ISGylation of EMD promotes its interaction with PDHA to inhibit aerobic oxidation in lung adenocarcinoma

Congcong Zhang¹ | Jiangtao Cui² | Leiqun Cao¹ | Xiaoting Tian³ | Yayou Miao³ | Yikun Wang⁴ | Shiyu Qiu⁴ | Wanxin Guo⁴ | Lifang Ma⁴ | Jinjing Xia⁵ | Xiao Zhang³,⁴

¹Anhui University of Science and Technology School of Medicine, Huainan, Anhui, China
²Department of Thoracic Surgery, Shanghai Chest Hospital, Shanghai Jiao Tong University School of Medicine, Shanghai, China
³Shanghai Institute of Thoracic Oncology, Shanghai Chest Hospital, Shanghai Jiao Tong University School of Medicine, Shanghai, China
⁴Department of Clinical Laboratory Medicine, Shanghai Chest Hospital, Shanghai Jiao Tong University School of Medicine, Shanghai, China
⁵Department of Pulmonary Medicine, Shanghai Chest Hospital, Shanghai Jiao Tong University School of Medicine, Shanghai, China

Abstract
Abnormal nuclear structure caused by dysregulation of skeletal proteins is a common phenomenon in tumour cells. However, how skeletal proteins promote tumorigenesis remains uncovered. Here, we revealed the mechanism by which skeletal protein Emerin (EMD) promoted glucose metabolism to induce lung adenocarcinoma (LUAD). Firstly, we identified that EMD was highly expressed and promoted the malignant phenotypes in LUAD. The high expression of EMD might be due to its low level of ubiquitination. Additionally, the ISGylation at lysine 37 of EMD inhibited lysine 36 ubiquitination and upregulated EMD stability. We further explored that EMD could inhibit aerobic oxidation and stimulate glycolysis. Mechanistically, via its β-catenin interaction domain, EMD bound with PDHA, stimulated serine 293 and 300 phosphorylation and inhibited PDHA expression, facilitated glycolysis of glucose that should enter the aerobic oxidation pathway, and EMD ISGylation was essential for EMD-PDHA interaction. In clinical LUAD specimens, EMD was negatively associated with PDHA, while positively associated with EMD ISGylation, tumour stage and diameter. In LUAD with higher glucose level, EMD expression and ISGylation were higher. Collectively, EMD was a stimulator for LUAD by inhibiting aerobic oxidation via interacting with PDHA. Restricting cancer-promoting role of EMD might be helpful for LUAD treatment.

Keywords
glucose metabolism, glycolysis, ISG15, proteasome, skeletal proteins, ubiquitination
1 | INTRODUCTION

Skeletal proteins are the main support structures of cells. In addition to its important role in the stability of cell morphology, and the ability to withstand external forces and the orderliness of internal structures, it also participates in tumorigenesis. The cytoskeleton may induce cell proliferation through the activity of specific proteins that control cell stiffness, and oncogenes become active throughout this process. Alterations in cytoskeletal proteins also lead to nuclear abnormalities common in a wide range of cancer types. It has long been recognized that tumour cells frequently exhibit abnormal nuclear architecture, including nuclear size and structure, nucleolar size and number and chromatin texture, and that these alterations are critical risk factors for tumour formation and developments.

Emerin (EMD) is 254 amino acids in length and with an N-terminal globular LAP2-EMD-MAN1 (LEM, aa: 1–45) domain, followed by an β-catenin interaction (IT, aa: 168–186) domain, and anchored to the nuclear envelope. EMD is an integral nuclear membrane protein that is expressed in most human tissues and is involved in the maintenance of nuclear structure. EMD also has important effects on nuclear assembly cell cycle progression. Furthermore, EMD has been implicated in the regulation of gene expression, cellular signalling, and nuclear and genomic structure. Although EMD was highly expressed in various types of cancers including ovarian epithelial cancer and breast cancer, how EMD exerts its ontogenetic role in cancer progression remain largely uncovered.

Interferon-stimulated gene 15 (ISG15) is a highly inducible interferon-stimulated gene first discovered by Haas et al. ISG15 covalently binds to the target protein through the glycine–glycine motif at the C-terminus and modifies the target protein. This binding is called ISGylation. ISGylation is similar to ubiquitination, but it does not function to degrade proteins. It is closely related to stress response, transcriptional regulation and protein translation. On the contrary, ISGylation could be an antagonistic process for ubiquitination and the following degradation via proteasomes. For example, through in vitro studies, it is known that ISG15 and ubiquitin (Ub) can form mixed chains. Lysine (K) residues 29 and 48 on Ub are the substrates for ISG15, and ISG15 may alter the binding of Ub to its target proteins and inhibit their subsequent degradation. Moreover, an increase in the ubiquitination levels in breast cancer cells is promoted by the reduction of ISG15 or UBCH8 (ubiquitin-conjugating enzyme H8, an ISG15 conjugating E2 enzyme), suggesting that ISGylation avoids protein ubiquitination and confers stability. In other studies, it has been demonstrated that the ISGylation of the interferon regulatory factor 3 (IRF3) transcription factor decreases its interaction with peptidylprolyl cis/trans isomerase, NIMA-interacting 1 (PIN1) to avoid its ubiquitination. However, whether the protein ISGylation can inhibit ubiquitination in other ways needs further elucidation.

Glucose metabolism is usually abnormal in tumour. Aerobic oxidation is inhibited, while glycolysis, pentose phosphate pathway and hexosamine biosynthesis pathway (HBP) are activated. Glycolysis, pentose phosphate and HBP pathways produce less adenosine triphosphate (ATP), but can provide more internal products for tumour cell proliferation compared with aerobic oxidation. Catalytic alpha subunit of PDHc (PDHA) is an enzyme that acts as a key link between glycolysis and the tricarboxylic acid cycle, converting acetyl coenzyme A (acyetyl-CoA) to pyruvate. PDHA is phosphorylated and inhibited by pyruvate dehydrogenase kinase (PDK). Studies have pointed out that abnormal PDHA expression is closely related to the occurrence and development of various tumours, including glioma, cholangiocarcinoma, breast cancer and so on. However, whether PDHA links with lung adenocarcinoma (LUAD) and its relationship with skeletal proteins in tumour formation and development remain elusive.

Here, we identified that EMD was highly expressed in LUAD and stimulated LUAD malignant phenotypes. In addition, the ubiquitination level of EMD was lower, but its stability was higher than that of other skeleton proteins. EMD ISGylation inhibited its ubiquitination and the following proteasome degradation. Moreover, we noticed that EMD promoted glucose uptake in LUAD cells and inhibited aerobic oxidation, allowing glucose to enter the glycolysis pathway. Mechanistically, EMD IT domain was essential for interaction and inhibition to PDHA expression and phosphorylation. In clinical LUAD specimens, EMD was negatively associated with PDHA, while positively associated with EMD ISGylation, tumour stage and diameter. LUAD with higher glucose level was associated with higher EMD expression and ISGylation level compared with that with lower glucose level. Therefore, EMD was a stimulator for glucose metabolism in LUAD, new treatment strategies for LUAD could be proposed considering the impact of EMD and its ISGylation.

2 | MATERIAL AND METHODS

2.1 | Cell culture

A549, H1299, PC9, Calu-1, H1975 and H2030 cell lines were purchased from Fuheng Biotechnology (Shanghai, China). All cells were maintained in Dulbecco’s modified eagle’s medium (DMEM, WILEY)
(A) Cohort #1

- Non-paired normal
- LUSC
- LUAD

IHC score graph with ** annotations.

(B) Cohort #2

- n=20/group
- Adjacent normal
- LUAD

EMD concentration (pg/ml) distribution.

(C) Cohort #3 (n=14/group)

- EMD
- GAPDH

Western blot analysis with NT and T marks.

(D) Cohort #4

- n=60/group
- NT

EMD concentration (pg/ml) distribution.

(E) Cohort #5 (n=193/group)

IHC score graph with ** annotations.

(F) Western blot analysis of EMD and Actin protein intensities.

Protein intensity (%) graph for A549, H1299, PC9, Calu-1, H1975, and H2030.
HyClone, Logan, UT, USA) supplemented with 10% foetal bovine serum (FBS), 100 U/ml penicillin and 100 mg/ml streptomycin.

2.2 | Mouse experiments and tissue samples

For xenograft experiments, H2030 cells (initially 5 × 10⁶) with EMDWT-FLAG or EMDDel-IT-FLAG overexpression with or without ISG15 knockout were subcutaneously injected into 8-week-old athymic nude mice. For patient-derived xenograft (PDX) generation, fresh LUAD tissues in the size of 2–3 mm³ were subcutaneously implanted into 8-week-old athymic nude mice. The PDX-planted mice could be used for further study after successful passage. The tumour volume was calculated as 0.5 × L × W², with L indicating length and W indicating width. All the tissue samples (mean age ± SD, 61.16 ± 10.35 years; male: female ratio, 1.14:1) were recruited in Shanghai Chest Hospital (Shanghai, China) from May 2013 to January 2022. Cohort #1 were collected from March 2015 to May 2017. Cohort #2 were collected from January 2021 to May 2021. Cohort #3 were collected from May 2021 to July 2021. Cohort #4 were collected from May 2021 to September 2021. Cohort #5 were collected from May 2013 to March 2019. Cohort #6 were collected from September 2021 to January 2022. Cohort #1 was stored in the form of tissues slices in 4°C. Cohort #5 was stored in the form of tissue microarrays in 4°C. Cohort #2, #3, #4 and #6 samples were stored in the form of tissue blocks in −80°C. Other detailed information was summarized in Table S1–6.

2.3 | Regents and plasmids

For regents, cycloheximide (CHX, Sigma, St Louis, MO) and MG132 (MedChemExpress, Monmouth Junction, NJ) were used to treat LUAD cells.

For plasmids, ISG15 and ISG15-KO plasmids were acquired from previous studies.EMDWT-FLAG and PDHAWT-HA were purchased from Zorinbio (Shanghai, China). LentCRISPR v2-based constructs were used for knocking out EMD. All the mutation plasmids were constructed using overlapping PCR and cloned into pcDNA3.1(+) plasmids (Biolink, Shanghai, China). The sequences for small guide RNA (sgRNA) were summarized in Table S7.
FIGURE 3  EMD ISGylation inhibited ubiquitination. (A) EMD expression in ISG15 overexpression or knockout H2030 and H1299 cells as measured by IB. (B) EMD ISGylation and ubiquitination were measured in H2030 cells with or without EMD overexpression or knockdown. The EMD level in each co-IP sample was adjusted to the same protein content. (C) EMD and PSMB5 colocalization in GFP−/− or ISG15−/− H2030 cells was measured by IF. Scale bar, 20 µm. (D) Association of EMD and PSMB5 analysed by IB in proteasomes isolated from ISG15 overexpression or knockout H2030 cells in the presence of MG132 (8 µM, 24 h). Samples from affinity or control beads were analysed in parallel. (E) ISGylation of EMDWT-FLAG and EMDK36A-FLAG were measured in H2030 cells. (F) The location of the potential EMD ISGylation site on total protein. (G) ISGylation and ubiquitination of EMDWT-FLAG, EMDK37A-FLAG, EMDK78A-FLAG, EMDK79A-FLAG and EMDK88A-FLAG were measured in H2030 cells. (H) EMDWT-FLAG or EMDK37A-FLAG was measured by IB in H2030 cells with CHX (10 µg/ml) treatment for indicated hours. (I) Association of EMD and PSMB5 analysed by IB in proteasomes isolated from EMDWT-FLAG, EMDK36A-FLAG or EMDK37A-FLAG overexpressed H2030 cells in the presence of MG132 (8 µM, 24 h). Samples from affinity or control beads were analysed in parallel. (J) Relationship between EMD ISGylation and expression were measured in indicated LUAD cell lines. The EMD level in each co-IP sample was adjusted to the same protein content. IB images were selected from three biological replicates. Data in J were analysed by a Spearman rank-correlation analysis.

2.5 | Co-immunoprecipitation

Co-immunoprecipitation was performed as described previously.27,28,30,31 Cell lysates were mixed with protein A/G-magnetic beads (Novex, Oslo, Norway) and incubated at 4°C overnight with the selected antibodies. The beads were washed using Western/IP lysis buffer (Beyotime, Haimen, China, composition: 20 mM Tris [pH 7.5], 150 mM NaCl, 1% Triton X-100, and inhibitors containing sodium pyrophosphate, β-glycerophosphate, EDTA, Na3VO4 and leupeptin), and eluted using the Acid Elution buffer (Beyotime). Next, total protein amount of each group was measured by BCA protein quantification, and eluted using the Acid Elution buffer (Beyotime). Next, total protein amount of each group was measured by BCA protein quantification. After the first quantification, the samples were diluted according to the difference in protein amount, and the second quantification was carried out for confirmation. Finally, the samples were suspended in SDS-PAGE loading buffer and then measured by IB. The antibodies used for co-IP were as follows: anti-FLAG (CST, #14793 and #8146), anti-PDHA (Abcam, #ab168379 and #ab110330), anti-antibodies used for co-IP were as follows: anti-FLAG (CST, #14793 and #8146), anti-PDHA (Abcam, #ab110330), anti-Lamin A (Abcam, #ab226198), anti-Nesprin1 HA (Abcam, #ab9110 and #ab1424), anti-EMD (Abcam, #ab156871) and anti-EMD (Abcam, #ab204987).

2.6 | Measurements of cell viability and anchorage-independent colony formation

Cell viability was measured using a 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide (MTT)-dependent method. As for colony formation assay, LUAD cells were seeded in a 6-well plate containing 0.3% agarose in DMEM at a density of 6 × 103 cells per well. After 2 weeks, the numbers of colonies were counted under microscope.

2.7 | Quantitative RT-PCR

Total RNA was extracted using Trizol- (Ambion, Carlsbad, CA, USA) dependent method and reverse-transcribed into complementary DNA using the PrimeScript™ RT reagent kit (Takara, Dalian, China). Next, the SYBR premix Ex Taq (Takara) kit was used for real-time qPCR. The primers are listed in Table S7.

2.8 | Proteasome isolation

Proteasomes were isolated by proteasome isolation kit (Sigma, #539176). Isolation was performed in accordance with manufacturer guidelines. Affinity and control beads were used to isolate the proteasome and serve as a negative control, respectively.

2.9 | Extracellular acidification rate and oxygen consumption rate assays

ECAR and OCR were analysed using the extracellular flux analyser XF96 (Seahorse Bioscience, Billerica, MA, USA) with the glycolysis stress test kit (Agilent, Wilmington, DE, USA, # 103020–100) and mitochondrial stress test kit (Agilent, # 103015–100), respectively.

2.10 | Measurements of metabolites and O-GlcNAcylated proteins

Glucose, acetyl-CoA, lactic acid, citrate, isocitrate, α-ketoglutarate, succinate, fumarate, malate, oxaloacetate, pyruvate and phosphoenolpyruvate (PEP, upstream of pyruvate) concentrations were measured by kits purchased from Sigma. 6-phosphogluconate (6-PG) concentration was measured by kit purchased from Abcam. ribose-5-phosphate (Rib-5-P) and uridine diphospho-N-acetylglucosamine (UDP-GlcNAc) concentrations were measured by kits purchased from Yingxin. O-GlcNAcylated proteins were measured using the kits from Abcam. All the experiments were performed strictly according to the instruction provided by the manufacturer.

2.11 | Protein ligation assay experiments

PLA was performed to identify the direct interactions between two proteins using the Duolink in situ Red Starter Kit (mouse/rabbit) (Sigma). The detailed procedure was described in a previous study.28 The antibodies used were anti-EMD (Abcam, #ab156871) and anti-PDHA (Abcam, #ab110330).
2.12  |  Statistical analysis

Tests for this study included Student’s t-test, one-way, two-way ANOVA, χ² test and the Spearman rank-correlation analysis. A P < 0.05 was considered statistically significant.

2.13  |  Ethics statement

This study was approved by the Ethics Review Committee of Shanghai Chest Hospital, Shanghai Jiao Tong University School of Medicine (permit number: KS[Y]1922). All patients completed informed consent forms.

3  |  RESULTS

3.1  |  EMD protein was highly expressed in LUAD

Firstly, we explored EMD expression in LUAD by analysing five cohorts. In cohort #1 (including 20 lung squamous cell carcinoma [LUSC] tissues, 20 LUAD tissues and 20 normal tissues adjacent to tumour), we found that the expression intensity of EMD in normal tissues was lower than that in LUSC tissues, which in turn was lower than that in LUAD tissues (Figure 1A). After analysing tissues from cohort #2 (including 60 paired of LUAD and adjacent normal tissues), cohort #3 (including 14 paired of LUAD and adjacent normal tissues) and cohort #4 (including 50 paired of LUAD and adjacent normal tissues), we found that EMD protein was highly expressed in LUAD tissues compared with adjacent normal tissues, whereas EMD mRNA level was not changed between LUAD and adjacent normal tissues (Figure 1B-D and Figure S1A-C). Additionally, in cohort #5 (including 193 paired of LUAD and adjacent normal tissues), we also observed that EMD expression was higher in LUAD tissues compared with adjacent normal tissues (Figure 1E). These data suggested that EMD protein expression was elevated in LUAD. We further screened six LUAD cell lines and selected the cell line with highest EMD expression (H2030) and lowest EMD expression (H1299) (Figure 1F and Figure S1D-E).

3.2  |  EMD was hypoubiquitinated in LUAD cells

Since EMD protein level was elevated in LUAD while mRNA level was unchanged, we ruled out the possibility that RNA modification and transcription contributed to the EMD elevation. Ubiquitination is a mechanism that inhibits protein expression by reducing protein stability. We hypothesized that the increased expression of EMD in LUAD might be related to its ubiquitination status. We found that the stability of EMD was higher than that of several other common cytoskeleton proteins, including Lamin A, Nesprin 1, Nesprin 3, Sun1, Sun2, in either the H2030 cell line with highest EMD expression or the H1299 cell line with lowest EMD expression (Figure 2A and Figure S2A-C). In H2030 cells, compared with other cytoskeleton proteins, EMD had the lowest ubiquitination level (Figure 2B and Figure S2D-E). These data suggested that high expression of EMD might be because of its hypoubiquitinated station.

3.3  |  EMD was ubiquitinated in K36

We used Prediction of Ubiquitination sites with Bayesian Discriminant Method (BDM-PUB) database (http://bdmpub.biocuckoo.org/prediction.php) to predict potential EMD ubiquitinated sites, and found its K36 position has the highest possibility of being modified (Figure 2C-D). After K36 was mutated to alanine (A), the ubiquitination of EMD was significantly decreased (Figure 2E), resulting in an increase in the stability of EMD (Figure 2F) and a significant decrease in the ability of EMD to enter the proteasome (Figure 2G). In addition, K36A mutation of EMD enhanced cell viability and colony formation ability of H2030 and H1299 cells (Figure 2H-I and Figure S2F). These data suggested that K36 was an important ubiquitination site for EMD.

3.4  |  EMD ISGylation inhibited its ubiquitination to increase protein stability

Since ISGylation and ubiquitination both occurs at K residue, we subsequently investigated whether ISGylation influenced EMD stability and expression via its ubiquitination. We found that in both H2030 and H1299 cells, overexpression of ISG15 increased, whereas knockout of ISG15 decreased EMD expression (Figure 3A). In addition, ISG15 elevated the ISGylation of EMD while inhibited its ubiquitination (Figure 3B and S3A), Knockout of ISG15 significantly promoted colocalization of EMD and one active site of the proteasome, PSMB8 (Figure 3C and S3B), and ISG15 also inhibited the ability of EMD to enter the proteasome (Figure 3D). Unfortunately, the mutation of ubiquitination site K36 into A did not inhibit the
ISGylation of EMD, indicating that K36 is not the ISGylation site of EMD (Figure 3E and S3C). We mutated the Ks near K36 (including K37, K78, K79, and K88) into A, and found that only K37A enhanced ubiquitination of EMD and inhibited its ISGylation (Figure 3F-G and S3D). In addition, K37A significantly decreased the stability of EMD (Figure 3H). We also found the ability of EMD to enter the proteasome was weakened by K36A and strengthened by K37A (Figure 3I). Moreover, the ISGylation of EMD was positively correlated with its expression in LUAD cell lines (Figure 3J and S3E). Collectively, these data suggested that ISGylation at the K37 site of EMD inhibited its ubiquitination at the K36 site and upregulated EMD protein stability.

3.5 | EMD inhibited aerobic oxidation and stimulates glycolysis

The above data have confirmed that EMD could promote the malignant phenotype of LUAD cells (Figure 2H-I), indicating that EMD could promote LUAD. Abnormal glucose metabolism is an important factor in tumour development. We observed that EMD overexpression induced, while EMD knockout suppressed glucose consumption and intracellular glucose level (Figure 4A and S4A-B), and EMD inhibited the biomarkers of aerobic oxidation including OCR (Figure 4B), acetyl-CoA, and products of tricarboxylic acid cycle (citrate, isocitrate, α-ketoglutarate, succinate, fumarate, malate and oxaloacetate) (Figure 4C-D). In addition, we found that EMD stimulated ECAR, the biomarker of glycolysis (Figure 4E) and increased lactic acid level (Figure 4F). However, EMD did not affect pentose phosphate pathway products 6-PG and Rib-5-P (Figure 4G-H), as well as HBP products UDP-GlcNAc and O-GlcNAcylated proteins (Figure 4I-J). These data suggested EMD inhibited aerobic oxidation and stimulates glycolysis.

3.6 | EMD interacted and degraded PDHA

To further investigate how EMD regulated aerobic oxidation and glycolysis, we overexpressed or knocked out EMD to evaluate the effects of EMD on the expression of aerobic oxidation and glycolysis-related metabolic enzymes including HK, G6P, PFK, ALDO, GAPDH, PKG1/2, PGAM1, ENO1, PKM, LDH, PDH and PDKs. We observed that EMD only affected the protein expression of PDHA but did not affect the protein expression of other enzymes (Figure 5B and S5A). In addition, EMD had no effect on mRNA expression of all enzymes (Figure 5C and S5B). EMD and PDHA protein colocalization and interaction were also validated (Figure 5D-F and S5C). These data suggested that EMD bound to PDHA and inhibited PDHA expression.

3.7 | Molecular mechanism for EMD-PDHA interaction

EMD protein has two key protein interaction domains LEM and IT, and PDHA protein has three important phosphorylation sites serine (S) 232, 293 and 300 which are phosphorylated by PDK1 (Figure 5G). Deletion of IT domain blocked EMD WT-FLAG and PDHA WT-HA interaction, whereas deletion of LEM domain did not have this effect (Figure 5H and S5D). In addition, we found that EMD could promote PDHA phosphorylation and interaction with PDK1, and deletion of IT domain abolished this effect (Figure 5I and S5E). Deletion of the 290–310 region (where S293 and S300 phosphorylation sites are located) inhibited the binding of PDHA to EMD, whereas deletion of the region 220–240 (where S232 phosphorylation site is located) did not affect the EMD-PDHA interaction (Figure 5J and S5F). EMD-PDHA interaction was inhibited by mutations of S293A and S300A, but not influenced by S232A mutation (Figure S5G). EMD inhibited the production of acetyl-CoA catalysed by PDHA, resulting in an increase in the upstream pyruvate content, and these effects would be abolished by the depletion of IT domain (Figure 5K and S5H). However, PEP was not regulated by EMD (Figure 5K and S5G). In addition, PDHA S232A, S293A and S300A mutations did not influence EMD expression, indicating phosphorylation of PDHA had no effect on EMD (Figure 5L). These data suggested that IT domain of EMD and...
S293 and S300 phosphorylation of PDHA were essential for EMD and PDHA interaction.

3.8  |  EMD ISGylation was essential for aerobic oxidation inhibition

Subsequently, we investigated whether EMD ISGylation was related to its interaction with PDHA and regulation on aerobic oxidation. ISG15 knockout abolished the EMD inhibitory effect on PDHA expression and promotion effect on PDHA phosphorylation (Figure 6A and S6A) and disrupted the interaction between EMD and PDHA (Figure 6B and S6B). ISG15 knockout also abolished the suppression role of EMD on products of tricarboxylic acid cycle (Figure 6C and S6C) and acetyl-CoA (Figure 6D and S6D), as well as the stimulation role of EMD on pyruvate (Figure 6E and S6E). However, ISG15 knockout did not significantly influence PEP level (Figure 6F and S6F). In xenograft formed by H2030 cells, we observed that deletion of IT domain or ISG15 knockout both abolished EMD stimulation on tumour growth (Figure 6G-H), inhibition on PDHA expression (Figure 6I) and acetyl-CoA level (Figure 6J), and blocked the interaction between EMD and PDHA (Figure 6J). These data elucidated the essential role of EMD ISGylation for its inhibition on PDHA and aerobic oxidation.

3.9  |  EMD was negatively associated with PDHA in PDX

We established two series of PDX and used immunohistochemistry (IHC) to identify that EMD expression was higher in PDX#1 compared with PDX#2 (Figure 6K). Interestingly, we found that PDHA expression was higher in PDX#2 compared with PDX#1 (Figure 6L). EMD level was also positively associated with its ISGylation level (Figure 6M) and negatively associated with acetyl-CoA level (Figure 6N). The above data could further validate that EMD was negatively associated with PDHA and acetyl-CoA, and positively associated with its ISGylation level.

3.10  |  Clinical correlation between EMD and PDHA

After tested 60 paired of LUAD and adjacent normal tissues from cohort #6, we found that EMD, EMD ISGylation and global ISGylation raised, while PDHA declined in LUAD tissues compared with adjacent normal tissues (Figure 7A-D), and EMD level was negatively correlated with PDHA level in LUAD tissues (Figure 7E). In addition, we found that EMD ISGylation was positively correlated with EMD (Figure 7F-G) and global ISGylation level (Figure S7A), and negatively correlated with PDHA level (Figure 7H). The level of EMD and its ISGylation, as well as global ISGylation increased with the tumour stage progression (Figure 7J-I, Figure S7I), whereas PDHA level decreased in stage II and III tissues compared with those of stage I (Figure 7K). Furthermore, EMD level, its ISGylation and global ISGylation was higher in LUAD tumours with ≥3 cm diameter (Figure 7L-M, Figure S7C), whereas PDHA level was higher in LUAD tumours with <3 cm diameter (Figure 7N). Since EMD and PDHA were related with glucose metabolism in LUAD (Figure 5-6),21 we evaluated that EMD expression, EMD ISGylation, global ISGylation and PDHA expression in high and low glucose LUAD tissues, and found that EMD expression, EMD ISGylation and global ISGylation increased, and PDHA expression decreased in high glucose tissues compared with low glucose tissues (Figure S7D-H). These data suggested that EMD expression was positively correlated with its ISGylation, and negatively correlated with PDHA expression in clinical LUAD tissues.

4  |  DISCUSSION

EMD is an important integral protein of the nuclear inner membrane, but the mechanism that promotes its expression is not fully understood to a great extent. Existing studies confirmed that EMD transcription is controlled by transcription factor LIM domain.7,36 and its expression is inhibited by Semaphorin 3A.37 In this study, we explored that EMD expression was simultaneously controlled by ubiquitination and ISGylation. The ubiquitination occurred at K36 and ISGylation occurred at K37, and there was a dynamic balance between them, which
controlled the expression of EMD; ubiquitination promoted the degradation of EMD, while ISGylation inhibited this process (Figure 2-3). Given that EMD has more than 30 phosphorylation sites and more than five known O-GlcNAcylated sites,9 whether phosphorylation and O-GlcNAcylation would affect the ubiquitination and ISGylation of EMD, and even other post-translational modifications are also involved in their dynamic balance, which can be further explored in subsequent studies. Based on the known high expression of EMD in breast cancer and ovarian epithelial cancer,12,13 we further supplemented the evidence of high expression of EMD in LUAD and clarified the relevant mechanism to a certain extent. These results provide basis for individualized treatment of cancer.

The LEM-domain is a ~40-residue helix–loop–helix fold conserved both in eukaryotes and in prokaryotic DNA/RNA-binding proteins.38 LEM domain in EMD can directly bind to lamin9 and barrier-to-autointegration factor (BAF),30 together forming a major component of nucleoskeletal structure known as the nuclear "lamina."41 However, binding to PDHA was another domain of EMD, the IT domain (Figure 5). The IT domain also has the function to bind proteins and is known to bind β-catenin.8 A deeper understanding of the properties of LEM and IT domains may lead to a more systematic understanding of the proteins they bind and their physiological effects. Followed by binding to PDHA, EMD promoted the phosphorylation and led to the degradation of PDHA (Figure 5), which is consistent with the previous study reported that PDHA phosphorylation leads to its inhibition.22

ISGylation modification is a kind of ubiquitin-like modification, and both ISGylation and ubiquitination occurs at the K residue of the protein. Therefore, there are important connections between ISGylation and ubiquitination functions.42 ISGylation can inhibit ubiquitination by destroying the ubiquitin chain and inhibiting the connection between the target protein and the ubiquitination-promoting protein, thereby inhibiting the degradation of the target protein.16,18 Here, we found that ubiquitination and ISGylation of EMD occurred at adjacent sites (K36 and K37) and were completely different between metastatic subpopulations compared with the primary tumour. Here, we proposed a hypothesis that EMD induces hypoxia in tumours by inhibiting PDHA, and hypoxia can lead to activation of activating transcription factor 4 (ATF4), which maintains tumour cells in high levels of autophagy.22 Subsequently, EMD is degraded by autophagy, leading to the occurrence of pro-invasive effects such as EMT. However, this hypothesis needs further experiments to validate. The possible dynamic changes of EMD poses a challenge to the treatment of tumours with high EMD expression. However, a deep understanding of EMD expression regulations and cancer-promoting roles would provide a basis for the proposal of new tumour treatment strategies.
AUTHOR CONTRIBUTIONS
Congcong Zhang: Data curation (equal); formal analysis (equal); investigation (equal); methodology (equal); validation (equal); visualization (equal); writing – original draft (equal). Jiatao Cui: Data curation (equal); formal analysis (equal); investigation (equal); methodology (equal); validation (equal); visualization (equal). Leiqun Cao: Data curation (equal); formal analysis (equal); investigation (equal); methodology (equal); validation (equal); visualization (equal). Xiaoting Tian: Resources (lead). Yayou Miao: Resources (lead). Shiyu Qiu: Data curation (supporting); formal analysis (supporting); investigation (supporting); methodology (supporting); validation (supporting); visualization (supporting). Yikun Wang: Data curation (supporting); formal analysis (supporting); investigation (supporting); methodology (supporting); validation (supporting); visualization (supporting). Wanxin Guo: Data curation (supporting); formal analysis (supporting); investigation (supporting); methodology (supporting); validation (supporting); visualization (supporting). Lifang Ma: Conceptualization (equal); formal analysis (equal); funding acquisition (equal); project administration (equal); supervision (equal). Jinjing Xia: Conceptualization (equal); formal analysis (equal); funding acquisition (equal); project administration (equal); supervision (equal). Xiao Zhang: Conceptualization (equal); formal analysis (equal); funding acquisition (equal); project administration (equal); supervision (equal); writing – original draft (equal).

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CONFLICT OF INTEREST
The authors have no conflicts of interest to declare.

DATA AVAILABILITY STATEMENT
The data that support the findings of this study are available from the corresponding author upon reasonable request.

ORCID
Xiao Zhang https://orcid.org/0000-0001-9076-5995

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