Deficient DNASE1L3 facilitates neutrophil extracellular traps-induced invasion via cyclic GMP-AMP synthase and the non-canonical NF-κB pathway in diabetic hepatocellular carcinoma

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Received 24 March 2021; Revised 28 February and 25 March 2022; Accepted 26 March 2022

doi: 10.1002/cti2.1386

Original Article

Abstract

Objective. Diabetic hepatocellular carcinoma (HCC) patients have high mortality and metastasis rates. Diabetic conditions promote neutrophil extracellular traps (NETs) generation, which mediates HCC metastasis and invasion. However, whether and how diabetes-induced NETs trigger HCC invasion is largely unknown. Here, we aimed to observe the effects of diabetes-induced NETs on HCC invasion and investigate mechanisms relevant to a DNA sensor cyclic GMP-AMP synthase (cGAS).

Methods. Serum from diabetic patients and healthy individuals was collected. Human neutrophil-derived NETs were isolated for stimulating HCC cell invasion. Data from the SEER and TCGA databases were used for bioinformatics analysis. In HCC cells and allograft models, NETs-triggered invasion was observed.

Results. Diabetic HCC patients had poorer survival than non-diabetic ones. Either diabetic serum or extracted NETs caused HCC invasion. Induction of diabetes or NETosis elicited HCC allograft invasion in nude mice. HCC cell invasion was attenuated by the treatment with DNase1. In TCGA_LIHC, an extracellular DNase DNASE1L3 was downregulated in tumor tissues, while function terms (the endocytic vesicle membrane, the NF-κB pathway and extracellular matrix disassembly) were enriched. DNASE1L3 knockdown in LO2 hepatocytes or H22 cell-derived allografts facilitated HCC invasion in NETotic or diabetic nude mice. Moreover, exposure of HCC cells to NETs upregulated cGAS and the non-canonical NF-κB pathway and induced expression of metastasis genes (MMP9 and SPP1). Both cGAS inhibitor and NF-κB RELB knockdown diminished HCC invasion caused by NETs DNA. Also, cGAS inhibitor was able to retard translocation of NF-κB RELB.

Conclusion. Defective
DNASE1L3 aggravates NETs DNA-triggered HCC invasion on diabetic conditions via cGAS and the non-canonical NF-κB pathway.

Keywords: cyclic GMP-AMP synthase, diabetes, hepatocellular carcinoma, metastasis, neutrophil extracellular traps, non-canonical NF-κB pathway

INTRODUCTION

Primary liver cancer is an increasing global health challenge with higher mortality and limited treatment options. Hepatocellular carcinoma (HCC) is the most common type of primary liver cancer and accounts for ~90% of cases.1 Intrahepatic invasion and distant metastases are the major cause of HCC-related deaths.2 Recent studies have demonstrated that neutrophils were able to promote HCC metastasis through forming neutrophil extracellular traps (NETs) during the process of NETosis.3,4 However, detailed molecular mechanisms underlying NETs-induced metastasis of HCC cells remain largely unknown.

Neutrophils usually undergo NETosis under multiple pathological conditions, such as pathogen infections,5 and pathophysiological states including diabetes mellitus (DM)6–9 and surgical stress.4 A growing body of evidence supports the notion that diabetic HCC patients had significantly higher rates of postoperative recurrence, metastasis and mortality than those non-diabetic HCC ones.10–17 Nevertheless, it is not clear whether diabetes-related HCC invasion or metastasis is affected by excessive NETs. As known, the secreted NETs consist of multiple components, such as granule proteins, like neutrophil elastase (NE) and myeloperoxidase (MPO), as well as chromatin double-stranded DNA (dsDNA). It was demonstrated that these extracellular NETs were able to trap tumor cells in the lungs and liver to fuel their distant metastasis18,19 and that such effects partially depended on dsDNA.3 However, whether dsDNA in extracellular NETs is responsible for diabetes-related HCC invasion or metastasis is rarely reported. Normally, there is only a trace amount of free dsDNA in the body and it is precisely maintained at a relatively stable level by DNases in/out the cells. A recent analysis of the TCGA and GEO databases surprisingly showed that expression of DNASE1L3, an extracellular DNase, is very low in HCC tissues,20 which may create a NETs DNA-rich microenvironment, thereby promoting invasion and/or metastasis.

In the cytosol, DNA can be sensed by pattern recognition receptors, among which cyclic GMP-AMP synthase (cGAS) is a key member. Once binding dsDNA, cGAS will be activated and synthesise the second messenger cyclic guanosine monophosphate-adenosine monophosphate (cGAMP).21 Activation of cGAS can initiate two distinctive effects: anti-inflammation/anti-cancer effect depending on stimulator of interferon genes (STING) and interferon,22 as well as pro-metastasis effect associated with the non-canonical NF-κB pathway.23 Hence, it is still necessary to elucidate whether and how cGAS is involved in NETs DNA-induced HCC invasion on diabetic conditions.

In the present study, we observed the effects of diabetes-induced NETosis on HCC invasion. With bioinformatics analysis and molecular biology assays, we investigated the mechanisms underlying NETs DNA-induced HCC invasion.

RESULTS

Diabetic conditions foster HCC cell invasion

Survival statistics of SEER*Stat showed that the survival rate of diabetic HCC patients was decreased when compared with that of non-diabetic HCC patients (Figure 1a). Since metastasis is a prominent cause of death in HCC patients, we speculated that diabetic conditions probably induced HCC cell metastasis or local invasion. In the transwell assay, it was found that treatment of HCC SMMC7721 cells with diabetic serum dramatically promoted invasion (P < 0.01), while remarkable invasion was not observed in the cells treated with healthy serum (Figure 1b). In addition, the effect of diabetic serum on invasion of LO2 hepatocytes was weaker than that on HCC SMMC7721 cells (Figure 1c). Moreover, we
established a diabetic nude mouse model, which was verified by high blood glucose levels (14.8 ± 4.2 vs. 7.6 ± 0.6 mM, P < 0.01, Figure 1d). In the two groups of nude mice, mouse HCC H22 cells were engrafted. After a 14-day growth of the allografts, the mice were sacrificed, and then the body and the allografts were weighed. As shown in Figure 1e, the body weight of diabetic nude mice was significantly elevated (P < 0.01). Importantly, the weight of HCC allografts in diabetic nude mice was also increased as compared with that in control nude mice (P < 0.05, Figure 1f and g). These data suggest that diabetic conditions can enhance HCC cell invasion.

DNA mediates NETs-induced HCC cell invasion

Given that diabetic conditions can increase NETs content, it was expected that the aforementioned enhancement of HCC invasion may be attributed to NETs generation. A cellular model of NETosis was established by treating human peripheral blood neutrophils with PMA. As shown in Figure 2, the treatment with 20 nM PMA for 2 h released a massive amount of net-like structure from the neutrophils, which was verified to be DNA by staining with Hoechst (Figure 2a) and PicoGreen (Figure 2b). The release of NETs from neutrophils is shown in the schematic diagram.
NETs promote diabetic hepatocellular carcinoma invasion

NL i et al

Figure 2. Roles of extracellular DNA in NETs-induced HCC cell invasion. (a–e) Neutrophils isolated from human peripheral blood were treated with 20 nM PMA for 2 h. DNA was labelled using Hoechst 33324 staining followed by fluorescence photography (a). Extracellular DNA was collected and quantified with a fluorescence microplate reader following PicoGreen staining (b). A schematic diagram showing NETs release from neutrophils (c). Granule proteins (MPO, MMP9 and NE) in the nets from PMA-treated or control cell supernatant were measured by Western blot assay after total proteins were quantified with a BCA kit (d), and the band intensity was quantified with the ImageJ software (e). (f–i) HCC SMMC7721 cells were stimulated with increasing concentrations of NETs (f). SMMC7721 cells or LO2 hepatocytes were treated with 0.2 μg mL⁻¹ NETs (g). SMMC7721 cells were treated with 0.2 μg mL⁻¹ NETs (h) or diabetic serum (i) in the absence or presence of 1.5 U DNase1. The cell invasion was observed with the transwell assay (mean ± SD, n = 4). *P < 0.05, **P < 0.01 vs. Control group. ##P < 0.01 vs. DNase1-free group.
Deficient DNASE1L3 facilitates NETs DNA-induced HCC cell invasion

To understand why HCC SMMC7721 cells were sensitive to NETs DNA or diabetic serum, we firstly investigated gene expression of various DNA-degrading enzymes in the TCGA_LIHC cohort. Among intracellular DNA-degrading enzymes (DNASE1L1, DNASE2, DNASE2B, DFFA, DFFB, TATDN1, TATDN2, TATDN3, TREX1 and FEN1), remarkable changes in gene expression were not observed (Figure 3a). Among extracellular DNA-degrading enzymes, the expression of ENDD1, DNASE1 and DNASE1L2 was also not markedly different between HCC and normal tissues (Figure 3b–d). Additionally, the survival could not be predicted or stratified by the expression of these enzymes (Figure 3e–g).

In various TCGA cancers, the expression of DNASE1L3 was downregulated in tumor tissues (Figure 4a and b). From the TCGA_LIHC cohort, it could be seen that the 5-year survival of HCC patients with low DNASE1L3 was dramatically poor (P < 0.0001, Figure 4c). The ROC curve of DNASE1L3 showed that the AUC score was 95%, indicating good performance in distinguishing HCC from normal tissues (Figure 4d). The examination of protein levels showed that DNase1L3 was also downregulated in HCC SMMC7721 cells compared with LO2 hepatocytes (Figure 4e). Additionally, such downregulation was observed in SMMC7721 cell supernatant (Supplementary figure 1). Importantly, after knocking down DNASE1L3 in LO2 hepatocytes (Figure 4f), NETs DNA-induced invasion was significantly enhanced (P < 0.01, Figure 4g). The results indicate that deficient DNASE1L3 in HCC tissues can exacerbate tumor invasion by reducing extracellular DNA degradation.

In the nude mice, induction of NETosis with LPS was able to significantly promote the development of mouse HCC H22 cell-derived allografts (P < 0.01). The knockdown of DNASE1L3 in H22 cells (Supplementary figure 2) was able to facilitate the effects of NETosis (P < 0.01) (Figure 5a–c). In the diabetic nude mice, the levels of NETs DNA were increased (P < 0.01) (Figure 5d). Similarly, the knockdown of DNASE1L3 in H22 cells promoted the invasive growth of the allografts (Figure 5e and f).

Bioinformatics analysis of differentially expressed genes and functional enrichment in tumor tissues of HCC patients

To dissect the possible mechanism underlying NETs DNA-induced HCC invasion, the TCGA_LIHC cohort was downloaded and analysed for differential gene expression. As shown in Figure 6a, under a criteria of P < 0.001 and fold of change (FC) > 1.8 (i.e. log2FC > 0.85), 2270 upregulated genes were identified. Under a criteria of P < 0.001 and log2FC < −0.85, 917 downregulated genes were found. The roles of upregulated genes in HCC development were investigated through screening enriched terms, which were shown in the bubble plot (Figure 6b). These terms (endocytic vesicle membrane, NF-κB signal pathway and extracellular matrix disassembly) were enriched in HCC tissues, indicating that deficient DNASE1L3 is likely to reduce extracellular DNA degradation. Once the accumulated DNA is endocytosed by HCC cells, the NF-κB signal will be activated and extracellular matrix will decompose, thereby boosting tumor invasion.

Non-canonical NF-κB pathway-induced MMP9 and SPP1 expression is responsible for NETs DNA-mediated HCC invasion

To verify the roles of the NF-κB pathway in HCC invasion, the analysis of differentially expressed
genes was performed. Firstly, the genes coding the NF-κB pathway subunits (NFKB2 and RELB) were found to be upregulated (Figure 7a and c). Additional survival analysis showed that the HCC patients with higher levels of NFKB2 or RELB had poorer survival status (Figure 7b and d). Furthermore, the in vitro experiments manifested that the exposure of HCC SMMC7721 cells to NETs DNA significantly enhanced the nuclear translocation of RelB proteins (Figure 7e and f). After knocking down the RELB gene (Figure 7g), NETs DNA-induced cell invasion was remarkably attenuated ($P < 0.01$) (Figure 7h). The results support the notion that the upregulation of the non-canonical NF-κB pathway is detrimental to HCC patients’ survival and metastasis may be a potential cause.

Next, we looked into how the non-canonical NF-κB pathway facilitated HCC invasion. Further
Figure 4. Roles of DNA-degrading enzyme DNASE1L3 in HCC cell invasion. (a) Expression of DNASE1L3 in various TCGA cancers. Data are shown as log2 (tumor median/normal median). (b) Expression of DNASE1L3 in HCC \((n = 371)\) and adjacent normal tissues \((n = 50)\) in the TCGA_LIHC cohort. Data are shown as median ± quartile in the violin plot. (c) Survival analysis of HCC patients between high and low DNASE1L3 expression groups. (d) Receiver operating characteristic (ROC) curve showing false-positive fraction and true-positive fraction of DNASE1L3 expression in the TCGA_LIHC cohort. The AUC is reported. (e) Expression of DNASE1L3 in LO2 hepatocytes and HCC SMMC7721 cells was measured by Western blot assay \((n = 3)\). (f) RNA interference was applied to knock down the expression of DNASE1L3 in LO2 hepatocytes \((n = 3)\). (g) Observation of NETs DNA-induced invasion between DNASE1L3 knockdown and LO2 hepatocytes \((n = 4)\). Data are shown as mean ± SD (e–g).
analysis of the relationship between the non-canonical NF-κB pathway and extracellular matrix disassembly suggested that NFKB2 and RELB interacted with the extracellular matrix disassembly members. In the interaction network, MMP9 was a bottleneck hub gene (Figure 8a). Moreover, during various stages of HCC, the expression of MMP9 was upregulated and the patients with higher MMP9 expression had a relatively lower survival rate (\( P < 0.0001 \)) (Figure 8b and c). In the network, SPP1 was found to be closely related to MMP9 (Figure 8a). Moreover, its expression and effects on survival were similar to MMP9 (Figure 8d and e). Importantly, the treatment with NETs DNA significantly raised MMP9 (Figure 8f) and SPP1 (Figure 8g) protein expression in HCC SMMC7721 cells. Further in vivo study showed that the allografts in NETotic mice had higher MMP9 and SPP1 expression than that in control nude mice. Notably, these effects of NETosis were fostered by the knockdown of DNASE1L3 (Figure 8h).

cGAS as a DNA sensor mediates NETs DNA-induced HCC invasion

To uncover how NETs DNA activated the non-canonical NF-κB pathway and promoted MMP9 and SPP1 expression, intracellular DNA sensor cGAS was investigated. The analysis of the TCGA_LIHC cohort indicated that HCC patients with higher cGAS levels had a poorer survival outcome (\( P < 0.004 \)) (Figure 9a). The treatment of HCC SMMC7721 cells with NETs DNA elevated intracellular cGAMP content in a time-dependent manner (Figure 9b). In addition, this treatment caused the perinuclear translocation of STING protein (Figure 9c). Importantly, the inhibition of cGAS with 20 \( \mu \)g mL\(^{-1} \) RUS521, a cGAS inhibitor, was able to suppress NETs DNA-induced activation of the non-canonical NF-κB pathway, characterised by the reduced RelB nuclear translocation (Figure 9d), as well as HCC cell invasion (Figure 9e). Notably, in the diabetic nude mice, the administration of cGAS inhibitor could
also prevent the allografts from invading (Figure 9f and g). These findings suggest that cGAS is a potential mediator in NETs DNA-induced HCC invasion.

**DISCUSSION**

In the present study, we found that diabetes-induced neutrophil NETosis boosted HCC invasion in a NETs DNA-dependent manner. The deficient DNASE1L3 expression in tumor tissues is a key cause for the impairment of NETs DNA degradation. The resultant accumulation of NETs DNA primed HCC cells to invade by activating the cGAS-ncNF-κB signalling pathway.

In diabetic rodent models, the increased NETs DNA has been demonstrated to promote HCC development. With the SEER*Stat database, the present study shows that diabetic HCC patients have poorer survival than non-diabetic HCC patients. It is surmised that such poor survival is probably attributed to tumor cell metastasis and
or local invasion elicited by diabetic conditions. Using cellular and mouse experiment models, it was found that the invasive growth of HCC cells was indeed enhanced under diabetic conditions. Actually, these findings can be supported by a variety of original articles or data analyses.\textsuperscript{10,12–15,25}

The previous studies showed that diabetic conditions prime neutrophils to undergo NETosis, thereby releasing DNA-containing NETs.\textsuperscript{6–8} In this study, we found that scavenging DNA dramatically suppressed the pro-inflammatory effects of diabetic serum. It is thus believed that the DNA ingredient is necessary for NETs-induced HCC invasion. The treatment of HCC cells with the extracted NETs was also able to induce cell invasion. Therefore, DNA in NETs is necessary and significant during HCC invasion induced by diabetic conditions. A recent study from Song and colleagues also suggested that the high NETs DNA level was a risk for cancer metastasis including HCC.\textsuperscript{18}

Under normal circumstances, extracellular DNA can be degraded to keep balance in the body. However, presently, in HCC tissues one of the extracellular DNA-degrading enzymes, DNASE1L3, was dramatically downregulated. Such downregulation is likely to attenuate DNA degradation, thereby creating a NETs DNA-rich microenvironment.\textsuperscript{20} Interestingly, the knockdown of DNASE1L3 was able to promote invasive growth of HCC allografts both in diabetic and NETotic nude mice. In the cellular experiment, this knockdown was also found to enhance the invasion. Therefore, the present evidence indicates that the effects of NETs DNA will be amplified, when endogenous DNA-degrading enzymes, like DNASE1L3, are defective, for example, in some HCC tissues. Nevertheless, this study did not provide evidence to demonstrate whether diabetes could affect the expression of DNASE1L3, which should be focused on in future studies.

Using the TCGA\_LIHC database, we investigated the mechanisms underlying NETs DNA-triggered HCC cell invasion. The functional analysis of differentially expressed genes showed that DNA metabolism, endocytic vesicle membrane and extracellular matrix disassembly were enriched. Such enrichment indicates that DNA degradation is defective and that the accumulated DNA may be endocytosed by tumor cells, thereby disassembling the extracellular matrix. Additionally, the NF-\textsuperscript{xB} signal pathway was enriched in HCC tissues and the high expression of non-canonical NF-\textsuperscript{xB} could predict poor survival. In the cell imaging assay, the RelB subunit was
translocated to the nucleus after exposure to NETs DNA. Furthermore, the knockdown of RELB attenuated NETs DNA-stimulated cell invasion. The findings provide evidence for the mediation of the non-canonical NF-κB pathway in NETs DNA-induced HCC invasion.\textsuperscript{23,26,27} Furthermore, the
interaction network between the non-canonical NF-κB pathway and extracellular matrix disassembly term revealed that cancer migration genes (MMP9 and SPP1) were involved in NETs DNA-induced invasive growth.28,29

Last but not least, to figure out how NETs DNA promoted HCC invasion, DNA sensors, pattern recognition receptors (PRRs)30 were investigated. As an important member of PRRs, cGAS is traditionally thought to suppress inflammation and cancer by activating dendritic cells.31–33 However, in this study, the survival analysis indicated that HCC patients with higher cGAS had poorer survival. The treatment of HCC cells with NETs DNA raised cGAMP generation and STING protein translocation around the nuclei. Importantly, the inhibition of cGAS attenuated NETs DNA-induced HCC invasion and allograft growth in the diabetic nude mice. These findings appear not to be consistent with those recent reports.32–34 Actually, other researchers have also pointed out that chronic or sustained activation
of cGAS in cancer cells, instead of dendritic cells, could induce inflammation and tumor metastasis.\textsuperscript{23,27,35} It was reported that chronic induction of nuclear DNA leakage into the cytosol activated STING-dependent cytokine production and tumorigenesis. The knockout of STING was almost completely resistant to 7,12-dimethylbenz[a]anthracene (DMBA)-induced skin carcinogenesis,\textsuperscript{36} which supports the present findings. Therefore, we believe that diabetes-induced NETs DNA could constantly activate cGAS and induce perinuclear translocation of STING in HCC cells, thereby promoting local invasion and/or metastasis. Therefore, in future immunotherapy of HCC patients, activating cGAS-STING in dendritic cells should be carefully handled to avoid arousing tumor cells nearby.\textsuperscript{37}

In summary, diabetes-induced NETs release is an important cause for HCC invasion. Such effects depend on non-canonical NF-κB pathway-mediated MMP9 and SPP1 expression. The activation of DNA sensor cGAS induced by deficient DNASE1L3 is involved in the above process. This study provides a significant clue for the treatment of HCC metastasis in diabetic patients.

METHODS

Human blood samples

Blood samples were drawn from four clinically diagnosed diabetic patients and four healthy individuals, who were enrolled from the 2nd Affiliated Hospital of Guangzhou Medical University. Serum was separated from the blood samples by centrifugation at 1800 g for 3 min. The experimental operation was approved by the Medical Ethics Committee of Guangzhou Medical University.

Isolation of neutrophils and purification of NETs

Neutrophils were isolated using Histopaque 11119 (Sigma-Aldrich, St Louis, MO, USA) and Percoll\textsuperscript{39} Plus Density Gradient Media, as described previously.\textsuperscript{5,8} The isolated neutrophils were incubated with 25 nm phorbol-12-myristate-13-acetate (PMA) (Sigma-Aldrich) for 2 h to induce NETosis.\textsuperscript{8} After incubation with PMA, the cell culture supernatant of neutrophils was collected. The cell bottoms were rinsed with Ca\textsuperscript{2+}/Mg\textsuperscript{2+}-free PBS. The collected supernatant and the rinsed PBS were transferred into a centrifuge tube for centrifugation at low speed (450 g, 10 min, 4°C). The supernatant was collected and then suffered from further centrifugation at high speed (20 000 g, 20 min, 4°C), and the pellet was collected. TE buffer was used to re-suspend the pellet, and the content of NETs DNA was quantified using NanoDrop\textsuperscript{TM} 2000. The NETs solution was stored at −20°C for the following experiments. The protocol was approved by the Ethics Committee of Guangzhou Medical University, China.

Detection of NETs DNA content

NETs DNA was measured as described previously.\textsuperscript{8} Briefly, after treatment with 20 nm PMA for 2 h, the neutrophils in 24-well plate were stained with Hoechst 33324 followed by photofluorography. The released DNA in cell culture supernatant was tested with Quant-iT\textsuperscript{TM} PicoGreen\textsuperscript{TM} dsDNA Assay Kits (Invitrogen, MD, USA). The fluorescence intensity was measured with a fluorescent microplate reader (Thermo, MA, USA).

Animal experiments

Six-week-old nude mice were bought from Nanjing Biomedical Research Institute (Nanjing, China) and maintained under SPF conditions. All experiments were performed under the approval of the Animal Ethics Committee of Guangzhou Medical University.

After a 1-week adaptation, 80 nude mice were randomly divided into four groups. Half of the mice were treated intraperitoneally with 1 mg kg\textsuperscript{−1} lipopolysaccharide (Sigma-Aldrich) and the other half were given normal saline (NE). When serum NETs DNA was increased in the lipopolysaccharide group, DNASE1L3 knockdown or wild-type mice HCC H22 cells (1 × 10\textsuperscript{6} cells/side/mice) were subcutaneously transplanted. These cells have highly proliferative and invasive abilities. After 3 weeks, all of the mice were sacrificed, and the allografts and protein expression were measured.

Diabetic animal models were established as described previously.\textsuperscript{17,38} Firstly, the nude mice were randomly divided into two groups: diabetes mellitus (DM, n = 30) and control (n = 10). In the DM group, the mice were fed a high-fat and carbohydrate diet for a week, followed by a combination of a single intraperitoneal injection of streptozotocin (1% w/v solution in fresh cold sodium citrate buffer, pH 4.4) at a dose of 150 mg kg\textsuperscript{−1}. Before the administration, the mice were deprived of food for 12 h. The control mice were injected with sodium citrate buffer.

Glucose levels were detected 3 days after streptozotocin injection using blood samples collected from the tail vein. Blood glucose > 9.5 mmol\textsuperscript{−1} was considered diabetic, and such mice were eligible for further experiments. One week after the onset of hyperglycaemia, four mice were randomly selected from either group for testing blood NETs. The rest of the diabetic nude mice were divided into three groups: in two groups, DNASE1L3 knockdown H22 cells (1 × 10\textsuperscript{6} cells/side/mice) were subcutaneously transplanted, and the other group and control group just received wild-type H22 cells. Four days later, the mice in one of the DNASE1L3 knocked-down groups were treated with RU521 (1.5 mg kg\textsuperscript{−1}) by intraperitoneal injection every day,\textsuperscript{39} and those in the other group were given normal saline. Three weeks later, all the mice were sacrificed, and the allografts were measured.
Data collection and analysis

The survival data of 62 952 HCC patients and 220 diabetic HCC patients were downloaded and analysed using SEER*Stat 8.3.8 software. For the Cancer Genome Atlas (TCGA) data, the heat map of genes was obtained from UALCAN (http://ualcan.path.uab.edu/analysis.html). The analysis of differentially expressed genes was performed using the TCGAbiolinks R package. The protein-protein interaction (PPI) was analysed with STRING v11.0 (https://string-db.org/), and the network of core genes was constructed using the Cytoscape software.

Cell culture

Human HCC SMMC7721 cells were purchased from Cell Bank of Type Culture Collection of Chinese Academy of Sciences (Shanghai, China). Human LO2 hepatocytes were supplied by Alexan Biotechnology Company (Guangzhou, China). Mouse HCC H22 cells were bought from Procell Life Sciences (Shanghai, China). Human LO2 hepatocytes were supplied by Alexan Biotechnology Company (Guangzhou, China). Mouse HCC H22 cells were bought from Procell Life Sciences (Shanghai, China). Human HCC SMMC7721 cells were purchased from Cell Bank of Type Culture Collection of Chinese Academy of Sciences (Shanghai, China).

Gene knockdown

Small interfering RNAs (siRNA) against DNASE1L3 or RELB were synthesised by Santa Cruz Biotechnology, Inc. (TX, USA). SMMC7721 cells or LO2 hepatocytes were seeded into 6-well plates at a density of $2 \times 10^5$ cells/well and cultured overnight. The constructed siRNA-DNASE1L3, siRNA-RelB or siRNA-control was transfected into the cells using Lipofectamine 2000 (Invitrogen). To raise transfection efficiency, the cells were incubated with 20 nm siRNA for 6 h followed by a 24-h culture. In mouse HCC H22 cells, lentivirus-mediated shRNA delivery system was used to knock down DNASE1L3 gene. Briefly, H22 cells were seeded into 6-well plates at a density of $2 \times 10^5$ cells/well to culture overnight, and then, the medium containing the lentiviral particles (GeneChem Co., Ltd, Shanghai, China) was supplemented. After a 12-h infection, the viral medium was replaced with a fresh medium followed by a 72-h growth. Subsequently, the cells were selected with 1.5 $\mu$g/mL puromycin for 48 h. The interference efficiency was evaluated by Western blot assay.

Cell invasion assay

The cell invasion was observed with a transwell assay according to a previous report. Briefly, 200 $\mu$L of SMMC7721 or LO2 hepatocyte suspension was placed in the upper chambers coated with matrigel (1:7 with cell medium). Into the lower chambers, 800 $\mu$L of medium containing NETs or diabetic serum was added as a chemo-attractant. After a 48-h culture, the upper chambers were washed with $Ca^{2+}$-free PBS, fixed in methanol for 30 min and stained with 0.1% crystal violet for 30 min. The cells remaining on the membrane were carefully wiped off with cotton swabs. The cells that migrated through the membranes were counted and captured under a microscope in four random visual fields.

Western blot assays for protein expression

Western blot assays were performed to analyse the expression of proteins in the cultured cells or allografts. After the indicated treatments, the cells or grafts were collected and split at 4°C, and the protein concentration was quantitated with a BCA kit. Equal protein content in each sample was loaded on 10% SDS-PAGE gels. After the electrophoresis, proteins in the gels were transferred to PVDF membranes. The membranes were blocked with 5% fat-free milk in Tris-base buffered saline containing 0.1% Tween-20 for 1 h at room temperature and incubated with primary antibody against RelB (No. AF6380; Affinity Biosciences, OH, USA), DNASE1L3 (No. 36420; Signalway Antibody LLC, MD, USA), SPP1 (No. 25715-1-AP) or MMP9 (No. 10375-2-AP; ProteinTech, Chicago, IL, USA) with gentle agitation overnight at 4°C. After three washes, horseradish peroxidase-conjugated secondary antibodies were applied and incubated for 1.5 h. The signals were visualised with an enhanced chemiluminescence detection system. The band intensity was quantified with ImageJ software.

Enzyme-linked immunosorbent assay

SMMC7721 cells were treated with 0.25 $\mu$g mL$^{-1}$ NETs DNA for the indicated times. The content of intracellular cGAMP was measured with a commercial ELISA kit provided by AmyJet Scientific Inc. (Wuhan, China) and followed the manufacturer’s protocol. The value of optical density was probed with a microplate reader (Thermo) at 450 nm.

Immunofluorescence

SMMC7721 cells were exposed to 0.25 $\mu$g mL$^{-1}$ NETs DNA for 48 h. The cells were fixed in ice-cold methanol for 15 min and soaked in 0.3% Triton X-100 for 10 min. After three washes with PBS, the cells were blocked with 5% BSA in PBS for 1 h, incubated with primary antibody against STING (No. 19851-1-AP; ProteinTech, Chicago, IL, USA), or RelB (No. AF6380; Affinity Biosciences, OH, USA) overnight at 4°C. After washes, the corresponding horseradish peroxidase-conjugated secondary antibodies were applied and incubated for 1 h. The cell nuclei were counterstained with DAPI. The cells were viewed and captured under a confocal microscope (Leica, Wetzlar, Germany).

Statistical analysis

The experiment data are presented as mean ± standard deviation (SD). Significance between groups was evaluated by one-way analysis of variance followed by the Student–Newman–Keuls test using the GraphPad Prism 8 software (San Diego, CA, USA). A probability < 0.05 was considered to be statistically significant.
Acknowledgments
This work was supported by the Natural Science Foundation of Guangdong Province (2021A1515011365 and 2017A030313495), Guangzhou Education Bureau Key Medical Discipline Construction Project (201851839) and National Natural Science Foundation of China (81201919). We thank the patients and healthy donors for supporting the present work. We also thank the reviewers for their careful and valuable work.

Conflict of Interest
The authors declare no conflict of interest.

Author Contributions
Na Li: Investigation; Methodology; Validation. Xue Zheng: Data curation; Methodology; Visualization. Mianrong Chen: Data curation; Investigation. Li Huang: Funding acquisition; Writing – review & editing. Li Chen: Methodology; Visualization. Rui Huo: Investigation; Methodology. Xiaotong Li: Data curation; Investigation; Software. Mingwen Sun: Investigation; Methodology. Suiqing Mai: Data curation; Visualization. Zhuoyi Wu: Data curation; Software; Visualization. Hui Zhang: Data curation; Methodology. Jinbao Liu: Funding acquisition; Investigation; Writing – review & editing. Chun-tao Yang: Funding acquisition; Project administration; Supervision; Writing – review & editing.

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**Supporting Information**

Additional supporting information may be found online in the Supporting Information section at the end of the article.