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Gene activation of CEBPA using saRNA: Preclinical studies of the first in human saRNA drug candidate for liver cancer

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Abstract

Liver diseases is a growing epidemic worldwide. If unresolved, liver fibrosis develops and can lead to cirrhosis and clinical decompensation. Around 5% of cirrhotic liver diseased patients develop

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hepatocellular carcinoma (HCC), which in its advanced stages has limited therapeutic options and negative survival outcomes.

CEPBA is a master regulator of hepatic function where its expression is known to be suppressed in many forms of liver disease including HCC. Injection of MTL-CEBPA, a small activating RNA oligonucleotide therapy (CEBPA-51) formulated in liposomal nanoparticles (NOV340-SMARTICLES®) upregulates hepatic CEBPA expression. Here we show how MTL-CEBPA therapy promotes disease reversal in rodent models of cirrhosis, fibrosis, hepatosteatosis and significantly reduces tumour burden in cirrhotic HCC.

Restoration of liver function markers were observed in a carbon-tetrachloride induced rat model of fibrosis following 2-weeks of MTL-CEBPA therapy. At 14-weeks animals showed reduction in ascites and enhanced survival rates. MTL-CEBPA reversed changes associated with hepatosteatosis in non-alcoholic methionine and cholic deficient diet induced steaotic liver disease. In diethylnitrosamine induced cirrhotic HCC rats, MTL-CEBPA treatment led to a significant reduction in tumour burden.

The data included here and the rapid adoption of MTL-CEBPA into a Phase 1 study may lead to new therapeutic oligonucleotides for undruggable diseases.

Keywords

HCC; Cirrhosis; NASH; Liver fibrosis; CEBPA; small activating RNA; Oligonucleotides Therapy

Introduction

Liver disease is caused by a multitude of components including obesity, alcoholism and viral hepatitis. These synergistically culminate in an asymptomatic 'compensated cirrhosis' stage where the average life expectancy of patients is 10 to 13 years. This is usually followed by a rapidly progressive phase of liver dysfunction termed 'decompensated cirrhosis' where the average life expectancy is reduced to as low as 2 years¹.

In the United States, there were 658,000 hospitalization associated with cirrhosis in 2011, which culminated in \$9.8 billion dollars in cost². Despite recent medical advances, liver cirrhosis remains a poorly treated stage of liver disease and in 2013 claimed 1.2 million lives globally. Liver cirrhosis is now the 8th leading cause of death in the US and the 13th leading cause of death globally, with a 46% increase in worldwide mortality over the past two decades¹.

Progression of cirrhosis can be accelerated by the development of other co-morbidities such as renal impairment, sepsis, hepatitis viral load and development of hepatocellular carcinoma³. According to long-term outcome studies, the severity of liver fibrosis is the most important predictor for patient outcome⁴.

Unresolved fibrosis ultimately leads to a decompensated liver where the only treatment at this stage is liver transplantation. The scarcity of organ donors, coupled with the increasing rates of incidents worldwide against a disease intractable to conventional therapeutic

compounds, poses a major threat for patient survival with chronic liver disease. Patients who progress to HCC end up with a severely compromised liver due to the natural progression of the disease which decreases their opportunity to meet eligibility for curative surgery, transarterial chemoembolization (TACE) or sorafenib treatment.

An improvement of liver function would be expected to benefit the general condition and Child Pugh status of patients; their quality of life and more significantly allow them to reenter eligibility criteria for surgical interventions, which would ultimately enhance chances of survival.

A treatment that reverses fibrosis, improves liver function whilst inhibiting tumour growth would be a major therapeutic breakthrough and here we present data for a novel RNA based drug candidate in the form of small activating RNAs (saRNAs), which is currently in a phase I clinical trial for patients with advanced cirrhotic HCC (ClinicalTrials.gov – NCT0271601).

saRNAs differ from other RNA-based therapeutics in that they up-regulate, rather than inhibit the expression of their therapeutic target⁵. The mechanism of action involves loading of saRNA into Ago2 protein, translocation into the nucleus, binding to the genomic target locus and promoting increased transcriptional activity by RNA Polymerase 2⁶.

Here we report an saRNA therapy that was designed to increase production of CEBPA (CCAAT/enhancer-binding protein alpha), a master transcriptional factor in the liver, which resets the natural gene regulatory mechanism of hepatocytes to reduce fibrosis and reverse liver dysfunction.

The idea of increasing C/EBP-α expression in diseased liver is long established. Downregulation of CEBPA is well known across many different liver disease models including fibrosis and cirrhosis^{7,8}. Furthermore the importance of CEBPA activity in normal liver function is observed in 'old age' related liver disease models, where hyperphosphorylation of CEBPA at serine residue 193 diminished transcriptional activity⁹. In transgenic CEBPA knock-out models, animals appear to develop liver tumours at an increased rate^{10,11}.

In order to validate the therapeutic benefit of increasing CEBPA expression in liver disease, we developed a therapeutic oligonucleotide (CEBPA-saRNA) for intravenous delivery using nanoparticles with high tropsim to the liver¹². The successful outcome of this study led us to further develop CEBPA-saRNA into a GMP grade manufactured clinical candidate (CEBPA51) with early indications to show high target specificity and reliance on Argonaute 2 and components of the RNA interacting transcriptional activating complex¹³. This therapeutically viable saRNA has been encased in liposomal nanoparticles (SMARTICLES [®])¹⁴ for the first in-human trial to study the outcome of targeting CEBPA activation in patients with HCC (clinicaltrials.gov - NCT0271601)

Here we report the preclinical data obtained from treating various liver diseased models with the new therapeutic candidate MTL-CEBPA. We show how restoration of CEBPA expression in the liver dramatically reverses markers of diseases in a fibrotic and steatotic model, whilst improving liver function tests and ultimately increasing survival rates in long-

term CCl_4 treated rats. The data included here and the rapid adoption of MTL-CEBPA into a Phase 1 study may lead to new therapeutic oligonucleotides for undruggable diseases.

Results

In vitro activation of C/EBP-a by CEBPA-51

The ability of CEBPA-51 to increase CEBPA mRNA expression *in vitro* was demonstrated in murine (AML12); rat (Clone 9) liver cell line, human (HEPG2) liver cells and primary human hepatocytes, (Fig. 1A). In each case, transfection of the newly GMP developed CEBPA-51 saRNA significantly increased CEBPA mRNA levels by about 2-fold while the non-specific control RNA oligonucleotide (siFLUC) was inactive. In human primary hepatocytes, transfection of CEBPA-51 showed increased expression of CEBPA mRNA and significant changes to expression of downstream targets of CEBPA. These included key liver function factors including albumin, HNF4A and CYP3A4 (Fig. 1B). Increased expression of these factors were also measured at the protein level by Western blotting (Fig. 1C).

In vivo delivery of MTL-CEBPA reduces tumour burden in DEN induced cirrhotic HCC

The newly GMP manufactured oligonucleotide formulated in SMARTICLES®; MTL-CEBPA was tested in the same DEN induced cirrhotic HCC rat model as previously reported with CEBPA-saRNA¹⁵. This was compared to treatment with an unspecific oligonucleotide also formulated in SMARTICLES® (NOV340/FLUC) (Fig. 2A). Briefly, Diethylnitrosamine (DEN) was administered for 9 weeks in male Wistar rats followed by a 2 week wash out period. At week 11, animals were randomized for each experimental arm. Tumour bearing animals were treated with 3 doses of either NOV340/FLUC control or MTL-CEBPA at days 1, 3 and 5. Animals were sacrificed one week later to assess the impact of MTL-CEBPA treatment on liver CEBPA levels and size of the tumour nodules.

Compared to NOV340/FLUC control, animals treated with MTL-CEBPA showed a significant increase in CEBPA mRNA expression in the liver (Fig. 2A). Although there was no significant increase in albumin transcript levels at day 12 across all the treated groups, we observed an increasing trend in the MTL-CEBPA treatment group. Animals treated with NOV340/FLUC control, after 12 days, showed larger sized tumour nodules at time of termination $(2,204 \pm 1,031 \text{ mm}^3)$, when compared to the nodule size in the MTL-CEBPA group $(401 \pm 181 \text{ mm}^3 \text{ (p=0.107 vs NOV340/FLUC control)}$ (Fig. 2B). Based on the mean tumour weight, this represented an 80% reduction in tumour size. In addition to this dramatic reduction in tumour size, liver function parameters for AST, ALT, bilirubin and ammonia levels all decreased between 30-50% over this short course of MTL-CEBPA treatment. The measured ammonia decrease was statistically significant (p<0.02) given the small group size and variability in tumour burden (Fig.2. C).

Activity of MTL-CEBPA (the CEBPA51 clinical candidate) improves liver parameters in carbon–tetrachloride (CCl₄) model of liver failure

Having recapitulated the same effects of CEBPA-saRNA as previously published, our new clinical candidate MTL-CEBPA was then investigated across different models of liver diseases. MTL-CEBPA was injected into a liver failure rat model exposed to the hepatotoxin

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carbon-tetrachloride (CCl₄). Sprague Dawley (SD) rats were treated with CCl₄ for 8 weeks and then randomized into four treatment groups (n=9 in each group). This comprised of MTL-CEBPA at three doses of 0.3mg/kg; 1.0 mg/kg and 3.0 mg/kg bi-weekly for 2 weeks co-administered alongside constant exposure to CCl₄. As control treatment, NOV340/FLUC was used at 3.0 mg/kg. To ascertain disease progression, a CCL₄ -only exposed group was maintained as ('Path control') and compared with normal animals ('Sham control') treated with olive oil.

Treatment with MTL-CEBPA showed an increase liver expression of CEBPAmRNA and protein levels (Fig. 3A). Increased expression of CEBPA culminated in a significant reduction in fibrosis as measured by hydroxyproline levels, when compared to the pathological control group ('Path control') while the NOV340/FLUC control had no impact (Fig. 3B). The reduction in hydroxyproline following MTL-CEBPA treatment was dose dependent. Liver tissues from animals in the MTL-CEBPA groups showed decreased histological fibrotic lesions when compared to the control group as shown here at 1.0mg/kg of MTL-CEBPA (Fig 3C and D).

Liver function tests demonstrated that MTL-CEBPA treatment reversed several disease parameters including ALT, AST and bilirubin despite continued exposure to CCl₄ (Fig. 3E). Ammonia levels significantly reduced after 2 weeks of treatment with MTL-CEBPA at all dose levels after just one week (Fig. 3F). Albumin, and total protein changes were minor in this model but appeared to be stabilized by MTL-CEBPA treatment (Fig. 3F). Whilst examining liver pathology at the end of the study, it became apparent that MTL-CEBPA drastically reduced lipid accumulation as observed by a significant decrease in red-oil-O staining and liver triglycerides shown here at 1.0mg/kg of MTL-CEBPA (Fig. 3G). Based on this initial study, the next strategy was to investigate the survival impact of MTL-CEBPA treatment in advanced cirrhotic animals where oligonucleotide treatment would be extended for 14 weeks of bi-weekly injectionS. In addition, based on our observation on reduction in liver triglycerides, the effect of MTL-CEBPA would also be studied on fatty liver disease in a non alcoholic fatty liver disease model.

Treatment of MTL-CEBPA in advanced stages of liver disease prolongs survival and improves liver function parameters

The long-term benefit of MTL-CEBPA on liver function and survival was assessed in animals chronically exposed to CCl_4 for 36 weeks. Animals were divided into two groups (n=9 in each group) comprising of a CCl_4 treatment control only and a $CCl_4 + MTL-CEBPA$ treatment group. Oligonucleotide treatment commenced after animals were exposed to CCl_4 for 8 weeks where MTL-CEBPA was maintained at biweekly injections for 14 weeks. After the last injection of MTL-CEBPA, animals were continually exposed to CCl_4 for a further 14 weeks until cessation of the study.

Upon treatment withdrawal, at week 22, fewer than half of the animals in the untreated control group remained alive. There was a significant 78% increased survival benefit (p<0.001 – log rank test) in the MTL-CEBPA treated group (Fig. 4A). Although all animals developed ascites; those in the control group rapidly escalated to grade 3 (visibly marked distension of the abdomen); all animals in the MTL-CEBPA treatment group displayed

ascites scores that were limited to Grade 1/2 (with only mild to moderate symmetrical distension of the abdomen) (Fig. 4B).

Albumin and total protein levels remained elevated in the MTL-CEBPA treated groups, when compared to untreated control (Fig. 4C and D). All animals treated with MTL-CEBPA also demonstrated reduction in liver damage markers including ALT, AST and ammonia (Fig. 4E-G). MTL-CEBPA treated animals also displayed an improvement in obstructive liver function as indicated by reduced accumulation of total bilirubin, GGT and ALP, (Fig. 4H-J). All MTL-CEBPA treated animals displayed normal body weight parameters and more efficient blood coagulation profile (prothrombin time), when compared to the untreated control group (Fig. 4K-L). Collectively, the evidence here suggests that the hepatoprotective effects of MTL-CEBPA against CCl₄-induced liver toxicity lasted for at least 6 to 8 weeks following withdrawal of treatment.

Activity of MTL-CEBPA in a MCD diet model of NASH

Having observed significant reduction in liver triglyceride levels from the CCl₄ liver failure model and when over-representation analysis of protein lysates from CEBPA51 transfected hepatocellular carcinoma lines (HepG2; Hep3B and PLCPRF5) were analysed by mass spectroscopy; significant protein changes in pathways that regulated lipid metabolism were observed (Supplementary Fig. 1). We therefore investigated the short term beneficial outcome of MTL-CEBPA in non alcoholic fatty liver disease. Here the well established methionine and choline deficient diet (MCD) induced non alcoholic steatohepatitis model was used. C57/BL6 mice were fed an MCD diet for 6 weeks to induce pathology. Liver sections confirmed distinct disruption to liver architecture with punctate steatotic lesions observed by week 4, (Fig. 5A). These represented accumulation of simple fat inclusions in the liver cells. MTL-CEBPA treatment commenced at week 6 where NASH showed the most severe histopathological progression (Fig. 5A). MCD fed animals were divided into 5 groups (n=10 in each) to follow increasing doses of MTL-CEBPA treatment (bi-weekly for 2 weeks) at 0.3, 1.0 and 3.0 mg/kg. This was compared to untreated MCD diet group and a NOV340/FLUC control treatment group. Treatment with MTL-CEBPA after two weeks caused a significant reduction in ALT and AST to near normal levels at all dose levels tested (Fig. 5B). Although bilirubin levels did not decrease significantly, they showed a trend towards reduction at higher doses of MTL-CEBPA treatment (Fig. 5B). When we investigated H&E and Masson's trichrome stained tissue sections of the liver, the MCD fed group treated with MTL-CEBPA at 0.3mg/kg; 1.0mg/kg and 3.0mg/kg showed major improvement in liver architecture at all concentrations (Fig. 5C). When the presence of lipid deposits was assessed by Oil red O staining, we observed a drastic reduction in positive staining only in the MTL-CEBPA treated groups to near normal levels when compared to the control group (Fig. 5D). This observation was also confirmed by quantifying liver triglyceride levels (Fig. 5E). This was similar to the CCl_4 induced liver disease model where we also observed improved clearance of fat deposits in MTL-CEBPA treated animals, (Fig. 3G).

Since infiltration of macrophages and activation of myofibroblasts within the liver are triggered during liver damage, we also measured for the presence of both by staining FFPE

fixed liver tissue sections with anti-α-smooth muscle actin and anti-F4/F80 (Fig. 5G and H, respectively). Animals fed with MCD diet showed a significant increase in F4/80 positive macrophage staining, when compared to normal diet control, and treatment at all doses of MTL-CEBPA significantly decreased the percentage of positively stained macrophages.

Discussion

The new findings presented here show development of a novel treatment for reversing liver fibrosis and liver steatosis through activation of CEBPA, a key liver transcription factor. Loss of CEBPA expression is a common feature in many forms of liver disease. This essential liver enriched transcription factor is responsible for regulating the normal function and differentiation of hepatocytes and this is illustrated in many transgenic knock-out animal models that report development of severe liver injury; liver steatosis and hyperammonemia in the absence of CEBPA⁹. In a more clinically relevant setting, published reports demonstrate that suppressed CEBPA mRNA levels is followed by a reduction in CEBPA target genes in tumour vs adjacent normal tissues of HCC patients¹⁶. Lower levels of CEBPA mRNA, moreover, correlated with shorter survival in HCC patients¹⁷. Paradoxically, there are also reports suggesting that higher CEBPA mRNA are observed in tumour vs adjacent normal tissue sections¹⁸. It must however be noted that C/EBP-a has 2 main isoforms (a p42 active and a p30 inactive form) that can be post-transcriptionally regulated¹⁹. Therefore it is possible that these observed discrepancies are due to the detection of inactive forms of C/EBP-a and in this regard the down-regulation of C/EBP-a target genes is a convincing reflection of the biological consequences in suppression of functional C/EBP-a.

Gain of function studies, where CEBPA expression is exogenously enhanced, show significant reduction in steatosis and reversal of liver damage²⁰. Gene therapy approaches for increasing C/EBP-a levels using either hydronamic liver delivery or adenoviral systems show reduction in liver fibrosis after CCl₄ treatment^{7,8}; whilst CEBPA overexpression via adenovirus eliminated orthotopic HCC xenograft nodules¹¹. The same observations are made where CEBPA overexpression using recombinant adenoviral cassettes reduces tumourogenicity of HCC tumour cell lines²¹. Collectively these previously published reports show data that is consistent with the in vivo data we have presented here using synthetic saRNA oligonucleotides in a nanoparticle formuation compatible for clinical use. Other studies have reported successful use of SMARTICLES® to deliver oligonucleotides (microRNA mimetic and DNAi) to achieve anti-tumour activity both preclinically and in the clinic^{14,22}, thus further validating use of SMARTICLES® as an attractive delivery vehicle to the liver. Liver activation of CEBPA through MTL-CEBPA therapy significantly improved liver function and enchanced survival in a CCL₄ induced advanced liver cirrhotic model. MTL-CEBPA drastically reversed steatosis in a diet induced NASH model and reduced tumour burden in a DEN induced cirrhotic HCC model. This was consistent with our earlier publication using CEBPA-saRNA delivered using PAMAM dendrimer nanoparticles¹².

After treatment with MTL-CEBPA, we confirmed up-regulation of CEBPA mRNA in the liver of both the DEN and CCl₄ treated models, which was also linked to up-regulation of albumin; a downstream target gene of CEBPA. Up-regulation of albumin by MTL-CEBPA

confirms directly that elevation of CEBPA mRNA leads to a functional co-activation in at least one of its target genes. In this report, the control oligonucleotide sequence encapsulated in the SMARTICLES® (NOV340/siFLUC) showed no anti-fibrotic, anti-steatotic or anti-tumour activities. NOV340/siFLUC furthermore showed no improvement in liver function consistent with the activity in the models being driven by the specific CEBPA-saRNA sequence and subsequent target up-regulation of CEBPA. This data also supports evidence of good liver delivery of MTL-CEBPA using SMARTICLES® although we have not examined the cellular distribution of MTL-CEBPA in these studies to fully define the liver cell populations impacted by the drug. Using histology we observed a reduction in macrophages and a decrease in stellate cell activity in the fibrosis models.

The mechanism of CEBPAsaRNA is still being unraveled. Currently, we show evidence of strand specificity and reliance on Argonaute 2^{13} . We also demonstrate it requires at least one component of the RNA interacting activating complex⁶. The exciting pre-clinical data reported here led us to progress MTL-CEBPA into a detailed one-month toxicology study in rats and cynomolgus monkeys that will be reported in detail elsewhere. The extended dosing in the CCl₄ model and the maintenance of activity for several weeks post cessation of dosing suggest a less frequent dosing schedule might be efficacious, but this will require additional studies in pre-clinical models. As far as we are aware, existing gene therapy approaches to modulate C/EBP- α (with their inherent complexity and safety challenges) have not yet progressed to the clinic.

MTL-CEBPA has successfully progressed into a Phase 1 dose escalation study in patients with primary HCC and liver metastases from other cancers (ClinicalTrials.gov – NCT0271601).

We believe that the unique combination of effects MTL-CEBPA has in enhancing liver function and preventing liver failure is suggestive of a potential breakthrough drug for chronic liver disease. We propose oligonucleotide therapy as a potentially novel and safe alternative to activate tumour suppressor genes.

Materials and Mehtods

Small activating RNA Oligonucleotides were synthesized with the following sequence

CEBPA-51

Sense strand: mGmCGmGUCAUUmGUCAmCUGGUCmUmU;

Anti-sense strand: GACCAGUGACAAUGACCGCmUmU.

siFLUC: The control oligonucleotide siFLUC targets firefly luciferase and has no complementarity in mammalian genome,

Sense strand: mCmUmUAmCGmCmUGAGmUAmCmUmUmCGAdTpsdT;

Anti-sense strand: UCGAAGmUACUmCAGCGmUAAGdTpsdT.

Oligonucletodies were synthesized by St Pharm at 99% HPLC purity. m=2'-O-Methyl and ps= Phosphorothioate are the strand modifications to the duplex RNA. All sequences are shown 5' to 3'.

Cell line studies

AML12—The AML12 (alpha mouse liver 12) cell line was established from hepatocytes from a mouse (CD1 strain, line MT42) transgenic for human TGF alpha and purchased from ATCC. Cells were cultured in a 1:1 mixture of Dulbecco's modified Eagle's medium and Ham's F12 medium with 0.005 mg/ml insulin, 0.005 mg/ml transferrin, 5 ng/ml selenium, 40 ng/ml dexamethasone and 10% fetal bovine serum (FBS).

Clone 9—Clone 9 (K-9) is an epithelial cell line isolated in 1968 from normal liver taken from a young male rat and was purchased from ATCC. Cells were cultured in F-12K Medium supplemented with 10% FBS.

HepG2—HepG2 cells are immortalized human liver carcinoma cells derived from the liver tissue of a 15 year old Caucasian male with a well differentiated hepatocellular carcinoma. Cells are culture in RPMI1640 medium supplemented with 10% FBS.

Primary human hepatocytes

Primary human hepatocytes (HMCPTS, Life technologies) were sourced from Donor ID: HU8200-A, a male African-American 52-year-old. Same batch numbered cells were purchased frozen and thawed following the manufacturer's protocol:

Primary hepatocyte thawing medium—Cryopreserved Hepatocyte Recovery Medium (CHRM) was used for thawing each vial (Life Technologies, CM7000)

Primary hepatocyte plating medium—Fetal bovine serum, heat inactivated 50ml (Life Technologies, 16140-071); Insulin-Transferrin-Selenium (100x)-5ml (Life Technologies, 41400-045); HEPES (1M)-5ml (Life Technologies, 15630-056); L-Glutamine-Penicillin-Streptomycin solution-5ml (Sigma, G1146); Dexamethasone-40ng/ml final concentration (Sigma, D8893); Phenol red free William's E Medium (Life Technologies, A12176-01).

Primary Hepatocyte Maintenance medium—Primary Hepatocyte Maintenance Supplement (Life Technologies, CM4000); Human Hepatocyte Growth Factor-40ng/ml final concentration (Sigma, H5791); Epidermal Growth Factor-20ng/ml final concentration (Sigma, E9644), Nicotinamide-2.5ug/ml final **concentration** (Sigma N0636); Phenol red free William's E Medium-500ml (Life Technologies, A12176-01).

Transfection studies (established cell lines)

For analyzing gene activation and protein expression, hepatocytes were seeded into 24-well plates at a density of 1×10^5 cells per well. Transfection was performed with lipofectamine 2000. CEBPA-saRNAs (CEBPA51) or scrambled saRNAs (siFLUC), were added to the cells at a final concentration of 20 nM, following the manufacturer's instructions

(Lifetechnologies, Cat number 11668019). The treatment was repeated 24 hours later and the cells were harvested at the 72-hour time point.

Transfection studies (primary human hepatocytes)

Cells were transfected at 1×10^5 cells per well in a 24-well collagen coated dish at a final oligonucleotide concentration of 50 nM with 3 µL of HiPerFect transfection reagent (Qiagen, 301704). The cells were incubated in plating media for 5 hours to allow monolayer formation before replacing with maintenance media. For the second (forward) transfection, the same conditions were used as for reverse transfection. Maintenance media was used for the remaining duration of the experiment.

Transfection studies (primary human hepatocytes)

Cells were transfected at 1×10^5 cells per well in a 24-well collagen coated dish at a final oligonucleotide concentration of 50 nM with 3 µL of HiPerFect transfection reagent (Qiagen, 301704). The cells were incubated in plating media for 5 hours to allow monolayer formation before replacing with maintenance media. For the second (forward) transfection, the same conditions were used as for reverse transfection. Maintenance media was used for the remaining duration of the experiment.

RNA extraction & Real-time PCR

Total RNA was extracted for reverse transcription (QuantiFast[®] Reverse transcription, Qiagen) and target cDNA amplification by real-time PCR (QuantiFast[®] SYBR[®]Green Master mix). The cDNA probes used we purchased as prevalidated QuantiTect[®] SYBR Probes from Qiagen. Real-time PCR was performed with validated QuantiTect[®] SYBR probe Qiagen or validated FAM probes from Applied Biosystems.

Western blot

Liver sections from three representative rats from the control and MTLCEBPA treated groups were processed for total protein extraction using Biosciences Tissue-PE-LBTM Buffer following manufacturer's instruction. 10μ g of protein were separated using SDS-PAGE. Samples were transferred onto PVDF membranes using a semi-dry transfer system (Biorad) for Western blotting using anti-CEBPA (Abcam: ab40764) and Lamin B (Santa-cruz: sc6216).

Statistical analysis

Statistical analysis was performed using one-way or two-way analysis of variance (ANOVA), followed by Dunnett's/Tukey's multiple comparison test wherever applicable. Kruskal Wallis test followed by Dunn's test was used for histological scores.

The following in vivo studies are included in the supplement

In vivo study 1: Effects of administering the clinical candidate MTL-CEBPA in Diethylnitrosamine (DEN) induced hepatocellular carcinoma (HCC).

In vivo study 2 and 3: Effects of administering the clinical candidate MTL-CEBPA following induction of chronic liver failure by carbon-tetrachloride (CCl4) – comparing short-term MTLCEBPA treatment and long-term MTL-CEBPA treatment.

In vivo study 4: The effects of administering the clinical candidate MTL-CEBPA following induction of NASH by methionine and choline deficient diet

Haematoxylin-eosin staining procedure (Mayer's H&E protocol)

Liver sections were processed and stained with H&E by following the standard procedures. Slides were deparaffinized and rehydrated by treatment through in series of 100% alcohol, 95% alcohol and 70% alcohol. Slides were washed with distilled water. To improve stain quality, formalin fixed tissue slides were re- fixed in Bouin's solution for 1 hour at 50°C. Slides were rinsed with tap water for 5 - 10 min to remove yellow color. Slides were stained with Weigert's iron hematoxylin working solution for 10 min. After staining, slides were rinsed with tap water for 10 min and washed with distilled water. After washing, slides were then stained with Biebrich scarlet-acid fuchsin solution for 10 - 15 min and washed with distilled water. Slides were differentiated with phosphomolybdic-phosphotungstic acid solution for 10 - 15 min or until collagen was not red. Slides were then transferred directly (without rinsing) to aniline blue solution for 2-5 min and washed with distilled water. Slides were quickly dehydrated through 95% ethyl alcohol, absolute ethyl alcohol and cleared with xylene. Finally, slides were mounted with resinous mounting medium.

Oil Red O Staining

Oil red O is a fat-soluble dye that can be used to stain neutral triglycerides and lipids on frozen sections. Frozen livers were cryomatrix (thermo scientific) fixed. Cryosections of 4-5 μ m in thickness were mounted on the glass slides and maintained at -20°C, until staining. These sections were air-dried and then fixed with formaldehyde for 5-10min. Then slides were placed in absolute propylene glycol for 2 - 5min and stained with 0.5% Oil red O stain. After staining, slides were rinsed with tap water for 10 min and washed with distilled water. The slides were then counterstained with Mayer's Hematoxylin for 1h, washed and mounted with glycerin. In this staining protocol, lipids/fat globules are stained bright red, while nuclei are stained blue.

F4/80 and a-SMA immunohistochemistry

Paraformaldehyde (PFA) fixed liver tissue sections were stained for F4/F80 (Thermo Scientific: PA5-MA1-91124 at 1:1000) and α smooth muscle actin (Thermo Scientific: PA5-19465 at 1:1000) to assess liver macrophages (Kupffer cells) and myofibroblast formation. Staining was performed following the standard procedure of antigen retrieval. Staining was quantified using Leica Qwin Software. From each slide 20x images were taken and the percentage area staining from each image was quantified from all the animals.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

Acknowledgments

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Abbreviations

ALT	Alanin transaminase
AST	Aspartate transaminase
CCL4	Carbon tetrachloride
СЕВРА	C/CAAT enhancer binding protein alpha
DEN	Diethylnitrosamine
НСС	Hepatocellular carcinoma
MCD	Methionine and choline deficient diet
MTL/CEBPA	CEBPA saRNA formulated in NOV340 SMARTICLES®
NOV340/FLUC	siRNA sequence to Luciferase as control formulated in NOV340 SMARTICLES [®]
NASH	Non alcoholic steato hepatitis
saRNA	small activating RNA

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Figure 1. Transcriptional acivity of CEBPA51 in vitro

(A) CEBPA-51 saRNA transfection increases CEBPAmRNA in murine AML12, Rattus (Clone 9), Human (HEPG2) and human (Primary hepatocytes). Key factors for liver function were screened in CEBPA-51 transfected primary human hepatocytes. These included (**B**) albumin, HNF4A (Hepatocyte nuclear factor 4-alpha) and CYP3A4 (Cytochrome P450 3A4). Data is expressed as mean \pm SEM. ¥=p(<0.00027) and *=p(0.0033). (**C**) A Western blot panel probed with antibody specific to HNF4A (Abcam; ab41898), CYP3A4 (Abcam; ab155029), Albumin (Abcam; ab106582) and CEBPA (CELL Signalling; 8178) showed

visible increase in protein expression in cells transfected with CEBPA51. Actin expression (Abcam: ab8226) was used as protein loading control across each of the lanes.

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Figure 2. MTL-CEBPA increases liver expression of CEBPA in DEN induced HCC Wistar rats (**A**) MTL-CEBPA showed a significant increase in CEBPAmRNA: **, p<0.001. Albumin showed a tedency to increase after MTL-CEBPA treatment, however this was not statistically significant (p=0.1769). **Tumour nodule size and liver function tests were assessed in the DEN induced cirrhotic HCC animals. (B**) Liver tumour nodule size was measured from the biopsies across the pre-treatment control and two treatment groups. The average tumour size in the DEN only group (pre-treatment) was 193 (\pm 63) mm³. Over the 12-day course of the study period, the tumour size of animals treated with NOV340/FLUC at time of termination increased to 2204 (\pm 1031) mm³. All animals treated with MTL-CEBPA showed a slower rate of increase with an average volume of 401 (\pm 181)mm³ (p=0.107 vs. NOV340/FLUC control). (**C**) Liver function parameters were measured for AST, ALT, albumin and bilirubin. All showed a 30-50% decrease over the short study period. Ammonia decreased significantly despite the small group size. *, p<0.02).



Figure 3. The effect of MTL-CEBPA treatment in CCl4 induced liver damage

(A) MTL-CEBPA administration in rats caused an increase in liver CEBPAmRNA expression and protein. Total RNA and protein were extracted from the liver of treated animals. Protein levels were assessed using Image J to perform band densitometry measurements. (B) MTL-CEBPA caused a dose dependent decrease in hydroxyproline at week 10 when compared to NOV340/Fluc control. Normal hydroxyproline levels were observed in the sham control group at week 10 compared to the pathological ('Path control') group at week 8. Data is expressed as mean \pm SEM vs. NOV340/FLUC. $\neq =p(<0.0002)$. (C)

Liver sections at week 11 ('Path control') and week 13 ('Sham control') were stained with H&E, Sirius red and Masson's trichrome. (Magnification – 20 X, Sham control, NOV304/ FLUC and MTL-CEBPA treatment groups). Sham control animals revealed normal architecture with hepatocytes arranged in the form of cords (light blue arrow) radiating away from the central vein (dark blue arrow). Normal distribution of sinusoidal spaces (yellow arrow) was observed. In the NOV340/FLUC treatment groups fibrous bands (green arrow) extended from one portal area to the adjacent or to the central vein. In the MTL-CEBPA treatment group, thinner fibrous bands (green arrow) were observed, when compared to NOV340/FLUC group. (D) The severity of hepatic necrosis was assessed. MTL-CEBPA treatment caused a significant reduction in Knodell score (activity index). ***p<0.001 compared to Nov340/FLUC. Statistical Analysis, one-way ANOVA followed by Tukey's multiple comparison test. The effect of MTL-CEBPA on liver function tests were assessed. (E) ALT, AST and bilirubin were measured and showed improvement at all doses of MTL-CEBPA. (F) Ammonia levels reversed despite continued exposure to CCl₄. Although minor changes in levels of albumin and total protein (globulin) were observed; this appeared to be stabilized over the course of treatment. Data is expressed as mean \pm SEM vs. NOV340/FLUC. *= p(0.0033); ¥=p(<0.00027); S=p(0.0006). The effect of MTL-CEBPA on fat deposit in the liver. (G) Cryosections of the liver were processed for Oil red O staining. Sham control animals showed absence of positively stained Oil red O positive fat globules. In the NOV340/FLUC group, there was an abundant number of Oil red O positive fat globules. This decreased in the MTL-CEBPA treated groups. Liver triglyceride levels were measured at week 13 to show a significant reduction in the MTL-CEBPA treated group. ###p<0.01 and \$\$\$p<0.001 compared to NOV340/siFLUC. One-way ANOVA followed by Tukey's multiple comparisons test.



Figure 4. The long-term effects of MTL-CEBPA on SD rats exposed to CCl₄ for 36 weeks (**A**). MTL-CEBPA treatment for 14 weeks caused a significant benefit to survival. (**B**) All animals had developed ascites which escalated to grade 3 in the PBS control group whereas in the MTL-CEBPA group this was limited to grade 1-2. Ascites scoring = 1: Mild ascites only detected by palpation; 2: moderate ascites causing moderate symmetrical distension of the abdomen and 3: large ascites cause visibly extended abdominal distension. **Liver function tests on blood collected from SD rats exposed to long term CCL₄ induced toxicity. (C**) Albumin and (**D**) total protein (globulin) remained raised when compared to the

untreated control group. (E) ALT, (F) AST, (G) ammonia, (H) total bilirubin, (I) GGT and (J) ALP, (K) Body weight and (L) prothrombin clotting time all collectively showed improvement in MTL-CEBPA treatment animals. All values represent mean ± SEM.

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(A) Histological analysis of liver sections from normal diet and MCD diet fed animals for (A, upper panels) 4 weeks and (A, lower panels) 6 weeks stained with H&E and Masson's trichome (magnification: 20x). (A, Left panel) Analysis of liver sections from normal diet control animals showed normal architecture with hepatocytes arranged in the form of cords (blue arrow) radiating away from the central vein. The portal trip (red arrow) contained an artery, vein and the bile duct. Normal distribution of the sinusoidal spaces (yellow arrow) was observed. (A, Right panel) Analysis of the liver sections from MCD diet animals

revealed steatosis (green arrow) and inflammatory cell infiltration (black arrow). (B) Serum levels of ALT and AST indicate reduction to near normal levels following two weeks of MTL-CEBPA treatment. Bilirubin levels do no continue to increase in the presence of 1 and 3mg/kg of MTL-CEBPA treatment. ¥=p(0.0002); *=p(0.002); \$=p(0.003). Histological analysis of liver sections from MCD diet fed animals treated with NOV340/FLUC (3mg/kg) and MTL-CEBPA at 0.3mg/kg; 1.0 mg/kg and 3mg/kg. (C) Liver sections stained with H&E and Masson's trichrome. (magnification: 20x). Analysis of liver sections from MCD + NOV340/FLUC revealed steatosis (green arrow) and inflammatory cell infiltration (black arrow) which was more prominent, when compared to MCD diet at week 6. The liver sections from all MTL-CEBPA treated groups showed reduced steatosis and inflammatory changes, when compared to MCD diet animals at week 6. The effect of MTL-CEBPA51 on fat deposits in the liver. (D) The liver cryosections from animals administered with all doses of MTL-CEBPA were stained with Oil red O. Staining pattern (**D**, left panel) at magnification: 20x showed less staining intensity in all animals treated with MTL-CEBPA. The scoring of Oil red O staining as percentage of the section area stained (**D**, **right panel**) showed significant reduction in staining from the treated groups. = p(<0.0002). (E) The effect of MTL-CEBPA treatment on liver triglyceride (TG) in MCD diet mice was also measured to consolidate the significant reduction seen from the Red Oil O staining. Data expressed as mean ±SEM when compared to NOV340/FLUC. ¥=p(<0.0002). The effect of MTL-CEBPA on myofibrolast and macrophage and formation. (F, left panel) Alpha SMA (a-SMA) staining of liver sections comparing animals fed on normal and MCD diet at week 6 and the effects of MTL-CEBPA treatment. (magnification: 20x). The liver sections from animals on normal diet revealed no sign of α -SMA staining. Animals fed on MCD diet showed prominent staining (green arrow), which was significantly reduced in MTL-CEBPA treatment at all concentrations. Scoring of the area as a percentage value (F, right panel) shows significant reduction in staining from animals treated with MTL-CEBPA. (G, left panel) F4/80 staining analysis of liver sections comparing animals on normal vs. MCD diet at week 6 and the effects of MTL-CEBPA treatment at increasing doses (magnification: 20x). The liver section revealed minimal F4/80 staining in the normal diet. More punctate staining was observed in the animals on MCD diet (black arrow). Scoring of the stained area (G, right panel) showed significant reduction in F4/80 at all doses of MTL-CEBPA groups compared with control. Data is expressed as mean \pm SEM. *=p(0.0026); **=p(0.0012); \$=p(0.0006).