N-glycosylation stabilizes MerTK and promotes hepatocellular carcinoma tumor growth

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\textbf{A B S T R A C T}

Despite the evidences of elevated expression of Mer tyrosine kinase (MerTK) in multiple human cancers, mechanisms underlying the oncogenic roles of MerTK in hepatocellular carcinoma (HCC) remains undefined. We explored the functional effects of MerTK and N-Glycosylated MerTK on HCC cell survival and tumor growth. Here, we show that MerTK ablation increases reactive oxygen species (ROS) production and promotes the switching from glycolytic metabolism to oxidative phosphorylation in HCC cells, thus suppressing HCC cell proliferation and tumor growth. MerTK is N-glycosylated in HCC cells at asparagine 294 and 454 that stabilizes MerTK to promote oncogenic transformation. Moreover, we observed that nuclear located non-glycosylated MerTK is indispensable for survival of HCC cells under stress. Pathologically, tissue microarray (TMA) data indicate that MerTK is a pivotal prognostic factor for HCC. Our data strongly support the roles of MerTK N-glycosylation in HCC tumorigenesis and suggesting N-glycosylation inhibition as a potential HCC therapeutic strategy.

\textbf{1. Introduction}

Hepatocellular carcinoma (HCC) is one of the most prevalent malignancies of liver and occurs predominantly in patients with underlying chronic liver disease and cirrhosis [1–3]. HCC is the third leading cause of cancer-related deaths worldwide in 2020, with over 900,000 people affected and 830,000 deaths, and the incidence is highest in Asia and Africa [1–3]. The incidence of HCC in the United States is rising because of the increased prevalence of hepatitis C virus (HCV) infection [4–6], and the overall 5-year survival rate for all stages of HCC combined is only 16% [7]. The poor prognosis of HCC is due to the fact that it is diagnosed at an advanced stage and lack of efficient therapy, as well as the highly aggressive and proliferative capacity of HCC cells [8,9]. HCC is a heterogeneous disease, that is usually not amenable to standard chemotherapy and is resistant to radiotherapy. Despite the recent advances in the diagnosis, staging and treatment of HCC, surgical resection and liver transplantation remain the only curative treatment options for advanced stage HCC patients. Therefore, understanding of molecular mechanisms of HCC and development of novel and effective HCC therapies is crucial as they have the potential to dramatically improve the outcomes of HCC patients.

MerTK is a member of the TYRO3, AXL, and MERTK receptors.
ligand GAS6 stimulates cell migration and induces procollagen expression in non-alcoholic steatohepatitis [24, 25]. Additionally, MerTK signaling in liver macrophages has been shown to promote liver fibrogenesis, MerTK is dramatically up-regulated [23]. Interestingly, liver disease patients with F2 gain-of-function capacity [22]. In the liver tissue of non-alcoholic fatty cytokine-independent growth in lymphocytes [14, 15]. MerTK is overexpression of MerTK leads to a transformed phenotype in fibroblasts and stored at already frozen in liquid nitrogen after surgical resection, and were then liver injury [26]. More importantly, allele-specific downregulation of MerTK was demonstrated to prevent HCV-induced liver fibrosis [27]. These evidences highlighted the essential function of MerTK in the process of liver pathological lesions. However, the role of MerTK in HCC tumorigenesis remains unclear. The importance of N-glycosylation of Tyrosine Kinase (TK) receptors were widely investigated such as epidermal growth factor receptor (EGFR) and ErbB2 [28, 29], VEGFR-2 [30], PDGF [31], PDGF [32], M-CSF [33], Src [34], etc. In HCC, N-glycosylation of CD147 is crucial for the tumor promotive activity of CD147 [35]. Here, we investigate a potential role of MerTK and mechanism in regulating HCC tumorigenesis with focus on the regulation of N-glycosylation of MerTK and stability of MerTK in facilitating HCC development.

2. Materials and methods

2.1. Patients and tissue specimens

Sixty-three paired HCC tissue samples including both tumor and matched normal tissue from the same patient were obtained from the First Affiliated Hospital of Wenzhou Medical University (Wenzhou, China) from February 2015 to August 2016. The tissues were immediately frozen in liquid nitrogen after surgical resection, and were then stored at −80 °C until use. The HCC tumors were graded and classified according to the criteria of the 2004 edition of the World Health Organization (WHO). The carcinomas were staged in accordance to the sixth edition International Union-American Joint Committee on Cancer (AJCC)/Union International Contre Cancer (UICC) tumor-node-metastasis (TNM) classification system. All patients who participated in this study provided written informed consents in accordance with the Declaration of Helsinki.

2.2. Tissue microarrays and immunohistochemistry assay

Tissue microarray (TMA) containing a total of 111 formalin-fixed, paraffin-embedded tissue sections were constructed as described previously [36, 37]. TMA specimens were subjected to immunohistochemistry (IHC) analysis as described previously [37].

2.3. Cell lines and cell culture

In this study, cell lines were used as follows: human HCC cell lines (MHCC97H, HCCLM3, HuH7), a human hepatoblastoma cell line HepG2, normal liver cell line (LO2) and low-passage HEK293T cells were purchased from the Cell Bank of Shanghai Institute of Cell Biology (Shanghai, China). The HLE cell line was kindly provided by Prof. Qichao Huang (Fourth Military Medical University, China). All cells were cultured in DMEM Medium (Life Technologies, Grand Island, NY, USA) supplemented with 10% fetal bovine serum (FBS, Life Technologies, Grand Island, NY, USA) and antibiotics (100U /ml penicillin and 100 µg/ml streptomycin) at 37 °C in a 95% air, 5% CO2 humidified incubator. All cell lines were confirmed mycoplasma free and authenticated by the Cell Bank of the Chinese Academy of Sciences before use. Cell lines were also routinely tested and confirmed for mycoplasma free during this study.

2.4. Reagents and antibodies

Diethylaminoamine (DEN, 73861), CCL4 (1601168), D-galactosamine (O4876), Carbonyl cyanide 4-(trifluoromethoxy) phenylhydrazide (FCCP, C2920), antismycin A (A8674), rotenone (R8875), 2-deoxy-o-glucose (2-DG, D8375), PNGase F (G5166) and Swainsonine (S9263) and D-(−)-Glucose (G7021) were obtained from Sigma (St. Louis, MO, USA). SYBR green was purchased from Bio-Rad (Hercules, CA, USA). Horseradish peroxidase (HRP)-conjugated anti-rabbit, anti-mouse immunoglobulin G were obtained from Beyotime (Haimen, Jiangsu, China). Glucose deficient DMEM medium and Trizol Reagent were obtained from Life Technologies (Grand Island, NY, USA). UNC-2250 (S7342) was purchased from Selleck Chemicals (Houston, TX, USA). The antibodies used in this study are: MerTK (abcam; ab52968), Phospho-MerTK (abcam; ab14921), Akt (Cell Signaling Technology; 4691), Phospho-Akt (Ser473) (Cell Signaling Technology; 4060), Phospho-GSK3β (Cell Signaling Technology; 55885), PKG1 (ZN; 501965), LDHA (ZN; 501146), PFKM (ZN; 505477), PKM2 (ZN; 505477), PDHK1 (Cell Signaling Technology; 38205), Lamin B (Proteintech; 12987-1-AP), GAPDH (Abmart; M20028) and β-Actin (Proteintech; 60008-1-1 g).

2.5. Western blot

Western blotting was performed as described previously [38].

2.6. RNA extraction and qRT-PCR

Total RNA was prepared from cells and tissues using Trizol Reagent according to the manufacturer’s protocol, and 2 μg of total RNA of each sample was used for reverse transcription using Primerscript™ RT reagent kit with gDNA Eraser (Takara, Dalian, China) according to the manufacturer’s instructions. qRT-PCR was performed using the CFX Connect™ real-time system (Bio-Rad, Hercules, CA, USA) according to the manufacturer’s protocol as follows: 95 °C for 10 min, 45 cycles of denaturation at 95 °C for 10 s and extension at 60 °C for 30 s. The threshold cycle number (CT) was recorded for each reaction. Primers for MerTK gene amplification were as follows: forward primer: 5’-GGAGGCTCGGATCTCTGTTCA-3’; reverse primer: 5’-GAGGGGGCATAATCTACTACCA-3’.

2.7. Plasmids

The human MERTK clone was obtained from the mRNA of 293T cell line by reverse transcription-Polymerase Chain Reaction (RT-PCR) subsequently cloned into pLVX-Puro lentiviral expression vector to establish MerTK overexpression cell line. Using the pLVX-Puro/MerTK expression vector as a template, MerTK mutants (N294Q, N395Q, N442Q, N454Q and 3Y) were developed by performing a site directed mutagenesis. PLKO.1-shRNA vector expressing shRNA targeting endogenous MERTK mRNA to interfere MERTK level. The asparagine and tyrosine point mutation primers and shRNA sequences of MERTK are shown in Tables S2 and S3.
2.8. Generation of MerTK knockdown and cDNA-overexpression cell lines

MerTK stable knockdown or overexpression of WT and various mutants of MerTK MHCC97H, HCCLM3, and Huh7 cells lines were constructed by lentivirus packaged plasmid infection. 24 h post infection, centrifugal sedimentation was used to increase the virus titer. Secondly, multiple rounds of transduction was employed to enhance cell infection ratio. Thirdly, polybrene (1:1000) was added into the culture medium to reduce the repulsion force between cells and viruses and thereby to promote the binding of retroviral particle to the cell surface resulting in a higher efficiency of transduction.

2.9. Cell proliferation assay

MerTK stable knockdown or overexpression MHCC97H, HCCLM3, and Huh7 cells were seeded into 12-well cell culture plates, and cultured at 37 °C, 5% CO2. Cell numbers were counted at following time points: 1, 2, 3, 4, 5, 6 days.

2.10. Tumor xenograft assay

Male BALB/c mice were purchased from Shanghai Laboratory Animal Center, CAS (Shanghai, China) and housed in a specific pathogen-free environment. For in vivo tumorigenesis analysis, control and MerTK stable knockdown or overexpression MHCC97H, HCCLM3, Huh7and HepG2 cells (2.5 × 10⁶ cells in 50 μl medium mixed with 50 μl Matrigel) were injected subcutaneously into the flanks of 5-week-old nude mice (n = 5–8). Tumor length (L) and width (W) were measured every 2 days with calipers, and the tumor volume (V) was calculated (V = 1/2(L × W²)).

2.11. DEN and CCl₄-induced HCC mouse model

The DEN and CCl₄-induced HCC mouse model was established as previously described [39,40]. Briefly, a single dose of DEN (25 mg/kg body weight in PBS) were injected into 15-day-old male mice intraperitoneally (i.p.) to initiate tumor formation. At 4 weeks of age, mice were injected i.p. Once a week with CCl₄ (0.5 mL/kg body weight in corn oil) for an additional 16 weeks. All mice were sacrificed at 30 weeks of age. Livers were removed and the number and size of HCC were recorded, and individual lobes were snap frozen in paraffin embedded and subjected to histological and immunohistochemical analyses.

2.12. Colony formation assay

MerTK stable depleted or overexpressed MHCC97H and HCCLM3 cells were counted, and a total of 400 cells per well were seeded evenly into 6-well plates and incubated at 37 °C for 14 days. Cells were washed with pre-warmed PBS three times, fixed with 4% paraformaldehyde, stained with Giemsa solution for 15 min, then washed with pre-warmed PBS three times, and counted by two independent investigators.

2.13. Metabolic flux assays

Real time intact cellular oxygen consumption rate (OCR) and extracellular acidification rate (ECAR) were measured using the Seahorse XF96 Extracellular Flux Analyzer (Seahorse Bioscience, North Billerica, MA, USA) as previously described [41]. Briefly, MerTK stable knockdown or overexpression MHCC97H and HCCLM3 cells were seeded into the seahorse 96-well cell culture plate at a density of 2.5 × 10⁶ and 2 × 10⁶ cells/80μl/well, respectively, and cultured at 37 °C, 5% CO₂ incubator overnight. Prepare the assay medium for OCR (primary medium 49 mL, 1 mL sodium pyruvate, 0.225 g glucose, pH7.4) and ECAR (primary medium 49 mL, 0.0164 g l-glutamine, pH 7.4). Add 140 μl calibrate solution into the calibrate plate and incubated at non-CO₂ incubator overnight. The next day, replace the calibrator microplate with cell culture plate and start the protocol as we optimized previously to measure the OCR and ECAR [41].

2.14. Apoptosis and mitochondrial membrane potential analysis by FACS

For apoptosis analysis, cells were collected and incubated with Annexin V-FITC/PI (BD, San Jose, CA), followed by incubation in the dark at room temperature for 20 min. Apoptosis rate was measured immediately using a BD Accuri™ C6 Plus flow cytometer (BD, Franklin Lakes, NJ).

Mitochondrial membrane potential was measured by FACS as previously described [41].

2.15. Glucose uptake

MerTK stable knockdown or overexpression MHCC97H, HCCLM3 Cells were collected and incubated for 20 min at 37 °C with 100 nM 2-NBDG (2-(1-(7-nitrobenz-2-oxa-1,3-diazol-4-yl) amino)-2-deoxyglucose), prior to FACS analysis.

2.16. Lactate production

Lactate levels secreted into the media were measured by Fluorescent colorimetric assays. The assay was performed as per the manufacturer’s instructions utilizing Amplite™ Fluorimetric L-Lactate Assay Kit (AAT Bioquest).

2.17. Preparation of nuclear extracts

To isolate nuclear component from cytoplasm, cells were collected and treated with a nuclear protein extraction kit (Beyotime Biotechnology, Haimen, Jiangsu, China) following the manufacturer’s instructions. Briefly, the cell samples were harvested and homogenated in the plasma protein extraction reagent A containing 1 mM PMSF, vortexed in the highest speed for 5s and then bathed on ice for 10–15 min, after which plasma protein extraction reagent B were added. The samples were vortexed in the highest speed for 5 s and then centrifuged at 13,000 rpm at 4 °C for 5 min. The supernatants, corresponding to the cytosolic fraction, were transferred to fresh tube, and measured for protein content by using Pierce BCA Protein Assay Kit.

Further, the nuclei pellet was suspended in nuclear protein extraction reagent, vortexed in the highest speed for 15–30 s. After which the nuclear components were bathed on ice for 1–2 min, vortexed in the highest speed for 15–30 s for every 1–2 min lasting 30 min and centrifuged at 12,000–16,000 × g at 4 °C for 10 min. The supernatants, corresponding to the nuclear fraction, were transferred to a fresh tube, and measured for protein content by using Pierce BCA Protein Assay Kit.

2.18. Blue native PAGE (BNG) gel

BNG was performed as previously published protocol with slight modifications [42]. Briefly, 40 μl of Buffer A (50 mM NaCl, 50 mM imidazole, 2 mM 6-aminohexanoic, and 1 mM EDTA, pH 7.0) was added to 10 mg of pelletized cells. Then 12 μl digitonin was added (20% (v/v)) and allowed to solubilization for 10–20 min at on ice. After centrifugation for 60 min at 20,000 g, supernatants were collected. Then 50% (v/v) glycerol and 5% (w/v) Coomassie blue-G-250 dye stock was mixed...
to yield a glycerol/dye ratio of 2. Next, the mixture was added to the samples to yield a samples/mixture ratio of 5 (v/v). Finally, the samples were run on 3–11% acrylamide gradient gels at 4 °C. The gel was run 60 min at 45 V for spacer gel and 150 min at 180 V for separation gel.

2.19. Immunoprecipitation and ubiquitination assay

To detect ubiquitination status of MerTK in vivo, MHCC97H and HCCLM3 cells were lysed with lysis buffer containing protein inhibitor for 30 min on ice. Then cell lysates were centrifuged at 13,000 rpm for 10 min at 4 °C and the supernatant was transferred to a fresh tube. The content of these cell lysates was measured as per the manufacturer’s instructions by using Pierce BCA Protein Assay Kit. 30 μl Protein A beads were added in 1 mg cell lysates for 4 h with rotation at 4 °C and then centrifuged for 5 min at 12,000 rpm. Supernatant was incubated with 4 μg MerTK primary antibody overnight, after which 50 μl Protein A beads were added for an additional 4 h. Immunoprecipitates were washed three times with PBS. Proteins were eluted in 1.25 × SDS-Loading buffer and analyzed by Western blot with anti-ubiquitin antibody.

2.20. Statistical analysis

Statistical significance was determined using GraphPad Prism software. The χ² test was performed to evaluate the relationship between the clinical-pathological features and MerTK expression in the IHC (TMA) results. Kaplan–Meier log-rank tests and Cox regression analysis were used for survival analysis. All data were presented as mean ± SD unless specified. Student’s t-test was used for two-group comparisons. One-way ANOVA with Bonferroni post-test was used to analyze multiple groups with only one variable tested. Statistical significance is displayed as *P < 0.05, **P < 0.01, ***P < 0.001, n. s., not significant.

3. Results

3.1. MerTK is overexpressed in HCC tissues and MerTK high expression is associated with poor outcomes in HCC patients

To understand the extent of MerTK protein comparative expression in human liver cancer patients as compared with adjacent normal liver tissue, we assessed 63 HCC patient biopsies by Western blot analysis and 37 HCC patient biopsies by qRT-PCR (Fig. 1A–C). Of the 63 or 37 HCC patients, 57 (92%) and 35 (95%), respectively, showed increased MerTK expression compared with normal liver tissues. Western blot analysis of MerTK expression correlated with an increased β-Actin expression (n = 63). Representative photomicrographs are shown in Figs. 1D and 1E. Kaplan–Meier survival analysis of HCC patients (n = 63) revealed that MerTK expression was associated with poor outcomes in HCC patients. Kaplan–Meier analysis of the OS of HCC patients with high (n = 35) or low MerTK (n = 76) expression. (E) Kaplan–Meier overall survival analysis of HCC patients diagnosed with I-II (n = 58) or III-IV (n = 53) stage according to MerTK protein expression. (G) Kaplan–Meier analysis of the OS of HCC patients with Cirrhosis (n = 87) according to MerTK protein expression.
3.2. MerTK is critical for HCC cell proliferation and migration

To better understand the role of MerTK in human liver cancer, we performed loss- and gain-of-function experiments. First, we assessed MerTK expression by Western blotting in four HCC cell lines HLE, Huh7, HCCLM3 and MHCC97H, as well as HepG2 cell, a human hepatoblastoma cell line, and normal liver cell line LO2. Our results revealed that MerTK protein was predominantly expressed in indicated human liver cancer cell lines, while it was barely detected in LO2 cells (Fig. 2A). As uncontrolled cell proliferation leads to tumor initiation and progression, we further investigate the role of MerTK in HCC cell proliferation. We constructed lentivirus packaged shRNA plasmid targeting MerTK (shMerTK#1 and shMerTK#2), as well as the negative control plasmid (shCont). HCC cell lines MHCC97H, HCCLM3 and Huh7 were infected with lentivirus packaged plasmids and the endogenous MerTK was successfully silenced (Fig. 2B). MerTK ablation caused significant inhibition of cell proliferation and migration in HCC cells (Fig. 2C; Fig. S2A). Conversely, over-expression of exogenous MerTK enhanced cell proliferation in both LO2 cells and HCC cells as well as cell migration in HCC cells (Fig. 2D and E; Fig. S2A). MerTK ablation also increased cell apoptosis (Figs. S2B and C) as well as reduced mitochondrial membrane potential in HCC cells (Fig. S2D). Moreover, we increased cell apoptosis (Figs. S2B and C) as well as reduced mito- 

Metastasis is a complex and multistage process. The first step is the primary tumor development (Fig. 2 F and G). The second step is the migration and invasion of cancer cells (Fig. S1A). The third step is the attachment and extravasation of cancer cells (Fig. S1B). The fourth step is the colonization of cancer cells at distant sites (Fig. S1C). The fifth step is the establishment of metastatic tumor lesions (Fig. S1D). Moreover, we found that MerTK knockdown inhibited HCC cell colony formation (Fig. 2F and G). Reversely, overexpressed MerTK increased colony number of HCC cell (Fig. 2F and G). As it is wildly demonstrated that MerTK is a pivotal regulator of ERK, Akt, NF-κB, and Src family kinases [43–47], we then investigated the Akt-GSK3α/β signaling in MerTK depleted or overexpressed HCC cells. We found that MerTK ablation and overexpression reversibly regulates the phosphorylation of Akt, GSK3β (Fig. 2H and I). Together, these results indicate a role for MerTK in conferring proliferative and migratory advantage in HCC partially by regulating Akt, GSK3β.

3.3. MerTK modulates the warburg effect in HCC cells

Metabolic reprogramming is a critical feature of cancer cells [48]. The liver serves as a central metabolic organ being responsible for detoxification, storing glycogen, biosynthesis of urea and several secreted protein and biotransformation. Consequently, patients suffering with HCC also have an altered metabolic pattern of liver displaying enhanced addiction to glucose, and enhanced aerobic glycolysis, like many tumor cells undergoing the Warburg effect [49]. To decipher the role of MerTK in regulating cellular metabolism, we measured Extracellular Acidification Rate (ECAR) and Oxygen Consumption Rate (OCR) in the indicated HCC cells. MerTK depletion caused repression of aerobic glycolysis in MHCC97H and HCCLM3 cells (Fig. 3A; Fig. S3A), while OCR was increased (Fig. 3B; Fig. S3B). In contrast, overexpression of MerTK enhanced the aerobic glycolysis (Fig. 3C; Fig. S3C) and repressed mitochondrial respiration (Fig. 3D; Fig. S3D). We also found MerTK depletion caused excessive ROS generation while overexpression of MerTK partially eliminated ROS (Fig. S3E). To further confirm the effect of MerTK ablation in switching of the Warburg effect, we measured cell glucose uptake and lactate production in MerTK depleted or overexpressed HCC cells, and found that relative glucose uptake and lactate production were significantly decreased in MerTK knockdown cells while increased by MerTK overexpression, respectively (Fig. 3E and F). Similarly, we found that MerTK regulated the expression of glycolytic enzymes PGK1, LDHA, PFKM, PKM2 and PDHK1, in a manner that knockdown or overexpression reversibly regulated the expression levels of these indicated proteins (Fig. 3G and H). Collectively, our data implicate that MerTK facilitates the Warburg effect through regulating key glycolytic enzymes.

3.4. MerTK promotes HCC cell growth in vivo

To further demonstrate the essential role of MerTK in promoting HCC cell growth, we subcutaneously injected control and MerTK knockdown MHCC97H, HCCLM3, Huh7 and HepG2 cells into BALB/c nude mice, respectively. MerTK downregulation significantly suppressed HCC cell growth in vivo (Fig. 4A–C). Stable Knockdown of MerTK protein in tumor of nude mice was confirmed by Western blot (Fig. 4D). We further assessed the levels of key enzymes of Akt-GSK3α/β signaling which involve in regulating glycolysis in cancer cells. Strikingly, Akt and GSK3β were all inactivated and PKG1, LDHA, PFKM, PKM2 and PDHK1 were significantly decreased due to MerTK knockdown (Fig. 4E). Consistently, overexpression of MerTK in MHCC97H cells promoted tumor growth in vivo (Fig. 4F–H). MerTK overexpression in tumor of nude mice was also confirmed by Western blot (Fig. 4I). Moreover, we demonstrated that MerTK expression was significantly increased in the tumor tissue of DEN and CCl4-induced primary HCC mice model (Figs. S4A and B). In addition, the phosphorylation of key enzymes of Akt-GSK3α/β signaling (Akt and GSK3β) were remarkably upregulated (Figs. S4B and C). The glycolytic enzyme PGK1, LDHA, PFKM, PKM2 and PDHK1 were also significantly increased (Figs. S4B and D). These data further confirmed an indispensable role of MerTK in promoting HCC cell growth through regulating the Warburg effect.

3.5. Mertk deletion suppresses chemical-induced liver tumorigenesis

To investigate the physiological and pathological functions of Mertk deletion suppresses liver tumorigenesis, we generated a Mertk knockout
Fig. 2. MerTK is crucial for HCC cell proliferation. (A) Western blot analysis and quantification of MerTK expression in total cell extracts in LO2, HLE, HepG2, HCCLM3, and MHCC97H cells. β-Actin was used as a loading control. Data are presented as mean ± SD of three independent experiments, statistical significance was assessed by One-way ANOVA with Bonferroni post-test, ***P < 0.001. (B) Western blot analysis demonstrating MerTK stable knockdown in the indicated HCC cells transfected with shRNA specific to MERTK mRNA. (C) Cell proliferation of MerTK stable knockdown MHCC97H, HCCLM3, and Huh7 cells lines was measured by cell counting. Data are presented as mean ± SD of three independent experiments, statistical significance was assessed by One-way ANOVA with Bonferroni post-test, ***P < 0.001. (D and E) Western blot analysis demonstrating MerTK overexpression in LO2 cells and the indicated HCC cells (D). Cell proliferation of MerTK overexpression LO2, MHCC97H, HCCLM3, and Huh7 cells lines were measured by cell counting (E). Cont, refers to cells transduced with empty vector and hereafter; MerTK OE, refers to cells transduced with plasmid overexpressing MerTK and hereafter. Data are presented as mean ± SD of three independent experiments, statistical significance was determined by Student’s t-test, *P < 0.05. (F and G) Representative images (F) and quantification (G) of colony formation assay of MerTK stable knockdown or overexpression MHCC97H and HCCLM3 cells. Data are presented as mean ± SD, statistical significance was assessed by One-way ANOVA with Bonferroni post-test and Student’s t-test, *P < 0.05, **P < 0.01. (H and I) Immunoblots of tumor growth related proteins p-GSK3β, p-Akt, Akt in MerTK stable knockdown (H) or overexpression (I) MHCC97H and HCCLM3 cells. β-Actin was used as a loading control.
Fig. 3. MerTK modulates bioenergetics metabolism through promoting the Warburg Effect and suppressing OXPHOS in HCC cells. (A and C) The intact cellular ECAR of control, MerTK stable knockdown (A) and overexpression (C) MHCC97H and HCCLM3 cells were measured in real time using the Seahorse XF96 Extracellular Flux Analyzer. Data were analyzed by Seahorse XF-96 Extracellular Flux Analyzer. ECAR is reported in mPH/minute, the results were normalized to cell number. Data are presented as mean ± SD, statistical significance was assessed by One-way ANOVA with Bonferroni post-test, *P < 0.05, **P < 0.01. (B and D) The intact cellular OCR of control, MerTK stable knockdown (B) and overexpression (D) MHCC97H and HCCLM3 cells were measured in real time using the Seahorse XF96 Extracellular Flux Analyzer. Data were analyzed by Seahorse XF-96 Extracellular Flux Analyzer. OCR is reported in pmols/minute, the results were normalized to cell number. Data are presented as mean ± SD, statistical significance was assessed by One-way ANOVA with Bonferroni post-test, *P < 0.05, **P < 0.01. (E) Glucose uptake of MerTK stable knockdown or overexpression MHCC97H, HCCLM3 by 2-NBDG incorporation by FACS. Data are presented as mean ± SD, statistical significance was assessed by One-way ANOVA with Bonferroni post-test and Student’s t-test, *P < 0.05, **P < 0.01. (F) Lactate production of MerTK stable knockdown or overexpression MHCC97H, HCCLM3 cells were measured following manufacturer’s instructions. Data are presented as mean ± SD, statistical significance was assessed by One-way ANOVA with Bonferroni post-test and Student’s t-test, *P < 0.05, **P < 0.01. (G and H) Immunoblots of glycolysis related enzymes PGK1, LDHA, PFKM, PKM2, PKDHK1 in MerTK stable knockdown (G) or overexpression (H) MHCC97H and HCCLM3 cells. β-Actin was used as a loading control.
Fig. 4. MerTK promotes HCC cell growth in vivo. (A) Representative dissected tumors derived from control and MerTK knockdown MHCC97H, HCCLM3, Huh7 and HepG2 cells are shown. (B) Growth of tumors derived from control and MerTK knockdown cells are shown. n = 8 (MHCC97H and HCCLM3 cells) or 5 (Huh7 and HepG2 cells). Data are presented as mean ± SEM, statistical significance was determined by Student’s t-test, **p < 0.01, ***p < 0.001. (C) Tumor weights at the end of the experiment are shown. Data are presented as mean ± SEM, statistical significance was determined by Student’s t-test, ***p < 0.001. (D) MerTK expression in the lysates of tumor tissue (A) are shown. (E) Immunoblots of tumor growth related proteins p-GSK3β, p-Akt and glycolysis related enzymes PGK1, LDHA, PFKM, PKM2, PDHK1 in tumor tissues derive from nude mice. β-Actin is used as a loading control. (F) Representative dissected tumors derived from control and MerTK overexpression MHCC97H cells are shown. (G and H) Tumors growth were measured once in two days (G) and tumor weights are measured at the end of the experiment (H). n = 8. Data are presented as mean ± SEM, statistical significance was assessed by Student’s t-test, *p < 0.05, **p < 0.01. (I) MerTK expression in the lysates of tumor tissue (F) are shown.
(Mertk\(^{-/-}\)) mice by crossing Mertk\(^{+/+}\) with Mertk\(^{-/-}\) mice. Western blotting analysis revealed that Mertk was knockout in all tissues, including heart, liver, spleen, lung, kidney, and skeletal muscle (Fig. 5A). To further investigate the contributions of MerTK to HCC development, the mouse liver tumors were induced by DEN and CCl\(_4\) in male mice of two genotypes, Mertk\(^{+/+}\) and Mertk\(^{-/-}\) (Fig. 5B). The tumor burden (number and size) was then recorded at 30 weeks of age (Fig. 5C and D). Strikingly, our results showed that the tumor numbers detected in the Mertk\(^{-/-}\) mice were much less than those in the Mertk\(^{+/+}\) control littermates, and the tumor size was also significantly smaller. Immunohistochemical analysis confirmed that MerTK was expressed in livers of the Mertk\(^{-/-}\) and Mertk\(^{+/+}\) mice (Fig. 5E). Histological examination revealed extensive liver pathology associated with severe steatosis in the DEN and CCl\(_4\)-treated Mertk\(^{-/-}\) mice (Fig. 5F). Moreover, the tumor lesions were positive for alpha-fetoprotein (AFP), an established liver tumor marker (Fig. 5G and H). These findings suggest that Mertk deletion suppresses chemical-induced liver tumorigenesis.

3.6. N-glycosylated MerTK is essential for HCC cell growth

MerTK was found to be heavily glycosylated on multiple asparagines residues which may regulate MerTK activity [50]. Thus, we asked whether N-glycosylation of MerTK is pivotal for the proto-oncogenic property in HCC. We used tunicamycin (TM), a GlcNAc phosphotransferase inhibitor which blocks the formation of N-glycosidic linkages, and swainsonine which can completely inhibit mammalian Golgi α-mannosidase II which is critical in the formation of N-glycosylation of proteins, as well as PNGase F which cleaves between the innermost GlcNAc and asparagine residues of high mannose, hybrid, and complex oligosaccharides. Expectedly, due to TM, swainsonine and PNGase F treatment, we found the molecular weight of MerTK was largely reduced which indicates that the higher molecular weight form is indeed N-glycosylated MerTK. We further treated the cells with or without glucose and found the molecular weight of MerTK was gradually reduced with time, which suggests that N-glycosylation was reduced due to glucose starvation (Fig. 6A). Next, we asked which asparagine site(s) of MerTK is/are glycosylated. According to the bioinformatics analysis, we mutated the N294, N395, N442, N454 (N was replaced by Q) and Y747, 753 and 754 (hereafter named 3Y, Y was replaced by A) in MERTK, and found N294 and N454 were crucial sites for N-glycosylation, but not N395 and N442 residues. Intriguingly, we found triple tyrosine mutant also deprived N-glycosylation of MerTK (Fig. 6B and C). To further understand the N-glycosylation of MerTK on HCC cell growth, LO2, MHCC97H and HCCLM3 cells were transfected with control, N294Q, N395Q, N442Q, N454Q and 3Y mutant plasmids. Cell growth was not enhanced by expression of N294Q, N454Q and 3Y mutants in
Fig. 6. N-Glycosylation of MerTK is Essential for HCC Cell Growth. (A) Western blot analysis of MHCC97H, HCCLM3 treated with or without Tunicamycin (TM), PNGase F, Swainsonine and glucose starvation. (B) Structure of MerTK and the modification site of N294Q, N454Q and 3Y. (C and D) MerTK expression was detected by Western blot in LO2, MHCC97H and HCCLM3 cells transfected with the N294Q, N395Q, N442Q, N454Q and 3Y mutant plasmids, respectively (C). Cell proliferation was analyzed by cell counting (D). Data are presented as mean ± SD of three independent experiments, statistical significance was assessed by One-way ANOVA with Bonferroni post-test, *P < 0.05. (E and F) The intact cellular ECAR and OCR of WT and indicated MerTK mutant LO2 Cells were measured in real time using the Seahorse XF96 Extracellular Flux Analyzer. Data are presented as mean ± SD, statistical significance was assessed by One-way ANOVA with Bonferroni post-test, *P < 0.05. (G) Glucose uptake of WT and MerTK indicated mutant LO2 Cells by 2-NBDG incorporation by flow cytometry. Data are presented as mean ± SD, statistical significance was assessed by One-way ANOVA with Bonferroni post-test, *P < 0.05, n. s., not significant.
vitre, whereas N395Q and N442Q mutants in which MerTK were still glycosylated exhibited pro-growth activity in these three cell lines (Fig. 6D). Moreover, we determined the ECAR and OCR alteration due to MerTK point mutation. N395Q- and N442Q-MerTK could effectively upregulate glycolytic rate, while mitochondrial oxidative phosphorylation (OXPHOS) was suppressed (Fig. 6E and F; Figs. S5A and B). The impacts of MerTK mutants on cellular bioenergetics were confirmed by glucose uptake only increased in N395Q- and N442Q-MerTK overexpressed LO2 cells (Fig. 6G). Together, the results indicate that MerTK is exclusively N-glycosylated at N-294 and N454, which is 3′ dependent, and N-glycosylated MerTK promotes both HCC cells and normal liver cells (LO2) growth.

3.7. N-glycosylation is crucial for MerTK stability

Next, we asked whether deglycosylated MerTK is sensitive to ubiquitination-dependent degradation. We treated LO2, MHCC97H and HCCLM3 cells with protease inhibitor MG132. Deglycosylated MerTK was increased in MG132 treated LO2 cells, but not in MHCC97H and HCCLM3 cells (Fig. 7A). However, MerTK was increased upon MG132 treatment when MerTK was deglycosylated by TM (Fig. 7B and C). We co-immunoprecipitated MerTK in TM or both TM and MG132 treated MHCC97H and HCCLM3 cells, and the immunoprecipitates were examined by Western blot with an anti-ubiquitin antibody. The data showed that ubi-MerTK level in MHCC97H and HCCLM3 cells was significantly increased following TM treatment (Fig. 7D). Importantly, MG132 treatment caused much more ubi-MerTK accumulation than the control (Fig. 7D), suggesting N-glycosylation is crucial for MerTK stability. To understand whether these mutants (N294Q, N454Q and 3Y) would affect MerTK stability, we treated MHCC97H cells with Cycloheximide (CHX, 125 μg/ml) at indicated intervals to examine MerTK turnover. N294, N454 and 3Y mutants of MerTK exhibited shorter half-life than G-MerTK (Fig. 7E). These data indicated that deglycosylated at N294 and N454 of MerTK is more sensitive to degradation. Intriguingly, dephosphorylation at Y749, Y753 and Y754 is crucial for N-glycosylation of MerTK which could also regulate the ubiquitin-dependent degradation. Previous work has demonstrated that ligand activated MerTK homodimerization is essential for autophosphorylation and subsequently stimulated multiple signals transduction as well as enhanced MerTK stability. We further asked whether N-glycosylation is indispensable for stabilizing MerTK. To clarify our hypothesis, we analyzed the dimer status of wild type MerTK and various mutant MerTK. MerTK mutant of N294Q, N454Q and 3Y impaired the MerTK dimerization (Fig. 7F). Strikingly, glucose deprivation and TM treatment caused deglycosylation of MerTK and almost completely abolished the ability to form dimer (Fig. 7G). Taken together, these data indicated that N-glycosylation and phosphorylation are essential for MerTK stability.

3.8. Nuclear located non-glycosylated MerTK is indispensable for HCC cell survival under stress

To further investigate the essential role of MerTK in promoting HCC cell survival, we examined cell death in response to glucose starvation. Intriguingly, we found glucose starvation induced gradually deglycosylation of MerTK as well as upregulated MerTK protein expression (Fig. 8A, upper panel). Furthermore, glucose starvation increased MERTK mRNA level in a time dependent manner (Fig. 8A, lower panel). To investigate pro-survival activity of MerTK under stress, we cultured LO2 cells that express low MerTK in glucose-free medium and collected the cells at 0, 6, 12 and 24 h followed by Western blot analysis. We found that MerTK expression was not affected under glucose starvation (MHCC97H used as a MerTK positive control to know the location of MerTK in blot) (Fig. 8B, left panel). We compared cell apoptosis rate of LO2, MHCC97H and HCCLM3 with or without glucose for 24 h, and found apoptosis was significantly induced in LO2 but only modestly in MHCC97H and HCCLM3 cells (Fig. 8B, right panel). Previous studies have demonstrated that prolonged Gas6 addition can promote MerTK translocation from cell surface to nucleus, which is related to the MerTK deglycosylation [51]. Hence, we further evaluated the sub-cellular location of N-glycosylated and deglycosylated MerTK. Our data shows a modest nucleus translocation of deglycosylated MerTK in both LO2 and MHCC97H cells transfected with N294Q- and N454Q-MerTK (Fig. 8C). Furthermore, we found that TM treatment induces a higher nuclear translocation of total and phosphorylated MerTK in MHCC97H cells (Fig. 8D, lane 8). Indeed, when phosphorylation was inhibited by UNC-2250, nucleus located MerTK was suppressed (Fig. 8D, lane 9), and cytoplasmic MerTK was also reduced significantly (Fig. 8D, lane 6). We also found UNC-2250 treatment reduced MerTK level (Fig. 8E) as well as facilitated cell apoptosis initiated by glucose starvation or TM treatment in MHCC97H cells (Fig. 8F). Further work is required to validate the downstream signaling induced by nucleus-located MerTK. Collectively, these results suggest that phosphorylation of deglycosylated MerTK is also required for nucleus translocation to facilitate HCC cell survival under stress condition.

4. Discussion

In this study, we report the pro-tumorigenic activity of MerTK in HCC through regulating Akt-GSK3α/β signaling as well as cellular bioenergetics. We also demonstrate that N-glycosylation is critical for MerTK homodimerization and stability. Importantly, we found tyrosine phosphorylation is indispensable for nucleus translocation of deglycosylated MerTK. These findings renovate current understanding of MerTK promoted tumorigenesis.

Aberrantly upregulated MerTK was widely found in multiple cancer, and MerTK inhibition probably caused cell growth arrest as well as increased cell death. Previous work has demonstrated that MerTK is a pivotal regulator of certain tumor associated signal transduction, such as Akt, Src, p38, ERK1/2, GSK3/β, MAPK, STAT5. CHK-2, focal adhesion kinase (FAK) and STAT6 [49,50]. Consistently, we show that MerTK depletion dramatically suppresses HCC cell growth in vitro and in vivo. In HCC, we found MerTK functions as a metabolic modulator by integrating aerobic glycolysis and OXPHOS. PGK1, LDHA, PKFM, PKM2 and PDHK1 are key downstream factors of MerTK in regulating the Warburg effect (Fig. 9). Our study unravels a novel role of MerTK in modulating cellular bioenergetics which enriches the metabolic regulator property of MerTK. However, further investigation will be needed to understand whether Akt may also regulate PKFM phosphorylation and stabilization in HCC cells.

Several studies have demonstrated and highlighted the glycosylation (both O-glycosylation and N-glycosylation) in regulating crucial TK receptors in carcinogenesis [52]. N-glycosylation of EGFR is essential for the stability and resistant to TK inhibition and N-glycosylation is also indispensable for the oncogenic property of c-Met [53,54]. N-linked glycosylation is a highly conserved post-translational modification via the addition of carbohydrates to regulate functions of various proteins, including protein folding, assembly, intracellular trafficking. As a transmembrane molecule, N-glycosylation commonly occurs at certain asparagine of MerTK. We identified two asparagines at N294 and N454 are important for MerTK N-glycosylation. Intriguingly, 3Y mutants also disturbed MerTK glycosylation. We demonstrated deglycosylation of MerTK deprived pro-tumor growth and metabolic reprogramming properties, which emphasized the significance of MerTK N-glycosylation. Additionally, partial deglycosylated MerTK could be stabilized in response to MG132 treatment in LO2 cells, but not in MHCC97H and HCCLM3 cells. However, TM caused non-glycosylated MerTK could also be stabilized by MG132 in MHCC97H and HCCLM3 cells. We validated that N-glycosylation is indispensable for MerTK to form dimmers which contributes to the increased stabilization. We also found under glucose starvation or TM treatment, MerTK fails to form homodimerization efficiently. Since MerTK, similar to other RTKs, was found to form dimer and activate auto-phosphorylation upon ligand stimulation which is
Fig. 7. N-glycosylation modification stabilizes MerTK. (A) Western blot analysis of LO2, MHCC97H and HCCLM3 treated with indicated concentration of MG132 for 24 h followed by Western blot analysis with indicated antibodies. (B) MHCC97H and HCCLM3 cells were treated with TM (1 μg/ml) and/or indicated concentration of MG132 for 24 h followed by Western blot analysis with indicated antibodies. (C) Western blot analysis of MHCC97H and HCCLM3 cells treated with TM (1 μg/ml) and/or MG132 (125 μg/ml) (left panel) and relative DG-MerTK expression level was quantified by Image J and normalized to the β-Actin (right panel). Data are presented as mean ± SD of three independent experiments, statistical significance was assessed by Student’s t-test, **P < 0.01. (D) MHCC97H and HCCLM3 cells with TM (1 μg/mL) and/or MG132 (125 μg/mL) treatment were subjected to MerTK immunoprecipitation (IP) followed by Western blot analyses with anti-ubiquitin. (E) WT and indicated MerTK mutants expressing in MHCC97H cells were treated with 125 μg/mL CHX at indicated intervals and detected by Western blot analysis. The intensity of MerTK protein was quantified using Image J and normalized to the β-Actin. Data are presented as mean ± SD of three independent experiments, statistical significance was assessed by One-way ANOVA with Bonferroni post-test, **P < 0.01. (F) SDS-PAGE and Blue Native PAGE of WT and indicated MerTK mutants expressing in MHCC97H and HCCLM3 cells to detect the MerTK dimerization. (G) SDS-PAGE and Blue Native PAGE of endogenous MerTK in MHCC97H cells treated with or without TM or in presence or absence of glucose (left panel). Proposed model of N-glycosylation and phosphorylation modification modulating MerTK homodimerization (right panel).
crucial for several downstream signals’ activation. N-glycosylation modification enhanced MerTK stability maybe the reason why MerTK protein level was increased in HCC tissues while the mRNA transcripts remained unchanged. In this context, we found that MerTK depletion increased the sensitivity of HCC cells to cellular stress such as glucose starvation. Under glucose starvation, increased MERTK transcription was exhibited in a time-dependent manner. Previous work demonstrated that in certain instances, TAM may heterodimerize or cross-phosphorylate [55], which may serve as the signaling for the nuclear translocation of MerTK. We proposed that TM treatment caused MerTK deglycosylation, therefore, MerTK may interact with other TAM or kinases to cross-phosphorylate or be phosphorylated. Furthermore, our results showed that decreased nucleus-located MerTK was more sensitive to glucose starvation or TM treatment which highlighted the pro-survival roles of nuclear MerTK. But whether nucleus located MerTK could bind to chromatin subsequently to activate MERTK transcription or regulate other potential transcriptional factors to promote MERTK transcription need more evidences. Complete understanding of molecular mechanism of MerTK in promoting HCC tumorigenesis maybe important for future developing effective therapeutic strategies in the

Fig. 8. MerTK is indispensable for HCC cell survival under glucose starvation stress or N-glycosylation blocked by TM. (A) MHCC97H and HCCLM3 cells were cultured in the presence or absence of glucose at indicated intervals. MerTK protein (upper panel) and mRNA (lower panel) levels were analyzed by Western blot and qRT-PCR, respectively. (B) Western blot analysis of MerTK expression in LO2 cells under glucose starvation at indicated intervals (left panel). LO2, MHCC97H and HCCLM3 cells were cultured with or without glucose for 24 h, and cell apoptosis was measured by FACS (right panel). Data are presented as mean ± SD of three independent experiments, statistical significance was assessed by One-way ANOVA with Bonferroni post-test, **P < 0.01. (C) Western blot analysis of nuclear and cytoplasmic located MerTK in LO2 cells transfected with WT and various MerTK mutants. (D) Western blot analysis of nuclear and cytoplasmic located MerTK in MHCC97H cells treated with TM (1 μg/mL) and/or UNC-2250 (16 μM) for 24 h. (E and F) Western blot analysis of MerTK and p-MerTK in MHCC97H cells treated with TM (1 μg/mL) and/or UNC-2250 (16 μM) or in presence or absence of glucose (E). Apoptosis rate was analyzed by FACS (F). Data are presented as mean ± SD, statistical significance was assessed by One-way ANOVA with Bonferroni post-test, **P < 0.01, ***P < 0.001.
HCC treatment.

5. Conclusions

In summary, we elucidated the critical role of MerTK in regulating HCC cell growth and revealed the N-glycosylation is essential for protein stability and tumor-promoter activity of MerTK. Moreover, phosphorylated MerTK could translocate to nucleus that renders HCC cells a cell survival response. Our present study provides insights into the molecular mechanism of MerTK in promoting HCC tumorigenesis that maybe important for targeting MerTK drug development and developing effective therapeutic strategies in the treatment of HCC.

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Author contributions

B.L conceived the project and designed the experimental protocols; Y.L., L.L., Y.-J.L., J.L., L.H., Y.D., M.F., J.-W. L., F.S., J.-P.L., J.W., and K.H. performed most of the experiments; Y.-J.L., and L.H. performed the DEN and CCl₄-induced HCC mouse model; L.H. and F.S. assisted the animal experiments; Y.L., L.L., Y.-J.L., L.H., J.-W. L., and B. L. analyzed the data. L.L., Y.-J.L., and J.-W. L. contributed to the statistical analyses of the data. B.L. supervised the study and wrote the manuscript.

Declaration of competing interest

The authors declare that they have no competing interests.

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Appendix A. Supplementary data

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