Introduction

Cap-dependent translation initiation is the most regulated step of protein production and is activated by multiple oncogenic signaling pathways. Previous studies show targeting this convergence point of signaling promotes strong antitumor activities. Previous studies show targeting this step of protein production and is activated by multiple oncoproteins, leading to growth arrest and apoptosis. In contrast to other eIF4A1 inhibitors, elatol induces markers of an integrated stress response, likely an off-target effect, but these effects do not mediate its cytotoxic activities. Elatol is less potent in vitro than the well-studied eIF4A1 inhibitor silvestrol but is tolerated in vivo at approximately 100× relative dosing, leading to significant activity against lymphoma xenografts.

Conclusions: Elatol’s identification as an eIF4A1 inhibitor with in vivo antitumor activities provides proof of principle for target-based screening against this highly promising target for cancer therapy. Clin Cancer Res; 24(17); 4256–70. ©2018 AACR.
Translational Relevance
Clinical activity of targeted signaling inhibitors is limited by acquired and de novo resistance, which stems from redundancies in signaling pathways, providing multiple ways for tumor cells to maintain activation of key downstream targets. Cap-dependent translation initiation is a key biologic process downstream from signaling, a convergence point of multiple pathways, that tumor cells depend on more than nonmalignant cells to maintain constitutive expression of key oncoproteins. A therapeutic window for targeting the cap-initiation complex eIF4F is well established preclinically, but no drug with this activity has entered evaluation in patients. The DEAD-box RNA helicase eIF4A1 is the enzymatic core of the eIF4F complex and is its most promising avenue for therapeutic targeting. Our study shows a way forward for high-throughput identification and characterization of eIF4A1 inhibitors to bring this highly promising therapeutic avenue to evaluation in patients.

eIF4A1 has proven a better drug target, being an ATP-dependent catalyst of mRNA unwinding (7).

Anti-eIF4A1 compounds identified through functional screening can work by increasing eIF4A1’s affinity for RNA, which sequesters it out of the eIF4F complex, and in the case of silvestrol possibly promoting particular binding to mRNA species containing polypurine sequences (8).

Alternatively, hippuristanol works by blocking the RNA interaction of both free and complexed eIF4A1 (5). We previously reported two compounds that inhibit eIF4A1 ATPase activity leading to detectable activity against tumor cell lines in vitro (9), but by comparison, this mode of action is minimally explored. Meanwhile, no eIF4A1 inhibitor described to date has led to a drug suitable for evaluation in clinical trials, and thus, the preclinical potential of this therapeutic approach remains untested in patients. A proof of principle for target-based screening versus eIF4A1 could help address the need for additional lead compounds suitable for development as cap-dependent translation inhibitors in cancer therapeutics.

The basis of in vitro therapeutic window for inhibiting cap-dependent initiation is thought to be differential dependence between cancer and nonmalignant cells for constitutive expression of cap-regulated proteins. Oncoproteins including MYC, MCL1, CYCLIN D3, and others are lost from cancer cells of multiple different cancer types upon eIF4A1 inhibition or knockdown (2, 10–12), and several different mRNA structural and sequence elements have been identified as potentially mediating these effects (reviewed in ref. 13). Regardless, cancer cells are more prone to enter apoptosis than nonmalignant cells upon treatment with eIF4A1 inhibitors, and animal studies demonstrate antitumor activities in vivo at doses tolerated by the host (2, 4).

Translationally regulated proteins are mediators of multiple cancer hallmarks, and their loss through eIF4F inhibition is seen as a way to help overcome both upstream signaling redundancies and tumor heterogeneity (14). Moreover, although gene expression changes at the level of mRNA are often reported as the key outputs of oncogenic signaling and epigenetic reprogramming, protein expression is what actually mediates cellular phenotype. Studies in several systems show poor correlation between mRNA and protein expression, highlighting translation as the step at which the proteome is primarily regulated (15).

In cancer, overexpression of components of the translational machinery, including eIF4F subunits, has been reported in a variety of diseases (reviewed in ref. 16). All of this evidence points to targeting translation directly as a highly promising avenue for cancer therapy for a wide range of cancer types, and a number of different approaches for translational inhibition are under preclinical investigation (16).

Proof of principle that inhibiting translation is a tolerable option in humans comes from the drug omacetaxine, which inhibits elongation at the ribosome and is FDA approved for patients with chronic myelogenous leukemia resistant to tyrosine kinase inhibitors (17). Inhibition at the more regulated and arguably more promising step of initiation, however, remains untested in patients.

eIF4A1 and other DEAD-box helicases are highly conserved enzymes that use the cooperative binding of ATP and RNA to cycle through conformational changes, allowing duplex unwinding (18–20). Inhibition of ATP hydrolysis in these proteins by RNA aptamers or protein-interacting partners has been shown to clamp them in place while bound to RNA and prevent their function, but optimized small molecules that work in this way have not yet been discovered (21, 22). Here, we report that the marine natural compound elatol is a novel eIF4A1 inhibitor successfully identified through target-based screening. Although off-target effects and synthetic complexities of this particular compound limit its utility in further development, we establish proof of principle for target-based screening against eIF4A1 based on inhibition of ATP hydrolysis, yielding a compound with in vivo antitumor activity.

Materials and Methods

Cell-free in vitro studies of eIF4A1
Protein purification and assessment of ATPase activity by malachite green were as described previously (9). Helicase assays were performed as per ref. 23. In isothermal titration calorimetry, eIF4a was dialyzed against buffer A (20 mmol/L MES-KOH, pH 6.0, 10 mmol/L potassium acetate, 2.5 mmol/L MgCl2, 1% glycerol, and 1 mmol/L DTT) for 12 hours. eIF4a was supplemented with 2% DMSO to match the ligand solution, degassed, and loaded in the cell of a nano-isorotermal titration calorimeter (TA Instruments). A total of 12 to 20 injections of 0.2 mmol/L elatol in buffer A were made every 200 seconds over a 3,000-second timeframe. NanoAnalyze software (TA Instruments) was used to integrate the peaks of the isotherm. The peaks were then integrated from injection start to 75 seconds postinjection and fit to an independent binding model. Replica experiments were done using 25 and 10 μmol/L eIF4A4 to add power to the stochiometric given by the NanoAnalyze software.

ATPase activity assays
Proteins were prepared as follows: 1 μmol/L eIF4A1 in buffer A (20 mmol/L MES-NaOH, pH 6.0, 100 mmol/L potassium acetate, 2.5 mmol/L MgCl2, 1% glycerol, and 1 mmol/L DTT), 500 nmol/L eIF4A1-K282R in buffer A, and 500 nmol/L eIF4A1-K238E in buffer A. ATP was added to generate samples of each protein containing a gradient of ATP (2 mmol/L, 1 mmol/L, 500 μmol/L, 250 μmol/L, 125 μmol/L, 62.5 μmol/L, 31.25 μmol/L, 0). The assay was carried out at 37°C, and after 1 hour, a 20 μL aliquot of the reactions was added to 40 μL of malachite green solution (9.3 μmol/L malachite...
green, 53 mmol/L (NH₄)₂MoO₄, 1 mol/L HCl, 0.04% Tween 20). After 5 minutes, the OD₅₆₂ was read on a GENs plate reader (BioTek Synergy 2). This was repeated after 2, 3, and 4 hours. The Michaelis–Menten curves were plotted and the Michaelis–Menten values were calculated (GraphPad Prism Software).

Molecular modeling
Modeling of elatol with eIF4A1 (PDB code: 2ZU6) was performed using Glide docking program (Schrodinger). Initially, the docking grid was created around the binding site and 1 was docked using extra precision (XP) glide docking. Resulting poses were evaluated using docking score and hydrogen bonding interaction with the active site residues.

Cell lines and reagents
All cell lines were routinely verified by STR fingerprinting and confirmed mycoplasma negative using the PlasmoTest Kit (Invivogen: REP-PT1). The mouse embryonic fibroblasts (MEF) eIF2α Serine 51 A/A and corresponding wild type were a kind gift from Dr. Randal Kaufman at Sanford Burnham Prebys Medical Discovery Institute (San Diego, CA). The MEFs ATF4 knockout and wild-type were a kind gift from Dr. Peter Johnson at the NCI (Rockville, MD). HBL1, TMD8, U2392, Riva, Toledo, OZ, SU-DHL-4, WSI-DLCL-2, MD901, and SNU-398 cell lines were grown in RPMI culture media (Corning) supplemented with 10% FBS and penicillin/streptomycin (P/S, RPMI). OCI-Ly2, OCI-Ly3, OCI-Ly10, and OCI-Ly19 were grown in IMDM with 20% FBS and P/S. DB, Farage, SU-DHL-6, SU-DHL-6 and Karpas-422 were grown in RPMI with 20% FBS and P/S. MDA-MB-468 were grown in DMEM with 10% FBS (D10). ATF4 wild type and knockout MEFs were cultured in D10 additionally supplemented with nonessential amino acids (Thermo Fisher Scientific) and 50 μmol/L β-mercaptoethanol. eIF2α wild-type and Ser37/A mutant MEFs were cultured in D10 media supplemented with NEAA. Silvestrol was purchased from MedChem Express (HY-12351). The PERK inhibitor GSK2606414 was purchased from Millipore/Calbiochem (516535). Tunicamycin was acquired from Millipore (L300015) following the manufacturer's protocol. Silvestrol, elatol, or tunicamycin at 48 hours and collected for analysis 56 hours following transfection with siRNA.

RT-PCR
RT-PCR was performed as described in ref. 25. TaqMan probes were purchased from Thermo Fisher Scientific: 18s rRNA 4319413E-0810041, GAPDH Hs00275899_g1, eif4a1 Hs00426773, eif4a2 Hs00756996_g1, Ccnd3 Hs00236949_m1, Mcl1 Hs01050896_m1, Myc Hs00153408_m1.

Viability assays
Cells were plated at 3–5 × 10³ cells per well in serial dilutions of drug ranging two logs with the top concentration for silvestrol 1 μmol/L and the top concentration for elatol 10 μmol/L. Viability was measured after 72 hours using CellTiter-Glo (Promega G7573) following the manufacturer’s protocol. Luminescence was detected on the BioTek HT Synergy plate reader, and LD₅₀ values calculated using nonlinear regression fit in GraphPad Prism7.

Apoptosis assays and Annexin staining
A total of 1–3 × 10⁵ cells were plated with indicated drug treatments and washed once with ice-cold PBS at the indicated time point and stained with PE-conjugated Annexin V and 7-AAD (BD Biosciences) according to the manufacturer’s protocol. Stained cells were analyzed by flow cytometry on the Attune NxT (Thermo Fisher Scientific). Data were analyzed using FlowJo v9.9.6 (FlowJo).

Dual-luciferase reporter assay
The dual-luciferase plasmids were a kind gift from Jerry Pelletier at McGill University (Montreal, Canada). A total of 7 × 10⁵ HEK-293 cells were plated and transfected with 400 ng of plasmid using Lipofectamine 3000 (Thermo Fisher Scientific L300015). Thirty-six hours after transfection, cells were treated as indicated before they were lysed using the Dual-Luciferase Reporter Assay Kit (Promega) and 100 μL added in triplicate to a white bottom 96-well plate for the detection of luciferase following the manufacturer’s protocol. Luciferase units were normalized to the DMSO-treated control.

O-propargyl puromycin assay
O-propargyl puromycin assay was performed as in ref. 26, but 1 × 10⁵ cells were plated and treated as indicated and pulsed with 50 μmol/L O-propargyl puromycin (OPP, Jena Biosciences) before being collected and washed twice with PBS. Alexa-488 azide (Thermo Fisher Scientific A10266) was used for conjugation to C-terminally labeled proteins and samples ran on Attune NxT.
Novel Screen for Translation Inhibitors Targeting eIF4A1

Polysome profiling
A total of 2 × 10^6 cells were treated as indicated and washed with ice-cold PBS with 100 g/mL cycloheximide and 100 U/mL RNasin. Lysates were cleared and equal A260 units (measured using NanoDrop 2000, Thermo Fisher Scientific) were loaded onto 10% to 50% sucrose gradients and centrifuged at 260,343 × g rpm in a SW-41 Ti rotor for 1.5 hours at 4°C. Samples were fractionated and 254 nm absorbance recorded using the gradient fractionation system (Brandel).

CLL patient cells
Chronic lymphocytic leukemia (CLL) patient cells were used as described previously (26).

Retroviral complementation experiments
Generation of the control and eIF4A stable knockdowns in NIH-3T3 cells was performed as described previously (27).

In vivo experiments
All in vivo experiments were performed following protocols approved by the relevant Institutional Animal Care and Use Committee. The initial non-tumor-bearing MTD studies and SU-DHL-6 xenografts were done with female 8- to 10-week-old SCID mice from Charles River Laboratories. Complete blood counts were collected from nontumor-bearing mice dosed as indicated and evaluated on a Hemavet950. For SU-DHL-6 xenografts, 2 × 10^6 cells were washed with ice-cold PBS and mixed 1:1 with Matrigel (Corning 354234) and injected subcutaneously into the right flank of SCID mice. Once the tumor reached 60 mm^3, mice were pair-matched and divided into two treatment groups: vehicle (sterile water with 5.2% Tween-80, 5.2% PEG-400) or 20 mg/kg elatol i.p. daily for 5 days. Mice were weighed and tumor volume calculated twice weekly. The MTD study of a single intraperitoneal injection of elatol up to 100 mg/kg was done on cohorts of 5 CD1 mice. H&E pathology slides were prepared using standard techniques and analyzed at the University of Miami (Miami, FL). For the OCI-Ly3 xenograft study, 10 × 10^6 cells were first washed with ice-cold PBS and mixed 1:1 with Matrigel and implanted subcutaneously on the flank of SCID mice. Once tumors reached 500 mm^3, tumors were dissected and dissociated using the gentleMACS Dissociator (Miltenyi Biotec) and 1 × 10^6 serially transplanted tumor cells were implanted subcutaneously into the flank of SCID mice. Once tumors reached 50 mm^3, mice were pair matched and split into two treatment groups: vehicle or 40 mg/kg elatol twice weekly. Mouse weight and tumor volume were monitored twice weekly.

Statistical analysis
All numerical data are based off of biological replicates represented as mean ± SEM. For Annexin staining and OPP incorporation, unpaired, two-tailed t tests were performed, with the addition of the two-stage linear setup procedure of Benjamini, Krieger, and Yekutieli to compare treatments over time, with P < 0.05 as significant. The OCI-Ly3 xenograft experiment was compared using a two-way ANOVA with P < 0.05 significant. All statistical comparisons were performed using GraphPad Prism7.

Results
Elatol is an eIF4A1 inhibitor with broad antitumor activity
We recently reported a screen for compounds able to inhibit eIF4A1's ATP hydrolysis and characterized the ATP-competitive inhibitors elatasib A and allolaurinterol (9). Despite inhibiting eIF4A1 ATP hydrolysis, when we evaluated the initial screen hits against non-Hodgkin lymphoma cell lines, where other eIF4A inhibitors show strong antitumor activities (2, 24, 28), elatasib A and allolaurinterol were inactive up to 10 μmol/L, indicating poor cell membrane permeability or metabolic liabilities (Fig. 1A). However, the brominated marine terpene elatol (Fig. 1B) identified on the same screening platform was toxic at concentrations less than 1 μmol/L in the preliminary screen of cancer cell lines and warranted further characterization. In the malachite-green assay used to identify eIF4A1 inhibitors in the initial screen, elatol is active against eIF4A1's ATP hydrolysis at 16.4 μmol/L (Fig. 1C). Counter-screening showed no activity against additional purified ATP-hydrolyzing enzymes, including the bacterial chaperonin GroEL, the human chaperone HSP70 (both considered unlikely to be affected by eIF4A1 modulators, as they have the least similar ATP pockets), the classical Walker A/B protein p97 and three other DEAD-box helicases: DDX3, DDX17, and DDX39A (Fig. 1C). In addition, in a commercial kinome screen, elatol had no activity against 97 oncogenic and related kinases, further validating its specificity (Supplementary Fig. S1A). To broadly interrogate elatol's antitumor activities, we tested it against the large collection of validated cancer cell lines maintained cooperatively by investigators at Harvard University (Boston, MA) and the Wellcome Trust (24). In this screen for growth inhibition, we found a wide range of sensitivity to elatol (Fig. 1D) with 37% (344/924) of cell lines having IC₅₀ < 1 μmol/L. Leukemia and lymphoma cell lines were the most sensitive tumor type overall, followed by breast and lung, all of which have been previously identified as candidates for eIF4A or eIF4F complex inhibition (12, 29). With lymphoma cell lines identified as a sensitive group, we proceeded to test three cell lines derived from diffuse large B-cell lymphoma (DLBCL), an aggressive lymphoma and the most common hematologic malignancy overall. A single elatol treatment completely halted proliferation in these lines (Fig. 1E). Elatol also induced apoptosis in a time- and concentration-dependent manner, with greater than 50% apoptosis by 24 hours in the most sensitive line (Fig. 1F), and the compound was less potent against normal human peripheral blood mononuclear cells (Supplementary Fig. S1B). Elatol therefore inhibits eIF4A's ATP hydrolysis in vitro and is highly toxic in a variety of cancer cell lines.

Elatol binds in a 2:1 ratio to eIF4A1 and disrupts helicase activity
To further characterize the interaction between elatol and eIF4A1, we first performed isothermal titration calorimetry. These results confirmed binding (Kᵦ = 1.98 ± 0.31 μmol/L) and revealed an unexpected 2:1 elatol:eIF4A1 stoichiometry (Fig. 2A). As has been the experience with other eIF4A1-interacting compounds,
our attempts to obtain a cocrystal structure with elatol have been unsuccessful. Molecular modeling studies, however, also predicted 2:1 stoichiometry, suggesting binding by two adjacent elatol molecules in the helicase core of eIF4A, with each elatol molecule interacting with a lysine residue from either the amino or carboxy RecA-like domains of eIF4A (Fig. 2B). The in silico model therefore predicts mutation of either lysine, K82 or K238, should lessen drug activity in vitro. K82 is part of the Walker A domain of eIF4A.
Figure 2.

Elatol binds to the N- and C-termini of eIF4A in a 2:1 stoichiometry. 

A, Isothermal titration calorimetry of eIF4A1 and elatol. Data fit to an independent binding model using NanoAnalyze software from TA instruments. Mean ± SEM, n = 3. 

B, Putative elatol binding with eIF4A1 based on molecular modeling experiments shows two elatol molecules (blue and pink) interacting with key lysines (yellow) in the RNA-binding groove between the two helicase domains of eIF4A (gray and purple). 

C, Malachite green assay for ATP hydrolysis showing IC50 for elatol treatment against wild-type eIF4A1 or proposed lysine mutants in the proposed binding sites. Mean ± SEM, n = 3. 

D, eIF4A helicase activity measured following treatment with various known eIF4A inhibitors and elatol. Mean ± SEM, n = 3.
ATPase motif conserved in all DEAD-box helicases (18, 19), and we found the nonconservative substitution K82E resulted in catalytically inactive protein (not shown). More conservative K82R, however, retained full ATPase activity of the wild-type protein but was significantly less sensitive to elatol (IC_{50} 85.5 ± 31.8 μmol/L; Fig. 2C; Supplementary Fig. S2A). K238E also retained full activity of wild type and was also less sensitive to elatol, although the effect was less pronounced (IC_{50} 34.2 ± 12.8 μmol/L; Fig. 2C; Supplementary Fig. S2A). Therefore although elatol's interactions with both residues in the helicase core likely contribute to inhibition, the interaction with K82 is likely more important. Moreover, these results highlight the ATP-binding pocket as a promising drug-binding site for additional development of eIF4A1 inhibitors. Finally, we assessed whether elatol affected eIF4A1's helicase activity in vitro using a fluorescence-based assay in which successful RNA unwinding results in quenching (23). Here, we found elatol results in inhibition in a manner similar to the previously characterized eIF4A1 inhibitors hippuristanol, silvestrol, and a silvestrol-derived roscaglate (Fig. 2D; Supplementary Fig. S2B). Elatol's inhibition of eIF4A1 ATPase therefore depends on binding two distinct pockets in the eIF4A1 core with 2:1 stoichiometry. In addition, elatol inhibits eIF4A1 helicase activity in a cell-free context.

Elatol is a translation inhibitor in cells with reduced potency compared with silvestrol

Silvestrol's mechanism as an eIF4A inhibitor and potent activity against a variety of tumor types in vitro are well described (2, 4, 30). In a panel of lymphoma cell lines, we found elatol was potent than silvestrol, with LD_{50} in a metabolic viability assay ranging from 130 to 5,756 nmol/L compared with 2.7 to 213 nmol/L for silvestrol (Fig. 3A). The 2:1 stoichiometry of elatol's interaction with eIF4A1 would only partially explain this disparity, and additional differences are likely to exist between the compounds. In particular, silvestrol might promote binding by free eIF4A1 to polypurine stretches of mRNAs that contain them (8), while its resulting reduced availability causes ribosome stalling on G-quadruplex structures found in complex 5' UTRs (30). These unique mechanistic properties of silvestrol would not be expected to apply to elatol, based on its simpler chemical structure and its predicted binding to the target. We next compared elatol and silvestrol in a variety of assays to assess elatol's effects on protein translation. At concentrations relative to the respective LD_{50} of either drug in each cell line, elatol, like silvestrol, causes a global decline of protein synthesis measured by incorporation of OPP, in a concentration- and time-dependent manner (Fig. 3B and C). OPP incorporation is measured on live gated cells, but to ensure these effects on translation are not observed with general cytotoxic agents, we compared the DNA-damaging agent carboplatin, again at similar concentrations relative to its LD_{50}, and found no significant decline in OPP incorporation (Fig. 3C; Supplementary Fig. S3A). Elatol also moderately reduced polystyrene translation at a 2-hour treatment compared with silvestrol, but by 16 hours showed complete elimination of polysome translation (Fig. 3D). Western blotting showed elatol, like silvestrol, causes loss of well-established translationally regulated oncoproteins in lymphoma cells (Fig. 3E; Supplementary Fig. S4A). Elatol also downregulated cap-dependent protein translation more strongly than cap-independent IRES-driven expression in a dual-luciferase reporter assay (Fig. 3F). Decreased eIF4A1 function either through knockdown or pharmacologic inhibition results in increased expression of the eIF4A2 transcript (31), a potentially useful marker of drug specificity against eIF4A1. In contrast to silvestrol treatment, which consistently induced eIF4A2 mRNA upregulation in sensitive lymphoma cell lines, we found a mixed response to elatol treatment. eIF4A2 increased in the two more sensitive cell lines OCI-Ly3 and SU-DHL-6 but did not in the less sensitive RIVA cell line (Supplementary Fig. S3B). In addition, we observed some decreases in both transcript levels at higher concentrations of elatol, suggesting elatol may have effects on mRNA transcription in addition to translation. We therefore examined mRNA expression of genes whose protein products are lost in response to elatol as in Fig. 3E and again found a mixed response across cell lines, with the most sensitive, OCI-Ly3, showing a decrease in mRNA levels following elatol treatment (Supplementary Fig. S3C). We therefore interrogated protein and mRNA expression in parallel across a range of elatol concentrations in this cell line and found that although both are affected, protein expression changes begin as low as 100 nmol/L below the LD_{50} of this cell line, while mRNA reductions are not significant until 500 to 1,000 nmol/L, well above the LD_{50} (Supplementary Fig. S3D). These results show elatol works more potently as a translation inhibitor but at higher concentrations, well above the LD_{50}, also affects transcription, raising questions about potential cellular off-target effects. Finally, silvestrol has been shown to inhibit deregulated protein translation in B-cell malignancies (25). We found that elatol, like silvestrol (Yeomans, Wilmore, Packham unpublished data) significantly reduced protein translation measured by OPP labeling in primary chronic lymphocytic leukemia (CLL) cells stimulated via the B-cell receptor using anti-IgM (Fig. 3G, ref. 26). Addition of caspase inhibitor Q-VD-OPh prevents induction of apoptosis, and therefore, the loss of protein translation in the CLL cells is due to inhibition of protein translation. In contrast, elatol did not significantly inhibit OPP labeling in unstimulated primary CLL cells, or normal malignant T cells, which were also present in the patients' blood samples (Supplementary Fig. S4B).

Integrated stress mediators induced by elatol do not result from an unfolded protein response

Elatol inhibits protein translation, but because of differences compared with silvestrol, we wanted to test for factors in addition to inhibition of eIF4A1 that might mediate elatol's toxicity in cells. We first employed the hepatocellular carcinoma (HCC) cell line SNU-398, which has constitutive mTORC1 activity due to biallelic loss of its negative regulator TSC2 and therefore high dependence on downstream translational activation (32). In contrast to the cytoplastic effect of mTORC1 inhibition by rapamycin, both silvestrol and elatol are highly cytotoxic in SNU-398 (LD_{50} 41 and 219 nmol/L, respectively, Fig. 4A), consistent with previous findings that eIF4A1 inhibition is more apoptotic than the cytoplastic effects of decreased eIF4E availability downstream of mTORC1 inhibition (33). Western blotting shows as expected that rapamycin potently inhibits phosphorylation of ribosomal protein S6 and 4EBP1 downstream of mTORC1 (Fig. 4B). Silvestrol and elatol both show no activity against mTORC1 targets, while still strongly downregulating expression of cap-dependent oncogenes. We noted, however, that elatol in contrast to silvestrol induced expression of the transcription factor ATF4, which is translationally upregulated downstream of eIF2α phosphorylation on serine 51 during an integrated stress response (ISR; ref. 34). Although eIF2α S51 phosphorylation was not notably induced by elatol in this experiment, these data point
Figure 3.

Elatol is toxic to non-Hodgkin lymphoma cell lines and inhibits protein translation. **A**, Sensitivity of a collection of non-Hodgkin lymphoma cell lines to known eIF4A inhibitor silvestrol or elatol. Viability measured after 72-hour treatment using Promega CellTiter-Glo reagent. LD50 calculated using nonlinear regression fit analysis in GraphPad Prism 7. Mean ± SEM, n = 4. **B**, Histogram plot showing OPP labeling in DLBCL cell lines treated with DMSO (green), 100 nmol/L silvestrol (red), or 10 μmol/L elatol (blue) for 24 hours. Gray, unlabelled cells. **C**, Mean fluorescent intensity of live cells labeled with OPP in DLBCL cells treated with indicated concentrations of silvestrol, elatol, or carboplatin for 4, 16, or 24 hours. Normalized to DMSO control. Mean ± SEM, n = 3; *P < 0.05; **P < 0.001. **D**, Polysome profiling of OCI-Ly3 cells treated with DMSO or 100 nmol/L silvestrol for 2 hours or 2 μmol/L elatol for 2 or 16 hours. **E**, Western blot showing protein expression of translationally regulated genes in DLBCL cells treated with DMSO (D), 50 nmol/L silvestrol (S), or 5 μmol/L elatol (E) for 16 hours. Representative images, n = 3. **F**, Dual luciferase reporter assay measuring cap-dependent versus IRES-mediated luciferase expression following an 8-hour treatment with the indicated translational inhibitors. Relative luciferase units normalized to DMSO-treated cells. Mean ± SEM, n = 3. **G**, Translation measured by OPP incorporation in CD19+CD5+ CLL patient cells following anti-Ig-M stimulation and elatol treatment. Normalized to anti-Ig-M stimulated but untreated cells. CpG-ODN stimulation used as a control. Mean ± SEM, n = 4; *P < 0.05; **P < 0.001.
Figure 4.
Elatol treatment induces ATF4 expression, not mediated by the UPR. A, Cell viability of SNU-398 cells measured after 72-hour treatment with translation inhibitors silvestrol, rapamycin, or elatol. Mean ± SEM, n = 4. B, Western blot showing protein expression of SNU-398 cells treated with DMSO (D), 200 nmol/L rapamycin (R), 100 nmol/L silvestrol (S), or 1 μmol/L elatol (E) for 16 hours. Representative images. C, Time course showing protein expression in DLBCL cell lines following elatol treatment. Representative images. n = 2. D, Cell death measured by flow cytometry following Annexin V staining in DLBCL cell lines treated with indicated inhibitors with or without the combination of the PERK inhibitor GSK2606414 for 24 hours. Mean ± SEM, n = 3. * P < 0.05; ** P < 0.001. E, Western blot showing protein expression in cells treated for 4 hours with DMSO, 100 nmol/L silvestrol, 5 μmol/L elatol, or 5 μmol/L tunicamycin with or without the combination with the PERKi. Representative images. n = 3.
to possible ISR induction by elatol that did not occur in response to silvestrol.

In lymphoma cells, as in the TSC2-deficient HCC cells, ATF4 is rapidly and strongly induced by elatol exposure, while eIF2α S51 phosphorylation is variable (Fig. 4C). The breast cancer cell line MDMB-MB-468, an elatol-sensitive cell line in the Harvard/Wellcome cell line collection, also showed ATF4 induction in response to treatment, indicating this is a general effect of elatol treatment, not cell type specific (Supplementary Fig. S4C). The ISR is a cytoprotective pathway that is activated in response to various cellular insults, and the end result of activation temporarily halts protein synthesis through eIF2α phosphorylation, allowing the cell to attempt to respond to the cellular stress or induce apoptosis if it is too severe (35). The unfolded protein response (UPR) is a type of ISR activated in response to unfolded proteins in the endoplasmic reticulum, leading to eIF2α phosphorylation by the Protein Kinase R-like Endoplasmic Reticulum Kinase (PERK), along with two additional branches indicated by expression of spliced XBP1 (XBP1s) and cleavage of ATF6, respectively. To evaluate whether activation of the ISR through the PERK kinase was causing the upregulation of ATF4 expression following elatol treatment, we tested the combination of the PERK inhibitor GS2606414 (PERKi; ref. 36) with drug treatment. PERKi treatment resulted in no significant reversal of elatol’s potency versus cultured cells, measured at 24 hours by Annexin V staining, in contrast to its antagonism of the UPR-inducing compound tunicamycin (Fig. 4D). In a short 4-hour treatment where elatol induction of ATF4 was strong, PERKi did not reverse elatol’s activation of ATF4 protein expression, and there was no evidence of overall UPR induction indicated by either XBP1s expression or ATF6 cleavage (Fig. 4E; Supplementary Fig. S4D). Elatol’s induction of ATF4 protein expression therefore is not a result of an unfolded protein response, but its importance to the drug’s activities in cells remains unclear from these results.

ISR induction by elatol does not mediate its toxicity to cells, but elf4A1 knockdown increases sensitivity to drug effects

To further assess ISR induction by elatol and determine its role in the drug's toxicity to cells, we first confirmed that the drug’s induction of ATF4 is downstream of eIF2α S51 phosphorylation. Murine embryonic fibroblasts (MEF) with both elf2α alleles mutated to S51A (elf2α-S51 A/A; ref. 37) show no induction of ATF4 in response to elatol, in contrast to littermate control elf2α-S51 S/S MEFs (Fig. 5A). There is no significant difference, however, in elatol sensitivity between these cells [Fig. 5B, IC_{50} = 1.029 nmol/L (A/A), 1.399 nmol/L (S/S); P = 0.9845]. We also employed MEFs with biallelic Atf4 deletion (38) in comparison with MEFs from wild-type littermates and found the Atf4-deficient cells are somewhat more sensitive [Fig. 5B, IC_{50} = 2.785 nmol/L (Atf4−/−), 4.038 nmol/L (WT); P = 0.0138]. For confirmation in tumor cells, we used siRNA to knock down ATF4 in elatol-sensitive MDA-MB-468 breast cancer cells and found no change in induction of apoptosis by elatol (Supplementary Fig. S5A and S5B). Similarly, the drug ISRIB (39), which ameliorates the short-term translational effects of eIF2α S51 phosphorylation by promoting eIF2B complex assembly (40), also showed no effect on elatol’s toxicity to OCI-Ly3 cells (Fig. 5C), despite strongly limiting its induction of ATF4 (Fig. 5D; Supplementary Fig. S5C). ISR induction by elatol therefore does not mediate its toxic effects and may in fact be a cytoprotective response to either elf4A1 ATPase inhibition or an off-target effect because MEFs lacking ATF4 were actually more sensitive.

In contrast to these results, knockdown of elf4A1 in NIH/3T3 cells, similar to methods employed previously to establish that elf4A1 is the key target of silvestrol (27), showed a significant shift in the elatol viability curve, similar to silvestrol [Fig. 5E and F; IC_{50} = 6.0 nmol/L (shRluc silvestrol), IC_{50} = 3.8 nmol/L (sh4A1 silvestrol)] P < 0.0001; IC_{50} = 3.547 nmol/L (shRluc elatol), IC_{50} = 2.382 nmol/L (sh4A1 elatol), P = 0.0026]. To confirm elf4A1 loss is not a general sensitizer to cytotoxic agents, we tested carboplatin in the same cells and found no change in drug sensitivity (Supplementary Fig. S5E). Manipulation of elf4A1 levels therefore adds to the toxic effects of elatol in a manner similar to the established elf4A1 inhibitor silvestrol, whereas multiple manipulations of the ISR pathway did not. Taken together with the in vitro and in vivo studies, binding and mutational studies, and effects on translationally regulated proteins, these data show elf4A1 functions as an elf4A1 inhibitor in cells, although with induction of an ISR for unclear reasons and effects on transcription at doses well above IC_{50}, making cellular off targets likely. elf4A1’s closely homologous but biologically distinct paralogs elf4A2 and elf4A3 are high-priority off targets based on structural similarity. Interestingly, we note in MDB-MB 468 cells that knockdown of either A2 or A3 results in strong induction of ATF4 (Supplementary Fig. S5E), but difficulties purifying these proteins from bacteria in a state that preserves their enzymatic activity have prevented cell-free assessment of elatol inhibition (see Discussion).

Antitumor activity and toxicity of elatol in vivo

Elatol has been given in vivo in only one prior study, resulting in antitumor activity against engrafted B16 murine melanoma cells, with the compound dosed at 10 mg/kg daily by intraperitoneal or oral administration (41). We tested elatol in vivo in nontumor-bearing SCID mice at 10, 20, and 30 mg/kg daily for 5 days intraperitoneally. Dosing at 30 mg/kg had to be stopped after 3 days due to weight loss and reduced animal activity, but neither 10 nor 20 mg/kg resulted in signs of toxicity (Supplementary Fig. S6A). Day 5 complete blood counts of the animals dosed at 20 mg/kg showed no differences from those of vehicle-dosed animals (Fig. 6A). We therefore treated SCID mice engrafted with SJ-DHL-6 flanking tumors at 20 mg/kg. We initially planned multiple 5-day cycles, but we found the first cycle resulted in weight loss and overt signs of toxicity, and treatment had to be discontinued (Supplementary Fig. S6B). The initial cycle halted tumor growth for 12 days from the start of treatment, relative to vehicle-treated animals but afterward tumors resumed enlarging (Fig. 6B). Elatol dosed at 20 mg/kg daily therefore halts tumor growth but was not well tolerated in tumor-bearing animals.

To optimize elatol dosing, we performed an MTD study in female CD-1 mice using a single-dose strategy with 5 mice per group (Fig. 6C). Group 1 tolerated 50 mg/kg without issue, but all 5 animals in group 2 died within five days of receiving 100 mg/kg. Dose reduction to 80 mg/kg (group 3) still showed unacceptable toxicity, with 3 of 5 mice dying or moribund requiring sacrifice within 5 days, but animals in group 4 tolerated 65 mg/kg without issue. Organ pathology on animals suffering toxicity in group 2 was compared with those that tolerated elatol without issue in group 1 (Fig. 6D; Supplementary Fig. S6C). Changes were seen in multiple organs, with damage to heart and/or liver considered...
most likely cause of morbidity/mortality. Pathology for animals treated at 65 mg/kg was similar to 50 mg/kg (not shown). Elatol’s MTD in nontumor-bearing animals is therefore 65 mg/kg, dramatically higher than for silvestrol, which is typically dosed at 0.2 to 0.5 mg/kg (4), potentially offsetting the potency difference between the compounds seen in vitro.

Figure 5.
Elatol induction of ATF4 is mediated by eIF2α phosphorylation, but toxicity is dependent on eIF4A inhibition. A, Western blot showing protein expression in eIF2α wild type and Ser51 A/A mutant MEFs treated with DMSO, 5 μmol/L elatol, or 5 μmol/L tunicamycin for 8 hours. B, Cell viability in MEF cells wild type or homozygous mutant for alanine at serine 51 of eIF2α and wild-type or homozygous knockout of ATF4 measured using CellTiter-Glo following 72-hour treatment with elatol. Mean ± SEM, n = 4, *, P < 0.05; **, P < 0.001. C, Western blot showing protein expression in cells treated for 4 hours with DMSO, 100 nmol/L silvestrol, 5 μmol/L elatol, or 5 μmol/L tunicamycin with or without the combination with 200 nmol/L ISRIB. Representative images, n = 3. D, Cell death measured by flow cytometry following Annexin V staining in DLBCL cell lines treated with indicated inhibitors with or without the combination of the 200 nmol/L ISRIB. Mean ± SEM, n = 3, *, P < 0.05; **, P < 0.001. E, Western blot showing eIF4A1 and eIF4A2 protein levels in NIH-3T3 cells stably expressing shRLuciferase (shRLuc) control or eIF4A1 shRNA transfected with the indicated proteins. Representative images, n = 2. F, Viability of NIH-3T3 cells stably expressing control or eIF4A1 shRNA measured by CellTiter-Glo after being treated with the indicated compounds for 5 days. Mean ± SEM, n = 4, *, P < 0.05; **, P < 0.001; ***, P < 0.0001.
We next engrafted OCI-Ly3 cells to SCID mice for additional assessment of elatol’s therapeutic window. Again, initial MTD for nontumor-bearing animals was not tolerated in tumor-bearing animals, with most engrafted animals experiencing morbidity and weight loss after a single dose at 65 mg/kg (not shown). Moving forward, we determined 40 mg/kg dosed twice per week was well tolerated and led to significant reduction of tumor growth compared with vehicle-treated controls ($P = 0.0117$).

**Figure 6.**
Elatol is well tolerated in mice up to 65 mg/kg and treatment slows tumor progression in vivo, but high doses show liver and cardiac toxicity. **A,** Complete blood counts for white blood cells, hemoglobin, and platelets of SCID mice following treatment with 20 mg/kg elatol daily for 5 days. Mean ± SEM, $n = 17$ and 7 vehicle and elatol, respectively. **B,** SCID mice were engrafted with $2 \times 10^6$ SU-DHL6 cells. Once tumors reached 60 mm$^3$, treatment began with 20 mg/kg i.p. daily for 5 days, and tumor volume was measured twice per week. Mean ± SEM, $n = 8$. **C,** MTD study in normal CD1 mice. Cohorts of 5 mice were given a single indicated dose of elatol and observed daily for signs of morbidity. At the time of death due to toxicity or by day 14 if no toxicities were observed, mice were sacrificed for organ pathology. **D,** H&E staining of the heart and liver of mice treated with 50 or 100 mg/kg elatol. Representative images, $n = 5$. **E,** SCID mice were implanted with $1 \times 10^6$ serially transplanted OCI-Ly3 xenograft cells. When tumors reached at least 50 mm$^3$, mice were pair matched and treatment began with either vehicle or 40 mg/kg of elatol i.p. twice weekly. Mean ± SEM, $n = 8$. *, $P < 0.05$. 

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Fig. 6E; Supplementary Fig. 5E) Elatol therefore shows proof of principle of a pipeline to identify inhibitors of the cap-dependent translation core enzyme eIF4A1 based on target-based cell-free 
in vitro screening for ATPase activity inhibition, taken through to 
in vivo therapeutic window.

Discussion

Target-based screening for small-molecule inhibitors of the DEAD-box RNA helicase eIF4A1 is a rational approach for discovery of novel cancer therapeutics for several reasons. First, translational activation is a nearly ubiquitous output of deregulated oncogenic signaling (16). Drugs targeting signaling molecules, typically kinases, have made substantial contributions to cancer therapeutic options, but short-lived responses in most clinical scenarios illustrate the ease with which messenger molecules may be bypassed by parallel or redundant signals (1, 2, 16). Translation, in contrast, particularly at the level of the eIF4F initiation complex, is a convergence point for these pathways, a bottleneck whose activation is necessary for transcriptional and epigenetic oncogenic outputs to be expressed as cellular phenotype. Both molecular heterogeneity of signaling leading to kinase-inhibitor resistance tracts and the vexing problem of tumor heterogeneity within individual patients may be bypassable with drugs against translation (42). Second, within the eIF4F complex, eIF4A1 has emerged as the most promising pharmacologic target. Innovative functional screening for interrupters of cap-dependent initiation revealed the initial natural compounds with this activator function. In addition, comparison of translatomes sensitive to inhibitors of eIF4A1's RNA-dependent ATPase activity can identifying compounds for further development but should clarify the reasons for elatol's mechanistic differences with silvestrol.

Importantly, using retroviral knockdown, we find increased sensitivity to elatol treatment upon loss of eIF4A1 similar to silvestrol and in contrast to the cytotoxic agent carboxatin. These results further establish that elatol is acting on eIF4A1 in a cellular context and that this is the cause of IRS-independent protein translation inhibition we observed. The eIF4A paralogs have increasingly well-characterized divergent biological roles. eIF4A1 is most strongly linked to the eIF4F cap-initiation complex, where it is necessary for efficient translation of most eukaryotic mRNAs (45). eIF4A2 can replace eIF4A1 in cell-free systems but is dispensable for cell survival (31). This factor is implicated instead in miRNA-mediated translational repression (46), although CRISPR deletion of the gene from murine fibroblasts showed it is not necessarily required for this (47). eIF4A3, meanwhile, is a critical component of the exon-junction complex and regulates nonsense-mediated mRNA decay (48). Because the three paralogs are so homologous at the amino-acid level, however, especially in their helicase cores, pan-inhibition by elatol could explain elatol's effects on ATF4 induction and could also mediate effects on mRNA expression, although as noted, we were unable to directly assess them as targets. Regardless, we believe evaluation of the specificity of eIF4A4 inhibitors for each paralog should become important steps for eIF4A1 inhibitor evaluation when technically feasible.

The eIF4A1 inhibitor space has stalled at a preclinical stage, and the only well-defined compounds have complex mechanisms and chemical structures. In over a decade, no derivative of silvestrol has made it to clinical evaluation, and its susceptibility to ABCB1-mediated drug efflux poses pharmacologic challenges (50). Elatol's interesting binding stoichiometry with eIF4A1 suggests immediate ideas for rational derivatization through covalent

typically dosed at 0.2 to 0.5 mg/kg. Silvestrol's effects on non-malignant host cells therefore may be significantly higher than elatol's, suggested also by silvestrol's 500× increased potency versus 3T3 fibroblasts in Fig. 5F. Overall, however, elatol is 5- to 10-fold less potent than silvestrol versus sensitive tumor cells in vitro, and is impotent against some lines that are silvestrol sensitive. A variety of mechanisms could account for these differences, and an intriguing question is whether elatol's novel mode of action against the target, being an ATPase inhibitor, is part of the reason. Alternately, elatol's off-target effects leading to induction of the cytotoxic PERK/UPR pathway, but otherwise, it is not clear from these experiments which eIF2α-phosphorylating kinase may be responsible. Importantly, however, we find elatol toxicity is not dependent on this in MEFs deficient for Atf4 or in breast cancer cells with ATF4 knockdown. Full interrogation of this question would be beyond our current scope, particularly because our initial derivatization efforts with elatol have not been successful raising questions about whether or not the compound is a good starting point for further development. We plan intensive additional screening using the approach we have established here, and identification of additional inhibitors with elatol's mode of action versus eIF4A1 may not only provide better lead compounds for further development but should clarify the reasons for elatol's mechanistic differences with silvestrol.

Published OnlineFirst May 29, 2018; DOI: 10.1158/1078-0432.CCR-17-3645
dimerization that might address some off-target effects, but as mentioned above, such efforts were not initially successful. Instead, we see the identification and characterization of elatol as laying the groundwork for a pipeline of novel eIF4A1 inhibitor discovery, including establishment of ATPase measurement as a basis for screening, along with multiple downstream steps for assessment of potency and specificity. Our initial screen was a relatively small one, assessing 500 natural compounds. With much larger scale efforts now underway, we hope to identify novel eIF4A1 inhibitors suitable for optimization with high antitumor potency to go after this promising target for cancer therapy.

Disclosure of Potential Conflicts of Interest

C.H. Benes reports receiving commercial research grants from Amgen, Alexza, and Novartis. No potential conflicts of interest were disclosed by the other authors.

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Acknowledgments

The technical assistance of Reynaldo Morales and Loreal del Valle during the isolation and purification of (+)-elatol is gratefully acknowledged. The authors wish to thank Ariel Ramirez Labrada for technical assistance in the OCI-Ly3 xenograft experiment. We thank Francesco Forconi, Isla Henderson, Ian Tracy, and Kathy Potter for kind help in collecting, storing, and characterizing CLL blood samples. Research reported in this publication/press release was supported by the NCI of the NIH under award number F30 CA021074 through the Experimental Mouse Shared Resource at the University of Arizona. J.H. Schatz was supported by startup funds from the University of Arizona and University of Miami Sylvester Comprehensive Cancer Center. A.D. Rodriguez was supported by NIH SC1 Program [grant 1SC1GM086271-01A1]. C.H. Benes was supported by The Wellcome Trust [grant 10926].

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Received December 7, 2017; revised April 20, 2018; accepted May 21, 2018; published first May 29, 2018.

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doi:10.1158/1078-0432.CCR-17-3645

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