Mitochondrial Dysfunction Due to Lack of Manganese Superoxide Dismutase Promotes Hepatocarcinogenesis

Anja Konzack,1 Mirza Jakupovic,2 Kateryna Kubaichuk,1 Agnes Görlach,3 Frank Dombrowski,4 Ilkka Miinalainen,5 Raija Sormunen,5 and Thomas Kietzmann1

Abstract

Aims: One of the cancer hallmarks is mitochondrial dysfunction associated with oxidative stress. Among the first line of defense against oxidative stress is the dismutation of superoxide radicals, which in the mitochondria is carried out by manganese superoxide dismutase (MnSOD). Accordingly, carcinogenesis would be associated with a dysregulation in MnSOD expression. However, the association studies available so far are conflicting, and no direct proof concerning the role of MnSOD as a tumor promoter or suppressor has been provided. Therefore, we investigated the role of MnSOD in carcinogenesis by studying the effect of MnSOD deficiency in cells and in the livers of mice.

Results: We found that loss of MnSOD in hepatoma cells contributed to their conversion toward a more malignant phenotype, affecting all cellular properties generally associated with metabolic transformation and tumorigenesis. In vivo, hepatocyte-specific MnSOD-deficient mice showed changed organ architecture, increased expression of tumor markers, and a faster response to carcinogenesis. Moreover, deficiency of MnSOD in both the in vitro and in vivo model reduced β-catenin and hypoxia-inducible factor-1α levels.

Innovation: The present study shows for the first time the important correlation between manganese superoxide dismutase presence and the regulation of two major pathways involved in carcinogenesis, the Wnt/β-catenin and hypoxia signaling pathway.

Conclusion: Our study points toward a tumor suppressive role of MnSOD in liver, where the Wnt/β-catenin and hypoxia pathway may be crucial elements.

Introduction

Mitochondrial dysfunction, one of the hallmarks of tumor cells, is associated with an imbalance of the antioxidant defense and an increased production of reactive oxygen species (ROS) resulting in oxidative stress. Among the first line of defense against oxidative stress is the dismutation of superoxide radicals to hydrogen peroxide and molecular oxygen by superoxide dismutases (SODs); in mitochondria, this task is carried out by manganese superoxide dismutase (MnSOD).

Accordingly, carcinogenesis may be expected to be associated with a dysregulation in the expression of antioxidant enzymes such as MnSOD. Indeed, first reports showed low levels or absence of MnSOD in many tumor types (3, 47), whereas forced expression of MnSOD could prevent the tumorigenic phenotype in melanoma, breast cancer, and glioma cells (8, 38, 57, 70). In addition, recent studies have reported that reduced MnSOD expression in primary tongue...
squamous cell carcinoma is associated with lymph node metastasis (40). In contrast, increased levels of MnSOD were found in progressive and metastatic breast, pancreatic, gastric, colorectal prostate, and lung carcinomas (12, 42), and findings from mouse experiments propose that MnSOD expression switches between early and advanced stages of skin cancer (13).

Interestingly, the direct role of MnSOD in hepatocellular carcinoma has not been widely addressed, although from all organs investigated so far, MnSOD activity was shown to be highest in the liver (32). Recent studies indicate that MnSOD promoter mutations may contribute to the carcinogenic process in hepatocarcinoma cells (65), and studies in mice suggested that lack of MnSOD may affect liver development (25, 36), likely without affecting the integrity of mitochondrial DNA (10). Together, these findings suggest that MnSOD might be linked to a specific type and stage of cancer, but they leave open whether a lack of MnSOD in hepatocytes would be linked to a carcinogenic process. Therefore, it was the purpose of the present study to investigate whether cellular deficiency of MnSOD can promote a carcinogenic process in the liver.

Thus, we studied the effect of MnSOD deficiency in a hepatocyte-derived cell line in vitro and in vivo in the liver of mice with hepatocyte-specific lack of MnSOD. Focusing on cellular properties generally associated with metabolic transformation and tumorigenesis, we found that absence of MnSOD in the cellular model contributed to their conversion to a malignant phenotype. In line, lack of MnSOD in mouse liver strongly promoted tumor formation in a chemically induced liver carcinogenesis model. Moreover, we found that expression of the Wnt/β-catenin pathway components as well as the hypoxia-inducible transcription factor (HIF-1α) was altered in both the cell as well as in the mouse model. Taken together, our study shows that lack of MnSOD exerts tumorigenic activity, which may involve the Wnt/β-catenin pathway as a crucial element.

Results

Loss of MnSOD results in mitochondrial dysfunction, formation of ROS, and changes in cell morphology and enhances cell proliferation

To investigate the role of MnSOD during hepatocarcinogenesis, we generated a MnSOD-knockdown (kd) cell line from human HepG2 hepatoma cells. The kd was confirmed by Western blot analysis; no MnSOD protein could be detected in the MnSOD-kd cells in contrast to the HepG2 scrambled (HepG2-sc) cells (Fig. 1A, B). As expected, MnSOD deficiency in MnSOD-kd cells resulted in a 40% increase in dihydroethidium (DHE) fluorescence, indicating an enhanced level of superoxide (Fig. 1C, D). In addition, a 2.3-fold increase in dichlorodihydrofluorescein (DCF) fluorescence in MnSOD-kd cells indicated an enhanced level of ROS.

FIG. 1. Knockdown of MnSOD induces ROS levels. (A) HepG2 cells were stably transfected with a scrambled (sc) shRNA or shRNA against MnSOD (kd). Protein lysates were prepared and analyzed by Western blotting with antibodies against MnSOD, Sod-1, Catalase, GPx-1, Prdx1, Prdx3, Trn2, Nrf-2, HO-1, and α-tubulin. Data are mean ± SD of fold induction normalized to HepG2-sc cells (n = 3, *p < 0.05) (B) Representative Western blots. α-tubulin is shown for equal loading of proteins. (C–E) The redox state of HepG2-sc and MnSOD-kd cells was determined by measuring (C) DHE fluorescence (n = 3), (D) DHE fluorescence microscopy showing mitochondrial localization of superoxide (n = 3), (E) DCF fluorescence (n = 10), and (F) GSH levels (n = 3) (c.f. section “Materials and Methods”). Data are mean ± SD of fold induction normalized to HepG2-sc (*p < 0.05). (G) Nrf2 localization analyzed by confocal fluorescence immunohistochemistry. (H) Representative Western blot showing Nrf2 localization after cell fractionation; Lamin A/C is shown to indicate nuclear fractions, and α-tubulin is shown for cytosolic fraction. DCF, dichlorodihydrofluorescein; DHE, dihydroethidium; Gpx-1, glutathione peroxidase-1; GSH, reduced glutathione; HO-1, heme oxygenase-1; kd, knockdown; MnSOD, manganese superoxide dismutase; Nrf2, NF-E2-related factor 2; Prdx, peroxiredoxin; ROS, reactive oxygen species; SD, standard deviation. To see this illustration in color, the reader is referred to the web version of this article at www.liebertpub.com/ars
total ROS (Fig. 1E). Furthermore, the reduced glutathione (GSH) level was decreased by about 45% in MnSOD-kd cells pointing toward a more oxidized state in these cells (Fig. 1F).

To examine whether lack of MnSOD may be compensated by enhanced expression of other antioxidative enzymes, Western blot analysis for Sod-1 (Cu/ZnSOD), catalase, glutathione peroxidase-1 (Gpx-1), thioredoxin-2, and peroxiredoxin (Prdx1) and Prdx3 protein levels was performed along with determination of GSH levels. We found that only Sod-1 showed a 25% compensatory increase in the absence of MnSOD, whereas catalase and Prdx3 levels were decreased by 30% and 20%, respectively. Other proteins involved in antioxidant reactions; among them, the NF-E2-related factor 2 (Nrf2) targets heme oxygenase-1 (HO-1), Prdx1, and the master regulator of antioxidant genes Nrf2 itself were not affected by MnSOD deficiency (Fig. 1A, B).

However, since Nrf2 changes its localization in response to redox changes, we performed cytosol/nuclear fractionation assays and confocal microscopy to assess Nrf2 differences with respect to cytosolic or nuclear presence. We detected Nrf2 almost exclusively in the cytosolic fraction of the wild-type cells, whereas Nrf2 was present in the nuclear fraction of the MnSOD-kd cells; In addition, confocal microscopy revealed that Nrf2 is primarily cytoplasmic in the control cells, whereas it is found much stronger in the nucleus of MnSOD-kd cells (Fig. 1G, H).

To assess whether loss of MnSOD changes mitochondrial morphology, we performed transmission electron microscopy. While the control cells displayed the intercellular networks, microvilli-like protrusions, and lipid droplets typical for hepatocytes, these features were lost in the MnSOD-kd cells. In addition, MnSOD deficiency altered mitochondrial structure and number; the mitochondria were smaller in size and contained fewer cristae, although MnSOD-kd cells appeared to contain a higher number of mitochondria compared to the control cells (Fig. 2A).

We next investigated whether these alterations in mitochondrial structure affect mitochondrial function and analyzed the mitochondrial integrity and membrane potential by labeling active mitochondria with the fluorescent dye tetramethylrhodamine ethyl ester (TMRE). In addition, we examined alterations in cellular respiration by high-resolution respirometry. Fluorescence microscopy of TMRE-labeled cells revealed that depletion of MnSOD increased fragmentation of the mitochondrial tubes, as indicated by shortened tube structures (Fig. 2B). In addition, MnSOD-deficient cells showed about 20% reduction in the mitochondrial membrane potential compared to the control cells (Fig. 2C).

**FIG. 2.** MnSOD knockdown alters mitochondrial function. (A) Transmission electron microscopy images of HepG2-sc and MnSOD-kd cells at 1200× and 4800× magnification. (B) Comparison of intracellular mitochondrial networks in HepG2-sc and MnSOD-kd cells visualized by TMRE (200 nM) staining and fluorescence microscopy. (C) Mitochondrial membrane potential (ΔΨm) was assessed by TMRE fluorescence using DNP as an uncoupler. Results are shown as percentage normalized to the HepG2-sc control and presented as mean±SD; *Significant difference sc versus kd, p<0.05. (D) Oxygen consumption of HepG2-sc and MnSOD-kd cells was measured by high-resolution respirometry. Measured were routine (R) respiration, proton leak (L) respiration after oligomycin-initiated inhibition of ATP synthase (5 μM oligomycin), ETC capacity (E) at FCCP-uncoupled respiration (5–7 μM FCCP), and nonmitochondrial respiration (rox) after inhibition of respiratory chain complexes I and III by rotenone (1 μM) and antimycin A (5 μM). Data were normalized to cell number, and the relative changes are presented. Abbreviations: R/E= ratio of routine and noncoupled respiration, L/E= ratio of oligomycin-inhibited and noncoupled respiration, rox/E'= rate of oxidative phosphorylation-independent oxygen consumption (*Significant difference p<0.05). DNP, 2,4-dinitrophenol; ETC, electron transport chain; FCCP, carbonyl cyanide-p-trifluoromethoxyphenylhydrazone; TMRE, tetramethylrhodamine, ethyl ester. To see this illustration in color, the reader is referred to the web version of this article at www.liebertpub.com/ars
Using high-resolution respirometry, we found that the noncoupled respiratory capacity of the electron transport system (ETS) was decreased and the proton leak after inhibition of ATP synthase with oligomycin was elevated due to MnSOD deficiency; however, the oxygen consumption during routine respiration was not affected by the loss of MnSOD. These changes in cellular respiration became very clear when comparing the flux control ratios of HepG2-sc and MnSOD-kd cells (Fig. 2D).

While control HepG2-sc cells utilized only about 50% of the ETS’s total respiratory capacity, MnSOD-depleted cells required about 65% for routine respiration. In addition, proton leakage after inhibition of ATP synthase with oligomycin indicated an increased potential of superoxide production (19%) for MnSOD-kd cells compared to the 12% of total respiratory capacity observed for HepG2-sc cells. Interestingly, after excluding the superoxide producing respiration, about 45% of the total ETS capacity was needed for ATP production in MnSOD-kd cells, whereas only 38% was used in control cells (Fig. 2D). Together, these results indicate a mitochondrial dysfunction, where the MnSOD-kd cells leak protons and operate significantly closer to the total respiratory capacity.

In addition to the changed mitochondrial morphology and function, we also detected changes in cell morphology upon staining of cells with crystal violet. While HepG2-sc cells were spherical in shape, their MnSOD-kd derivatives showed a fibroblast-like morphology with branches and an elongated shape. Furthermore, 4',6-diamidino-2-phenylindole (DAPI) staining revealed an increased degree of DNA condensation in the nuclei of MnSOD-kd cells compared to the scrambled control cells (HepG2-sc) (Fig. 3A).

Since increased ROS may affect proliferation and cell survival, we next determined whether kd of MnSOD alters proliferation. Therefore, bromodeoxyuridine (BrdU) incorporation and MTT assays were performed along with counting of cells; they show that loss of MnSOD caused an increase in proliferation by about 50% (Fig. 3B). Apoptotic as well as necrotic cell deaths, as assessed by flow cytometry using a combination of the DNA-binding dye propidium iodide (PI) and the phospholipid-binding protein annexin V, were not affected in MnSOD-kd cells (Fig. 3C, D). Together, these results indicate that lack of MnSOD promotes proliferation but not survival of hepatoma cells.

**FIG. 3.** Loss of MnSOD alters cell morphology and increases proliferation in HepG2 cells. (A) Representative images of HepG2-sc (sc) and MnSOD-kd (kd) cells stained with crystal violet and DAPI. Images were taken using a 60×/1.00 oil immersion objective in bright-field to visualize crystal violet staining and fluorescence at an appropriate filter set for DAPI. (B) The effect of MnSOD deficiency on the proliferative ability was measured by counting of cells, BrdU assay, and MTT assay. Data are presented as fold induction (mean ± SD) normalized to HepG2-sc cells (*p < 0.05). (C) FACS analysis of apoptotic and necrotic cells after double staining with Annexin-V-FLOUS (AV) and PI at a sampling rate of 1 ml/min and wavelengths of λ_em = 488 nm, λ_ex = 518 nm for AV and λ_em = 488 nm, λ_ex = 617 nm for PI. Representative plot displaying AV versus PI shows three clearly separated populations of cells: R2 = live (AV neg, PI neg), R3 = early apoptotic (AV pos, PI neg), and R4 = late apoptotic/necrotic (AV pos, PI pos). (D) Alive (h), early apoptotic (ea), and late apoptotic/necrotic (a/n) cells analyzed by FACS. Data are mean ± SD of the fold induction normalized to HepG2-sc cells. BrdU, bromodeoxyuridine; DAPI, 4',6-diamidino-2-phenylindole; PI, propidium iodide. To see this illustration in color, the reader is referred to the web version of this article at www.liebertpub.com/ars
MnSOD deficiency enhances malignant transformation of hepatoma cells

Malignant cell transformation is characterized by the ability of cells to overcome contact inhibition and to grow in a semisolid environment. Indeed, soft agar assays revealed that loss of MnSOD led to growth and formation of a few large colonies (>100 μm in diameter), which displayed about fivefold increase in colony volume, whereas HepG2-sc cells formed many small colonies (<50 μm in diameter) and stopped growing soon after plating (Fig. 4A, B). In addition, after 6 days in a monolayer colony formation assay, MnSOD-deficient cells covered a 40-fold larger area with loose widespread colonies compared to HepG2-sc cells that grew under the same conditions as dense overlaying colonies (Fig. 4A, B).

Next, migration and invasion of both the HepG2-sc and MnSOD-kd cells through a semipermeable membrane were studied in a transwell assay in response to serum starvation. Cell migration increased by about ninefold due to loss of MnSOD (Fig. 4C). Since tumor invasiveness is highly dependent on cell adhesion, the strength of cell attachment to the matrix of a culture plate was quantified in an adhesion assay. Properly attached cells were rotated for 2 h on an orbital shaker, while a control plate was kept motionless. MnSOD deficiency decreased cell adhesion significantly; about 60% detachment was observed for MnSOD-kd cells, whereas only 25% of HepG2-sc cells detached in response to motion (Fig. 4C, D). Together, these data underline that lack of MnSOD plays a role in malignant cell transformation.

Lack of MnSOD in hepatocytes contributes to liver failure and precancerosis in mice

To further investigate the role of MnSOD in liver tumorigenesis on an in vivo level, we created a hepatocyte-specific MnSOD-knockout (KO) mouse model. Although these mice appeared phenotypically normal at birth, they had a significantly decreased liver to body weight ratio (wild type [WT]: 1.90 ± 0.26 g, KO: 0.99 ± 0.10 g). The absence of MnSOD was confirmed on the level of protein expression. MnSOD protein was absent in the livers of these animals, whereas the levels in the brain and kidney were unchanged (Fig. 5A).

In line with the in vitro data, the loss of MnSOD in liver should cause an increase in ROS levels resulting in enhanced formation of 8-isoprostanes due to free radical...

FIG. 4. MnSOD depletion affects anchorage-independent and anchorage-dependent cell growth. (A) Anchorage-independent and anchorage-dependent growth of HepG2-sc and MnSOD-kd cells was assessed by colony formation and soft agar assay, respectively. Data are presented as sum of colony volume [Σ(V)] and sum of area [Σ(A)], respectively. (B) Representative images from soft agar and colony formation assay are shown. (C) Migration was determined by a transwell assay (8μm pore size) under serum starvation. Data are mean ± SD of the fold induction normalized to HepG2-sc cells. Representative images and quantification of migrated cells are shown. (D) Cell adhesion was assessed by rotating cells at 250 rpm on an orbital shaker for 2 h at 37°C, while keeping a control plate motionless. The number of adherent cells in the motionless control plates was set to 100%; data are mean ± SD (*Significant difference, p < 0.05). (E) Representative images of crystal violet stained cells still attached to the plate are shown. To see this illustration in color, the reader is referred to the web version of this article at www.liebertpub.com/ars
catalyzed peroxidation of arachidonic acid (Fig. 5B). To analyze this, we stained liver sections with antibodies against 8-isoprostane. Immunostaining for 8-isoprostane was absent in the livers of control mice, whereas hepatocytes from MnSOD-KO mice showed strong (40-fold increase) 8-isoprostane signal, present in almost all hepatocytes (Fig. 5C). In addition, the enhanced generation of $O_2^-$ due to lack of MnSOD may, via formation of peroxynitrite, lead to protein tyrosine nitration (Fig. 6A). Indeed, liver sections from MnSOD-KO mice showed about 20-fold increase in 3-nitrotyrosine staining, which was almost absent in the livers of control mice (Fig. 6B, C).

To determine the pathological changes caused by loss of MnSOD on a histological basis, hematoxylin–eosin-stained liver tissue sections were examined. Sections from control mice were morphologically normal and displayed no signs of liver damage or failure (Fig. 7A, B). In contrast, livers from MnSOD-KO mice showed neoductular proliferation in the portal triads as well as periportal and intra-acinar inflammatory infiltrates of lymphocytes, leucocytes, and macrophages. Many hepatocytes exhibited intracellular lipid droplets or contained more than two nuclei (multinuclear hepatocytes). Mitotic activity was significantly increased compared with the WT livers (Fig. 7A, B). In addition, the fibrosis index as measured by the Knodell score (33) was significantly increased.
Furthermore, evidence of liver failure due to hepatocyte injury was found on the biochemical level when the serum of MnSOD-KO and control mice was examined. The levels of albumin, a useful marker of the hepatic synthetic and secretory function, and cholinesterase, a common indicator for chronic and advanced liver disease, were markedly decreased in the MnSOD-KO mice (Fig. 7C, D). In contrast, the aminotransferases, aspartate aminotransferase and alanine aminotransferase, as well as the levels of alkaline phosphatase were increased in MnSOD-KO mice indicating hepatocellular damage. Moreover, bilirubin levels were not affected by loss of MnSOD (Fig. 7C, D).

To further examine whether the lack of MnSOD initiates neoplastic changes, we analyzed the expression of placental glutathione S-transferase (GST-P), a member of the GST family known as a reliable marker for hepatocarcinogenesis (52). We found that livers of control mice were completely devoid of GST-P. In contrast, MnSOD-KO mice displayed a large number of GST-P-positive cells (Fig. 8A, B), indicating structural changes within the liver acinar architecture. To further investigate this, we studied distribution of glutamine synthetase (GS), an enzyme known to be present in the most distal perivenous cells of the acinus, which thus represents a marker for acinar integrity (17). In control mice, GS was found only in the hepatocytes surrounding the central veins. In contrast, the MnSOD-KO mice displayed a scattered pattern of GS-positive cells without a clear zonal distribution, indicating that loss of MnSOD initiated structural changes within the liver acinus, which could over an extended time frame contribute to the formation of liver tumors (Fig. 8A, B). However, during the 3-month observation period, the animals remained free of tumors (Table 1).

**Loss of MnSOD in hepatocytes increases diethylnitrosamine-induced tumor formation in mice**

To determine whether MnSOD deficiency affects hepatocarcinogenesis, the formation of preneoplastic lesions and adenomas in livers of hepatocyte-specific MnSOD-deficient and control mice was studied at different time points after induction of hepatocellular carcinogenesis with diethylnitrosamine (DEN). The carcinogenic effect of DEN involves bioactivation by cytochrome P450 enzymes and formation of highly carcinogenic DNA adducts through nitrosamine alkylation (58). The MnSOD-deficient DEN-injected mice displayed an increased likelihood and earlier appearance of neoplastic lesions. The malignant phenotype included increased liver size, formation of invasive neoplasms, and increased metastases (58). The MnSOD-KO mice displayed an increased likelihood and earlier appearance of neoplastic lesions. The malignant phenotype included increased liver size, formation of invasive neoplasms, and increased metastases (58).

**FIG. 7. Lack of MnSOD in hepatocytes causes liver damage. (A) Livers from 3-month-old, male MnSOD<sup>flox/flox</sup> control mice (WT) and MnSOD-KO mice (KO) were prepared and analyzed for morphological changes. Cells showing mitotic signs, inflammatory infiltrates, and lipid droplets were counted using a ×400 magnification (0.15 mm<sup>2</sup> field), and the respective index (nuclei with visible chromosomes/total number of nuclei with observations at least 5000 counted nuclei) as well as the Knodell fibrosis score (33) compared to the WT animals, was determined. Data presented as mean ± SD (*p < 0.05). (B) Representative liver sections from wild-type MnSOD<sup>flox/flox</sup> (WT) and hepatocyte-specific MnSOD-KO mice stained with hematoxylin/eosin. The livers of hepatocyte-specific MnSOD-KO mice displayed signs of inflammation (inflammatory foci in the middle panel in KO). In addition, an enhanced number of mitotic cells (arrow in left KO) and lipid droplet containing cells (right KO) were present in these animals. (C) Blood was drawn from 3-month-old, male control MnSOD<sup>flox/flox</sup> mice (WT) and MnSOD-KO mice (KO); the individual samples of serum were analyzed for different parameters indicating liver injury. The respective levels or activities in the WT animals were set to 1 (*Significant difference p < 0.05). (D) Serum parameters in the WT and KO mice, values represent mean ± SD (*p < 0.05). Alb, albumin; ALP, alkaline phosphatase; ALT, alanine aminotransferase (formerly GPT, glutamate pyruvate aminotransferase); AST, aspartate aminotransferase (formerly GOT, glutamate oxaloacetate aminotransferase); Bili, bilirubin; Che, choline esterase. To see this illustration in color, the reader is referred to the web version of this article at www.liebertpub.com/ars
FIG. 8. Loss of MnSOD in hepatocytes promotes tumorigenesis. (A) Livers from 3-month-old, male MnSOD<sup>flox/flox</sup> control mice (WT) and MnSOD-KO mice (KO) were prepared and immunostained for GST-P and GS, and positive cells were counted using a 40× magnification. Data represented as positive cells per 100 hepatocytic nuclei; at least 5000 nuclei have been counted (n=6/group; *p<0.05). (B) Representative liver sections from wild-type MnSOD<sup>flox/flox</sup> (WT) and hepatocyte-specific MnSOD-KO mice immunostained with antibodies against GST-P or GS as early tumor markers. Dark brown precipitates indicate the GST- or GS-positive cells. 40× magnification. (C) Diethylnitrosamine induced liver tumors in mice. (i) Massive DEN-induced liver tumorigenesis in hepatocyte-specific MnSOD-KO mice 12 months after DEN injection (Table 1). (ii) Large areas of perivenular chronic inflammation in the livers of KO mice compared to normal morphology of WT animals 6 months after DEN treatment; 40×. (iii) Preneoplastic lesions in WT and KO mice 9 and 6 months after DEN injection, respectively; 200×. (iv) Oval cells in WT compared to liver adenoma in KO liver tissue 12 months after DEN treatment; 400×. Hematoxylin–Eosin staining. DEN, diethylnitrosamine; GS, glutamine synthetase; GST-P, placental glutathione S-transferase. To see this illustration in color, the reader is referred to the web version of this article at www.liebertpub.com/ars

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Number of preneoplastic lesions and adenoma (per cm<sup>2</sup> liver area) found in the livers of wild-type and MnSOD-deficient male mice at 3, 6, 9, and 12 months of age. Animals are grouped into DEN treated [DEN (+)] and untreated [DEN (−)]. Mean values±SDs are shown; the number of lesion presenting animals versus the number of examined mice in each group is given in brackets.

* Differences between WT and KO p < 0.05 were considered significant; ** p < 0.01.

DEN, diethylnitrosamine; KO, knockout; MnSOD, manganese superoxide dismutase; SD, standard deviation; WT, wild type.
hepatic preneoplastic lesions and adenomas (Table 1),
pointing toward a role for MnSOD as a tumor suppressor.

Signs of preneoplastic lesions in MnSOD-KO mice were
apparent already 3 months after injection of DEN, whereas in
control animals, the first preneoplastic lesions appeared af-
after 6 months. Hepatocellular adenomas appeared first in
MnSOD-KO mice 9 months after DEN treatment. Twelve
months after DEN injection, the livers of WT mice also
displayed adenomas; however, they were 10 times more
frequent in MnSOD-deficient animals. In this study, three out
of nine DEN-treated KO mice showed liver carcinogenesis in
its terminal stage, with five or more tumors of 2–5 mm di-
diameter (Fig. 8C[i]). In addition, a few preneoplastic lesions
and adenomas were found even in the livers of untreated
MnSOD-KO mice, but not in the untreated wild-type ani-
mals, further emphasizing the role of MnSOD as a tumor
suppressor.

In line, histological analyses of the hematoxylin–eosin-
stained liver tissues revealed large areas of perivenular in-
flammatory infiltrates in DEN-injected MnSOD-KO mice
after 6 months (Fig. 8C[ii]). At 12 months, all of the DEN-
treated KO mice displayed an advanced stage of liver ade-
noma. In contrast, the livers of DEN-treated wild-type ani-
mals were almost free of tumors, but contained large numbers
of oval cells, which are involved in liver regeneration after
injury (Fig. 8C[iiv]). Together, these data show that lack of
MnSOD in hepatocytes contributes to tumor formation in
the liver.

Loss of MnSOD in vitro and in vivo leads to inactivation of Wnt signaling and
to interruption of the hypoxic response

It is well known that the Wnt/β-catenin pathway is one of
the most important pathways associated with carcinogenesis
and malignancy (51). Moreover, loss of hepatic zonation and
pericentral tissue rearrangement has recently been associated
with improper Wnt signaling (4, 56). In light of this, we
determined whether MnSOD deficiency would affect ex-
pression and distribution of Wnt/β-catenin pathway compo-
nents using antibodies against β-catenin and the adenoma
polyposis coli (APC) protein. In line with previous findings
(5), we found that APC was highly expressed in hepatocytes
surrounding the central veins of wild-type mice, whereas this
perivenous zonation was almost absent in MnSOD-KO ani-
mals (Fig. 9A, B).

In line, APC protein levels of MnSOD-kd cells were sig-
nificantly decreased compared to HepG2-sc cells (Fig. 9C).
Since APC is responsible for directing the phosphorylation
and ubiquitination of β-catenin (67), the reduction in APC
due to loss of MnSOD would be expected to stabilize
β-catenin. In contrast, the levels of β-catenin were also found to

FIG. 9. Lack of MnSOD decreases APC, β-catenin, and
HIF-1α levels in vivo in mice and in vitro in HepG2 cells.
(A) Representative liver sections from 3-month-old, male
wild-type MnSODlox/lox (WT) and hepatocyte-specific
MnSOD-KO mice immunostained for APC, β-catenin, and
HIF-1α. Dark areas indicate high APC, β-catenin, and HIF-
1α expression. (B) Quantification of immunostained tissue
sections of 3-month-old animals (n = 4/group; *p < 0.05).
(C) HepG2 cells stably transfected with a scrambled (sc)
shRNA or shRNA against MnSOD (kd) were exposed to
normoxia 16% O2 or hypoxia 5% O2 for 4 h. Thereafter,
protein lysates were prepared and analyzed by Western
blotting with antibodies against APC, β-catenin, HIF-1α,
and α-tubulin, respectively. α-Tubulin is shown for equal
loading of proteins. Arrows indicate areas of changed ex-
pression. APC, adenoma polyposis coli; HIF-1α, hypoxia-
inducible transcription factor. To see this illustration in
color, the reader is referred to the web version of this article
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be reduced when MnSOD was knocked out in mice or knocked down in HepG2 cells (Fig. 9A–C).

Another important aspect during carcinogenesis is hypoxia signaling, where the HIF-1α plays a critical role (63). Since HIF levels can be influenced by ROS (21), we were interested to see whether lack of MnSOD may have an effect on HIF-1α. To do this, we first analyzed HIF-1α levels in the liver tissue of the wild-type and MnSOD-KO animals. We found that low levels of HIF-1α could be detected around the low oxygen containing central veins of WT animals. In contrast, HIF-1α was absent in the livers of MnSOD-deficient mice (Fig. 9A). Next, we exposed MnSOD-kd cells and HepG2-sc cells to normoxic and hypoxic conditions. We found that hypoxia mediated a strong upregulation of HIF-1α in the control cells, whereas hypoxia failed to induce HIF-1α levels in MnSOD-kd cells.

To further prove the interrelation between lack of MnSOD and the β-catenin and HIF-1α pathway, we transfected HepG2-sc and MnSOD-kd cells with a β-catenin responsive TopFlash luciferase (Luc) reporter construct as well as a HIF-responsive HRE-Luc construct in combination with shRNAs against β-catenin and HIF-1α, respectively. We found that, in line with the Western blot analyses, MnSOD-kd cells displayed reduced TopFlash and HRE-Luc activity similar to the ones of HepG2-sc cells cotransfected with the corresponding shRNA (Fig. 10A). In addition, we tried to rescue the reduced TopFlash and HRE-Luc activity levels by treating the transfected MnSOD-kd cells with the cell permeable MnSOD mimic Mn(III) tetrakis (4-benzoic acid) porphyrin chloride (MnTBAP). Both TopFlash and HRE-Luc expressions were induced upon treatment of MnSOD-kd cells with MnTBAP. This seemed to be a specific superoxide-driven event since treatment of cells with N-acetylcysteine (NAC) did not induce TopFlash and HRE-Luc activity (Fig. 10A), but treatment with MnTBAP was able to restore the GSH content of MnSOD-kd cells to the level of the control (Fig. 10B).

In addition, we found that kd of HIF-1α in HepG2-sc cells induced TopFlash activity, whereas kd of β-catenin did not affect HRE-Luc activity pointing to an interdependency of both pathways.

To estimate whether the downregulation of the Wnt/β-catenin or the HIF-1α signaling pathways may be involved in the transformation of HepG2-sc cells toward a malignant phenotype, HepG2-sc cells were transfected with shRNAs against β-catenin and HIF-1α and proliferation was measured.
by BrdU incorporation. Indeed, kd of β-catenin induced proliferation by about 20%, and kd of HIF-1α promoted proliferation of HepG2-sc cells by about 15%. Together, these data suggest that lack of MnSOD can have a promoting effect on hepatocarcinogenesis via affecting β-catenin and HIF-1α signaling.

Discussion

The present study showed that MnSOD deficiency promotes carcinogenesis in vitro in human hepatoma cells and in vivo in liver of mice. On the cellular level, loss of MnSOD caused an increased proliferation, reduced apoptosis, decreased contact inhibition and cell adhesion, as well as enhanced migration. This was in line with the in vivo findings showing that hepatocyte-specific loss of MnSOD in mice causes liver failure, 8-isoprostanate and 3-nitrotyrosine formation, as well as initiation of malignant transformation and tumor formation. In addition, MnSOD deficiency strongly promotes tumor formation in a model of chemically induced carcinogenesis. The mechanistic analyses of the in vivo and in vitro models indicate a loss of APC and β-catenin in the MnSOD-deficient cells and livers of MnSOD-deficient mice, indicating that MnSOD is a critical component necessary to sustain proper Wnt/β-catenin signaling.

Loss of MnSOD contributes to carcinogenesis

First evidence that MnSOD deficiency contributes to carcinogenesis and malignancy came from numerous reports showing a relative reduction of MnSOD in many solid tumors (54, 66). In addition, two MnSOD variants (23), showing either reduced mitochondrial translocation or enzyme activity (2, 55), were associated with a higher incidence of cancer (44, 62), which points to the role of MnSOD as a tumor suppressor, and of oxidative stress as a promoter of liver tumors.

Supporting the concept that loss of MnSOD leads to a transformed cell type is the observed change in morphology in MnSOD-kd HepG2 hepatoma cells toward a more fibroblast-like shape. Although the branched and elongated shape of MnSOD-deficient cells would lead to the assumption that they adhere stronger to the extracellular matrix, the data showed the opposite. Loss of MnSOD decreased adhesion and increased migration of cells. The potential of MnSOD to affect migration appears not to be restricted to liver-derived cells. Recent reports indicated that changed MnSOD levels affect also migration of certain cancer cell lines as well as of vascular smooth muscle cells in vitro and in vivo (9, 37, 61).

Changes toward a more malignant phenotype due to the loss of MnSOD were also observed in cell proliferation. Like in the present study, suppression of MnSOD by siRNA resulted in increased superoxide levels and stimulation of proliferation in ovarian cancer cells (24). Furthermore, overexpression of MnSOD has shown to inhibit cellular proliferation of numerous tumor cell types both in vitro and in vivo (31, 50, 61). These findings are in line with studies demonstrating changes in MnSOD during the cell cycle. MnSOD levels were decreased during the S phase (48), while increased MnSOD levels were observed in nonproliferating cells in the G0 phase (7).

Although the ROS generated due to MnSOD deficiency do not necessarily need to be carcinogenic, they can initiate or participate in reactions that contribute to transformation of cells and to a carcinogenic process. The concept that loss of MnSOD leads to a transformed cell type is supported by our findings from hepatocyte-specific MnSOD-KO mice displaying an enhanced number of GST-P- and GS-positive cells. Enhanced levels of GST isoforms as precancerous markers were also found in hepatic preneoplastic nodules (7), suggesting that hepatocyte-specific loss of MnSOD induces development of hepatocellular carcinomas. The appearance of GST-positive cells is in line with the role of ROS as tumor promoters since H2O2 was shown to induce GST in different cell types (1, 53). Moreover, GSTs also detoxify some of the carbonyl-, peroxide-, and epoxide-containing metabolites produced within xenobiotic metabolism, where ROS are produced as byproducts. Interestingly, similar neoplastic changes were also observed in livers of Cu/Zn-SOD−/− mice (14). Like our hepatocyte-specific MnSOD-KO mice, the nonconditional Cu/Zn-SOD−/− mice were born without a significant phenotype, but exhibited an increased incidence of nodular hyperplasia or hepatocellular carcinoma as well as neurodegeneration, and reduced fertility later in life (14). In addition, other antioxidant enzymes may contribute to the development of human hepatocellular carcinoma; for example, Gpx activity was found to be reduced in hepatocellular carcinoma patients compared to histologically normal livers (6).

Carcinogenesis is also accompanied by structural reorganization or remodeling of tissue. In the liver, various enzymes are distributed in a zonated manner, that is, they can be found in the periportal, intermediate, or perivenous zone (26, 27). In the healthy liver, GS is found only in distal perivenous hepatocytes (16, 17). However, our MnSOD-KO mice clearly showed GS-positive cells in other areas. GS-producing cells may possess a growth advantage (49), as well as a higher metastatic potential (19), which coincides with the phenotypic changes due to MnSOD deficiency in HepG2 cells. Since glutamine is an energy source of tumors, GS positivity indicates that these cells produce their own glutamine, independent from that supplied by the blood. Together, the lost GS zonation and the presence of GST-positive cells (Fig. 8A, B) appear to represent early hepatocellular carcinogenesis.

MnSOD deficiency in the livers of male humans and rodents has been associated with nonalcoholic fatty liver disease (NAFLD) (35). While NAFLD does not necessarily lead to liver cancer, it is a known cause of chronic liver disease, which possesses a risk for progression to cirrhosis and hepatic carcinogenesis (43). Although not enough epidemiological data are available to show whether loss of MnSOD is cause or consequence of NAFLD, our findings provide a link indicating that loss of MnSOD may be carcinogenic. In line with this, the additional increase in oxidative stress due to chemical induction of hepatocarcinogenesis with DEN resulted in an early-onset and increased incidence of hepatic preneoplastic lesions and liver adenomas in hepatocyte-specific MnSOD-deficient mice. This together with the presence of enhanced oxidative stress seen as 8-isoprostanate and 3-nitrotyrosine immunohistochemical staining (Figs. 6 and 7) indeed supports the role of oxidative stress as a tumor promoter in the liver.
The involvement of Wnt and hypoxia signaling in hepatocarcinogenesis

The Wnt/β-catenin pathway, as one of the most important pathways associated with cancer, has been specifically implicated with liver tumors. Mutations leading to accumulation of β-catenin were found in 17% of hepatocellular carcinomas (64). Therefore, we determined whether MnSOD deficiency affects expression and distribution of the Wnt pathway components β-catenin and APC. Surprisingly, both were decreased in MnSOD-deficient HepG2 hepatoma cells as well as in MnSOD absent mouse livers, indicating a dysregulation of β-catenin on expression rather than on stability. In line with our study, hepatocyte-specific β-catenin KO mice showed increased susceptibility to DEN-induced carcinogenesis (69). Paradoxically, hepatocyte-specific disruption of the APC gene in mice also resulted in activation of Wnt/β-catenin signaling and hepatocarcinogenesis (69), indicating that any imbalance in Wnt signaling may promote tumor development in the liver.

Since hypoxia plays an important role in tumor progression and the ability of cells to survive under low oxygen conditions is a key feature of malignant tumors, it was one of the aspects of this work to investigate HIF-1α protein levels. Indeed, induction of HIF-1α was disrupted both in vitro and in vivo due to the loss of MnSOD. These findings are partially in line with previous observations from MnSOD overexpressing human breast carcinoma MCF-7 cells, where hypoxia-dependent induction of HIF-1α was lost when MnSOD activity was moderately increased but restored when under highly increased MnSOD activity. Furthermore, overexpression of H2O2 metabolizing enzymes in these high MnSOD-expressing cells prevented HIF-1α induction under hypoxia. Thus, it was believed that the increase in HIF-1α is mediated by mitochondrial hydrogen peroxide (63).

Overall, these observations as well as the data presented here may follow the same regulatory mechanism of HIF-1α degradation. VHL-dependent degradation of HIF-1α is known to require the presence of Fe^{2+} as a cofactor for the prolyl hydroxylases (PHDs), which hydroxylate HIF-1α at conserved proline residues, mediating proteasomal degradation (63). While Fe^{2+} in general causes reduction of iron, H2O2 oxidizes it to Fe^{3+}. Thus, it is tempting to speculate that kd of MnSOD and accumulation of O2_2− would increase PHD activity and therefore HIF-1α degradation, whereas expression of MnSOD and accumulation of H2O2 would decrease PHD activity and prevent HIFα degradation.

Although our data do not unravel all mechanistic aspects of the carcinogenic process in response to MnSOD deficiency, they indicate that the interplay between HIF-1α and β-catenin contributes to this effect. The β-catenin/HIF-1α interplay appears to be complex, partially conflicting, and is far from being completely understood. Initial reports indicated that β-catenin is able to interact with HIF-1α to mediate hypoxic adaptation (28), which would imply that HIF signaling may negatively affect β-catenin signaling under emerging hypoxic conditions to allow immediate adaptation of a tumor. Indeed, several reports indicate that hypoxia and HIF-1α can block and even downregulate β-catenin protein levels (11, 39, 59, 60). These findings would be in line with our data indicating that kd of HIF-1α increases β-catenin-dependent TopFlash activity.

However, other reports showed that hif-1α deletion can also impair Wnt-dependent processes, including neuronal stem cell proliferation, differentiation, and neuronal maturation (41). As a consequence, loss of the β-catenin-mediated differentiating effect promotes a dedifferentiated state of cells, which in terms of cancer represents a more malignant status than a well-differentiated one. This is in line with our findings showing loss of β-catenin due to lack of MnSOD as well as with a study from 97 hepatocellular carcinoma patients where 76% of the patients displayed a reduced expression of β-catenin correlating with poor survival (68).

Although hypoxia usually correlates with a malignant behavior of cancer cells, it has been recently shown that HIF-1α is also responsible for the induction of cell cycle inhibitors such as p21 and p27 (20). Thus, lack of HIF-1α and β-catenin in MnSOD cells would promote progression into the S phase and at the same time inhibit cell differentiation with the consequence to have a procarcinogenic effect.

Although not enough epidemiological data are available to show whether loss of MnSOD is a general carcinogenic phenomenon, our study shows that hepatocyte-specific loss of MnSOD contributes to the induction of hepatotoxicity and liver cancer, which appear to involve the Wnt/β-catenin pathway and HIF-1α signaling, as well as their interplay, as crucial elements.

Materials and Methods

All biochemicals and enzymes used were of analytical grade and were obtained from commercial suppliers.

Cells

Stable transfectants of human HepG2 hepatoma cells were established using a MnSOD miRNA-expressing vector and a respective scrambled control vector. Both cell lines, a mixed population of MnSOD-kd and HepG2-sc, were cultured under normoxia (16% O2, 5% CO2, and 79% N2 \([v/v]\)) or when indicated under hypoxia (5% O2, 5% CO2, and 90% N2 \([v/v]\)) at 37°C in minimal essential medium (MEM) supplemented with 10% fetal bovine serum (FBS), 1% nonessential amino acids, and 0.5% ciprofloxacin hydrochloride.

ROS detection

The total ROS content of cells was assessed using 2',7'-dichlorodihydrofluorescein diacetate (H2DCF-DA), whereas DHE was used to measure superoxide levels. Cells (10^4 cells/well) cultured on 96-well plates were washed with 1× phosphate-buffered saline (PBS) and incubated with 10 μM H2DCF-DA (Molecular Probes) or 10 μM DHE (Sigma) in 1× PBS for 15 min at 37°C. After replacement of the reagents with 1× PBS, DCF formation was recorded at λex/λem: 495/529 nm and DHE fluorescence at λex/λem: 530/610 nm using a Tecan infinite M1000Pro microplate reader (Tecan). Values were corrected for H2DCF-DA and DHE auto-oxidation in 1× PBS. For live cell imaging of DHE, fluorescence cells were plated onto 35 μm μ-dishes (Ibidi) and treated with 20 μM DHE (Sigma) for 15 min, and after washing with PBS, images of cells were taken using a confocal microscope LSM700 (Zeiss) at the appropriate filter set for DHE.
GSH assay

GSH levels were measured using o-phthaldehyde (OPT) as a fluorescent agent (22). HepG2-sc and MnSOD-kd cells were washed with 1× PBS, resuspended at 10^6 cells/ml in a lysis buffer (5% trichloroacetic acid, 1 mM ethylene-diaminetetraacetic acid [EDTA], 0.1 M HCl, 1:1:1, v/v/v), and incubated on a roller for 20 min at 4°C. Following lysis, insoluble material was spun down at 5600 g for 20 min at 4°C. After mixing 18 parts of 0.1 M Na3HPO4/5 mM EDTA (pH 8.0), one part of OPT (0.1 mg/ml in methanol), and one part of sample supernatant, fluorescence was measured at λex/λem: 350/420 nm using a Tecan infinite M1000Pro microplate reader (Tecan).

Transmission electron microscopy

Cells were fixed in 1% glutaraldehyde and 4% formaldehyde mixture in 0.1 M phosphate buffer for 10 min. Cells were detached, and fixation was continued for 1 h. After fixation, cells were centrifuged, immersed in 2% agarose in distilled water, postfixed in 1% osmium tetroxide, dehydrated in acetone, and embedded in Epon LX 112 (Ladd Research Industries). Thin sections were cut with a Leica Ultracut UCT ultramicrotome, stained in uranyl acetate and lead citrate, and examined in a Tecnai G2 Spirit transmission electron microscope (FEI Europe). Images were captured by using a Que mesa CCD camera (Olympus Soft Imaging Solutions GMBH).

TMRE mitochondrial staining

TMRE is a cell membrane permeant, positively-charged fluorescent dye that accumulates in the active mitochondria. Treatment with carbonyl cyanide-p-trifluoromethoxyphenylhydrazone (FCCP) or 2,4-dinitrophenol (DNP) uncouples the ETS from the oxidative phosphorylation, thereby decreasing the proton gradient of mitochondria. For live cell imaging of TMRE, fluorescence cells were plated onto 35-μm μ-dishes (ibidi) and treated with 200 nM TMRE (Sigma) for 30 min. Cells treated with 20 μM FCCP (Sigma) 10 min before TMRE staining were included as a depolarization control. After washing with PBS, images of cells were taken using an Observer Z1 live cell analyzing microscope (Zeiss) at the appropriate filter set for TMRE (λex/λem: 549 nm/575 nm). The mitochondrial membrane potential was determined after treating cells (10^4 cells/well) on 96-well plates with 1 μM TMRE for 30 min. Treatment with 2 mM DNP (Sigma) for 10 min before TMRE staining served as a control for depolarized mitochondria. Controls for fluorescent TMRE autofluorescence were included. After washing with 0.2% bovine serum albumin (BSA) in PBS, the fluorescence was measured using a Tecan infinite M1000Pro microplate reader (λex/λem: 549 nm/575 nm).

Oxygen consumption

Oxygen consumption measurements were performed using an Oxygraph-2k (Oroboros). Before each measurement, a background calibration of MEM supplemented with HEPES (10 mM) was conducted under contact to the gas phase (air). Cells (~10^6 cells/ml) in MEM-HEPES were measured in parallel at 37°C and under stirring at 750 rpm and oxygen exclusion. The oxygen flux was allowed to stabilize till the aerobic metabolic activity under routine conditions (ROU-TINE respiration) was reached. The proton leakage caused by ATP synthase inhibition (LEAK respiration) was determined by 1–2 μl injections of oligomycin (5 mM). Then, the maximal capacity of the ETS (respiration) was measured by sequential injections of FCCP (1 mM) until no further increase in O2 flow could be achieved. Finally, single injections of rotenone (1 mM) and antimycin A (5 mM) were implemented to completely inhibit complex I and III for the determination of oxidative phosphorylation-independent oxygen consumption (rox). The rox flux was subtracted from the other respiratory states for data analysis.

DAPI and crystal violet staining

Cells were grown on glass coverslips within six-well plates at low confluence. After washing, cells were fixed with 4% paraformaldehyde and stained with 0.0075% crystal violet. Excess stain was removed, and glass coverslips were placed with a drop of Mounting Medium containing DAPI (Duolink II) onto a microscope slide. In addition, single stain controls were prepared. Images were taken at the Olympus BX51 microscope using an NPL (normal field of view plan) 60×/1.00 oil immersion objective.

Cell proliferation

Determination of cell number. Cells (4×10^5) were seeded in 100-mm culture dishes, and cell number was determined by counting with a hemocytometer after 5 days.

BrdU assay. Incorporation of BrdU into newly synthesized DNA was detected by immunolabeling according to the manufacturer’s (Merck Millipore) instructions using 2×10^4 cells per well in a 96-well culture plate.

MTT assay. Cell viability was assessed by measuring conversion of the yellow (3-[4,5-dimethylthiazol-2-yl]-2,5-diphenyl tetrazolium bromide) into dark blue water-insoluble formazan crystals (46). The extent of MTT reduction was quantified by spectrophotometry at 570 nm.

Flow cytometry analysis

Fluorescence double staining with Annexin-V-FLOUS and PI was performed according to the instructions provided with the Annexin-V-FLUOS Staining Kit (Roche). Approximately, 10^6 cells were washed with 1× PBS before being resuspended in 100 μl Annexin-V-FLUOS labeling solution and incubated for 20 min at room temperature. Two sets of controls were included: single stain controls for electronic compensation of the plot and necrosis-positive controls using 0.01% Nonidet P40. Data were collected using a CyFlow Space flow cytometer (Partec) at a sampling rate of 1 ml/min. Excitation wavelength was 488 nm, and emission wavelength was 518 nm for Annexin-V-FLUOS and 617 nm for PI.

Colony formation

Anchorage-independent growth was analyzed in a three-layer soft agar assay, containing a basal agar layer followed by a cell and a top agar layer. The agar layers were prepared by mixing prewarmed (37°C) 2× MEM (containing 20% FBS, 2% nonessential amino acids, and 1% ciprofloxacin) with an equal volume of agarose (DNA grade). For the basal
and top layer, 1.2% agar was used, while only 0.8% agar was used for the cell containing layer with 2000 cells/well. A fifty microliter suspension was used for each layer per well of a 96-well plate. The plates were incubated at 37°C for 14 days before the colonies were visualized with resazurin (50 μl/well, 0.2 μg/ml) and colonies were counted and measured for quantification. Anchorage-dependent growth was assessed in a monolayer colony formation assay by seeding cells at different densities (500–3000 cells/well in steps of 500) in six-well plates and growing them for 6 days. Afterward, cells were fixed and stained with crystal violet and analyzed as described above.

Migration assay

Migration of cells was assessed in vitro in a transwell assay, where the cells migrate through an 8 μm polycarbonate membrane of 24-well transwell chambers (Becton Dickinson Labware) in the course of serum starvation. Cells were seeded into the top chamber at a density of 10^5 cells/well in 100 μl of serum-free MEM; the bottom chamber was filled with 600 μl MEM containing 10% FBS. After 16 h in a humidified incubator, cells were fixed and stained with crystal violet as described above. The nonmigrated cells were removed from the upper surface of the Transwell membrane with a cotton swab.

Cell adhesion

Cells (4 × 10^5) were seeded onto 24-well culture plates, and after attachment, one plate was placed in an orbital shaker and rotated for 2 h at 250 rpm and 37°C, while a control plate was kept motionless. Adherent cells were stained with crystal violet as described above as well as quantified by the MTT assay described above.

Western analysis

Western blot analysis was carried out as described (30). In brief, 100 μg of protein was loaded onto sodium dodecyl sulfate–polyacrylamide gels and after electrophoresis blotted onto a nitrocellulose membrane. The primary antibodies against MnSOD (SOD2) (#574596; Calbiochem), SOD1 (#574597; Calbiochem), Nrf-2 (#ab62352; abcam), HO-1 (#SPA-895; assay designs), Catalase (#8841; Cell Signaling), GPx1 (#3206; Cell Signaling), Prdx1 (#8499; Cell Signaling), Prdx3 (#sc-59661; Santa Cruz), Trx2 (#14907; Cell Signaling), HIF-1α (#610959; D Biosciences), APC (#sc-896; Santa Cruz), Lamin A/C (#sc-7292; Santa Cruz), and β-catenin (#sc-7963; Santa Cruz) were used in a 1:1000 dilution. The secondary antibody was either an anti-sheep, anti-mouse, or anti-rabbit immunoglobulin G horseradish peroxidase (1:5000; Santa Cruz Biotechnology). The enhanced chemiluminescence system (Amersham) was used for detection.

Plasmid constructs, cell transfection, and luciferase assay

The β-catenin responsive Topflash Firefly luciferase (Luc) construct (45), the HIF-responsive HRE-Luc (29) construct, and the expression vectors for shHIF-1α (15), as well as for shβ-catenin (34), have been previously described. Cells (4 × 10^5) per 60-mm dish were transfected essentially as described (30). In brief, 2 μg of the appropriate β-catenin-responsive Topflash Firefly Luc or HIF-responsive HRE-Luc construct was cotransfected in duplicate and, to control transfection efficiency, with 0.25 μg of a Renilla luciferase expression construct (pRLSV40) (Promega). After 12 h, the medium was changed and the cells were further cultured for 24 h, where indicated also in the presence of MnTBAP (100 μM) or NAC (1 mM). The detection of luciferase activity was performed with the Dual-Glo Luciferase Assay Kit (Promega).

Animals

The hepatocyte-specific MnSOD-KO mice (MnSOD^−/−, KO) used in this study were described (30). Control mice (MnSOD^+/−/−) are referred to as WT. Animals were housed in specific pathogen-free facilities on a 12-h light/12-h dark cycle, at a constant temperature of 22°C, and were fed a standard diet and received water ad libitum. Hepatocellular carcinogenesis in these mice was induced by a single intraperitoneal injection of DEN (20 μg/g body weight) 2 weeks after birth. A total of 122 animals were included in this experiment—62 wild-type and 60 MnSOD-KO mice, of which 32 and 30 received DEN injections, respectively (Table 1). Three, six, nine, and twelve months following the DEN treatment, animals were euthanized by CO2 inhalation and livers were examined. All protocols for animal use and experiments were approved by the Animal Care Committee of local institutions (30).

Histology and immunohistochemistry

Mouse livers fixed in 10% neutral buffered formalin were embedded in paraffin, and 5 μm sections were stained with hematoxylin and eosin according to standard techniques. Inflammatory infiltrates and cells undergoing mitosis were counted and the respective index was estimated (observations per 100 hepatocytes). For immunohistochemical staining, 5 μm paraffin sections were prepared by deparaffinization and rehydration with xylene and a graded ethanol series, respectively. Tissue sections were boiled in 0.1 M citric acid, pH 6.0, endogenous peroxidase activity was quenched with 3% H2O2 in methanol and nonspecific binding blocked with 5% BSA in T-PBS. The primary antibodies and the secondary peroxidase-conjugated antibodies were applied in 1% BSA in PBS overnight at 4°C or 60 min at room temperature, respectively. Antibodies against 8-isoprostane (#IS20; Oxford Biochemical Research), 3-nitrotyrosine (#N0409; Sigma), APC (#sc-896; Santa Cruz), Lamin A/C (#sc-7292; Santa Cruz), and β-catenin (#sc-7963; Santa Cruz) were used in a 1:1000 dilution. The secondary antibody was either an anti-sheep, anti-mouse, or anti-rabbit immunoglobulin G horseradish peroxidase (1:5000; Santa Cruz Biotechnology). The enhanced chemiluminescence system (Amersham) was used for detection.

Statistical analysis

Densitometry data were plotted as the fold induction of relative density units, with the zero value absorbance in each figure set arbitrarily to 1 or 100%. Statistical evaluation was conducted by the paired, two-tailed Student’s t-test, and differences of p < 0.05 were considered statistically significant. The results are presented as mean ± standard deviation.
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Author Disclosure Statement
No competing financial interests exist.

References
18. This reference has been deleted.


63. Wang M, Kirk JS, Venkataraman S, Domann FE, Zhang HJ, Schafer FQ, Flanagan SW, Weydert CJ, Spitz DR, Buettner GR, and Oberley LW. Manganese superoxide dismutase suppresses hypoxic induction of hypoxia-


Address correspondence to:
Prof. Thomas Kietzmann
Faculty of Biochemistry and Molecular Medicine, and Biocenter Oulu
University of Oulu
Aapistie 7B
Oulu 90230
Finland

E-mail: tkietzm@gwdg.de

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Abbreviations Used

Alb = albumin
ALP = alkaline phosphatase
ALT = alanine aminotransferase (formerly GPT, glutamate pyruvate aminotransferase)
APC = adenoma polyposis coli
AST = aspartate aminotransferase (formerly GOT, glutamate oxaloacetate aminotransferase)
Bil = bilirubin
BrdU = bromodeoxyuridine
BSA = bovine serum albumin
Che = choline esterase
DAPI = 4′,6-diamidino-2-phenylindole
DCF = dichlorodihydrofluorescein
d DEN = diethylnitrosamine
DHE = dihydroethidium
DNP = 2,4-dinitrophenol
EDTA = ethylenediaminetetraacetic acid
ETS = electron transport system
FBS = fetal bovine serum
FCCP = carbonyl cyanide-p-trifluoromethoxyphenyl-hydrazone
Gpx-1 = glutathione peroxidase-1
GS = glutamine synthetase
GSH = reduced glutathione
GST-P = placental glutathione S-transferase
H2DCF-DA = 2′,7′-dichlorodihydrofluorescein diacetate
HIF-1α = hypoxia-inducible transcription factor
HO-1 = heme oxygenase-1
kd = knockdown
KO = knockout
MnSOD = manganese superoxide dismutase
NAC = N-acetylcysteine
NAFLD = nonalcoholic fatty liver disease
Nrf2 = NF-E2-related factor 2
OPT = o-phthalaldehyde
PBS = phosphate-buffered saline
PHD = propyl hydroxylase
PI = propidium iodide
Prdx = peroxiredoxin
ROS = reactive oxygen species
SOD = superoxide dismutase
TMRE = tetramethylrhodamine, ethyl ester
WT = wild type