal cancer (CRC) tissues and a high amount of *F. nucleatum* was associated with an immunosuppressive tumor environment. PD-L1 is an important immune checkpoint expressed on tumor cells and promotes tumor immune escape. Whether PD-L1 is regulated by *F. nucleatum* is still unclear. We demonstrated that *F. nucleatum* promoted CRC progression and upregulated PD-L1 protein expression in CRC cell lines. Combined m^6^A-seq and RNA-seq identified m^6^A-modified IFIT1 mediating *F. nucleatum* induced PD-L1 upregulation. IFIT1 mRNA was modified with m^6^A modifications in 3'UTR and the m^6^A levels were altered by *F. nucleatum* treatment. Our results also indicated that IFIT1 served as a potential oncosine in CRC and regulated PD-L1 protein levels through altering PD-L1 ubiquitination. Clinical CRC data confirmed the correlation among *F. nucleatum* abundance, IFIT1 and PD-L1 expressions. Our work highlighted the function of *F. nucleatum* in stimulating PD-L1 expression through m^6^A-modified IFIT1 and provided new aspects for understanding *F. nucleatum* mediated immune escape.

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Introduction

Colorectal cancer (CRC) is the third most commonly diagnosed cancer as well as the third fatal cancer in the world [1,2]. Recent evidence has demonstrated that host-microbiota interactions play an important role in CRC progression [3,4]. *F. nucleatum* is an important cancer-associated bacterium contributing to cancer development especially CRC [5,6]. A high enrichment of *F. nucleatum* is associated with colorectal carcinogenesis, disease stages and overall survival. *F. nucleatum* coculture could stimulate cancer cell proliferation and *F. nucleatum* gavage resulted in more and larger tumors in AOM-DSS CRC model and Apc^m^A^m^A^−^ mice [7]. Besides, *F. nucleatum* also promotes resistance to chemotherapy and glycolysis in CRC [8,9].

PD-L1 is an essential immune checkpoint molecule, generating an inhibitory signal and establishing an immunosuppressive environment [10]. PD-L1 is highly expressed in various cancer cells and stromal immune cells, allowing tumors to evade attacks of infiltrated T cells [11,12]. Several population-based studies have reported a higher abundance of *F. nucleatum* associated with an immunosuppressive tumor environment, with lower T cell density [13,14]. The virulence factor derived from *F. nucleatum*, Fap2, inhibited NK cell cytotoxicity through interacting with TIGIT [15]. Whether *F. nucleatum* could regulate PD-L1 expression in CRC is still unclear.

m^6^A-methyladenosine (m^6^A) modification is abundant and prevalent in eukaryotic cells, involving in diverse biological processes in cancer [16,17]. PD-L1 expression could be directly modified with m^6^A modifications and also indirectly regulated by m^6^A-modified target genes [18,19]. Whether RNA m^6^A modification participated in *F. nucleatum* mediated immunosuppressive tumor environment was still unclear.

In this work, we found *F. nucleatum* promoted CRC cell proliferation in vitro and in vivo. *F. nucleatum* treatment could markedly stimulate PD-L1 protein expression without mRNA levels alteration. Through combined m^6^A-seq and RNA-seq in CRC cells cocultured with *F. nucleatum*, we identified m^6^A-modified IFIT1 and found IFIT1 knockdown could reverse *F. nucleatum* induced PD-L1 expression. Mechanically, IFIT1 overexpression...
increased PD-L1 protein levels through inhibiting its ubiquitination. Also, we found IFIT1 was higher expressed in CRC tissues than normal colon tissues. A higher expression of IFIT1 was associated with higher cancer stage and worse survival. In addition, METTL3 and METTL14 methylated IFIT1 in 3’UTR and promoted its mRNA stability in a IGF2BP2/3-dependent manner. In clinical samples, F. nucleatum abundance, IFIT1 and PD-L1 expressions were positively correlated. Our work indicated that F. nucleatum stimulated PD-L1 protein expression via m^1A modification of IFIT1 and provided new aspects for understanding F. nucleatum mediated immune escape.

**Materials and methods**

**Cell lines**

RKO cells and SW1116 cells were purchased from American Type Culture Collection (ATCC) and incubated with RPMI-1640 medium (Gibco) supplemented with 10% fetal bovine serum (FBS). HT29 cells and HCT116 cells were also purchased from ATCC and cultured with McCoy's 5A (Gibco) with 10% FBS (Gibco). All cell lines were grown at 37°C in a humidified 5% CO₂ atmosphere.

**Patient specimens**

We studied four cohorts of CRC patients having undergone surgery in Renji Hospital affiliated with Shanghai Jiao Tong University School of Medicine. The study was approved by the ethics committee of Renji Hospital and all patients have obtained the written informed consent. Respectively, there were 36 pairs of FFPE CRC tissues and non-cancerous normal colon tissues in cohort 1, 79 FFPE cancer tissues in cohort 2, 34 pairs of FFPE cancer tissues and non-cancerous normal tissues in cohort 3 and 180 FFPE cancer tissues in cohort 4. The clinical characteristics of these cohorts were shown in Supplementary Table 1, 2, 3 and 4.

**Animal studies**

Four-week-old male BALB/c nude mice were housed in Shanghai Model Organism Center under specific pathogen-free conditions. To investigate the effects of F. nucleatum on tumor progression, HT29 and HCT116 cells (5×10^3) were injected subcutaneously into the right axilla. Mice were randomly separated into three groups after 5 days’ inoculation, receiving intratumoral injection of PBS, F. nucleatum solution and E. coli solution (10^7 CFU per mouse) separately. The injection was carried twice one week. Mice were sacrificed 20 days later and subcutaneous tumors were harvested and weighed. To investigate the effects of IFIT1, HCT116 cells were stably transfected with shIFIT1 lentivirus, pcDNA3.1_IFIT1 lentivirus and their corresponding control. HCT116 cells (5×10^3) were also injected subcutaneously into the right axilla. Mice were sacrificed 20 days later and subcutaneous tumors were harvested and weighed. These experiments were performed following the Institute’s guidance on animal experiments.

**Bacterial strains and growth conditions**

*Fusobacterium nucleatum* ATCC 25586 and *Enterotoxigenic Bacteroides fragilis* were purchased from ATCC. *DH5α* E. coli was obtained from Tiangen Company. They were cultured as previously described [8].

**Detection of bacterial abundance**

QIAamp DNA FFPE Tissue Kit (QIAGEN) was used to extract the genome DNA from FFPE CRC tissue. The extracted DNA was assessed using qPCR and the *F. nucleatum* abundance was normalized using 16S genes. The primer for *Fusobacterium nucleatum* detection is “F-CAACCATTCTTTACCTACATGTTCA” and “R-GT TGACTTTACAGAGGAGATTATGTAAATATC”. The primer for 16S detection is “F- GGTGAATACGTTCCCGG” and “R- TACGCTACCTTGTACGACTT”.

**Cell proliferation assay**

Cell proliferation was assessed with Cell Counting Kit-8 assay (Dojindo), EdU assay (RiboBio) and colony formation assay. For Cell Counting Kit-8 assay, 3000 cells were planted into 96-well plates. Cell Counting Kit-8 (Dojindo) was mixed with medium at 1:10 dilution. 100ul of the above mixture was added into the wells. Incubated for 2 hours, the absorbance at 450nm was measured. For EdU assay, cells were incubated with EdU stain for 2 hours. Cells with positive stain were counted under microscope. For the colony formation assay, 600 cells were planted into 12-well plates. After 7 to 10 days’ culture, cells were washed with PBS twice, fixed with 4% paraformaldehyde and stained with 0.1% crystal violet. The colonies were counted after washing with PBS. The above experiments were conducted independently three times.

**Construction and transfection of small interfering RNAs, lentivirus shRNA and plasmids**

METTL3, METTL14, IGF2BP1/2/3 and IFIT1 small interfering RNAs (siRNAs) were obtained from GenePharma (Shanghai, China). IFIT1 lentivirus shRNAs and pcDNA3.1_IFIT1 overexpressed lentivirus were also obtained from GenePharma (Shanghai, China). The control plasmid, METTL3, METTL14, IFIT1 and PD-L1 overexpressing plasmids were constructed in Generay Technologies (Shanghai, China). Cells were seeded in six-well plates overnight. For siRNA transfection, we made the transfection mixture with 5ul siRNA (50nm), 5ul Dharma FECT 1 transfection reagent and 390ul opti-MEM medium. For plasmid transfection, we added 1ug plasmid and 3ul viaFECT for one well. After six hours of incubation, the medium was replaced by normal culture media. Cells were harvested after 48h. For lentivirus shRNA transfection, cells were incubated with lentivirus at the MOI of 100. After 24 hours’ incubation, cells were treated with puromycin for two weeks. The sequences of siRNAs and shRNAs were listed in Supplementary Table 7.

**M^4^A-seq and RNA-seq**

M^4^A-seq was conducted according to the previous studies [20]. Briefly, polyadenylated mRNA was isolated from total RNA with GenEluteTM mRNA Miniprep Kit. The quantity and quality of mRNA were tested with agarose gel electrophoresis and NanoDropTM instruments. The isolated mRNA was fragmented into short fragments and approximately 1/10 of the RNA fragments were conserved for RNA sequencing by oebiotech(Shanghai, China). The M^4^A-methylated mRNAs were immunoprecipitated with anti-M^4^A antibodies for further mRIP-seq. The differentially m^4^A-RP-enriched regions (peaks) were calculated using exomePeak.

**MeRIP-qPCR**

This experiment was conducted as a previous report [17]. Polyadenylated mRNA was fragmented with RNA fragmentation reagents (NEB). The anti-m^4^A antibody (NEB) was used to immunoprecipitate m^4^A-methylated mRNAs. The m^4^A-methylated mRNAs were extracted, purified and resolved with nuclease-free water as previously described [17]. Real-time PCR was performed to measure the m^4^A level in target gene transcripts. Primers in this experiment were listed in Supplementary Table 7.
Western blot assay and Immunoprecipitation

Cells were collected after lysis with RIPA added with a protease inhibitor cocktail. Cell supernatant was subjected to BCA Protein Assay Kit to measure the concentration. Same amount of protein samples was separately with 10% SDS-PAGE and then transferred to PVDF membranes (Bio-Rad). Membranes were blocked with 5% nonfat milk and then incubated with primary antibodies overnight at 4°C. Next day, membranes were incubated with secondary antibodies. The ECL Kit was used to detect the signals. Immunoprecipitation was performed as previously described. Cells were harvested with IP lysis buffer (Thermo Fisher). After centrifugation, 1/10 of the supernatant was added with 5× SDS-PAGE loading buffer, serving as an input control. The remaining was supplemented with 20ul Pierce Protein G Magnetic Beads (thermo fisher, 88847). After supplemented with 2ul primary antibodies and isotype-matched IgG, the mixture was rotated slowly at 4°C overnight. Subsequently, the precipitation was isolated using the magnetic rack (NEB), followed by washing with PBS five times. The precipitation was mixed with 50 µl 1× SDS-PAGE loading buffer and heated at 95°C for 10min. Subsequent steps are as above described. The antibodies used in the experiments are listed as follow: anti-PD-L1 (CAT#13684, CST); anti-IFIT1 (CAT#14769S, CST); anti-METTL3 (CAT#15073-1-AP, Proteintech); anti-METTL14 (CAT#51104, CST); anti-HA tag (CAT#3724, CST); anti-FLAG tag (CAT#2368, CST); anti-Pi3K (CAT#4249T, CST); anti-p-AKT (CAT#9271T, CST); anti-AKT (CAT#4691T, CST); anti-p-JAK2 (CAT#ab32101, abcam); anti-JAK2 (CAT#ab108596, abcam); anti-p-STAT1 (CAT#ab109457, abcam); anti-STAT1 (CAT#ab109320, abcam); anti-p-NF-kB (CAT#3033T, CST); anti-NF-kB (CAT#8242T, CST); anti-GAPDH (KANGCHENG Technology); anti-rabbit-HRP (KANGCHENG Technology); anti-mouse-HRP (KANGCHENG Technology).

Quantitative real-time PCR

Cells were lysed and extracted with RNaiso Plus (Takara). Reverse transcription was conducted with PrimeScriptRT Reagent Kit (Takara) and specific genes expressions were measured with SYBR Premix Ex Taq II (Takara). Primers were obtained from tisngle Company and the sequences were listed in Supplementary Table 7.

Immunofluorescence

Cells were seeded into eight-well chamber slides. After adherence and treatment, cells were fixed using 4% formaldehyde (Biohashp), permeabilized using Triton-X-100 for 10 minutes and blocked with 1% BSA for 1 hour. Following incubated with primary antibodies overnight at 4°C, cells were incubated with fluorescently-labeled conjugated secondary antibodies (Alexa Fluor 594 or 488; Invitrogen). Slides were washed with PBS, stained with DAPI and sealed with coverslips. Fluorescence was observed with Zeiss LSM710 confocal microscope. The antibodies used in the experiment are listed as follow: Rabbit anti-PD-L1 (CAT#17952-1-AP, Proteintech), 1:100 dilution; Mouse anti-IFIT1(CAT#ab118062, abcam), 1:100 dilution. The above experiments were conducted independently three times.

Immunohistochemistry

Paraffin-embedded colon tissues were deparaffinized and rehydrated. After antigen retrieval and endogenous peroxidase block, the sections were blocked with goat serum, incubated with primary antibodies overnight and incubated with secondary antibody for 30 minutes. DAB was used as chromogen and hematoxylin was used for nucleus counterstaining. Immunohistochemistry (IHC) staining score was assessed under 200X microscope. The antibodies used in the experiment are listed as follow: Rabbit anti-PD-L1 (CAT#17952-1-AP, Proteintech), 1:200 dilution; Rabbit anti- IFIT1 (CAT#ab236256, abcam), 1:50 dilution.

Flow cytometry assays

Flow cytometry assays were performed for cell surface PD-L1 detection. After harvested with trypsin, cells were stained with APC- conjugated anti-human PD-L1 antibody (1:200) (BioLegend) for 30min at 4°C followed by washing with PBS twice. The staining process was protected from light. And then cells were analyzed with flow cytometry. The above experiments were conducted independently three times.

Detection of PD-L1 protein stability

Cells were transfected with IFIT1 siRNAs and incubated with cycloheximide (beyotime) (50 μg/mL) for specific times. The proteins were extracted and analyzed using western blot. The intensity of bands was calculated with Image J and were normalized by GAPDH intensity.

RNA stability assay

Cells were seeded in 12-well plates overnight, followed by actinomycin D (5 μg/mL) treatment for 0h, 2h, and 4h. Total RNA was extracted and qPCR was performed to assess IFIT1 mRNA levels. The half-life of IFIT1 mRNAs was estimated using linear regression analysis.

RNA immunoprecipitation (RIP) and RIP-qPCR

RIP assays were performed with Magna RIP Kit (Millipore, New Bedford, MA). Anti-IGF2BP2 (proteintech, 11601-1-AP), anti-IGF2BP3(proteintech, 14642-1-AP), and isotype-matched IgG were used for RIP. Real-time PCR was conducted to determine the interaction between IGF2BP2/3 and IFIT1 mRNA.

Statistical analysis

GraphPad Prism (8.0.2) was used to plot column bar graphs and scatter plots. Values were showed as mean±SD with three independent experiments. Two independent samples were compared with two-tailed Student’s t test and more than two groups were compared using one-way ANOVA. Immunofluorescence and western blot were analyzed with ImageJ (2.1.0) software. Flow cytometry assays were analyzed with Flowjo 10.4 (Biosaccine). The correlation assays were analyzed using Spearman correlation test.

Results

F. nucleatum promoted tumor progression and upregulated PD-L1 expression in CRC

F. nucleatum is a well-known oncobacterium in cancers, especially in CRC. We measured F. nucleatum abundance in 36 pairs of formalin-fixed paraffin-embedded (FFPE) CRC tissues and normal colon tissues in cohort 1. The results showed a higher F. nucleatum abundance in cancer tissues (Fig. 1A). In vitro, F. nucleatum coculture (MOI=300, MOI, multiple of infection) remarkably enhanced HT29 and HCT116 cell proliferation compared with DH5a E. coli (MOI=300), heat-killed F. nucleatum and medium control through EdU assays (Fig. 1B-C, Supplementary Figure S1B-C). The similar results were also obtained through CCK8 assays and colony formation assays (Fig. 1D-E, Supplementary Figure S1A, D-F). In vivo, we inoculated CRC cells into nude mice and injected F. nucleatum solution intratumorally
and found *F. nucleatum* treatment stimulated tumor growth significantly (Fig. 1F-G, Supplementary Figure S1G-H). Ki-67 is an important proliferation marker in cancer progression. In clinical CRC samples, we observed a higher *F. nucleatum* abundance was associated with higher Ki-67 scores (Supplementary Figure S1I), however, the difference was not significant. Various factors including histological differentiation, tumor size, lymphatic metastasis and AJCC II/III/IV stage could also influence Ki-67 score.

Previous researches have suggested that *F. nucleatum* in colorectal tissues could modulate a tumor immunosuppressive environment, accompanied with lower density of CD3+ T cells and an expansion of myeloid-derived immune cells [13,14]. PD-L1 is an important immunosuppressive molecule, dampening cancer immunity through providing suppressive signals to CD8+ T cells [10]. Whether *F. nucleatum* treatment could stimulate PD-L1 expression is still unclear. We treated CRC cells with *F. nucleatum* for 3 hours and measured PD-L1 mRNA expression using RT-qPCR. The results showed that *F. nucleatum* coculture didn’t alter PD-L1 mRNA levels (Supplementary Figure S2A-B, D). However, western blot assays showed that *F. nucleatum* coculture (MOI=300) stimulated PD-L1 protein expression compared with DH5α *E. coli* (MOI=300), heat-killed *F. nucleatum* (MOI=300) and medium control in CRC cells, while mRNA levels were not altered (Fig. 2A-B). The similar results also existed in HCT116 and RKO cells (Supplementary Figure S2C, E). Altogether, our data indicated that *F. nucleatum* promoted CRC progression and simulated PD-L1 protein expression without significant mRNA alterations.

**F. nucleatum** promoted **PD-L1** expression via **m6A** modification of **IFIT1**

RNA m6A modification is a prevalent posttranscriptional mRNA modification in eukaryotic cells [20]. Previous reports have indicated that RNA m6A modification was an important regulator of PD-L1 and was also associated with tumor immune cell infiltration [21]. Whether m6A modification participated in PD-L1 expression induced by *F. nucleatum* was still unknown. To explore this question, we preformed combined m6A-seq and RNA-seq in HT29 cells treated with *F. nucleatum* (MOI=300) and attempted to find the potential genes regulating PD-L1 expression. Consistent with the previous findings [22], the m6A consensus motif GGAC was enriched in the detected peaks (Fig. 2C). The m6A peaks were mostly distributed in 5’-untranslated regions (5’UTR), coding regions and 3’-
Fig. 2. *F. nucleatum* promoted PD-L1 expression via m6A modification of IFIT1. A. PCR analysis of PD-L1 mRNA levels in HT29 cells treated with *F. nucleatum* (MOI=300), DH5α E. coli (MOI=300), heat-killed *F. nucleatum* (MOI=300) and medium control (n=3, mean± SD, One-way ANOVA). B. Western blot assays detecting PD-L1 protein expression in HT29 cells treated with *F. nucleatum* (MOI=300), DH5α E. coli (MOI=300), heat-killed *F. nucleatum* (MOI=300) and medium control. C. The top consensus m6A motif in HT29 cells. D. The distribution of m6A peaks. E. Distribution of genes with a significant change in both m6A level and gene expression level in HT29 cells treated by *F. nucleatum* (MOI=300) compared with medium control (p<0.05). F. The flow chart of selecting candidate target genes. G. PCR analysis of 5 candidate target genes in HT29 cells treated with *F. nucleatum* (MOI=300) and medium control (n=3, mean± SD, two-tailed Student’s t-test). H. IGV analysis representing m6A abundances in IFIT1 transcripts in HT29 cells in response to *F. nucleatum* (up panel) and the medium control (bottom panel). I. Agarose electrophoresis (up panel) and meRIP-qPCR assays (down panel) of IFIT1 3’UTR m6A levels in HT29 cells (n=3, mean± SD, two-tailed Student’s t-test). J. MeRIP-qPCR analysis of IFIT1 3’UTR m6A levels in HT29 cells treated with medium control, *F. nucleatum* (MOI=300), ETBF (MOI=300) and DH5α E coli (MOI=300) (n=3, mean± SD, One-way ANOVA). K. PCR analysis of IFIT1 mRNA levels in HT29 cells treated with different MOI of *F. nucleatum* (n=3, mean± SD, One-way ANOVA). L. Western blot assays detecting IFIT1 protein expression in HT29 cells treated with different MOI of *F. nucleatum*. M. PCR analysis of IFIT1 mRNA levels in HT29 cells treated with medium control, *F. nucleatum* (MOI=300), ETBF (MOI=300) and DH5α E coli (MOI=300) (n=3, mean± SD, One-way ANOVA). N. Western blot assays detecting IFIT1 protein expression in HT29 cells treated with medium control, *F. nucleatum* (MOI=300), ETBF (MOI=300) and DH5α E coli (MOI=300). O. Western blot assays detecting PD-L1 protein expression in IFIT1 knockdown and the control HCT116 cells treated with *F. nucleatum* (MOI=300). P. Western blot assays detecting PD-L1 protein expression in IFIT1 knockdown and the control RKO cells treated with *F. nucleatum* (MOI=300).
untranslated regions (3′UTR), especially abundant surrounding the stop codon (Fig. 2D, Supplementary Figure S2F). Through combined analysis, we have identified 112 genes with altered m^A modifications and expression (Fig. 2E) (Supplementary Table 5 and 6). The top five genes were listed as the candidate genes, including CLDN2, IFIT1, PHLD1B1, LONP1, and SLPI (Fig. 2F). Real-time PCR was performed to confirm which candidate genes were regulated by *F. nucleatum* treatment. The results showed IFIT1 was the most significantly altered in HT29 and HCT116 cells after *F. nucleatum* infection (MOI=300) (Fig. 2G, Supplementary Figure S2G). The Integrative Genomics View (IGV) data showed a higher m^A level in 3′UTR of IFIT1 after *F. nucleatum* treatment (Fig. 2H). MeRIP-qPCR assays confirmed the m^A peaks located in 3′UTR of IFIT1 transcripts and the m^A levels were specifically increased after *F. nucleatum* infection (MOI=300) in HT29 cells, compared with treatments with *ETBF* (MOI=300) and *DH5α E. coli* (MOI=300) (Fig. 2I). The IFIT1 mRNA and protein expressions were induced by *F. nucleatum* in a MOI-dependent manner (Fig. 2K-L, Supplementary Figure S2H-I). Additionally, IFIT1 upregulation was induced specifically by *F. nucleatum* (MOI=300) compared with *ETBF* (MOI=300) and *DH5α E. coli* (MOI=300) (Fig. 2M-N, Supplementary Figure S2J-K). To explore whether IFIT1 mediated *F. nucleatum* induced PD-L1 expression, we cocultured *F. nucleatum* with IFIT1 knockdown CRC cells and the control cells. The results showed that IFIT1 knockdown could partially reversed *F. nucleatum* (MOI=300) induced PD-L1 protein upregulation (Fig. 2O-P). Previous researches have also indicated that PI3K pathway is an important signal regulating PD-L1 expression in cancer progression [23,24]. And *F. nucleatum* treatment could activate PI3K pathway [25,26]. To investigate whether *F. nucleatum* induced PD-L1 expression also depended on activating PI3K pathway, we treated CRC cells with the PI3K inhibitor, LY294002, prior to *F. nucleatum* coculture. The western blot results showed that *F. nucleatum* treatment could activate PI3K pathway in CRC cells. However, PI3K inhibition could not reverse PD-L1 upregulation induced by *F. nucleatum* treatment (Supplementary Figure S2L-Q). Thus, our results indicated that *F. nucleatum* promoted PD-L1 expression partially via m^A modification of IFIT1.

**IFIT1** promoted CRC cell proliferation and participated in *F. nucleatum* induced CRC progression

IFIT1, interferon-induced proteins with tetratricopeptide repeats gene 1, is a member of interferon-stimulated genes (ISGs), which function as essential anti-viral proteins [27]. Emerging findings demonstrated its roles in tumor progression [28,29]. However, the roles of IFIT1 in CRC progression are still unknown. We performed CCK8 assays and colony formation assays in HT29 and HCT116 cells with IFIT1 knockdown. The results showed that IFIT1 knockdown dampened CRC proliferation and colony numbers (Fig. 3A-C, Supplementary Figure S3A-C). To confirm the results in *vivo*, we used lentivirus to establish HCT116 cells stably transfected with IFIT1 shRNAs. Knockdown of IFIT1 markedly reduced HCT116 tumor burden in nude mice models (Fig. 3D-E). Meanwhile, IFIT1 overexpression stimulated CRC cell proliferation and colony numbers (Fig. 3F-H, Supplementary Figure S3D-F). To confirm the results in *vivo*, we used lentivirus to establish HCT116 cells stably overexpressing IFIT1. Overexpression of IFIT1 dramatically increased HCT116 tumor volume and tumor weight in xenograft mice models (Fig. 3I-J). Since *F. nucleatum* could increase IFIT1 expression in CRC cells, whether IFIT1 participated in *F. nucleatum* induced CRC progression was still unknown. To explore this question, we decreased IFIT1 in HT29 and HCT116 cells using siRNAs and treated cells with *F. nucleatum* and performed CCK8 assay and colony formation assays. The results showed that IFIT1 knockdown could partially block *F. nucleatum* (MOI=300) induced cell proliferation (Fig. 3K-M, Supplementary Figure S3G-I). Altogether, IFIT1 promoted CRC proliferation in *vivo* and in *vivo*. Also, FIT1 upregulation participated in *F. nucleatum* induced CRC progression.

**IFIT1** altered PD-L1 protein expression

Since IFIT1 knockdown could reverse *F. nucleatum* stimulated PD-L1 protein expression, we guessed that IFIT1 may regulate PD-L1 protein expression. First, we measured PD-L1 mRNA and protein expressions in IFIT1 siRNAs-transfected RKO cells. The results suggested that IFIT1 knockdown impaired PD-L1 protein expression but not mRNA expression (Fig. 4A-B, Supplementary Figure S4A). Ectopically overexpressed IFIT1 markedly enhanced PD-L1 protein expression without mRNA alterations in RKO cells (Fig. 4C-D, Supplementary Figure S4B). Immunofluorescence assays and flow cytometry also displayed that IFIT1 knockdown induced a reduction of PD-L1 in RKO cells (Fig. 4E-G). These results indicated that IFIT1 regulated PD-L1 protein expression without mRNA level alterations.

**IFIT1** mediated PD-L1 stabilization via regulating PD-L1 ubiquitination

We sought to explore how IFIT1 regulated PD-L1 protein expression. NF-κB and JAK-STAT signals were two important pathways regulating PD-L1 expression [30–32]. Firstly, to investigate whether IFIT1 regulated these two pathways, we detected the expression of key components of NF-κB and JAK-STAT pathways in RKO cells with IFIT1 knockdown and overexpression using western blot assay. The results showed that IFIT1 knockdown and overexpression didn’t alter these two pathways (Supplementary Figure S5A-B). Previous studies have indicated that NF-κB and JAK-STAT pathways regulate PD-L1 expression in mRNA levels. Consistent with these results and previous studies, our data showed IFIT1 altered PD-L1 protein expression not mRNA expression. In consideration of the TPR motifs of IFIT1, we hypothesized that IFIT1 may exert its effects through interacting with PD-L1. We immunoprecipitated RKO cell lysates with primary anti-PD-L1 antibodies and detected IFIT1 using western blot. The results indicated endogenous interaction of IFIT1 and PD-L1 (Fig. 5A). Next, we transfected FLAG-IFIT1 and HA-PD-L1 into RKO cells and confirmed their interaction ectopically (Fig. 5B). Since IFIT1 didn’t affect PD-L1 mRNA levels, we guessed that IFIT1 affected PD-L1 protein degradation. CHX assays showed PD-L1 showed a more rapid decay after IFIT1 knockdown in RKO cells (Fig. 5C-E). Treatment with the proteasome inhibitor MG132 could reverse IFIT1 knockdown induced PD-L1 downregulation, while the lysosomal inhibitor chloroquine (CQ) and autophagy inhibitors 3-methyladenine (3-MA) displayed no such effects (Fig. 5F-H). These results indicate that IFIT1 may regulate PD-L1 expression via the ubiquitin-proteasome pathway. To investigate whether IFIT1 influenced PD-L1 ubiquitination, we treated RKO cells with PD-L1 and ubiquitin plasmids and transfected them with IFIT1 siRNA and IFIT1 plasmid separately. The results showed that knockdown of IFIT1 could increase PD-L1 ubiquitination and overexpression of IFIT1 could decrease its ubiquitination (Fig. 5I-J). These results suggested that IFIT1 upregulated PD-L1 expression via decreasing its ubiquitination and degradation.

**IFIT1** was methylated by METTL3 and METTL14 and recognized by IGF2BP2/3

The above results suggested obvious m^A peaks in 3′UTR of IFIT1 mRNAs. We attempted to investigate which m^A regulators mediated m^A modifications in IFIT1 mRNAs and what functions m^A modifications have. In biological validation assays, IFIT1 mRNA and protein levels were markedly dampened in METTL3 and METTL14 knockdown CRC cells (Fig. 6A-D, Supplementary Figure S6A-B). Overexpression
Fig. 3. IFIT1 promoted CRC cell proliferation and participated in *F. nucleatum* induced CRC progression. A. CCK8 assays detecting cell growth of HT29 cells after transfection with IFIT1 siRNAs (mean± SD). B. Representative images of colony formation assays of HT29 cells after transfection with IFIT1 siRNAs. C. The summarized data of colony formation assays of HT29 cells after transfection with IFIT1 siRNAs (n=3, mean± SD, One-way ANOVA). D. Representative images of tumors in nude mice bearing HCT116 cells stably transfected with shIFIT1 lentivirus (n=10). E. Statistical analysis of tumor weights in D (n=10, two-tailed Student’s t-test). F. CCK8 assays detecting cell growth of HT29 cells after transfection with pcDNA3.1_IFIT1 (mean± SD). G. Representative images of colony formation assays of HT29 cells after transfection with pcDNA3.1_IFIT1. H. The summarized data of colony formation assays of HT29 cells after transfection with pcDNA3.1_IFIT1 (n=3, mean± SD, One-way ANOVA). I. Representative images of tumors in nude mice bearing HCT116 cells stably transfected with IFIT1 plasmid (n=7). J. Statistical analysis of tumor weights in I (n=7, two-tailed Student’s t-test). K. CCK8 assays detecting cell growth of HT29 cells transfected with IFIT1 siRNAs and treated with *F. nucleatum* (MOI=300) (mean± SD). L. Representative images of colony formation assays of HT29 cells transfected with IFIT1 siRNAs and treated with *F. nucleatum* (MOI=300). M. The summarized data of colony formation assays of HT29 cells transfected with IFIT1 siRNAs and treated with *F. nucleatum* (MOI=300) (n=3, mean± SD, One-way ANOVA).
of METTL3 and METTL14 increased IFIT1 expression (Fig. 6E-F, Supplementary Figure S6C-D). Since METTL3 and METTL14 positively regulated IFIT1 expression, we guessed that METTL3 and METTL14 may increase IFIT1 mRNAs stability. RNA stability assays showed that IFIT1 transcripts were less stable when METTL3 and METTL14 knockdown and more stable when METTL3 and METTL14 overexpressed in HCT116 cells (Fig. 6G-J).

M6A readers recognize and bind methylated mRNAs and mediate corresponding functions. IGF2BP family, including IGF2BP1, IGF2BP2 and IGF2BP3, are responsible for mRNA stability while other readers including YTHDF2 and YTHDF3 are responsible for mRNA decay [33,34]. Considering that METTL3 and METTL14 positively regulated IFIT1 expression and increased IFIT1 mRNA stability, we guessed that IGF2BP family may recognize IFIT1 mRNA and increase mRNAs stability. To investigate this question, we designed separately two siRNAs targeting IGF2BP1, IGF2BP2, and IGF2BP3 to knockdown their expression and detected IFIT1 mRNA expression alterations. The results showed that IFIT1 mRNA expression was remarkably suppressed after IGF2BP2/3 knockdown, but not IGF2BP1 knockdown in CRC cells (Fig. 6K, Supplementary Figure S6E). Furthermore, RNA stability assays showed that the half-life of IFIT1 transcripts was shortened after IGF2BP2 and IGF2BP3 knockdown HCT116 cells (Fig. 6L-M). RIP assays validated that IGF2BP2 and IGF2BP3 could directly bind the 3'UTR of IFIT1 mRNA (Fig. 6N-O). These data demonstrate that METTL3 and METTL14 methylated IFIT1 mRNAs and increased its expression through IGF2BP2/3-dependent mRNA stability regulation.

**F. nucleatum abundance, IFIT1 and PD-L1 expression were positively correlated in CRC patients**

Our results have suggested that *F. nucleatum* stimulated PD-L1 protein expression via m6A-modified IFIT1. The function of PD-L1 has been extensively studied in clinical samples. The functions of IFIT1 have not been studied in CRC samples, we measured IFIT1 expression in 34 pairs of CRC tissues and non-cancerous normal colon tissues in cohort 3. The results showed that IFIT1 was significantly upregulated in CRC tissues (Fig. 7A-B). Also, we measured IFIT1 expression in clinical CRC tissues using immunohistochemistry and found that IFIT1 expression was more highly expressed in TNM stage III/IV CRC than in stage I/II (Fig. 7C). Kaplan–Meier analysis showed higher IFIT1 expression correlated with worse overall survival based on TCGA COAD database (Fig. 7D). In cohort 2, there was a strong correlation between IFIT1 and PD-L1 expression (r=0.5853, p<0.0001) (Fig. 7E-F). *F. nucleatum* abundance was also positively correlated with IFIT1 expression (Fig. 7G). Our data showed that IFIT1 was higher expressed in tumor tissues and *F. nucleatum* abundance, IFIT1 and PD-L1 expression were positively correlated in CRC patients.

**Discussion**

Increasing evidence suggests that microorganisms are strongly associated with colorectal carcinogenesis [3,35]. *F. nucleatum* is reported as a pathogenic bacterium in CRC progression [36]. Emerging evidence has indicated that higher *F. nucleatum* amount was associated with an immunosuppressive
Fig. 5. IFIT1 mediated PD-L1 stabilization via regulating PD-L1 ubiquitination. A. Co-immunoprecipitation (Co-IP) assay detecting IFIT1 in RKO cell lysates immunoprecipitated with anti-PD-L1. B. RKO cells were overexpressed with FLAG_IFIT1 and HA_PD-L1. Co-IP assay detecting exogenous IFIT1 (FLAG) in RKO cell lysates immunoprecipitated with anti-HA. C-D. RKO cells transfected with control siRNAs and IFIT1 siRNAs were treated with cycloheximide (CHX) (50 μg/mL) at the indicated time points. Immunoblot of PD-L1 at indicated time points. E. Quantification of relative remaining PD-L1 in C-D. F. Immunoblot of PD-L1 in RKO cells transfected with IFIT1 siRNAs and treated with MG132 (10μM, 6 hours). G. Immunoblot of PD-L1 in RKO cells transfected with IFIT1 siRNAs and treated with CQ (25μM, 6 hours). H. Immunoblot of PD-L1 in RKO cells transfected with IFIT1 siRNAs and treated with 3-MA (6 hours). I-J. Immunoblot of cell lysates immunoprecipitated with anti-PD-L1 in RKO cells transfected with the indicated constructs.
Fig. 6. IFIT1 was methylated by METTL3 and METTL14 and recognized by IGF2BP2/3. A. PCR analysis of IFIT1 mRNA levels in HCT116 cells after transfection with METTL3 siRNAs (n=3, mean± SD, One-way ANOVA). B. PCR analysis of IFIT1 mRNA levels in HCT116 cells after transfection with METTL14 siRNAs (n=3, mean± SD, One-way ANOVA). C. Western blot assays detecting IFIT1 protein expression in HCT116 cells after transfection with METTL3 siRNAs. D. Western blot assays detecting IFIT1 protein expression in HCT116 cells after transfection with METTL14 siRNAs. E. PCR analysis of IFIT1 mRNA levels in HCT116 cells after transfection with pcDNA3.1_METTL3 and pcDNA3.1_METTL14 (n=3, mean± SD, two-tailed Student’s t-test). F. Western blot assays detecting IFIT1 protein expression in HCT116 cells after transfection with pcDNA3.1_METTL3 and pcDNA3.1_METTL14.

G. The detection of half-life(t1/2) of IFIT1 mRNAs in HCT116 cells transfected with control siRNAs and METTL3 siRNAs using real-time PCR, linear regression analysis. H. The detection of half-life(t1/2) of IFIT1 mRNAs in HCT116 cells transfected with control siRNAs and METTL14 siRNAs using real-time PCR, linear regression analysis. I. The detection of half-life(t1/2) of IFIT1 mRNAs in HCT116 cells with and without METTL3 overexpression using real-time PCR, linear regression analysis. J. The detection of half-life(t1/2) of IFIT1 mRNAs in HCT116 cells with and without METTL14 overexpression using real-time PCR, linear regression analysis. K. PCR analysis of IFIT1 mRNA level in HCT116 cells with IGF2BP1, IGF2BP2 and IGF2BP3 siRNAs transfection (n=3, mean± SEM, One-way ANOVA). L. The detection of half-life(t1/2) of IFIT1 mRNAs in HCT116 cells transfected with control siRNAs and IGF2BP2 siRNAs using real-time PCR, linear regression analysis. M. The detection of half-life(t1/2) of IFIT1 mRNAs in HCT116 cells transfected with control siRNAs and IGF2BP3 siRNAs using real-time PCR, linear regression analysis. N. The direct binding of IGF2BP2 with IFIT1 3’ UTR using Agarose electrophoresis and RIP-qPCR assays in HCT116 cells (n=3, mean± SD, two-tailed Student’s t-test). O. The direct binding of IGF2BP3 with IFIT1 3’ UTR using Agarose electrophoresis and RIP-qPCR assays in HCT116 cells (n=3, mean± SD, two-tailed Student’s t-test).
Fig. 7. *F. nucleatum* abundance, IFIT1 and PD-L1 expression were positively correlated in CRC patients. A. Immunohistochemical staining for IFIT1 in colorectal cancer tissues and para-cancerous normal tissues in cohort 3, scale bar, 100μm in 200X, 50μm in 400X. B. Statistical analysis of IFIT1 IHC score in cohort 3 (paired two-tailed Student’s t-test). C. TNM stages of samples in cohort 2 grouped by IFIT1 expression. D. Overall survival analysis of TCGA_COAD samples grouped by IFIT1 expression. E. Representative images of IFIT1 and PD-L1 staining of samples in cohort 2. F. Statistical analysis of the correlation of IFIT1 and PD-L1 expression (r=0.5853, p<0.0001, two-sided Pearson’s correlation). G. Statistical analysis of the correlation among *F. nucleatum* abundance and the expression of IFIT1 (r=0.4016, p<0.0001, two-sided Pearson’s correlation).

tumor environment [14]. In *Apc*^min/+^ mice, *F. nucleatum* gavage resulted in an expansion of myeloid cells in colon tumors, which represented tumor-promoting functions including myeloid-derived suppressor cells (MDSC) and tumor-associated macrophages (TAM). A population-based study suggested that higher *F. nucleatum* amount in tumors was associated with a lower CD3+ T cell density [13,37]. Also, *F. nucleatum* could inhibit NK cell activity through its virulence factor Fap2, which directly interacted with the inhibitory receptor TIGIT [15]. PD-L1 is an important immune checkpoint expressed on tumor cells and promotes tumor immune escape [38]. Here, we found *F. nucleatum* treatment promoted CRC progression and stimulated PD-L1 protein expression. The PD-L1 upregulation didn’t depend on *F. nucleatum* treatment activated PI3K pathway. Combined m^6^A-seq and RNA-seq identified m^6^A modified IFIT1 mediating *F. nucleatum* induced PD-L1 upregulation. IFIT1 regulated PD-L1 expression without altering mRNA level through interacting with PD-L1 protein and influencing its ubiquitination.

The regulatory mechanisms of PD-L1 expression are complex, including genetic alterations, transcriptional regulations, post-transcriptional regulations and post-translational modifications [10,39]. Emerging researches have indicated RNA m^6^A modification could regulated PD-L1 expression directly and indirectly [18,19]. PD-L1 can be directly modified with m^6^A modifications and indirectly regulated by m^6^A modified target genes. Here, we performed combined m^6^A-seq and RNA-seq in *F. nucleatum*-treated HT29 cells and identified m^6^A-modified IFIT1 mediated *F. nucleatum* stimulated PD-L1 upregulation. IFIT1 altered PD-L1 protein expression without affecting its mRNA levels. IFIT1 could interact with
PD-L1 protein and its overexpression inhibited PD-L1 ubiquitination and decreased its degradation.

IFIT1 is a member of interferon-stimulated genes (ISGs) which function as essential anti-viral proteins [27]. Emerging evidence has indicated its roles in carcinogenesis. IFIT1 and IFIT3 were reported as oncogenes in oral squamous cell carcinoma with the function to promote epithelial-mesenchymal transition and activate EGFR signaling [28]. IFIT1 and IFIT3 were also overexpressed in hepatocellular carcinoma tissues and mediated cell migration [29]. Here, we found IFIT1 was higher expressed in CRC tissues and its higher expression indicated worse survival. Inhibition of IFIT1 dampened CRC progression in vitro and in vivo. IFIT1 was found to interact with PD-L1 protein and regulate its ubiquitination. In addition, IFIT1 was modified with m^3^A peaks located in 3’UTR and the m^3^A levels were altered by F. nucleatum treatment. IFIT1 was methylated by METTL3 and METTL14 and m^3^A-modified IFIT1 was more stable dependent on IGFBP2/3 recognition. Our data suggested that IFIT1 may be a potential oncogene in CRC and IFIT1 may participate in tumor immune escape.

In conclusion, F. nucleatum promoted CRC progression in vitro and in vivo and upregulated PD-L1 expression. Combined m^3^A-seq and RNA-seq identified m^3^A-modified IFIT1 mediated F. nucleatum induced PD-L1 upregulation. IFIT1 was modified with m^3^A in 3’UTR and the m^3^A levels were altered by F. nucleatum treatment. IFIT1 served as a potential oncogene in CRC and regulated PD-L1 expression through regulating PD-L1 ubiquitination. Our work suggested that F. nucleatum promoted PD-L1 protein expression via m^3^A modification of IFIT1 and provided new aspects for understanding F. nucleatum mediated immune escape.

**Author contributions**

Y.Q.G., TH.Z., PPX., J.H., H.Y.C., and J.Y.F. designed the experiments. Y.Q.G., TH.Z., PPX., Y.C.W., and Y.J. performed the experiments and analyzed the data. Y.Q.G. wrote the manuscript. Y.-X.C., H.J., H.Y.C. and J.-Y.F. reviewed and revised the manuscript.

**Data availability**

The raw sequencing data are available in the Gene Expression Omnibus database under the accession number GSE191257. All other data are available in the article and supplemental information.

**Declaration of Competing Interest**

The authors declare no potential conflicts of interest.

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**Supplementary materials**

Supplementary material associated with this article can be found in the online version, at doi:10.1016/j.neo.2022.100850.

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