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Lucidone Suppresses Hepatitis C Virus Replication by Nrf2-Mediated Heme Oxygenase-1 Induction

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Upon screening of plant-derived natural products against hepatitis C virus (HCV) in the replicon system, we demonstrate that lucidone, a phytochemical, isolated from the fruits of Lindera erythrocarpa Makino, significantly suppressed HCV RNA levels with 50% effective concentrations of 15 ± 0.5 μM and 20 ± 1.1 μM in HCV replicon and JFH-1 infectious assays, respectively. There was no significant cytotoxicity observed at high concentrations, with a 50% cytotoxic concentration of 620 ± 5 μM. In addition, lucidone significantly induced heme oxygenase-1 (HO-1) production and led to the increase of its product biliverdin for inducing antiviral interferon response and inhibiting HCV NS3/4A protease activity. Conversely, the anti-HCV activity of lucidone was abrogated by blocking HO-1 activity or silencing gene expression of HO-1 or NF-E2-related factor 2 (Nrf2) in the presence of lucidone, indicating that the anti-HCV action of lucidone was due to the stimulation of Nrf2-mediated HO-1 expression. Moreover, the combination of lucidone and alpha interferon, the protease inhibitor telaprevir, the NS5A inhibitor BMS-790052, or the NS5B polymerase inhibitor PSI-7977, synergistically suppressed HCV RNA replication. These findings suggest that lucidone could be a potential lead or supplement for the development of new anti-HCV agent in the future.

Approximately 170 million people across the world are infected with hepatitis C virus (HCV). The virus causes chronic inflammation of the liver that ultimately leads to severe consequences such as cirrhosis and hepatocellular carcinoma (1). The HCV genome is a 9.6-kb positive single-strand RNA molecule encoding a single polyprotein of 3,000 amino acids that is processed by viral and cellular proteases to produce structural (C, E1, and E2) and nonstructural (NS2, NS3, NS4A, NS4B, NS5A, and NS5B) proteins (2). Pegylated alpha interferon (PEG–IFN-α) combined with ribavirin is the only recommended standard therapy for hepatitis C patients at present. However, the current therapeutic regimens only achieve a 40 to 50% cure rate in genotype 1 HCV-infected patients (3). In addition, the severe side effects of the current treatments, including depression, fatigue, flu-like symptoms, and hemolytic anemia, often lead to treatment discontinuation (4). Despite recent advancement of therapeutics with the approval of NS5/4A protease inhibitors, telaprevir and boceprevir, in combination with PEG–IFN-α and ribavirin, viral resistance and side effect to both inhibitors was still observed in clinical studies (5–7). Therefore, it is essential to develop new anti-HCV agents for therapeutic purposes.

Heme oxygenase-1 (HO-1), the rate-limiting enzyme in the oxidative degradation of heme, protects against oxidative stress and liver inflammation (8). Its reaction products including biliverdin, carbon monoxide, and ferrous iron are regarded as potent cytostatic agents (9). Among the various intracellular molecules capable of inducing HO-1, BTB and CNC homolog 1 (Bach1) and nuclear factor erythroid-2 derived 2 related factor 2 (Nrf2) play crucial roles in the regulation of HO-1 expression for the maintenance of cellular redox homeostasis (10). Nrf2 is essential for the transcriptional induction of antioxidant response element (ARE)-mediated phase II detoxifying and antioxidant genes via heterodimerization with small Maf (sMaf) proteins. In contrast, Bach1 heterodimerizes with sMaf proteins and interferes with HO-1 expression by competing the binding of the Nrf2-sMaf complex to the ARE of the HO-1 promoter region, which serves as a transcription repressor (11). Recent advances have revealed that the upregulation of HO-1 expression or overproduction of its metabolite is a promising strategy for suppressing HCV replication by activating the HO-1 transcriptional activator Nrf2 or targeting its transcriptional repressor Bach1 (12–14). The HO-1 product biliverdin has been demonstrated to be a major effector against viral replication by increasing the antiviral IFN response and inhibiting HCV NS3/4A protease activity (12, 14). Consequently, the discovery of HO-1-inducible agents may offer an advantage of therapeutic strategy by simultaneously targeting both viral and host factors for future HCV therapies.

By screening many natural products from plant using cell culture-based HCV replicon system, we identified a phytochemical, lucidone (Fig. 1A), isolated from the fruit of Lindera erythrocarpa that specifically inhibited HCV RNA replication. L. erythrocarpa Makino, belonging to the family Lauraceae, is widely cultured in Asian countries such as Taiwan, Japan, Korea, and China, and its fruits are traditionally used in folk medicine because of their extensive pharmacological properties, such as analgesic, digestive, diuretic, antidiotal, and antibacterial properties (15). The lucidone derivatives, such as methyllynderone and methyllycludone, also exhibited anti-inflammation activity and anti-farnesyl protein.
transferase (16, 17). Recent in vitro and animal model studies have demonstrated that lucidone exerts anti-inflammatory activity with significant suppression of iNOS and COX-2 production (16, 18) and antimelanogenic activity (19). In the present study, we characterize the anti-HCV activity of lucidone and evaluate its possible mechanism of action against HCV replication.

MATERIALS AND METHODS
Cell culture and reagents. Ava5 cells (20), a human hepatoma cell line (Huh-7 derivative) harboring HCV subgenomic replicon RNA, were cultivated in Dulbecco modified Eagle medium (DMEM) with 10% heat-inactivated fetal bovine serum, 1% antibiotic-antimycotic, 1% nonessential amino acids, and 1 mg of G418/ml. IFN-α2a (Roferon-A) was purchased from Roche, Ltd. BMS-790052 and PSI-7977 were purchased from Shanghai Haoyuan Chemexpress Co., Ltd. Telaprevir was purchased from Legend Stat International Co., Ltd. These compounds were stored at a concentration of 10 mM in 100% dimethyl sulfoxide (DMSO). The final concentration of DMSO was maintained at 0.1% for all experiments.

Plasmid construction. The pHO-1-Luc vector, kindly provided by Anupam Agarwal (University of Alabama), was used to measure the transcription activity of HO-1 (21). p3xARE-Luc, a reporter vector containing triple repeats of the Nrf2-dependent ARE, was used to measure the translocation and transcription activity of Nrf2. The AREs (TGACTCAGC) flanked by XhoI were inserted into the promoterless firefly luciferase vector pMCS-Luc (Stratagene, La Jolla, CA) and are designated as p3xARE-Luc. pISRE-Luc, a reporter vector containing firefly luciferase under the control of an IFN-stimulated response element (ISRE), was used to mea-
TABLE 1 Oligonucleotides used for real-time RT-PCR

<table>
<thead>
<tr>
<th>Oligonucleotide</th>
<th>Sequence (5’–3’)</th>
</tr>
</thead>
<tbody>
<tr>
<td>5’NS5B</td>
<td>GGAACACAGCTGCCATCA</td>
</tr>
<tr>
<td>3’NS5B</td>
<td>CCTCAAGGATAGAAATTAAA</td>
</tr>
<tr>
<td>5’GAPDH</td>
<td>GTTCTACACCATGTGGAGGA</td>
</tr>
<tr>
<td>3’GAPDH</td>
<td>ATGGAGATGCTGTTGCAATA</td>
</tr>
<tr>
<td>5’OAS1</td>
<td>CAAGTTAAGGCCCTACCC</td>
</tr>
<tr>
<td>3’OAS1</td>
<td>TGGGCTGTGGTTGAAATGTTG</td>
</tr>
<tr>
<td>5’OAS2</td>
<td>ACAGCTGAGAACCTTTGGA</td>
</tr>
<tr>
<td>3’OAS2</td>
<td>GCTTTAAGGAGCAGACG</td>
</tr>
<tr>
<td>5’OAS3</td>
<td>CACTGACATCCCCGAGCATG</td>
</tr>
<tr>
<td>3’OAS3</td>
<td>GATGAGCTGCTTCTGGG</td>
</tr>
<tr>
<td>5’PKR</td>
<td>ATGATGGAAAGCGCAAGG</td>
</tr>
<tr>
<td>3’PKR</td>
<td>GAGATGATGGCATCCCGT</td>
</tr>
<tr>
<td>5’IFN-α2</td>
<td>GCAAGTCAAGCTGCTTGG</td>
</tr>
<tr>
<td>3’IFN-α2</td>
<td>GATGTTTACCCTTTGGA</td>
</tr>
<tr>
<td>5’IFN-α17</td>
<td>AGGGATTTAGGCAACAGG</td>
</tr>
<tr>
<td>3’IFN-α17</td>
<td>CATCAGGAGGCTTCCTTCA</td>
</tr>
</tbody>
</table>

*GAPDH, glyceraldehyde 3-phosphate dehydrogenase; OAS, 2,5-oligoadenylate synthetase; PKR, protein kinase R; IFN-α, alpha interferon.

sure IFN response-dependent transcription activity (Stratagene). HO-1 (NM_002133), Nrf2 (NM_006164), and enhanced green fluorescent protein (EGFP) shRNA (as a negative control) were purchased from the National RNAi Core Facility, Institute of Molecular Biology/Genomic Research Center, Academia Sinica, Taiwan. The cloned DNA fragments were verified by DNA sequencing.

Western blot assay. A standard procedure was used for Western blotting (22). Membranes were probed with either anti-NS5B (1:5,000; Abcam, Cambridge, MA), anti-GAPDH (1:10,000; GeneTex, CA), anti-HO-1 (1:3,000; GeneTex), anti-BVR (1:1,000; Abcam, Cambridge, MA), or anti-Nrf2 (1:3,000; GeneTex) antibody. Signals were detected using an ECL detection kit (Perkin-Elmer, CT).

Quantification of cellular mRNA and HCV RNA. Total RNA was extracted from cell lysates using a total RNA miniprep purification kit (GMBiolab Co., Ltd., Taiwan) according to the manufacturer’s instructions. Quantitative real-time RT-PCR (qRT-PCR) was performed using an ABI Step One real-time PCR system (ABI, Warrington, United Kingdom). Each sample was normalized by the endogenous glyceraldehyde 3-phosphate dehydrogenase (gadphl) gene expression. The primers used are listed in Table 1.

HCV JFH-1 infection assay. The inhibitory effect of compound on HCV infection was assayed as previously described (23). In brief, the Huh-7 cells were seeded at density of 4 × 10³ cells/well in 24-wells culture plate and infected with 100 μl of HCV JFH-1 particles at a multiplicity of infection of 0.1 for 6 h, followed by incubation with various concentrations of lucidone for an additional 3 days. The total RNAs were then extracted and subjected to qRT-PCR to measure the mRNAs of HCV and GAPDH as described above.

Cytotoxicity assay. Cells seeded in a 96-well plate at a density of 5 × 10³ cells/well were exposed to various concentrations of crude extracts and the pure compounds. The cells were incubated at 37°C in an atmosphere of 5% of CO₂ for 4 days. Cell viability was determined by the MTS assay using the CellTiter 96 Aqueous One solution cell proliferation assay system (Promega, WI) according to the manufacturer’s instructions. Absorbance was detected at 490 nm using a Bio-Rad 550 plate reader (Bio-Rad, Hertfordshire, United Kingdom).

Transfection and luciferase activity assay. To evaluate the regulation of HO-1 Nrf2, or IFN response by lucidone, Ava5 cells were transfected with 1 μg of pPHO-1-Luc, p3xARE-Luc, or pSRE-Luc using T-Pro reagent (Ji-Feng Biotechnology Co., Ltd., Taiwan) in accordance with the manufacturer’s instructions. To evaluate the role of HO-1 and Nrf2 in HCV replication by lucidone, Ava5 cells were transfected with increasing concentrations of the HO-1 or Nrf2 shRNA expression vector (pPHO-1-shRNA and pNrf2-shRNA: 0.25 to 2 μg) in the presence of 30 μM lucidone. Each transfection complex contained 0.1 μg of a secreted alkaline phosphatase (SEAP) expression vector (pSEAP), which served as a transfection control. After 3 days of incubation, cell lysates were prepared for luciferase activity assay with the Bright-Glo luciferase assay system (Promega, Madison, WI) in accordance with the manufacturer’s instructions and Western blotting with specific antibodies. The supernatants were harvested for SEAP activity assay with a Phospha-Light assay kit (Tropix, Foster City, CA). For each experiment, luciferase activity was normalized to the SEAP activity. The basal level of promoter activity was defined as 1 for comparison of the fold values after normalization of the luciferase activity.

Intracellular bilirubin measurement. Cells were seeded in six-well plates at a density of 4 × 10³ and then treated with lucidone at various concentrations. After 3 days of incubation, the cells were lysed using lysis buffer (25 mM Tris-phosphate [pH 7.8], 2 mM dithiothreitol [DTT], 2 mM DTTA [1,2-diaminocyclohexane-N,N,N’,N’-tetraacetic acid], 10% glycerol, 1% Triton X-100). The total bilirubin concentration was measured by using a MedHiPro direct bilirubin test kit (Fornos Biomedical Technology Corp., Taiwan) according to the manufacturer’s protocol. The assay is calibrated using a CFAS (calibrator for automated systems; Roche Diagnostics, LTD., USA). The absorbance was detected at 546 and 660 nm using an Epoch microplate spectrophotometer (Bio-Tek Instruments, Inc., United Kingdom).

Cell-based and in vitro transcription and translation (TnT) assay for NS3/4A activity. To evaluate the effects of NS3/4A protease activity by lucidone inside cells, the Huh-7 cells were transfected with 1 μg of NS3/4A reporter vector pEG(ΔDEA4B)SEAP and 0.5 μg of the NS3/4A expression vector pNS3/4A using the T-Pro reagent (Ji-Feng Biotechnolog Corp., Ltd., Taiwan). Each transfection complex contained 0.1 μg of a firefly luciferase expression vector (pFLuc), which served as a transfection control. Subsequently, the transfected cells were incubated with different concentrations of lucidone with or without 20 μM HO-1 inhibitor SnPP. The reporter activity assays were performed as described above. An in vitro-coupled TnT reactions for NS3/4A protease activity assay were performed as described previously (24). In brief, the NS3/4A protease and EGFP(ΔDEA4B)SEAP substrate protein were generated by respective incubation of 1 μg of pNS3/4A or pEG(ΔDEA4B)SEAP in the presence of 1 mM methionine or 355 Ci of [35S]methylionine (Institute of Isotopes Co., Ltd., Budapest, Hungary)/ml in a total volume of 50 μl of rabbit reticulocyte lysate solution, where incubation of pEGFP alone served as the identity of cleaved product EGFP derived from NS3/4A protease-mediated proteolytic processing of EG(ΔDEA4B)SEAP. Reactions were carried out at 30°C for 2.5 h and then terminated by adding 5 U of Tagetin RNA polymerase inhibitor (Epiconcentre, Madison, WI). The NS3/4A protease activity assay was performed by coincubation of 10 μl of non-radioisotope-labeled NS3/4A protease with 10 μl of 35S-labeled EGFP(ΔDEA4B)SEAP in the presence of various concentrations of lucidone in a total volume of 25 μl at 30°C for 15 min. Treatment with NS3/4A protease inhibitor telaprevir (0.3 μM) served as a positive control. The reactants were subsequently analyzed by SDS-PAGE and autoradiography. Densitometric quantification was analyzed with Quantity One one-dimensional analysis software (Bio-Rad Laboratories, Inc., USA).

Analysis of drug synergism. Ava5 cells were treated with serially diluted lucidone (3, 6.125, 12.5, 25, and 50 μM) in combination with serially diluted IFN-α (7.5, 15, 30, and 60 U/ml), the HCV NS3/4A protease telaprevir (0.075, 0.15, 0.3, and 0.6 μM) (25), the HCV NS5A inhibitor BMS-790052 (1, 2, 4, and 8 pM) (26), and the RNA-dependent RNA polymerase nucleoside inhibitor PSI-7977 (10, 20, 40, and 80 nM) (27). Three days later, the total RNA was extracted to quantify HCV RNA levels by qRT-PCR. The relative RNA levels were normalized by cellular gapdh mRNA expression. Combination index (CI) values were analyzed using the CalcuSyn software (Biosoft, Cambridge, United Kingdom), a computer program based on the method of Chou and Talalay (28, 29). The 95% confidence intervals for the dose-response values were used to deter-
mine the data statistically. According to the percent inhibition of HCV RNA, the CI value is calculated using the following formula: $CI = (Da + Db)/(Dxa + Dxb) + DaDb/DxaDxb$. Da and Db are the concentrations of drugs A (for example, lucidone) and B (for example, IFN-α), respectively, required to inhibit X% of HCV RNA as single agents, whereas Dxa and Dxb are the concentrations of A and B, respectively, required to inhibit X% of HCV RNA in combination, for which treatment 0.1% DMSO was considered as a negative control. The effect of multiple drug combinations is presented as antagonism ($CI > 1$), additivity ($CI = 1$), or synergism ($CI < 1$). In addition, traditional isobologram analysis was used to confirm the drug-drug interaction (30).

Preparation of nuclear extract. Nuclear extracts were prepared using hypotonic and high-salt buffer extraction as previously described (31). Briefly, Avα5 cells were seeded in a 6-cm dish at a density of $4 \times 10^5$ cells/dish and then treated with or without lucidone at the indicated dose. After 3 days and the indicated times of incubation, the cells were lysed using the ice-cold hypotonic buffer (10 mM HEPES, 1.5 mM MgCl$_2$, 10 mM KCl, 0.5 mM DTT, 10% Nonidet P-40 [pH 7.9]). After centrifugation at 7,000 × g for 15 min, the resulting nuclear pellets were extracted with high-salt nuclear extraction buffer (20 mM HEPES, 1.5 mM MgCl$_2$, 0.2 mM EDTA, 0.6 M KCl, 0.2 mM DTT, 0.5 mM EDTA [pH 7.9]) at 4°C for 30 min. The protease inhibitors and phosphatase inhibitors were added to hypotonic buffer and high-salt buffer immediately before use. Finally, nuclear proteins were collected after centrifugation at 20,000 × g for 15 min and stored at −80°C until use.

Statistical analysis. The results were analyzed and are presented as means ± the standard deviations (SD) for at least three independent experiments. The statistical significance was analyzed by using the Student t test.

RESULTS
Lucidone inhibits HCV replication. To assess the potential of lucidone in inhibiting HCV replication, Avα5 cells, a parent Huh-7-derived cell line harboring an HCV subgenomic RNA replicon (20), were treated with lucidone either at increasing concentrations for 4 days or at the single concentration of 50 μM for various times of incubation (1 to 4 days). The inhibitory effect of lucidone on HCV protein synthesis was then examined by Western blotting. The results indicated that lucidone markedly decreased the HCV NS5B protein levels in a concentration- and time-dependent manner (Fig. 1Ba and b). The inhibitory effect of lucidone on HCV RNA replication was examined by qRT-PCR in HCV replicon cells. As expected, HCV RNA levels were suppressed by lucidone in a concentration-dependent manner (Fig. 1C, left axis). The calculated 50% effective concentration (EC$_{50}$) of lucidone for reducing HCV RNA levels was 15 ± 0.5 μM in HCV replicon cells. The cell viability assay revealed a 50% cytotoxic concentration (CC$_{50}$) value of 620 ± 5 μM (Fig. 1C, right axis) for lucidone, which indicated that lucidone is not cytotoxic at effective antiviral concentrations. In addition, we performed HCV JFH-1 infectious assay to confirm the inhibitory effect of lucidone on viral RNA replication, with an EC$_{50}$ of 20 ± 1.1 μM (Fig. 1D). Because of an acceptable selectivity index (SI; CC$_{50}$/EC$_{50}$) of ~31, we suggested that lucidone could be potential as a promising lead compound for development of new anti-HCV agent.

Lucidone upregulates HO-1 expression in an HCV replicon system. Lucidone has been demonstrated to effectively down-regulate lipopolysaccharide (LPS)-induced inflammation by blocking iNOS and COX-2 (16, 18, 32). Because HO-1 is a critical regulator involved in the suppression of iNOS and COX-2 expression in response to inflammation (33, 34), we next examined whether lucidone could modulate HO-1 expression in HCV replicon cells. We first performed a HO-1 promoter-based reporter assay. Avα5 cells were transfected with the reporter plasmid pH0-1-Luc carrying the HO-1 promoter-driven firefly luciferase gene and then incubated with increasing concentrations of lucidone for 3 days, for which pH0-1-Luc-transfected-Huh-7 cells provided the basal level of HO-1 expression. As shown in Fig. 2A, the luminescent signal was markedly diminished in Avα5 cells compared to the signal in Huh-7 cells, indicating that HCV proteins strongly downregulated HO-1 gene transcription. Subsequently, lucidone concentration dependently enhanced the luminescent signal in Avα5 cells compared to that in lucidone-untreated Avα5 cells. Similar to the results obtained in the promoter activity assay, qRT-PCR analysis clearly revealed that lucidone increased HO-1 RNA...
levels in a concentration-dependent manner (Fig. 2B). Immunoblot analysis was performed to confirm the induction of HO-1 protein synthesis by lucidone in a concentration-dependent manner (Fig. 2C).

Lucidone augments antiviral IFN response and suppresses HCV NS3/4A protease activity through biliverdin production. Recently, Lehmann et al. and Zhu et al. have demonstrated that the HO-1 product biliverdin inhibits viral replication by increasing the antiviral interferon response and blocking HCV protease activity (12, 14). To verify whether lucidone could enhance biliverdin production due to the increase of HO-1 production, we first measured the amount of bilirubin, the biliverdin metabolite, to reflect the intracellular biliverdin levels following lucidone treatment. The results revealed that lucidone concentration-dependently increased the bilirubin levels (Fig. 3A). Biliverdin reductase (BVR) is an important catalase for biliverdin formation from biliverdin in HO-1-dependent reaction (35). To rule out the possibility that the accumulation of biliverdin was due to the induction of BVR by lucidone, we further detected the effect of lucidone on BVR expression. As shown in Fig. 3B, the BVR protein levels were not interfered by lucidone under increasing concentrations. Because the conversion of biliverdin to bilirubin is proportional, we concluded that lucidone treatment resulted in the increase of biliverdin, which may lead to anti-HCV activity. To verify whether lucidone inhibited HCV replication via its aforementioned anti-HCV effects, we performed a transient ISRE activity assay using the ISRE-mediated firefly luciferase expression vector. Ava5 cells were transfected with pISRE-Luc reporter plasmids and incubated with increasing concentrations of lucidone for 3 days. As shown in Fig. 4A, lucidone treatment significantly increased ISRE-mediated luciferase activity at effective antiviral concentrations (black columns). Conversely, treatment of the specific competitive HO-1 inhibitor tin protoporphyrin (SnPP) significantly abrogated the transcriptional induction of ISRE by lucidone (white column). Furthermore, our results confirmed that SnPP reversed the inductive effect of lucidone on ISRE-mediated activity due to completely blocking enzymatic activity of HO-1 because the lucidone-enhanced bilirubin levels were dramatically reduced by SnPP at tested concentration (Fig. 4Ba). In contrast, SnPP did not interfere concentration-dependent induction of HO-1 protein levels by lucidone (Fig. 4Bb). In addition, we further measured the expression of IFN-α2 and -α17 in lucidone-treated Ava5 cells (30 and 50 μM) by qRTPCR analysis. As shown in Fig. 4C and D, lucidone significantly induced the mRNA levels of both IFN genes compared to no lucidone treatment (lanes 1, 3, and 5). Subsequently, we examined the mRNA levels of critical IFN-mediated antiviral genes (36), including protein kinase R (PKR), 2′-5′-oligoadenylate synthetase 1 (OAS1), OAS2, and OAS3 under the same experimental conditions. As expected, lucidone significantly induced the expression of PKR, OAS1, OAS2, and OAS3 compared to their expression in lucidone-untreated cells (Fig. 4Aa to d, lanes 1, 3, and 5). Conversely, treatment of HO-1 inhibitor SnPP significantly abrogated the expression of IFN-α2 and -α17 (Fig. 4C and D, lanes 2, 4, and 6) and the induction of antiviral genes (Fig. 4Aa to d, lanes 2, 4, and 6). Treatment with SnPP alone served as a mock control (lack 2). For the cell-based HCV NS3/4A protease activity assay, Huh-7 cells were cotransfected with the NS3/4A expression vector pNS3/4A-myc and its substrate vector pEG(DEΔ4AB)SEAP (37) (Fig. 5Aa), which contained a NS3 cleavage site between EGFP and SEAP, in the presence of increasing concentrations of lucidone for 3 days. Treatment with telaprevir (25) or bilirubin (14) served as positive controls. As shown in Fig. 5B, a concentration-dependent reduction of SEAP activity was observed in lucidone-treated cells compared to its activity in lucidone-untreated cells (black columns), reflecting that NS3/4A protease activity was inhibited by lucidone inside cells. Conversely, treatment of SnPP attenuated the inhibitory effect of lucidone on NS3/4A protease activity by detection of SEAP activity (white columns). To examine whether lucidone directly reacted with NS3/4A protease, we performed a cell-free transcription and translation (TnT) of NS3/4A activity assay (24). As shown in Fig. 5C, the cleavage of the EG(DEΔ4AB)SEAP fusion protein by NS3/4A protease was observed in the absence or presence of increasing concentrations of lucidone compared to telaprevir treatment, a positive control of protease inhibition. Densitometric quantification of the uncleaved- and cleaved-specific bands showed that there was no difference in the residual EG(DEΔ4AB)SEAP levels, remaining from 43 to 49%, from TnT experiments with or without lucidone (Fig. 5D, white squares), indicating that the cleavage efficiency of NS3/4A protease was not interfered by lucidone. Compared to the findings in cell-based reporter assay described above, we indicated that lucidone did not directly target NS3/4A protease. Taken together, these findings are suggestive that lucidone treatment leads to substantial biliverdin production through HO-1 induction, which results in the effective inhibition of HCV replication via the combination of the antiviral interferon response and anti-NS3/4A protease activity. Therefore, despite the enhanced level of IFN-stimulated gene (ISG), the expression of lucidone-treated cells was lower than that of lucidone-untreated cells.
FIG 4 Induction of the antiviral IFN responses by lucidone in HCV replicon cells. (A) Concentration-dependent induction of ISRE activity by lucidone. (B) Restoration of lucidone-induced HO-1 activity by HO-1 inhibitor SnPP. (C to E) Concentration-dependent induction of gene expression of IFN-α2 (C) and IFN-α17 (D) and IFN-mediated gene expression (E, panels a to d) by lucidone. For reporter analysis, Ava5 cells were transiently transfected with 1 μg of the IFN response reporter vector pISRE-Luc. Subsequently, the pISRE-Luc-transfected cells were incubated with the indicated concentrations (0 to 50 μM) of lucidone with or without 20 μM HO-1 inhibitor SnPP for 3 days, and total cell lysates were analyzed for luciferase activity. Luciferase activity in lucidone-untreated cells was defined as 1. For the detection of bilirubin production and HO-1 expression, the total cell lysates of lucidone-treated Ava5 cells were harvested for the quantification of bilirubin concentration and Western blot analysis using the MeDiPro direct bilirubin test kit combined with CFAS (calibrator for automated systems) and anti-HO-1 antibody, respectively, under the same assay conditions. GAPDH protein levels confirmed equal loading of cell lysates. For gene expression analysis, the total RNA of lucidone-treated Ava5 cells was extracted to quantify the RNA levels of IFN-α2, IFN-α17, OAS1, OAS2, OAS3, and PKR by qRT-PCR analysis under the same assay conditions. The relative RNA levels were normalized by cellular gapdh mRNA expression. The RNA level in lucidone-untreated Ava5 cells was defined as 1. Each value represents the mean fold of normalized data ± the SD for triplicate experiments. The error bars denote the SD of the mean. *, P < 0.05; **, P < 0.01.
FIG 5 Effect of lucidone on the HCV NS3/4A protease activity. (A) Schematic representation of the NS3 response reporter vector pEG(ΔE4AB)SEAP. The decapeptide sequence, named 8×DEMEEC-ASHL, corresponding to the NS4A/B junction was inserted between egfp and seap. (B) Concentration-dependent reduction of NS3/4A protease activity by lucidone in cell-based analysis. Hub-7 cells were transiently cotransfected with 1 μg of the pEG(ΔE4AB)SEAP vector and 0.5 μg of the NS3/4A expression vector pNS3/4A. Subsequently, the transfected cells were treated with the indicated concentrations (0 to 50 μM) of lucidone with or without 20 μM SnPP. After 3 days, total cell lysates were analyzed for SEAP activity. Treatment with 1 μM telaprevir or 100 μM biliverdin served as the positive controls. (C) No significant inhibition of NS3/4A protease activity by lucidone in cell-free TnT analysis. The reaction mixtures contained non-radioisotope-labeled NS3/4A protein and 35S-labeled EG(ΔE4AB)SEAP substrate protein generated by a cell-free TnT system (Promega) in the absence or presence of increasing concentrations of lucidone. Treatment with 0.3 μM telaprevir served as a positive control. TnT product of EGFP alone served as an indicator of protease-mediated proteolytic product from EG(ΔE4AB)SEAP. After incubation for 15 min at 30°C, reactants were subjected to SDS-PAGE and autoradiography. Western blotting with anti-NS3 antibody was performed to reveal equal amounts of NS3/4A protein in each reaction. (D) Densitometric quantification was performed to present the relative cleavage of EG(ΔE4AB)SEAP in the absence or presence of lucidone. The arrowheads indicated the expected sizes of EG(ΔE4AB)SEAP, ASHL-SEAP, EGFP-8×DEMEEC, and EGFP. The relative intensity of EG(ΔE4AB)SEAP was determined as the ratio of EG(ΔE4AB)SEAP band to total predominant bands corresponding to the TnT products, expressed as the remaining percentage. Each value represents the mean fold ± the SD of triplicate experiments after normalization of luciferase activities. The error bars denote the SD of the mean. *, P < 0.05; **, P < 0.01.
of IFN-treated cells at the effective concentration of anti-HCV activity (data not shown), and lucidone was shown to act in an additive manner on anti-HCV replication by both of the antiviral actions described above.

Anti-HCV activity of lucidone is mediated by HO-1 induction and activity. Subsequent experiments were performed to further investigate whether the anti-HCV activity of lucidone is mediated through HO-1 expression. Ava5 cells were incubated with increasing concentrations of the SnPP (2.5, 5, 10, and 20 μM) in the presence of lucidone (30 μM) for 3 days. Western blotting and qRT-PCR were used to determine the inhibitory effect of combinational treatment on HCV protein and RNA expression, respectively. As shown in Fig. 6A, SnPP treatment reversed the inhibitory effect of lucidone on HCV protein synthesis (lanes 3 to 6) in a concentration-dependent manner compared to the effects of no lucidone treatment (lane 1) and lucidone treatment in the absence of SnPP (lane 2). Similarly, HCV RNA levels were gradually restored by increasing the concentrations of SnPP (Fig. 6B). To rule out the possibility of a nonspecific effect of a pharmacological inhibitor, we performed an RNA interference technique to disrupt HO-1 function. Lucidone-treated or untreated Ava5 cells were transfected with either specific HO-1 shRNA (0.25 to 2 μg) or nonspecific control EGFP shRNA vectors. The effect of HO-1 shRNA on HO-1 and HCV protein synthesis was examined by Western blotting. As shown in Fig. 6C, HO-1 shRNA markedly reduced lucidone-mediated HO-1 induction (middle panel, lanes 3 to 6), and simultaneously reversed the inhibitory effect of lucidone on HCV protein synthesis (upper panel, lanes 3 to 6), whereas control shRNA had no effect on HO-1 expression (middle panel, lane 2) and the recovery of HCV protein synthesis in the presence of lucidone (upper panel, lane 2). It is noteworthy that the recovery of HCV protein levels was comparable to that in lucidone-untreated Ava5 cells when HO-1 expression was efficiently silenced (lanes 1 and 6). Likewise, the percent recovery of HCV RNA levels was correlated to the increase in the amount of HO-1 shRNA compared to their recovery in untransfected Ava5 cells in the presence of lucidone (Fig. 6D). Taken together, these results clearly revealed that HO-1 upregulation contributes to the antiviral action of lucidone.

Lucidone triggers nuclear translocation of Nrf2 and stimulates Nrf2-mediated transcriptional activity for HO-1 induction. Nrf2 is one of the important nuclear factors that transcriptionally activate HO-1 expression. To examine whether Nrf2 is involved in HO-1 induction by lucidone, we first analyzed the effect of lucidone on Nrf2 expression in Ava5 cells. As shown in Fig. 7A, lucidone increased total Nrf2 protein levels and nuclear Nrf2 accumulation in a concentration-dependent manner. A time-dependent accumulation of nuclear Nrf2 in response to lucidone treatment was also observed (Fig. 7B). According to the qRT-PCR analysis, we found that lucidone caused the elevation of Nrf2 expression at the transcriptional level (data not shown). We next used a p3XARE-Luc luciferase reporter construct to verify the specificity of lucidone in the induction of ARE-mediated HO-1 upregulation in response to Nrf2 binding. p3 × ARE-Luc-transfected Ava5 cells were incubated with increasing concentrations of lucidone for 3 days. As expected, lucidone enhanced luciferase activity in a concentration-dependent manner (Fig. 7C). Compared to the findings in the lucidone-untreated cells, an ~8-fold
increase in luciferase activity was observed at a lucidone concentration of 50 μM. We next silenced gene expression of Nrf2 using shRNA expression vector to verify involvement of Nrf2-mediated HO-1 upregulation in anti-HCV activity of lucidone. As shown in Fig. 8A, the increasing gene silence of Nrf2 gradually reduced lucidone-mediated HO-1 induction (second and third panels, lanes 3 to 6) and simultaneously reversed the inhibitory effect of lucidone on HCV protein synthesis (first panel, lanes 3 to 6), whereas control shRNA had no effect on HO-1 expression and the recovery of HCV protein synthesis in the presence of lucidone (lane 2). Likewise, the percentage recovery of HCV RNA levels was correlated to the increase in the amount of Nrf2 shRNA (Fig. 8B). Taken together, these results clearly revealed that Nrf2-mediated HO-1 induction may contribute to the antiviral action of lucidone.

**Lucidone synergistically inhibits HCV replication in combination with IFN-α or viral enzyme inhibitors.** To evaluate combination treatments, we treated Ava5 cells with lucidone in combination with IFN-α, the NS3/4A protease inhibitor telaprevir (25), which is a drug currently approved by the U.S. Food and Drug Administration, the NS5A inhibitor BMS-790052 (26), or the NS5B polymerase inhibitor PSI-7977 (27), a prodrug of 2’-F-2’-C-methyluridine monophosphate, at various fixed ratios of concentrations as described in Materials and Methods. After 3 days, total RNAs were harvested and quantified by qRT-PCR analysis. The inhibitory effect of the combination treatments on HCV replication was calculated using the isobologram method and CalcuSyn software (28, 30). The results of the combination studies expressed as the mean of CI values at an effective dose of 50% (ED$_{50}$), 75% (ED$_{75}$), or 90% (ED$_{90}$) inhibition. By definition, a CI value of 1 denotes additivity, a CI value of <1 denotes synergism, and a CI value of >1 denotes antagonism. As shown in Table 2, the CI values for the ED$_{50}$, ED$_{75}$, and ED$_{90}$ ranged from 0.21 to 0.78, indicating that the combination treatments synergistically inhibited HCV replication according to the guidelines of the program. We also observed an enhanced induction of ISG expression upon combination treatment with lucidone (50 μM) and PEG–IFN-α (50 U/ml), compared to monodrug treatment (data not shown), which supported the results of synergistic effect for lucidone in combination with IFN-α. A traditional isobologram of each combination was presented in Fig. S1 in the supplemental material. There was no significant cytotoxic effect observed in each combination at the higher concentrations using MTS assay (see Fig. S2 in the supplemental material), which excluded the possibility of synergistic cytotoxicity upon combination treatment.
A previous report demonstrated that treatment with miRNA-196, which targets Bach1, a transcriptional repressor of HO-1, resulted in HO-1 upregulation and subsequent suppression of HCV replication (13). In addition to suppressing HCV replication, the induction or overexpression of HO-1 has been demonstrated to interfere with the replication of other viruses, such as HIV (38), hepatitis B virus (39), and entrovirus 71 (40). Therefore, targeting HO-1 may be a potential therapeutic approach to inhibit virus infection. The present study is the first time to identify a phytocompound as a protective agent against HCV replication via HO-1 induction. In addition to Nrf2, nuclear factor-κB (NF-κB) and activator protein-1 (AP-1) are also involved in the transactivation of HO-1 expression (10). We will further investigate the effect of lucidone on the NF-κB and AP-1 expression in terms of HO-1 upregulation. In addition, HO-1 gene expression can be transcriptionally mediated by a number of intracellular signaling molecules, such as extracellular signal-regulated protein kinase (ERK) (41), p38 mitogen-activated protein kinase (MAPK), c-Jun N-terminal kinase, protein kinase C, and phosphatidylinositol 3-kinase (8). Among them, the activation of MAPK/ERK kinase-ERK1/2 signaling has been reported to be involved in anti-HCV activity upon oxidative stress (42). Moreover, lucidone was previously reported to exert protective effects against LPS-induced inflammation through NF-κB/MAPK signaling pathways (18). Therefore, it is worthwhile to further investigate additional molecules or signaling pathways involved in HO-1 activation by lucidone to clearly demonstrate the detailed molecular mechanism by which lucidone blocks HCV replication.

A number of new direct-acting antivirals (DAAs) for treating HCV infection are currently under development. Telaprevir, a recently approved NS3/4A protease inhibitor, significantly improves the sustained virological response rate in combination with the standard of care of PEG-IFN and ribavirin among patients with chronic genotype 1 HCV infection (43). Although the antiviral efficacy of protease inhibitor triple therapy appears to be positive, this regimen still has several challenges, such as anemia, a well-recognized side effect of IFN-based regimens, and the emergence of DAA-related resistant variants due to the high replication rate of the virus and the low fidelity of the NS5B polymerase (44). Indeed, targeting host factors required for the viral life cycle has been considered a favorable strategy to overcome the genetic variability of the viral genome because the mutation rate of host genes is lower than that of viral genes (45). Accordingly, combinations of DAAs and host-targeted antivirals may provide alternative regimens to overcome these limitations. In addition to targeting host HO-1 expression, lucidone exhibits synergistic activity against HCV replication in combination with other promising inhibitors against distinct targets of HCV (Table 2). In regard to lucidone as a potentially clinically useful drug or adjuvant, assessment of anti-HCV activity assay in vivo animal mode is required. Recently, Kumar et al. have performed in vivo experiments to demonstrate anti-inflammatory activity of lucidone against lipopolysaccharide.

**Fig. 9** Model for the inhibitory action of lucidone on HCV replication.

**TABLE 2** Effects of a combination of lucidone and various inhibitors on HCV replication

<table>
<thead>
<tr>
<th>Combination compound</th>
<th>Mean CI ± SD at ED50 (CI)</th>
<th>ED25 (CI)</th>
<th>ED10 (CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td>IFN-α</td>
<td>0.61 ± 0.012</td>
<td>0.53 ± 0.034</td>
<td>0.47 ± 0.072</td>
</tr>
<tr>
<td>Telaprevir</td>
<td>0.78 ± 0.04</td>
<td>0.66 ± 0.047</td>
<td>0.56 ± 0.069</td>
</tr>
<tr>
<td>BMS-790052</td>
<td>0.48 ± 0.095</td>
<td>0.42 ± 0.069</td>
<td>0.37 ± 0.05</td>
</tr>
<tr>
<td>PSI-7977</td>
<td>0.45 ± 0.087</td>
<td>0.30 ± 0.1</td>
<td>0.21 ± 0.104</td>
</tr>
</tbody>
</table>

*Avv5 cells were treated with combinations of various concentrations of lucidone and IFN-α, telaprevir, BMS-790052, or PSI-7977 for 3 days. The anti-HCV activity was determined by qRT-PCR to analyze the HCV RNA levels. The combination index (CI) values for the effective dose for 50% (ED50), 75% (ED25), or 90% (ED10) inhibition were calculated using the CalcuSyn computer program. Results are expressed as mean values for three independent experiments. The CI values indicate the degree of interaction of the potential drugs; values of <1, 1, and >1 are indicative of synergistic, additive, and antagonistic effects, respectively.**
According to the formula for dose translation by the used in the adult human is approximated to be 16.22 mg/kg. The body surface area normalization method is more commonly used to safely predicate a suitable starting dose for a human clinical trial from animal toxicology data, which incorporates several physiologic parameters between different mammalian species, such as oxygen utilization, caloric expenditure, basal metabolic rate, blood volume, circulating plasma proteins, and renal function (46). For example, the safe starting dose of lucidone used in the adult human is approximated to be 16.22 mg/kg according to the formula for dose translation by the factor. In addition, predication of in vivo human metabolic drug clearance from in vitro metabolism data and/or in vivo animal experiments is an important assessment for clinical studies. For predicting the hepatic clearance of drug, several liver models, such as the well-stirred parallel tube and dispersion, have been used to analyze pharmacokinetic parameters using in vitro hepatic microsome and isolated hepatocytes from animals or humans (47, 48). These animal experiments will provide practical information on developing lucidone as an adjuvant in current regimens against HCV in the future.

In summary, we found that lucidone possesses anti-HCV activity. An investigation of the mechanism(s) disclosed that the inhibition of HCV replication by lucidone was because of biliverdin production through Nrf2-mediated HO-1 induction, which indicated the feasibility of using selective HO-1 inducers to improve the efficacy of cellular defense pathways against HCV infection and also inhibit viral protease activity.

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