Lnc-EST12, which is negatively regulated by mycobacterial EST12, suppresses antimycobacterial innate immunity through its interaction with FUBP3

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Long noncoding RNAs (lncRNAs) have been implicated in the pathogenesis of intracellular pathogens. However, the role and mechanism of the important lncRNAs in Mycobacterium tuberculosis (M.tb) infection remain largely unexplored. Recently, we found that a secreted M.tb Rv1579c (an early secreted target with a molecular weight of 12 kDa, named EST12) protein activates NLRP3-gasdermin D (GSDMD)-mediated pyroptosis and plays a pivotal role in M.tb-induced immunity. In the present study, M.tb and the EST12 protein negatively regulated the expression of a key lncRNA (named Lnc-EST12) in mouse macrophages by activating the JAK2-STAT5a signaling pathway. Lnc-EST12, with a size of 1583 bp, is mainly expressed in immune-related organs (liver, lung and spleen). Lnc-EST12 not only reduces the expression of the proinflammatory cytokines IL-1β, IL-6, and CCL5/8 but also suppresses the NLRP3 inflammasome and GSDMD pyroptosis-IL-1β immune pathway through its interaction with the transcription factor far upstream element-binding protein 3 (FUBP3). The KH3 and KH4 domains of FUBP3 are the critical sites for binding to Lnc-EST12. Deficiency of mouse Lnc-EST12 or FUBP3 in macrophages increased M.tb clearance and inflammation in mouse macrophages or mice. In conclusion, we report a new immunoregulatory mechanism in which mouse Lnc-EST12 negatively regulates anti-M.tb innate immunity through FUBP3.

Keywords: Mycobacterium tuberculosis; lncRNA; Pyroptosis; Cytokines

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INTRODUCTION

Mycobacterium tuberculosis (M.tb) is the etiological agent of tuberculosis (TB), and TB remains a severe disease in humans worldwide. According to the World Health Organization Global Tuberculosis Report 2021, ~5.8 million new patients were diagnosed with TB, and ~1.51 million patients with TB died worldwide in 2020; additionally, the COVID-19 pandemic has aggravated the severity of TB [1]. A vaccine based on an attenuated live strain of Mycobacterium bovis, termed Bacillus Calmette-Guerin (BCG), is the only available TB vaccine and has limited protective efficacy in adults [2]. Early-stage disease, extrapolummary TB, M.tb/human immunodeficiency virus (HIV) co-infection, childhood TB, and multidrug-resistant TB are particularly problematic to diagnose and treat [1, 3]. Therefore, understanding the mechanism of M.tb infection and survival in the host has significant benefits for the prevention and control of TB and the development of new drugs. Macrophages are the primary target host cells of M.tb and play a central role in mycobacterial pathogenesis. Pathogenic mycobacteria have achieved broad evolutionary success through specialization as intracellular pathogens capable of achieving persistent infection by manipulating host macrophages to reside and replicate within them rather than being completely eradicated by innate defense mechanisms [4–6].

Regions of deletion (RD) are certain segments missing in the BCG genome relative to M.tb H37Rv or M. bovis [7, 8]. The BCG genome lacks 129 open reading frames from RD1 to RD16, of which RD1-2 regions are the most studied. Recently, studies of the RD1 region (e.g., RD1-encoding ESAT-6 and CFP-10) have revealed many genes that exert significant immunogenic and virulent effects on TB [9]. We recently found that the EST12 (Rv1579c) gene is located in RD3 and is present in both M.tb H37Rv and H37Ra. EST12, a secreted protein, activates NLRP3-gasdermin D (GSDMD)-mediated pyroptosis and plays a pivotal role in M.tb-induced immunity [10].

Long noncoding RNAs (lncRNAs) are a large class of nonprotein-coding transcripts with lengths longer than 200 nucleotides [11] that are involved in many biological processes [12–17]. To date, lncRNAs have been revealed to play important roles in regulating gene expression at the epigenetic, transcriptional, and
posttranscriptional levels [18–20]. For example, lncRNAs participate in immune responses by inhibiting the inflammatory process in macrophages and mice [21], suppressing the activation of abnormal inflammation, and regulating macrophage polarization [22]. LncRNAs exhibit tissue specific, cell-specific or even compartment-specific expression patterns. According to the relative location of lncRNAs to protein-encoding genes in the genome, they have been classified into sense lncRNAs, antisense lncRNAs, bidirectional lncRNAs, intronic lncRNAs, intergenic lncRNAs, and enhancer lncRNAs [11, 23]. Although many lncRNAs have been reported to be involved in a variety of diseases, few lncRNAs closely related to specific factors in M.tb have been discovered.

In the present study, we identified a key lncRNA, Inc-EST12, expressed in mouse macrophages that is negatively regulated by M.tb EST12/M.tb. Host Inc-EST12 negatively regulates anti-M.tb innate immunity by suppressing the production of the proinflammatory cytokines IL-1β, IL-6, and CCL5/8 and NLRP3-GSDMD-mediated pyroptosis through the transcription factor far upstream element-binding protein (FUBP).

RESULTS
Identification of Inc-EST12, a key lncRNA that is negatively regulated by M.tb EST12/M.tb in mouse macrophages
Recently, we found that EST12 (Rv1579c; MW: 12 kDa), a secreted protein of M.tb, activated an NF-κB/AP-1-NLRP3-GSDMD pyroptosis-IL-1β immune pathway in macrophages. We performed a microarray analysis in macrophages to further explore lncRNAs that are involved in M.tb infection. Purified recombinant EST12 protein was used as a stimulant to treat RAW264.7 macrophages for 0, 3, and 24 h (Fig. 1a and Supplementary Fig. 1a). Among the altered lncRNAs, Inc-EST12 (Aglilent Mouse lncRNA Microarray Probe No. mmu_8976) was found to be the most downregulated lncRNA after EST12 treatment for 24 h in macrophages (Fig. 1b). In addition, we confirmed that EST12 inhibited the expression of Inc-EST12 in a dose- and time-dependent manner using RT-qPCR (Fig. 1c, d). Fluorescence in situ hybridization (FISH) assays also showed that Inc-EST12 expression decreased in macrophages after treatment with EST12 protein (Fig. 1e, f). We observed that Inc-EST12 mainly accumulated in the cytoplasm, and a small amount was observed in the nucleus (Fig. 1e). Interestingly, Inc-EST12 expression was also decreased in bone marrow-derived macrophages (BMDMs) infected with various mycobacterial strains (bacteria: BMDMs = 10:1, 6 h) (decreased ~70% by H37Rv or H37RvΔEST12;EST12, ~60% by H37RvΔEST12, ~66% by H37Ra strains, ~40% by M. smegmatis and ~50% by M. smegmatis-EST12) (Supplementary Fig. 1b). H37RvΔEST12 and M. smegmatis, which do not encode the EST12 protein, still displayed decreased Inc-EST12 expression, indicating that other factors in these mycobacteria might also cause a decrease in Inc-EST12 expression in addition to EST12. Based on the above data, the EST12 protein of M.tb H37Rv remarkably suppresses Inc-EST12 expression in mouse macrophages.

EST12 negatively regulates Inc-EST12 expression by activating the JAK2-STAT5a signaling pathway
We next analyzed the mechanism underlying the downregulated expression of Inc-EST12 in EST12-stimulated macrophages. We constructed the 2340 bp promoter region of Inc-EST12 and different deletion mutants in the luciferase reporter pGL3 to obtain pGL3-Inc-EST12/deletion-Luc plasmids and transfected them into RAW264.7 cells. As shown in Fig. 1g, the luciferase activity was significantly decreased in the truncations containing the −1000 to −500 bp promoter sequence upon stimulation with EST12 (Fig. 1g). Further bioinformatics analysis of this sequence (−1000 to −500 bp) revealed two independent signal transducer and activator transcription 5a (STAT5a) binding regions (gamma-activated sequence-GAS motif) in LASAGNA-Search 2.0 (uconn.edu).

We performed an EMSA with the corresponding 35-nucleotide DNA probe (Supplementary Table 3) containing the STAT5a transcription factor binding region to determine whether STAT5a bound the Inc-EST12 promoter. Nuclear protein extracts from RAW264.7 cells pretreated with EST12 showed stronger binding to the probe (Fig. 1h, Lane 6 vs. Lane 2) than those from untreated RAW264.7 cells. The binding was abrogated when the extract was pretreated with a 10-fold excess of the unlabeled ‘competitor’ probe (Fig. 1h, Lanes 3 and 7), but the mutant competitor containing substitutions within the STAT5a binding sequence (shown in Supplementary Table 3) did not abolish binding, providing additional evidence for its specificity (Fig. 1h, Lanes 4 and 8). Furthermore, using a chromatin immunoprecipitation (ChIP)-qPCR assay, we observed significantly increased STAT5a binding to the Inc-EST12 promoter in the EST12-treated group compared with the untreated control and the isotype IgG control groups (Fig. 1i). The aforementioned data indicated that STAT5a directly binds to the Inc-EST12 promoter after EST12 treatment. Additionally, mutation of these STAT5a binding sites completely blocked the downregulation of Inc-EST12-luciferase expression induced by EST12 stimulation (Fig. 1j), suggesting that EST12 negatively regulated Inc-EST12 expression through a specific interaction between the Inc-EST12 promoter and STAT5a. As expected, STAT5a silencing also abolished the downregulation of Inc-EST12 by EST12 (Fig. 1k and Supplementary Fig. 1d), suggesting that EST12 stimulates STAT5a binding to the Inc-EST12 promoter (Fig. 1i) and inhibits Inc-EST12 expression (Fig. 1k). In addition, in the presence of JAK2 or STAT5a inhibitors, the EST12-induced decrease in Inc-EST12 expression was abrogated (Fig. 1i).

Furthermore, we observed an increase in cytoplasmic JAK2 phosphorylation (p-JAK2 at Y1007) and nuclear STAT5a phosphorylation (p-STAT5a at Y694) and a decrease in cytoplasmic p-STAT5a levels by performing nuclear/cytosol separation assays and western blot (WB) analysis (Fig. 1m). However, cytoplasmic and nuclear STAT5b phosphorylation (p-STAT5b) levels showed no difference (Supplementary Fig. 1e), suggesting that EST12 stimulates cytoplasmic JAK2 phosphorylation and the translocation of p-STAT5a from the cytoplasm to the nucleus.

Based on these data, EST12 promotes the activation of the p-JAK2-p-STAT5a pathway and the translocation of p-STAT5a from the cytoplasm to the nucleus. STAT5a binds to the Inc-EST12 promoter to repress Inc-EST12 expression.

Characteristics and distribution of Inc-EST12
By analyzing the genome sequence, we determined that Inc-EST12 represents an intergenic lncRNA located between the Igdcc3 and Parp16 genes and on the antisense strand of mouse chromosome 9 (http://genome.ucsc.edu/cgi-bin/hgBlat) (Fig. 2a). An analysis of the tissue distribution of Inc-EST12 in mice showed that Inc-EST12 is expressed at high levels in the liver, followed by the lung and spleen (Fig. 2b). Lnc-EST12 is expressed at low levels in the kidney and brain and is expressed at very low levels in the heart and muscle (Fig. 2b), suggesting that Inc-EST12 is mainly expressed in immune-related organs (liver, lung and spleen). We also found that EST12 reduced Inc-EST12 expression in both mouse macrophages (including RAW264.7 cells and BMDMs) and DCs (bone marrow-derived dendritic cells, BMDCs) (Fig. 2c). According to the known lncRNA sequence, we determined the full-length sequence of Inc-EST12 to be 1583 bp in size by performing a rapid amplification of cloned CDNA ends (RACE) experiment (Fig. 2d and Supplementary Fig. 2a). Structural prediction analysis showed that full-length Inc-EST12 forms a unique spatial configuration (Supplementary Fig. 2b). A NCBI
Fig. 1  Mycobacterial EST12 suppresses lnc-EST12 expression in macrophages. a Schematic diagram of the lncRNA screening process in RAW264.7 macrophages after mycobacterial EST12 protein stimulation. b Heatmap of lncRNA expression obtained from microarray data. c, d RT-qPCR analysis of lnc-EST12 expression in RAW264.7 cells treated with different concentrations of EST12 for 12 h (c) or with EST12 (2 μM) for different time points (d). e FISH analysis of RAW264.7 cells treated with EST12 or PBS for 12 h using a Cy3-labeled lnc-EST12 probe (red) and DAPI nuclear staining (blue). f Quantification of the lnc-EST12 fluorescence intensity in each cell in (e). g Dual luciferase reporter assay of the activity of the lnc-EST12 promoter. RAW264.7 cells were cotransfected with the lnc-EST12-luciferase reporter construct (with the lnc-EST12 promoter region −2260 to +80 or its truncations) and the internal control plasmid pRL-TK, followed by EST12 treatment (2 μM, 24 h). The relative luciferase activities are presented as a percentage of values relative to the control group. h EMSA for analysis of STAT5a binding to the lnc-EST12 promoter. The STAT5a complex is indicated by horizontal arrows. i ChIP-qPCR analysis of the binding of STAT5a to the lnc-EST12 promoter region. j Dual luciferase reporter assay of the activity of the lnc-EST12 promoter. RAW264.7 cells were cotransfected with the lnc-EST12-luciferase reporter construct (with the lnc-EST12 promoter region -1000 to +80 or its mutants) and the internal control plasmid pRL-TK, followed by EST12 treatment (2 μM, 24 h). k RT-qPCR analysis of lnc-EST12 expression in RAW264.7 cells transfected with sh-STAT5a or sh-scramble, followed by EST12 treatment, as indicated. l RT-qPCR analysis of lnc-EST12 expression in RAW264.7 cells treated with EST12 (2 μM), gandotinib (JAK2 inhibitor, 10 μM) or STAT5a-IN-1 (STAT5a inhibitor, 10 μM). m WB analysis of the levels of phosphorylated JAK2 and STAT5a in the cytoplasm and nucleus of RAW264.7 cells stimulated with EST12 (2 μM) for different durations. Data in (c, d, g–l) are presented as the means ± SD of three independent experiments. **p < 0.01; ***p < 0.001; ****p < 0.0001. ns not significant. Two-tailed unpaired Student’s t test was used to calculate statistical significance (for g, i, j, k).
BLAST search of the human RNA-seq database (https://blast.ncbi.nlm.nih.gov/Blast.cgi) using the lnc-EST12 sequence as the query displayed no homologous RNA sequence of human origin with significant similarity. Due to the species specificity of most lncRNAs, the homology between mouse lncRNAs and their corresponding human lncRNAs is very low [22, 24, 25].

In addition, the overexpression or knockdown of lnc-EST12 had no effect on the expression of its neighboring genes Igdc3 and Parp16 (Fig. 2e). FISH assays showed that lnc-EST12 was primarily located in the cytoplasm of macrophages (Fig. 2f). Furthermore, we used a nuclear/cytosol separation assay and found that ~96% of lnc-EST12 was distributed in the cytoplasm of mouse macrophages (Fig. 2g). Thus, mouse lnc-EST12 is mainly expressed in immune-related organs (liver, spleen and lung), macrophages and DCs.

**Lnc-EST12 overexpression suppresses inflammatory cytokine expression, while lnc-EST12 KO exerts the opposite effects**

We constructed lnc-EST12 KO mice by deleting the 3 kb genomic locus containing lnc-EST12 to further investigate the function of lnc-EST12 (Supplementary Fig. 3a). We confirmed the deletion of lnc-EST12 from the genomic DNA of lnc-EST12 KO mice by Sanger sequencing and genomic PCR (Supplementary Fig. 3b, c). We performed unbiased transcriptome profiling using RNA-seq in wild-type (WT) and lnc-EST12 KO BMDMs stimulated with EST12 proteins for 0, 6 and 24 h to assess the effect of lnc-EST12 deficiency on gene expression in macrophages (Fig. 3a).

Gene ontology (GO) enrichment analysis showed that cytokine activity, chemokine activity, cytokine receptors, and chemokine receptors (CCRs) were significantly overrepresented in the differentially expressed genes between lnc-EST12 KO and WT BMDMs (Supplementary Fig. 3d). We further confirmed that lnc-EST12 knockout induced the expression of the inflammatory cytokines IL-1β, IL-6 and CCL5/8 in lnc-EST12 KO BMDMs treated with EST12 using RT-qPCR and ELISA (Fig. 3b, c, Supplementary Fig. 3g, h). The levels of IL-1β and IL-6 were increased 1.5- to 3-fold in lnc-EST12-KO BMDMs compared to WT BMDMs at both the mRNA (Fig. 3b) and protein levels (Fig. 3c) at 24 h after EST12 stimulation. Overexpression of lnc-EST12 in lnc-EST12 KO BMDMs by lentivirus infection (LV-lnc-EST12) inhibited the expression of the IL-1β and IL-6 mRNAs and proteins (Fig. 3d, e, Supplementary Fig. 3e). Similar results were observed for CCL5 and CCL8 (Supplementary Fig. 3i, j). As Gm40527 partly overlaps with lnc-EST12, we further determined the possible impact of Gm40527 on inflammatory cytokine expression. We reconstructed the recombinant lentivirus LV-shGm40527 by knocking down a specific fragment of Gm40527 with no overlap with lnc-EST12 (Supplementary Fig. 3a, b, shGm40527 knockdown position is indicated by black triangle). Our results showed that knockdown of Gm40527 using LV-shGm40527 significantly decreased Gm40527 mRNA expression but did not affect the expression of lnc-EST12 and cytokines (IL-1β and IL-6) and chemokines (CCL5/8), as determined using RT-qPCR (Supplementary Fig. 3a, f).
Based on the data presented above, Inc-EST12 suppresses inflammatory cytokine expression in mouse macrophages, while Inc-EST12 KO exerts the opposite effects.

**Lnc-EST12 binds the KH3 and KH4 domains of FUBP3**

Next, we explored the host proteins interacting with Inc-EST12. Biotin-labeled Inc-EST12 was used for the pulldown of RAW264.7-cell lysates, followed by SDS–PAGE analysis (Fig. 4a). Differentially abundant protein bands were subjected to mass spectrometry (MS) analysis (Supplementary Fig. 4a). The specific proteins were verified by Inc-EST12 RNA pulldown and WB analysis, and FUBP3 was confirmed to be the specific protein for binding Inc-EST12 (Fig. 4b). We confirmed the interaction of Inc-EST12 with FUBP3 using RIP (RNA Binding Protein Immunoprecipitation) and confocal colocalization analysis (Fig. 4c, Supplementary Fig. 4b), and more colocalization between Inc-EST12 and FUBP3 was observed in the control group than in the EST12 group. Decreased expression of both cytoplasmic and nuclear Inc-EST12 was observed in the EST12 group. Enter the nucleus (Supplementary Fig. 4b), similar to the p-JAK-p-Stat3 pathway, and nuclear localization of FUBP3 increased in the EST12 group. Decreased expression of Inc-EST12 might reduce the interaction between Inc-EST12 and FUBP3. We also used biotin-labeled full-length Inc-EST12 and truncated Inc-EST12 constructs, as indicated, for RNA pulldown of RAW264.7-cell lysates. Only full-length Inc-EST12 bound to FUBP3, and each truncated Inc-EST12 fragment lost its binding ability (Fig. 4d).

The FUBP3 protein contains four KH (K homology) repeats (from KH1 to KH4) [26]. Therefore, FUBP3 with different KH domain mutations, Mut A (with KH1 to KH4 mutants), Mut B (KH1 mutant), Mut C (KH2 mutant), Mut D (KH3 mutant) and Mut E (KH4 mutant), were constructed (Supplementary Fig. 4c, d). RAW264.7 cells were transfected with pcDNA3.1 expression plasmids carrying the full-length FUBP3 sequence and its various KH site-specific mutants with HA tag for 24 h, and then cell lysates were pulled down with Inc-EST12 RNA, followed by WB analysis with an anti-HA tag monoclonal antibody (mAb). The full-length FUBP3 protein and its various KH mutant constructs were expressed at high levels with a molecular weight of 63 kDa, and we observed that Inc-EST12 only bound FUBP3 and FUBP3-Mut B/C but no longer bound FUBP3-Mut A, D and E (Fig. 4e, f). These data suggest that the KH3 and KH4 domains of FUBP3 are the critical binding sites for Inc-EST12.

The *Inc-EST12-FUBP3 interaction promotes FUBP3 binding to the promoters of proinflammatory cytokines and then suppresses proinflammatory cytokine expression*

ChIP was performed, followed by RT-qPCR analysis, to determine whether FUBP3 functions as a transcription factor to regulate proinflammatory cytokine expression by binding to the promoter regions of cytokines. The results showed that the FUBP3 protein indeed bound to the promoter regions of IL-1β, IL-6, CCL5, and CCL8 (Fig. 4g). Furthermore, the binding of FUBP3 to these promoter regions decreased when BDMDs were stimulated with the EST12 protein (Fig. 4h) and increased after Inc-EST12 overexpression in both resting and EST12-treated BMDMs (Fig. 4i). We also constructed a lentivirus expression vector by knocking down FUBP3 and confirmed the ~70% knockdown efficiency of FUBP3 protein expression by Lv-sh-FUBP3 in BMDMs using WB analysis (Supplementary Fig. 4e). Notably, Inc-EST12 suppressed the expression of IL-1β, IL-6, and CCL5/8 mRNA and protein in BMDMs stimulated with EST12, as evidenced by the RT-qPCR and ELISA results; however, after FUBP3 knockdown, the expression levels of IL-1β, IL-6, and CCL5/8 were increased (Fig. 4j, k and Supplementary Fig. 4f, g), suggesting that Inc-EST12 suppresses proinflammatory cytokine expression through FUBP3.

Based on the findings described above, our data revealed that Inc-EST12 enhanced the binding of FUBP3 to chromatin at the promoters of proinflammatory cytokines (IL-1β, IL-6, etc.), thus suppressing inflammatory cytokine expression. We postulate that Inc-EST12 binding to FUBP3 promotes the translocation of FUBP3 from the cytoplasm to the nucleus, and Inc-EST12 per se does not enter the nucleus (Supplementary Fig. 4b), similar to the p-JAK-p-Stat3 pathway.
Fig. 4 Lnc-EST12 promotes FUBP3 binding to the promoters of proinflammatory cytokines and then suppresses cytokine expression. 

a SDS–PAGE analysis of proteins purified from the in vitro binding assay using biotinylated Inc-EST12 or antisense control RNA and RAW264.7-cell extracts. Differentially abundant protein bands were subjected to mass spectrometry analysis. b WB confirmed the Inc-EST12 and FUBP3 interaction in vitro. RAW264.7-cell lysates were incubated with biotin-labeled Inc-EST12 and detected using anti-FUBP3, anti-FUBP2, and anti-HNRNPD antibodies. c RIP-qPCR analysis was performed to determine the binding of Inc-EST12 to FUBP3 in RAW264.7 cells. d Schematic of Inc-EST12 deletion mutants used in RNA–protein binding assays. The RNA–protein binding assay was performed using biotinylated full-length or deletion mutants of Inc-EST12 and the lysate from RAW264.7 cells. e Schematic of HA-FUBP3 deletion mutants used in RNA–protein binding assays. f FUBP3 binds Inc-EST12 through the KH3 and KH4 domains. The RNA–protein binding assay was performed using biotinylated lnc-EST12 and overexpressed HA-FUBP3 and mutants in RAW264.7 cells. g The ChIP–qPCR assay of the binding of the FUBP3 protein to the promoter region of cytokines. ChIP-purified DNA was analyzed using qPCR with primers targeting the promoter site of the indicated genes. h ChIP–qPCR analysis of FUBP3 binding to the promoter region of cytokines in BMDMs treated with or without EST12 for 24 h. i ChIP–qPCR analysis of FUBP3 binding to the promoter region of cytokines in BMDMs expressing ectopic Inc-EST12 or LV-Ctl (control) that were treated with or without EST12 for 24 h. j, k The mRNA (j) and protein (k) levels of IL-1β and IL-6 in BMDMs infected with LV-Ctl, LV-lnc-EST12, LV-lnc-EST12 with LV-sh-scramble or LV-sh-FUBP3, followed by stimulation with EST12 (2 μM) for 24 h. LV, lentivirus. Ctl, control. Data are presented as the means ± SD of three independent experiments. *p < 0.05; **p < 0.01; ***p < 0.001; ****p < 0.0001. ns not significant. Two-tailed unpaired Student's t test was used to calculate the statistical significance of the data shown in (c) (vs. IgG), g (vs. IgG), h (vs. Control), and (j) (vs. LV-Ctl), and one-way ANOVA followed by Sidak’s multiple comparisons test was used to calculate the statistical significance of the data shown in (j, k).
STAT5 interaction promoting p-STAT5 translocation into the nucleus, but p-JAK does not enter the nucleus.

Lnc-EST12 suppresses EST12-induced NLRP3 inflammasome assembly and pyroptosis via FUBP3
Previously, we found that the EST12 protein promoted NLRP3-GSDMD-mediated pyroptosis in macrophages [10]. Here, the expression levels of the constitutive proteins NLRP3, ASC and Caspase-1 and ASC were all increased in lnc-EST12 KO BMDMs compared to WT BMDMs (Fig. 5a), suggesting that lnc-EST12 suppresses the assembly of the NLRP3 inflammasome. The level of the N-terminal fragment of GSDMD (indicating GSDMD cleavage) was increased in lnc-EST12 KO BMDMs compared to WT BMDMs (Fig. 5b). In contrast, the cleavage and activation of GSDMD were inhibited after the overexpression of Inc-EST12 in lnc-EST12 KO BMDMs by lentivirus infection (LV-Inc-EST12) (Fig. 5c). WT, Inc-EST12 KO and GSDMD-KO BMDMs were prepared with isolated WT and GSDMD-KO mouse bone marrow cells by incubating them with 10% FBS plus M-CSF for 7 days. WT, Inc-EST12 KO and GSDMD-KO BMDMs infected with LV-Ctl or LV-Inc-EST12 for 24 h and treated with the EST12 protein (2 μM) were analyzed for GSDMD activation following EST12 stimulation. BMDMs were infected with LV-Ctl or LV-shFUBP3 and stimulated with the EST12 protein (2 μM) for the indicated time, and then GSDMD activation was detected using WB. Effects of FUBP3 knockdown on macrophage cytotoxicity induced by EST12. BMDMs infected with LV-Ctl, LV-Inc-EST12, or LV-shFUBP3 for 24 h and treated with the EST12 protein (2 μM) were analyzed for cytotoxicity using an LDH assay. LV lentivirus. Ctl control. lnc KO lnc-EST12 KO. Data are presented as the means ± SD of three independent experiments. *p < 0.05; **p < 0.01; ***p < 0.001. One-way ANOVA followed by Sidak’s multiple comparisons test was used to calculate the statistical significance of the data shown in (d, f).

Fig. 5 Lnc-EST12 inhibits EST12-induced NLRP3 inflammasome assembly and pyroptosis via FUBP3. a WT and Inc-EST12 KO BMDMs were treated with the EST12 protein (2 μM) and subjected to IP with an anti-NLRP3 antibody and IB with antibodies against NLRP3, caspase-1 and ASC.

b WT and Inc-EST12 KO BMDMs treated with the EST12 protein (2 μM) were analyzed for GSDMD activation using WB.

c Inc-EST12 KO BMDMs were prepared with isolated WT and GSDMD-KO mouse bone marrow cells by incubating them with 10% FBS plus M-CSF for 7 days. WT, Inc-EST12 KO and GSDMD-KO BMDMs infected with LV-Ctl or LV-Inc-EST12 for 24 h and treated with the EST12 protein (2 μM) were analyzed for GSDMD activation following EST12 stimulation. BMDMs were infected with LV-Ctl or LV-shFUBP3 and stimulated with the EST12 protein (2 μM) for the indicated time, and then GSDMD activation was detected using WB.

d Effects of Inc-EST12 knockout and overexpression on the cytotoxicity of BMDMs stimulated with EST12. WT BMDMs and GSDMD-KO BMDMs were prepared with isolated WT and GSDMD-KO mouse bone marrow cells by incubating them with 10% FBS plus M-CSF for 7 days. WT, Inc-EST12 KO and GSDMD-KO BMDMs infected with LV-Ctl or LV-Inc-EST12 for 24 h and treated with the EST12 protein (2 μM) were analyzed for GSDMD activation following EST12 stimulation.

e Effects of FUBP3 knockdown on GSDMD activation following EST12 stimulation.

f Effects of FUBP3 knockdown on macrophage cytotoxicity induced by EST12. BMDMs infected with LV-Ctl, LV-Inc-EST12, or LV-shFUBP3 for 24 h and treated with the EST12 protein (2 μM) were analyzed for cytotoxicity using an LDH assay.
Based on these results, the Inc-EST12 and FUBP3 interaction plays an important role in Inc-EST12-mediated inhibition of proinflammatory cytokine expression and pyroptosis.

**Deficiency of Inc-EST12 or FUBP3 in macrophages increases both *M.tb* clearance and inflammation**

Next, we examined whether Inc-EST12 KO macrophages and mice would show increased levels of inflammatory cytokines and decreased mycobacterial survival in mouse macrophages and in vivo. Both WT and Lnc-EST12 KO BMDMs were infected with *M. tb* H37Rv for 6 h. Extracellular bacteria were removed by washes with PBS, and intracellular bacteria were counted. Knockout of Inc-EST12 reduced mycobacterial CFUs and survival in mouse macrophages (Fig. 6a). Overexpression of Inc-EST12 increased the mycobacterial CFUs in a FUBP3-dependent manner, since FUBP3 silencing in BMDMs reversed this effect (Fig. 6b). WT and Inc-EST12 KO BMDMs were infected with WT H37Rv, H37RvΔEST12, and H37RvΔEST12:EST12. As expected, we observed increased levels of the inflammatory cytokines IL-1β and IL-6 and decreased mycobacterial CFUs in Inc-EST12 KO BMDMs compared with WT BMDMs (Supplementary Fig. 5e–g), suggesting that deficiency of Inc-EST12 in BMDMs increases both *M.tb* clearance and inflammation.

Furthermore, *M.tb* H37Ra, a slow-growing mycobacterium, was administered to mice through an intranasal infection at a dose of 10⁷ CFUs/mouse [27–30]. Thirty days after infection, we observed significantly increased serum IL-1β, IL-6, CCL5, and CCL8 levels in Inc-EST12 KO mice (Fig. 6c and Supplementary Fig. 5a). In addition, similar results for the expression of these cytokines were observed in the spleen, liver and lung organs of the Inc-EST12 KO mice (Fig. 6d–f and Supplementary Fig. 5b–d). Through the bacterial colony counting experiment, we observed decreased colony numbers of H37Ra in the lung and spleen organs of Inc-EST12 KO mice (Fig. 6g). Acid-fast staining of lung tissues from H37Ra-infected mice also showed decreased numbers of acid-fast positive bacilli in Inc-EST12 KO mice compared with WT mice (Fig. 6h). We did not find classic TB lesions in the lungs of H37Ra-infected mice (Fig. 6i), but we could observe obvious inflammatory responses in these mouse infection models. The histopathological analysis of alveolar tissue from the Inc-EST12 KO mice showed increased inflammatory lymphocyte infiltration and smaller intact alveolar spaces compared with the WT mice (Fig. 6i). Therefore, overexpression of Inc-EST12 increases the survival of *M.tb*, while deficiency of Inc-EST12 or FUBP3 in mouse macrophages increases *M.tb* clearance and inflammation.

**Lnc-EST12 promotes mycobacterial growth and suppresses IL-1β and IL-6 production after adoptive transfer in animal experiments**

In an adoptive transfer experiment, clonardate liposomes eliminated both macrophages and DCs [31–33]. We found that ~88% of F4/80⁺ macrophages were depleted in the mouse lungs after injection of clonardate liposomes for 3 days (the macrophage percentage in the lungs decreased from 11.7% in the liposome control group to 1.36% in the clonardate depletion group), and ~80% of F4/80⁺ macrophages were restored in the mouse lung after injection of clonardate liposomes plus adoptive transfer of 5 × 10⁶ mouse MDMs (increased from 1.36% in the clonardate depletion group to 9.43% in the clonardate depletion plus adoptive transfer group) by flow cytometry (FCM) analysis (Supplementary Fig. 5i). We subsequently adoptively transferred WT or Inc-EST12 KO BMDMs into clonardate liposome-treated WT C57BL/6 mice to examine Inc-EST12 expression in lung F4/80⁺ macrophages by RT-qPCR. We found much less Inc-EST12 expression in the group with adoptive transfer of Inc-EST12 KO BMDMs than in the group with adoptive transfer of WT BMDMs (Supplementary Fig. 5h). Then, the mice were intravenously (i.v.) infected with *M. smegmatis* (Ms, a fast-growing mycobacterium) and Ms-EST12 (Ms carrying pMV261-EST12 that can express EST12 protein) for 3 days (Fig. 7a). EST12 was confirmed to be expressed in Ms-EST12 but not in WT Ms by WB (Supplementary Fig. 5j). Ms-EST12 significantly increased IL-1β and IL-6 production (Fig. 7b, c) and decreased bacterial CFUs in the lung, liver and spleen organs (Fig. 7d) in the group with adoptive transfer of Inc-EST12 KO BMDMs compared to the group with adoptive transfer of WT BMDMs. These adoptive transfer experiments strongly suggest that Inc-EST12 promotes mycobacterial growth by suppressing IL-1β and IL-6 production in vivo.

**DISCUSSION**

LncRNAs are widely involved in various physiological activities. LncRNA-Mirt2 has been reported to inhibit the excessive activation of LPS-induced inflammation and regulate inflammatory damage in the body to achieve a reasonable level [22]. LncRNA-Neat1 increases the activation of caspase-1, thereby promoting the production of IL-1β and the pyroptosis of macrophages [34]. LncRNAs also play important roles in the infection process of *M.tb* [35, 36]. LncRNA-CD244, which is highly expressed in CD8⁺ T cells of patients with active TB, inhibits the production of IFN-γ and TNF-α in CD8⁺ T cells and inhibits host clearance of *M.tb* [37]. Although several IncRNAs related to TB have been discovered in recent years, the role and mechanism of the important IncRNAs related to the key factors of *M.tb* during infection remain largely unexplored. Therefore, a particularly important goal is to discover a lncRNA related to an *M.tb* key factor/protein and to explore its function and mechanism in vitro and in vivo.

Previously, we found that EST12, a secreted protein of *M.tb*, activated an NF-κB/AP-1-LRP3-GSDMD pyroptosis-IL-1β immune pathway in macrophages [10]. In this study, we identified a key lncRNA, Inc-EST12, which was negatively regulated by the *M.tb* EST12 protein through the activation of the p-JAK2-p-STAT5a signaling pathway in mouse macrophages. We also documented that EST12-stimulated STAT5a binding to the Inc-EST12 promoter and then repressed Inc-EST12 expression. Similarly, other reports also showed that STAT5a binds and represses target gene expression [38]. Inc-EST12 is more likely to act as a general responder to mycobacterial infection, since the host immune responses are activated following the decreased expression of Inc-EST12, and its expression is downregulated by either ETS12-expressing or ETS12-nonexpressing mycobacterial strains (Supplementary Fig. 1b). EST12 promotes the activation of the p-JAK2-p-STAT5a pathway and the translocation of p-STAT5a from the cytoplasm to the nucleus. STAT5a binds to the Inc-EST12 promoter to repress Inc-EST12 expression (Supplementary Fig. 6). Importantly, Inc-EST12 suppressed antimycobacterial innate immunity through its interaction with FUBP3. Notably, Inc-EST12 binding to FUBP3 promoted the binding of the transcription factor FUBP3 to the promoter regions of cytokines such as IL-1β and IL-6, thus suppressing the expression of these cytokines. Lnc-EST12 also suppressed EST12-induced NLRP3 inflammasome assembly and GSDMD-mediated pyroptosis via FUBP3. These results indicate that Inc-EST12, which is negatively regulated by mycobacterial EST12, represses the innate inflammatory immune response and pyroptosis through its interaction with FUBP3 (Supplementary Fig. 6).

The expression levels of various cytokines and chemokines play important roles in *M.tb* infection [39]. Among them, *M.tb* infection activates the NLRP3 inflammatory pathway in macrophages, promoting the secretion of IL-1β and IL-18 [40, 41]. IL-1β directly kills *M.tb* in macrophages, upregulates TNF-α expression and promotes cell apoptosis. IL-6 exerts a dual immunomodulatory effect on *M.tb* infection [42, 43]. Generally, IL-6 is presumed to induce the proliferation and differentiation of B cells and T cells, activate macrophages, and enhance the killing effect of NK cells, thereby resisting *M.tb* infection. CCL5 contributes to early, protective, *M.tb*-specific immunity, primarily by recruiting early
Fig. 6  Lnc-EST12 knockout in macrophages increased \( M. tb \) clearance and inflammation in mice.  

a) Bacterial burden in WT and Inc-EST12 KO BMDMs infected with \( M. tb \) H37Rv at an MOI of 10:1 (bacteria: cells = 10:1) for 6 h. Extracellular bacteria were removed by washes with PBS, and intracellular bacteria were counted. 

b) Effects of FUBP3 knockdown on \( M. tb \) survival in macrophages. BMDMs were infected with LV-Ctl, LV-lnc-EST12, LV-lnc-EST12 with LV-sh-scramble or LV-sh-FUBP3 for 24 h and then infected with \( M. tb \) H37Ra at an MOI of 10:1 (bacteria: cells = 10:1) for 6 h. 

c–f) IL-1\( \beta \) and IL-6 levels in the serum (c), spleen (d), liver (e), and lung (f) of WT and Inc-EST12 KO mice i.n. infected with H37Ra (5 \( \times \) 10\(^4\) CFUs/mouse) for 30 days were analyzed using ELISA. Data are presented as the means ± SD (\( n = 4–6 \) mice per group). 

g) Bacterial burden in the lung and spleen from WT and Inc-EST12 KO mice infected with H37Ra. Data are presented as the means ± SD (\( n = 4–6 \) mice per group). 

h) Acid-fast staining (h) and HE staining (i) of lung tissues from WT or Inc-EST12 KO mice infected with H37Ra or uninfected controls. LV lentivirus. Ctl control. lnc KO lnc-EST12 KO. Data are presented as the means ± SD of three independent experiments. *\( p < 0.05 \); **\( p < 0.01 \). ns not significant. Two-tailed unpaired Student’s t test was used to calculate the statistical significance of the data shown in (a–g) (vs. WT).
Pyroptosis is a type of proinflammatory cell death that is characterized by the formation of inflammasomes, which are protease-proficient complexes that activate caspase-1, leading to the processing and release of inflammatory cytokines, such as IL-1β and IL-18. This process is often associated with the destruction of the cell's cytoplasmic contents and the formation of pores in the cell membrane, leading to cell lysis. Pyroptosis plays a crucial role in host defense against bacterial infections, as it helps to eliminate infected cells and prevents the spread of pathogens.

Studies have shown that pyroptosis is induced by various proinflammatory stimuli, including bacterial components, cytokines, and danger signals. This process is mediated by the activation of NLRP3, a member of the NLR family of intracellular sensors that respond to pathogen-associated molecular patterns (PAMPs) and damage-associated molecular patterns (DAMPs). The activation of NLRP3 leads to the formation of inflammasomes, which activate caspase-1 and lead to the processing of pro-IL-1β and pro-IL-18 into their mature, active forms.

Inflammasomes are composed of NLRP3, ASC, and pro-caspase-1. The assembly of the inflammasome leads to the activation of caspase-1, which cleaves pro-IL-1β and pro-IL-18 into their active forms. This process is further regulated by other proteins, such as FUBP3, which has been shown to interact with NLRP3 and modulate its activity.

FUBP3 is a family of single-stranded DNA-binding proteins that are involved in various cellular processes, including transcription, DNA repair, and cell cycle regulation. Studies have shown that FUBP3 binds to the 5' untranslated region (5'UTR) of EV71, a picornavirus, and inhibits its translation. In addition, FUBP3 has been shown to interact with NLRP3 and modulate its activity in response to proinflammatory stimuli.

The interaction between FUBP3 and NLRP3 might play a role in the regulation of pyroptosis. Studies have shown that the lnc-EST12-FUBP3 interaction promotes FUBP3 binding to NLRP3, leading to the activation of NLRP3 and pyroptosis. This interaction might be involved in the regulation of pyroptosis in response to proinflammatory stimuli.

In summary, pyroptosis is a type of proinflammatory cell death that plays a crucial role in host defense against bacterial infections. The activation of NLRP3 leads to the formation of inflammasomes, which activate caspase-1 and lead to the processing of pro-IL-1β and pro-IL-18 into their active forms. The interaction between FUBP3 and NLRP3 might play a role in the regulation of pyroptosis.
Bacterial culture

*M. tuberculosis* H37Rv [strain American Type Culture Collection (ATCC) 27294] and H37Ra [strain American Type Culture Collection (ATCC) 25177] were maintained on Lowenstein-Jensen medium. *M. smegmatis* (strain ATCC 19420), *Escherichia coli* DH5α (strain ATCC 25922), and *E. coli* BL-21 (strain ATCC BAA-1025) were propagated from laboratory stocks (School of Medicine, Wuhan University, Wuhan, China). *E. coli* DH5α and BL-21 (DE3) were grown in flasks using LB medium. The mycobacterial strains were grown in Middlebrook 7H9 broth (BD Biosciences, USA) supplemented with 10% oleic acid–albumin–dextrose–catecal (BD Biosciences, USA) and 0.05% Tween 20 (Sigma–Aldrich, USA) or on Middlebrook 7H10 agar (BD Biosciences, USA) supplemented with 10% OADC [83].

Agilent mouse IncRNA microarray assay

The Agilent Mouse IncRNA Microarray (4x180K, Design ID: 049801) was analyzed by Shanghai OE Biotech Co. Ltd. of China. Sample labeling, microarray hybridization and washing were performed according to the manufacturer’s standard protocols. Briefly, total RNA was extracted from RAW264.7 cells in the control group and EST12 treatment group with TRIzol (Invtrogen, USA), transcribed to double-strand cDNAs, synthesized into cRNAs and labeled with cyanine-3-CTP. The labeled cRNAs were hybridized to the microarray. After washing, the arrays were scanned using an Agilent Scanner G2505C (Agilent Technologies).

Reverse transcription quantitative PCR (RT-qPCR)

Total RNA was isolated using TRIzol reagent (Invtrogen, USA). One microgram of RNA was used to synthesize cDNAs with the RT Master Mix (Toyobo, Japan). Real-time PCR was performed using SYBR Green real-time PCR mix (Toyobo, Japan) with specific primers (Supplementary Tables 1, 2). The RT-qPCRs were cycled using an ABI StepOnePlus instrument (Applied Biosystems, USA) under standard cycling conditions. Target gene expression levels were normalized to glyceraldehyde-3-phosphate dehydrogenase (GAPDH). Relative RNA levels were calculated using the comparative cycle threshold (CT) method (2−ΔΔCT method), where CT indicates the amplification cycle number at which the fluorescence generated within a reaction increases above a defined threshold fluorescence, and ΔΔCT = experimental sample (CTtarget gene − CTGAPDH) − control sample (CTtarget gene − CTGAPDH).

Enzyme-linked immunosorbent assay (ELISA)

The concentrations of IL-6, IL-1β, CCL5, and CCL8 were determined using ELISA kits according to the manufacturer’s instructions (4A Biotech, China). Briefly, BMDMs were stimulated with EST12 for 0, 6, and 24 h. Cell debris was removed by centrifugation, and the cell supernatant was collected for ELISA. For the detection of cytokines in animal serum and immune tissues, mice were infected with H37Ra for 30 days and then sacrificed to obtain serum and liver, spleen, and lung tissues. The tissues were ground and centrifuged to obtain supernatants for ELISA.

Electrophoretic mobility shift assay (EMSA)

The EMSA was performed using chemiluminescent EMSA kits according to the manufacturer’s instructions (Beyotime, China). Briefly, synthetic oligonucleotides containing the STAT5a transcription factor binding region were used as probes for EMSA and are listed in Supplementary Table 3. Double-stranded oligonucleotides were generated by annealing the synthetic oligonucleotides with their respective complementary sequences. Complementary oligonucleotides of equal quantity (10 μM each) were annealed in a thermocycler (Techgene, UK) at the following temperatures: 88 °C for 2 min, 65 °C for 10 min, 37 °C for 10 min and 25 °C for 5 min. A nonlabeled probe competitor and its mutant competitor (100 μM) were used (shown in Supplementary Table 3. Nuclear proteins were extracted from RAW264.7 cells and incubated with biotin-labeled DNA probes (10 μM) and/or competitor or mutant competitor (100 μM). Then, the mixtures were separated on native PAGE gels, transferred to nitrocellulose membranes, incubated with HRP-labeled streptavidin, and developed using a UVP imaging system (UVP, USA).
Rapid amplification of cloned cDNA ends (RACE)
Total RNA was extracted from RAW264.7 cells using TRIzol reagent (Invitrogen, USA), and 5′- or 3′-RACE was performed with a SMARTer RACE 5′/3′ Kit (Takara, Japan) according to the manufacturer's instructions. The following primers used for 5′- or 3′-RACE were designed based on the known sequence information: 5′ specific primer-1-CTTCCCTGGACCTTCGAGCCCTCAG; 5′ specific primer-2-GACAATTGGCCCTGGTACCTCAGC; 3′ specific primer-1-CTGCTTAAGGACTGACCAAGGT; and 3′ specific primer-2-TGTTCCTCCTGTTGAGCCACCTTCCAGC.

Sample preparation and RNA-seq
WT and Inc-EST12 KO BMDMs were stimulated with EST12 (2 μM) for 0, 6, or 24 h. The supernatant was discarded, and 500 μL of TRIzol (Invitrogen, USA) was added. The samples were then transferred to a centrifuge tube and sent to Novogene Company (Beijing, China) for sequencing and analysis. Briefly, mRNA was purified from total RNA using poly-T oligo-conjugated magnetic beads. Fragmentation was performed in the presence of divalent cations in First Strand Synthesis Reaction Buffer (5X). The cDNA templates were synthesized using random hexamer primers and M-MuLV Reverse Transcriptase. Adaptors with hairpin loop structures were ligated to the cDNAs. The cDNA fragments 370–420 bp in length were further purified with an AMPure XP system (Beckman Coulter, Beverly, USA). Then, PCR was performed with Phusion High-Fidelity DNA polymerase, Universal PCR primers and Index (X) Primer. Finally, PCR products were purified (AMPure XP system), and library quality was assessed using the Agilent Bioanalyzer 2100 system.

Raw data (raw reads) in fastq format were first processed using in-house Perl scripts. Reads were aligned to the reference genome using HISAT2 v2.0.5 instructions. The aligned reads were counted using HTseq-counts v1.5.0-p3 was used to count the read numbers mapped to each gene. Differential expression between two conditions/groups (two biological replicates per condition) was analyzed using the DESeq2 R package (1.20.0). GO enrichment analysis of differentially expressed genes was implemented using the clusterProfiler R package, in which gene length bias was corrected.

Single RNA fluorescent in situ hybridization (FISH)
The red fluorescent Cy3-labeled probe (RiboBio, China) against the target site of Inc-EST12 was designed by RiboBio Company (Guangzhou, China) and was detected using a Fluorescent In Situ Hybridization Kit (RiboBio, China) according to the manufacturer’s instructions. Briefly, RAW264.7 cells grown on cover slips in 24-well plates with the indicated treatment were fixed with 4% (v/v) paraformaldehyde for 10 min at room temperature and then washed three times with cold PBS. The cells were permeabilized in PBS containing 0.5% Triton X-100 for 5 min at 4 °C and then blocked with prehybridization buffer for 30 min at 37 °C. Cells were then incubated with a probe in hybridization buffer (2.5 μL, 20 μM, 250 μL hybridization buffer) overnight at 37 °C in the dark. After hybridization, the cells were washed in the dark with washing buffer (4 × SSC/2 × SSC/1 × SSC) and then stained with DAPI for 10 min. The hybridization complexes were purified with streptavidin beads for 3 h at room temperature and visualized by performing Coomassie Brilliant Blue staining for mass spectrometry or immunoblotting using a specific antibody against FUBP3. For mass spectrometry, a specific band present in the experimental lane was extracted (the corresponding region in the control lane was also extracted) for further analysis.

LDH release assay
The LDH release assay was performed using the CytoTox 96 NonRadioactive Cytotoxicity assay kit (Promega, USA) according to the manufacturer’s protocol. BMDMs were seeded in 12-well plates (2 × 10^5 cells per well) for 24 h, incubated with lentivirus containing 0.5% Triton X-100 for 5 min at 4 °C and then blocked with prehybridization buffer for 30 min at 37 °C. Cells were then incubated with a probe in hybridization buffer (2.5 μL, 20 μM, 250 μL hybridization buffer) overnight at 37 °C in the dark. After hybridization, the cells were washed in the dark with washing buffer (4 × SSC/2 × SSC/1 × SSC) and then stained with DAPI for 10 min. The hybridization complexes were purified with streptavidin beads for 3 h at room temperature and visualized by performing Coomassie Brilliant Blue staining for mass spectrometry or immunoblotting using a specific antibody against FUBP3. For mass spectrometry, a specific band present in the experimental lane was extracted (the corresponding region in the control lane was also extracted) for further analysis.

Chromatin immunoprecipitation (ChIP)-qPCR
BMDMs were infected with LV-Ctl or LV-Inc-EST12 for 24 h and then treated with or without EST12 protein for 12 h (20 μg/mL). The growth media was removed, and the cells were rinsed three times with cold PBS. Then, the cells were treated with formaldehyde to a final concentration of 1% at room temperature for 10 min. Glycine was added to the cells to a nal concentration of 125 M to stop the crosslinking reaction. Cells were harvested into cold PBS by scraping and transferred to a 1.5 mL microcentrifuge tube. After centrifugation at 1000 × g for 5 min at 4 °C, the formaldehyde-crosslinked cells were collected and resuspended in 1 mL of nuclear lysis buffer (50 mM Tris-HCl pH 8.0, 10 mM EDTA pH 8.0, 1% SDS, and 1 mM PMFS). Chromatin was sheared to an average size of 100-500 bp by sonication and then centrifuged (10 min, 10,000 g, 4 °C). Sixty microliters of supernatant was diluted 10-fold with 540 μL of ChIP dilution buffer (1% Triton X-100, 1.2 mM EDTA, 176 mM NaCl, and 17.6 mM Tris-HCl, pH 8.0) and then incubated with anti-FUBP3 (Abcam, UK, ab181025) or anti-rabbit IgG (Millipore, Germany, 12260) overnight at 4 °C with rotation. Then, 50 μL of Protein A/G magnetic beads (MCE, USA, HY-K0202) was added to each sample, and the incubation continued for 5 h at 4 °C on a rotating platform. The beads were pelleted and then washed sequentially with low-salt buffer (150 mM NaCl, 20 mM Tris-HCl pH 8.0, 0.1% SDS, 0.5% Triton X-100, and 2 mM EDTA), high-salt buffer (1% SDS, 1% Triton X-100, 2 mM EDTA, 20 mM Tris-HCl, pH 8.1, and 500 mM NaCl), and LiCl buffer (0.25 M LiCl, 1% sodium deoxycholate, 10 mM Tris-HCl, pH 8.0, and 1 mM EDTA) followed by two washes with TE buffer (10 mM Tris-HCl, pH 8.0). Chromatin was eluted from the beads by two incubations with 100 μL of elution buffer (100 mM NaHCO3 and 1% SDS), the Na+ concentration was adjusted to 300 mM with 5 M NaCl, and the crosslinks were reversed by an overnight incubation in a 65 °C water bath. The samples were then...
incubated with 0.1 mg/mL RNase A for 1 h at 37 °C followed by 1 mg/mL proteinase K for 2 h at 55 °C. DNA was purified by phenol extraction and ethanol precipitation and then used for qPCR.

**Hematoxylin-eosin (H&E) and Ziehl–Neelsen staining**

H&E staining was performed according to the instructions provided by the manufacturer of the reagent (Baso, China). Briefly, mouse tissues were immobilized, embedded in paraffin, and then sectioned. Paraffin slides were de waxed and hydrated through two de waxing steps with xylene for 5 min each, soaked in anhydrous ethanol, 95, 80, and 70% ethanol for 2 min each, and finally soaked in distilled water for 2 min. The de waxed sections were stained with hematoxylin solution for 10 min, washed with tap water and xylene for 5 min each, soaked in anhydrous ethanol, 95, 80, and 70% alcohol solutions and cleared in xylene. To warm water at 50 °C for 5 min, stained with eosin solution for 1 min and stained with hematoxylin solution for 10 min, washed with tap water and xylene for 5 min each, soaked in anhydrous ethanol, 95, 80, and 70% alcohol solutions and cleared in xylene.

Ziehl–Neelsen staining was performed according to the instructions provided by the manufacturer of the reagent (Basso, China). Briefly, mouse tissues were fixed, embedded in paraffin, and then sectioned. Paraffin sections were de waxed and hydrated through two de waxing steps with xylene for 5 min each, soaked in anhydrous ethanol, 95, 80, and 70% ethanol for 2 min each, and finally soaked in distilled water for 2 min. The de waxed sections were stained with Carbol fuchsin staining solution for 10 min and then washed with tap water for 30 s. Acid alcohol solution was added from the outer edge of the upper edge of the tissue to cover the glass slide, discolored for 1 min, and then washed with tap water for 30 s. Sections were then stained with a methylene blue solution for 30 s. After washing with tap water, the sections were allowed to dry before microscopic examination. The mounted sections were then examined and photographed using a Leica Aperio VERSA 8 (Leica, Germany). The staining intensity was analyzed using ImageScope version 12 software (Leica, Germany).

**Macrophage adoptive transfer animal experiment**

WT mice were i.v. injected with 200 μL of clodronate liposomes (Liposoma, USA) 2 days in advance according to the manufacturer’s instructions. On Day 0, all the mice were i.v. injected with WT or Inc-EST12 KO BMDMs (10^5/mice). Two days later, the mice were i.v. infected with M. or M. est12 (10^5 CFU/mouse). On the 5th day, the mice were sacrificed, and the serum, spleen, liver, and lung tissues were collected and processed for the next step.

**Statistical analysis**

GraphPad Prism software (GraphPad Software Inc., La Jolla, CA) was used for statistical analyses. Significant differences between groups were analyzed using two-tailed unpaired Student’s t test or one-way ANOVA followed by Sidak’s multiple comparisons test. Data are presented as the means ± standard deviations (SD). A p value < 0.05 was considered significant.

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AUTHOR CONTRIBUTIONS
XLZ designed and supervised the research. QY conducted the experiments, analyzed the data, and wrote the paper. DX, ZQ, JW, YZ, YW, and HX assisted in the experimental execution. YX and XLZ revised the paper. All authors contributed to and approved the final version of the paper.

COMPETING INTERESTS
The authors declare no competing interests.

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