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Yandong Zhang
Washington University School of Medicine in St. Louis

Anthony J. Saporita
Washington University School of Medicine in St. Louis

Jason D. Weber
Washington University School of Medicine in St. Louis

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p19ARF and RasV12 Offer Opposing Regulation of DHX33 Translation To Dictate Tumor Cell Fate

Yandong Zhang, Anthony J. Saporita, Jason D. Weber

BRIGHIT Institute and Department of Internal Medicine, Division of Molecular Oncology, Siteman Cancer Center, Washington University School of Medicine, St. Louis, Missouri, USA

DHX33 is a pivotal DEAH-box RNA helicase in the multistep process of RNA polymerase I-directed transcription of the ribosomal DNA locus. We explored the regulation of DHX33 expression by RasV12 and ARF to determine DHX33’s role in sensing these opposing signals to regulate ribosome biogenesis. In wild-type primary fibroblasts, RasV12 infection induced a transient increase in DHX33 protein level, as well as an rRNA transcriptional rate that was eventually suppressed by a delayed activation of the ARF/p53 pathway. DHX33 expression was exclusively controlled at the level of translation. ARF caused a dramatic reduction in polysome-associated DHX33 mRNAs, while RasV12 led to a complete shift of existing DHX33 mRNAs to actively translating polysomes. The translation of DHX33 by RasV12 was sensitive to inhibitors of phosphatidylinositol 3-kinase, mTOR, and mitogen-activated protein and was pivotal for enhanced rRNA transcription and enhanced overall cellular protein translation. In addition, DHX33 knockdown abolished RasV12-induced rRNA transcription and protein translation and prevented both the in vitro and in vivo transforming properties of oncogenic RasV12. Our results directly implicate DHX33 as a crucial player in establishing rRNA synthesis rates in the face of RasV12 or ARF signals, adjusting ribosome biogenesis to match the appropriate growth or antigrowth signals.

Canerous cells frequently harbor genetic mutations that activate oncogenes or inactivate tumor suppressors, leading to uncontrolled cell growth, evasion of apoptosis, and other cellular properties (1). To accommodate the rapid proliferation of cancer cells, several associated biological activities are also augmented in cancer cells (2). Recently, increasing evidence has shown that cancer cells often increase ribosome production to improve protein translation and cell growth (3–7). Ribosome biogenesis is frequently targeted by activated oncogenes and repressed by tumor suppressors (as reviewed in references 3 and 8). In fact, the link between nuclear hypertrophy and tumorigenesis was recognized more than 100 years ago (8, 9). More recent data indicate that a marked increase in rRNA synthesis is a general attribute of many cancers (9, 10), which is consistent with the idea that changes in rRNA synthesis may be prerequisite alteration in the progression to cellular transformation. The rate of cancer cell proliferation in tumors is directly proportional to nuclear size and RNA polymerase I (Pol I) activity, with overexpression of pre-rRNA correlating with poor prognosis in many cancers (10–13).

Ribosome biogenesis largely occurs in the nucleolus and is a highly coordinated biological process that includes rRNA synthesis, modification, processing, and assembly into ribosome subunits (10, 14–16). It is tightly controlled and directly linked to cell cycle events; defects in ribosome biogenesis often lead to apoptosis or cell cycle arrest (17–19). The initial step of ribosome biogenesis, ribosomal DNA (rDNA) transcription, is subject to numerous layers of regulation (20–22). Human rDNA contains >400 copies of the rRNA genes, organized in tandem arrays on five different human chromosomes. Initiation of rDNA transcription requires assembly of a specific multigene complex including Pol I and numerous associated proteins (3, 10). Two of these proteins are upstream binding factor (UBF) and the promoter selectivity factor, SL1/TIF-IB. Interaction of these two proteins at rDNA promoter leads to assembly of the preinitiation complex and subsequent transcriptional activation at the promoter (15, 23). Given its extreme importance in initiating ribosome biogenesis, rDNA transcription is greatly influenced by the Ras, Myc, and NPM oncogenes, as well as the ARF, p53, and PTEN tumor suppressors (14, 16, 24–29).

We previously identified the nucleolar DHX33 DEAH-box RNA helicase as an important mediator of RNA Pol I transcription through its interaction with UBF at rDNA loci following serum stimulation (30). In the present study, we explored the mechanism underlying DHX33 regulation. We now report that DHX33 is positioned at the crossroads of opposing Ras and ARF activities; oncogenic RasV12 stimulates but ARF represses translation of existing DHX33 mRNAs. In this manner we show that DHX33 is used as an endpoint of contrasting signals to set ribosome biogenesis rates. Using xenograft models and established Ras mutant cancer cell lines, we demonstrate that DHX33 accumulation is pivotal for RasV12 to initiate tumor formation.

MATERIALS AND METHODS

Cell culture. Wild-type mouse embryonic fibroblasts (MEFs), Arf−/− ear fibroblasts, Arf+/− MEFs, p53−/− MEFs, p53−/+ MEFs, Mdm2−/+ MEFs, and p53−/−; Mdm2−/+; Arf−/− MEFs were isolated from C57BL/6/Sv129 mixed mice. BxPC-3, Capan-2, Miapaca-2, and Panc-1 pancreatic cancer cell lines were kindly provided by Andrea Wang-Gillam (Washington University), Beas2B cells were provided by Gregory Longmore (Washington University), and H441 and A549 lung cancer cells were provided by Steven Brody (Washington University). BxPC-3 and H441 cells were

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Address correspondence to Jason D. Weber, jweber@dom.wustl.edu.

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grown in RPMI 1640 containing 10% fetal bovine serum (FBS) with antibiotics and supplements (10 mmol of HEPES, 4.5 g of glucose, 2 mmol of l-glutamine, and 1 mmol of sodium pyruvate/liter). A549 cells were grown in F-12 medium supplemented with 10% FBS and 1% penicillin-streptomycin. All other cells were maintained in Dulbecco modified Eagle medium (DMEM) supplemented with 10% FBS and 1% penicillin-streptomycin. All cell lines were incubated at 37°C with 5% CO₂ in a humidified incubator. U0126 and LY294002 were purchased from Sigma. Rapamycin was purchased from LC laboratories.

**Western blotting and antibodies.** Whole-cell lysates were prepared by incubation with 1 × NP-40 buffer that included 0.5% NP-40 and 1% sodium dodecyl sulfate (SDS) supplemented with Halt protease and phosphatase inhibitors (Sigma). Lysates were cleared by centrifugation and protein concentration was tested by DC assay (Bio-Rad). Lysates were boiled with SDS sample buffer, separated by SDS-PAGE, and transferred to polyvinylidene difluoride membrane (Millipore). Membranes were blocked in 5% nonfat dry milk TBS-T buffer (10 mmol of Tris-HCl [pH 7.4]/liter, 150 mmol of NaCl/liter, 0.1% Tween 20) and incubated in primary antibodies diluted in blocking buffer at 4°C overnight. Blots were washed with TBS-T buffer and incubated with horseradish peroxidase-conjugated secondary antibodies (1:10,000; GE Healthcare) in blocking buffer at room temperature. Immunocomplexes were visualized with an enhanced chemiluminescence kit (GE Healthcare). Primary antibodies for immunodetection were sourced as follows: anti-ARF (rat; Santa Cruz), antitubulin (goat; Santa Cruz), anti-AKT (Cell Signaling), anti-pS6 (Cell Signaling), anti-NF1 (Santa Cruz), anti-Ras (Santa Cruz), anti-p53 (Cell Signaling), anti-akt (Cell Signaling), and anti-pS473-AKT (Cell Signaling).

**Quantitative reverse transcription-PCR (qRT-PCR).** Primers were all designed by Primer Express 2.0 software and purchased from Integrated DNA Technologies. Total RNA was extracted by Nucleospin II (Clontech) RNA isolation kit and was reverse transcribed into cDNA by SuperScript III first-strand synthesis kit (Invitrogen). PCRs were performed on Bio-Rad C1000 thermal cycler and managed with Bio-Rad MyCycler. SYBR green (Clontech) was used and transcript quantification was performed by comparison with standard curves generated from dilution series of cDNA of human 47S rRNA (cloned in pCR2.1Topo). Melting curve analysis confirmed that single products were amplified, indicating that rRNA transcription might be a focal point of opposing activities from these two pathways caused DHX33 levels from 2 days until 6 days postinfection (Fig. 1D).

**Focus assay.** Human cancer cell lines were infected by pLKO.1 lentivirus encoding shScrambled RNA or shRNA to knockdown DHX33, and cells were selected by puromycin for 2 days. Cells were then plated at a density of 10⁴ per 100-mm dish and grown for 10 to 20 days. Colonies were washed with cold phosphate-buffered saline twice and fixed with 100% methanol for 10 min at room temperature. Colonies were then stained with Giemsa stain for 1 h at room temperature and washed with water before air-dried and photographed.

**Soft agar assay.** A total of 10⁴ cells were mixed in 4.0 ml of 0.3% agar–1 × DMEM–10% FBS as the top agar and plated into 60-mm plates with 4.0 ml of 0.6% agar–1 × DMEM–10% FBS as the base agar. Plates were incubated at 37°C and checked every 3 days, and the cells were fed with 2.0 ml of 0.3% agar–1 × DMEM–10% FBS every week. The colonies were photographed and counted 2 to 3 weeks later.

**Polysome profiles.** Cells (3 × 10⁴) after transduction with the indicated virus for 96 h were treated with 10 μg of cycloheximide/ml prior to harvesting and counting. Cells were subjected to cytoplasmic ribosome fractionation as described previously using sucrose density gradient system ranging from 7 to 47% (Tedelyne ISCO). Fractions were collected, and RNA was extracted with TRIzol and converted into cDNA with superscript reverse transcriptase III (Invitrogen) before quantitative PCR analysis with the appropriate primers.

**[3H]uridine pulse-chase labeling.** Cells were first infected with the indicated virus. At 3 days postinfection, the cells were replated at a confluence of 60 to 70% per plate. On the following day, the cells were then pulsed with [3H]uridine at a concentration of 2.5 μCi/ml for 30 min and chased with unradioabeled uridine at a concentration of 5 mM for the indicated time points. Approximately 2 × 10⁶ cells were pelleted, and the total RNA was isolated after dissolving cells in RNASolv (Omega Biotek, Norcross, GA). Formaldehyde RNA denaturing gel was run to separate different species of rRNA and then transferred to nylon Hybrid Plus membrane. After UV cross-linking and spraying with Enhancer (Perkin Elmer), the membrane was exposed to film and subjected to autoradiography.

**[35S]methionine incorporation.** Cells were starved in cysteine-methionine-free medium for 4 h and then pulsed with [35S]methionine (50 μCi/ml)-containing medium for 30 min before being harvested. The cells were lysed, and supernatants were precipitated with trichloroacetic acid at a concentration of 10%. Protein pellets were subsequently dissolved by 1% SDS and analyzed for protein concentration. The samples were then analyzed for radioactivity by liquid scintillation counting. The data presented were normalized based on equal amount of protein in each sample.

**Mouse xenografts.** Animals were handled according to protocols approved by the Washington University Animal Studies Committee. Nude mice were purchased from Jackson laboratories. Arf-null cells after transduction with the indicated virus were injected subcutaneously with 10⁶ cells into the flanks of mice. Tumors were dissected after 2 weeks and photographed.

**RESULTS**

**p19ARF induction during oncogenic stress lowers DHX33 protein levels.** ARF is the principal tumor surveillance protein charge with preventing aberrant cell growth and proliferation during oncogenic insult (18, 31). In wild-type primary fibroblasts, oncogenic RasV12 induces ARF protein expression, resulting in subsequent p53 activation and cell cycle arrest. Numerous tumor suppressors and oncoenes are known to influence the levels and activities of key molecules involved in rRNA transcription, suggesting that RNA transcription might be a focal point of opposing signaling moieties (3). Previously, we have shown that the DHX33 DEAH RNA box helicase is a novel regulator of rRNA transcription (30). To test whether p19ARF and RasV12 could affect DHX33 expression, wild-type (WT) MEFs were infected with either control, RasV12, or ARF-expressing retroviruses. DHX33 protein levels were analyzed at 2, 3, or 5 days postinfection. As shown, the ectopic expression of ARF resulted in a significant decrease in the DHX33 protein expression 2 days postinfection (Fig. 1A). Reduction of DHX33 by ARF continued through 3 days postinfection, when the DHX33 protein levels decreased 10-fold (Fig. 1B). In contrast, oncogenic stress by RasV12 infection resulted in slightly more DHX33 protein than control cells at 2 and 3 days postinfection (Fig. 1A and B). Noticeably, upregulation of ARF and p53 were only modest at these early time points following ectopic RasV12 expression. However, after 5 days of RasV12 infection, wild-type primary cells expressed significant levels of ARF with a resultant decrease in DHX33 protein (2-fold reduction) compared to control cells (Fig. 1C). These results indicate that two major pathways might regulate DHX33: ARF/p53 and oncogenic RasV12. The opposing activities from these two pathways caused DHX33 levels to be transiently increased and then significantly decreased. In accordance with this hypothesis, RasV12 infection of fibroblasts deficient in ARF (Arf-null MEFs) resulted in a robust increase in DHX33 levels from 2 days until 6 days postinfection (Fig. 1D). This is in contrast to a 2-fold reduction of DHX33 in WT MEFs in...
which endogenous ARF function is intact and induced during RasV12 infection. Thus, our results indicate that endogenous ARF is a key regulator of DHX33 expression during oncogenic stress.

We next performed quantitative real-time PCR (qRT-PCR) to determine ribosome RNA transcriptional rates by analyzing 47S pre-rRNA transcript levels in both RasV12 and ARF-infected wild-type MEFs. As shown in Fig. 1E, ARF infection resulted in the downregulation of rRNA transcription in a time-dependent manner. At 5 days postinfection, 47S rRNA levels dropped to 30% of that in the control sample. Moreover, RasV12 infection first resulted in a transient increase in pre-rRNA synthesis at 2 and 3 days postinfection (up to 2.5-fold), but after 5 days postinfection, after endogenous ARF induction, the pre-rRNA levels dropped to 70% of empty vector control (Fig. 1E). This trend is in agreement with the increase in DHX33 followed by its decrease over time after RasV12 infection.

ARF regulation of DHX33 is dependent on Mdm2 and p53. The ARF tumor suppressor has p53-dependent and -independent
functions (32). Wild-type MEFs maintain an intact p53 pathway downstream of ARF, suggesting that regulation of DHX33 by ARF in WT MEFs could be p53 dependent. To study whether this regulation occurs in a p53-dependent manner, we infected p53-null MEFs, p53−/−; Mdm2−/− MEFs, and p53−/−; Mdm2−/−; Arf−/− MEFs with ARF-expressing retroviruses. As shown in Fig. 2, ARF overexpression in WT MEFs resulted in a significant reduction in DHX33 levels. However, the reduction of DHX33 was far less significant in p53-null MEFs and DKO MEFs. In TKO MEFs, we observed no reduction in DHX33 protein levels, indicating that reduction of DHX33 by ARF requires p53 (Fig. 2B). In addition, we found that the infection of ARF in wild-type MEFs resulted in a much greater inhibition of ribosome RNA synthesis than in TKO MEFs (Fig. 2C). Our results indicate that ARF inhibits ribosome biogenesis not only in a p53-independent manner but also in a p53-dependent manner. Knockdown of endogenous ARF only mildly enhanced DHX33 protein expression in p53−/− MEFs (Fig. 2D), suggesting that p53 is required for DHX33 induction following loss of ARF.

**ARF reduces DHX33 protein levels through a translational control mechanism.** To dissect the mechanism of DHX33 reduction by ARF, we first analyzed DHX33 mRNA levels.
was performed on total RNAs isolated from ARF- and RasV12-infected cells at 2, 3, and 5 days postinfection. Both GAPDH mRNA and actin mRNAs were used as internal controls. We observed no significant change in DHX33 mRNA expression at each time point after ARF or RasV12 infection of WT MEFs compared to the empty vector control (Fig. 3A). These results indicate that reduction of DHX33 by ARF does not occur by transcriptional regulation. ARF has been previously shown to influence the stability of several proteins (33, 34). To determine whether DHX33 protein reduction was due to accelerated protein degradation upon ARF induction, cells were treated with MG132, a 26S proteasome inhibitor, for 6 h. As shown in Fig. 3B, we found that DHX33 was not stabilized in the presence of MG132. As a positive control, p21CIP1 was stabilized to a significant degree with MG132 treatment, demonstrating that MG132 is functioning as expected to inhibit 26S proteasome. These results imply that reduction of DHX33 in the presence of ARF is not due to accelerated protein degradation.

To determine whether DHX33 reduction was due to translational repression of existing DHX33 mRNAs, we chose to analyze polysome-associated DHX33 mRNAs. We performed a polysome fractionation by sucrose gradient after lysis of WT MEFs that were transduced with vector control or ARF overexpressing retroviruses (Fig. 3C). We analyzed the mRNA distribution of DHX33 in monosome and polysome fractions by qRT-PCR. As shown in Fig. 3D, we found that in ARF-infected WT MEFs, a large portion of DHX33 mRNAs (up to 60% of total mRNA) had moved into the monoribosome fractions. Conversely, empty vector-infected WT MEFs exhibited a majority of their DHX33 mRNAs associated with polysomes (70%). These data clearly in-
Indicate that ARF induction causes a translational repression of DHX33 in the cytoplasm.

DHX33 protein reduction decreases protein translation, while DHX33 overexpression enhances protein translation. Our previous data has shown that DHX33 is an important regulator of rRNA transcription; DHX33 knockdown reduced rRNA production, while DHX33 overexpression enhanced rRNA synthesis (30). In TKO MEFs, we manipulated DHX33 levels by utilizing lentivirus infection to knockdown (Fig. 4A) or overexpress (Fig. 4E) DHX33 protein. As shown in Fig. 4B, knockdown of DHX33 nearly abolished all rRNA production. Since rRNA is the key component for ribosome assembly, we hypothesized that DHX33 knockdown should result in less available ribosomes and thus decrease overall protein translation. mRNAs undergoing active translation are bound to multiple ribosomes, forming polysomes. The level of polysomes is widely regarded as an indicator of overall protein translational activity. Therefore, we performed cytosolic ribosome fractionation using sucrose gradients to monitor polysome levels. Strikingly, we noted a significant reduction of polysomes in TKO MEFs infected with DHX33 knockdown lentiviruses (Fig. 4D). The cytosolic 40S and 80S ribosomes were also decreased dramatically. Interestingly, the 60S ribosome peak was enhanced, indicating a different dynamic regulation of 40S and 60S, even though all of the rRNA species were decreased significantly (Fig. 4C).

Previously, we have found that wild-type DHX33 overexpression enhanced rRNA synthesis, while helicase-defective mutant of DHX33 (K94R) inhibited it (30). To determine the effect of DHX33 overexpression on cell growth, we transduced wild-type DHX33 and helicase-dead K94R mutant of DHX33 in TKO MEFs.
by lentivirus infection. Wild-type DHX33 only slightly enhanced 80S formation and polysome formation, whereas the K94R DHX33 mutant resulted in decreased levels of polysomes (Fig. 4E). This suggests that DHX33 is important for translation, but its overexpression might not be sufficient to significantly enhance protein synthesis. Western blot analysis showed levels of overexpressed wild-type and K94R mutant DHX33 (Fig. 4F).

**RasV12 stimulates DHX33 mRNA translation.** We previously showed that RasV12 expression caused a significant increase in DHX33 protein expression in Arf-null MEFs, implying that Ras hyperactivation regulates DHX33 levels in the absence of Arf. The neurofibromin (Nf1 gene) tumor suppressor protein is an upstream regulator of Ras signaling; loss of Nf1 results in irreversible activation of Ras and results in subsequent heightened growth and proliferation in vitro and in vivo tumor formation (35–37). We isolated MEFs from Nf1<sup>fl/fl</sup> mice and used adenovirus to overexpress Cre recombinase, resulting in the successful deletion of Nf1 alleles (Fig. 5A). Compared to the control Lac Z adenovirus-infected Nf1<sup>fl/fl</sup> MEFs, we found that DHX33 was upregulated 2.7-fold following Nf1 loss (Fig. 5A). As a confirmation for the activation of Ras signaling, we also detected increased levels of phospho-S6 and phospho-AKT (Fig. 5A).

In order to dissect the mechanism of DHX33 induction by Ras signaling, we first confirmed the activation of several conserved signaling events downstream of Ras. As shown in Fig. 5B, RasV12 expression in Arf-null MEFs induced activation of the mitogen-activated protein kinase (MAPK) pathway as indicated by increased phospho-ERK1/2 and activation of phosphatidylinositol

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**FIG 5** Ras activity induces DHX33 protein expression. (A) Nf1<sup>fl/fl</sup> MEFs were infected with adenoviruses encoding either LacZ or Cre recombinase at a multiplicity of infection of 200. At 2 days postinfection, the cells were then serum starved for 72 h. Equal amount of cell lysates were subjected to Western blot analysis with the indicated antibodies. (B) Arf-null ear fibroblasts from 2-month-old mice were infected with retroviruses encoding either pBABE empty vector (EV) or pBABE-RasV12. At 3 days postinfection, infected cells were subjected to Western blot analysis with the indicated antibodies. (C) Total RNA was isolated from the above-mentioned cells and changes of DHX33 mRNA levels were analyzed by qPCR with GAPDH as a control. (D) Arf-null ear fibroblasts were infected with empty vector or RasV12. At 3 days postinfection, the cells were treated with U0126 (20 μM), wortmannin (100 nM), or LY294002 (50 μM) for 24 h. Cell lysates were subjected to Western blot analysis with the indicated antibodies. (F) Arf-null cells infected with empty vector or RasV12 were treated with rapamycin, wortmannin, or LY294002 as indicated for 24 h. Cell lysates were prepared and analyzed for DHX33 protein levels with GAPDH as a loading control. (G) RasV12-null MEFs were infected with retroviruses encoding myristoylated Akt (Myr-Akt), RasV12, or empty vector. Cell lysates were prepared at 4 days postinfection after puromycin selection and analyzed by Western blotting for DHX33, pAkt-473, Akt, and GAPDH protein levels. The fold change is indicated below identified blots.
3-kinase (PI3K)/AKT pathway as indicated by phospho-AKT-S473, as well as activation of mTOR pathway by increased phospho-S6. To check whether upregulation of DHX33 in this setting was due to protein stability or mRNA level changes, we performed protein half-life assays. As shown in Fig. 5C, there was no significant change in DHX33 stability in empty vector or RasV12-transduced cells. Next, we analyzed mRNA levels of DHX33 by RT-PCR and found no significant change in DHX33 mRNA in RasV12-infected cells (Fig. 5D). To dissect the mechanism of DHX33 induction by Ras, we treated RasV12-infected Arf-null cells with PI3K/AKT or MAPK pathway inhibitors. Upregulation of DHX33 was completely abolished by the PI3K pathway inhibitors wortmannin and LY294002 but only partially by MEK inhibitor U0126 (Fig. 5E), demonstrating that Ras/PI3K is the main signaling pathway that regulates DHX33 protein induction. To determine whether DHX33 upregulation was controlled by mTOR activation, we treated cells with rapamycin. As shown in Fig. 5F, rapamycin inhibited the induction of DHX33 in a dose-dependent manner to a similar extent as wortmannin and LY294002, indicating that the Ras/PI3K/mTOR pathway is primarily responsible for upregulating DHX33 translation. To further confirm these results, we infected Arf-null MEFs with a constitutively active myristoylated Akt (Myr-Akt) retrovirus and found that activation of Akt alone was able to induce DHX33 protein levels but not to the levels seen in RasV12-infected cells (Fig. 5G).

We also analyzed DHX33 mRNA distribution on polysomes. As expected, RasV12 infection significantly enhanced production of cytosolic ribosomes and polysome formation (Fig. 6A). Approximately 70% of DHX33 mRNA was not associated with polysomes in Arf-null cells (Fig. 6B). In contrast, a majority (75%) of DHX33 mRNAs associated with polysomes in Arf-null cells that were infected with RasV12 retroviruses (Fig. 6B). As a control, GAPDH mRNA distribution was also analyzed and showed no significant difference between empty vector and RasV12 infection (Fig. 6B). This significant difference shows that DHX33 mRNAs are selectively translated upon RasV12 infection in the absence of Arf. To confirm that the Ras/PI3K/mTOR pathway indeed translationally regulates DHX33, we further treated the cells with rapamycin and analyzed DHX33 mRNA distribution on polysomes. As shown in Fig. 6C and D, rapamycin treatment resulted in a reduction of DHX33 protein levels and global protein translational repression. A significant proportion of DHX33 mRNA was shifted from polysomes to monoribosomes following rapamycin treatment (Fig. 6E).

DHX33 upregulation is required for enhanced rRNA transcription during Ras activation. We have previously reported that DHX33 is an important factor in rRNA transcription (30). We hypothesized that elevated levels of DHX33 during Ras activation are important for Ras to promote rRNA synthesis. To test this hypothesis, we first detected pre-rRNA transcript levels by qRT-PCR in both empty vector and RasV12-infected Arf-null cells and saw a 2- to 3-fold increase in 47S rRNA levels (Fig. 7A). To test whether DHX33 was required for this observed increase in pre-rRNA levels, Arf-null fibroblasts were first infected with RasV12 retroviruses, followed by a second infection with lentiviruses expressing knockdown shRNAs for DHX33 (Fig. 7B). We performed pulse-chase labeling with [3H]uridine to detect ongoing RNA synthesis. We found that reduction in DHX33 resulted in significantly lower 47S rRNA transcript levels that mirrored those seen in uninfected Arf-null cells (Fig. 7C). We also measured global protein synthesis by [35S]methionine incorporation into newly synthesized proteins for RasV12-transformed Arf-nulls cells after DHX33 knockdown and found that DHX33 knockdown caused a significant reduction in protein synthesis (Fig. 7D).

DHX33 is required in RasV12-initiated tumor formation. Given that we have shown a requirement for DHX33 in RasV12-initiated 47S rRNA transcription, we next sought to determine the contribution of DHX33 to RasV12-driven cellular transformation. Arf-null MEFs infected with RasV12 retroviruses were subjected to a second infection with shSCR or shDHX33 lentiviruses. DHX33 protein knockdown efficiency was analyzed by Western blotting (Fig. 8A). Importantly, DHX33 knockdown did not reduce DHX33 levels below those seen in control cells (Fig. 8A, lanes 1 and 4). After DHX33 knockdown, cells were plated in soft agar and grown for 2 weeks and resultant transformed cell colonies were counted. We observed a significant decrease in soft agar colonies in RasV12+shDHX33-infected cells, underscoring the importance of heightened DHX33 expression in RasV12-mediated cellular transformation (Fig. 8B). We next determined whether DHX33 knockdown influenced RasV12-initiated mRNA translation. Again, Arf-null MEFs were infected with RasV12 retroviruses and subjected to a second infection with lentiviruses encoding shRNAs for DHX33. Western blot analysis confirmed successful overexpression of RasV12 and knockdown of DHX33 (Fig. 8C).

We measured significant decreases in cytosolic ribosome subunits and actively translating polysomes in the RasV12+shDHX33 cells (Fig. 8D), indicating that elevated DHX33 expression is required for enhanced ribosome production and mRNA translation following ectopic RasV12 expression. To assess the impact of DHX33 knockdown on RasV12-initiated tumor formation, we injected 106-infected cells into the flanks of immunocompromised mice. At 2 weeks postinjection, we detected significant tumor cell growth of the cells infected with RasV12+shSCR, while cells infected with RasV12+shDHX33 did not exhibit any measurable tumor formation (Fig. 8E). This striking difference indicates that DHX33 is a crucial target of oncogenic RasV12 and is required to enhance RasV12-mediated cell growth and tumor formation.

Correlation between DHX33 protein levels, 47S rRNA levels, and cell proliferation in K-Ras mutated human cancer cell lines. Ras gene mutation has been frequently observed in human cancers (23). To determine whether endogenous DHX33 is upregulated in human cancers harboring mutant Ras alleles, we performed Western blot analysis for endogenous DHX33 protein levels on a panel of human cancer cell lines. As shown in Fig. 9A, we found elevated DHX33 protein levels in three of five K-Ras mutant cancer cell lines using wild-type K-Ras cell lines as a comparison. DHX33 protein levels were significantly upregulated in MiaPaCa-2, Panc-1, and A549 cells compared to wild-type K-Ras human cancer cell line, BxPC-3 or normal immortalized human lung epithelial cell line, Beas-2B (Fig. 9A). Due to the pivotal role of DHX33 in rRNA transcription, we also measured 47S rRNA levels by qRT-PCR in a panel of K-Ras mutated cancer cell lines. We discovered that 47S rRNA transcript levels correlated with DHX33 protein levels (Fig. 9B). For example, 47S rRNA transcript levels were the highest in MiaPaCa-2 cells, where DHX33 protein level was also the highest. Although in Capan-2, where DHX33 levels were the lowest, 47S rRNA level were also the lowest (Fig. 9B). Moreover, we also noticed that cell proliferation rates were tightly correlated with DHX33 protein levels and 47S rRNA levels in these K-Ras mutated human cancer cell lines (Fig. 9C).
To study the importance of DHX33 protein upregulation in K-Ras mutated human cancer cells, we utilized two unique shRNAs to knock down endogenous human DHX33 protein levels and measured cell growth over time. The knockdown efficiency of DHX33 for all five different cancer cell lines is shown in Fig. 9D. All cells exhibited some dependency on DHX33 for sustained proliferation (Fig. 9E). However, the negative impact of DHX33 on long-term proliferation was the most dramatic in the highly proliferative Miapaca-2 and A549 cells. The p53 mutational status might influence the different outcomes we observed for DHX33 knockdown. DHX33 knockdown in Miapaca-2 (mutant p53) resulted in significant cell death, while in p53 wild-type A549 cells, DHX33 knockdown resulted in a G2/M arrest (Fig. 9F). Taken together, our results show that elevated DHX33 protein expression in mutant K-Ras cancer cell lines is pivotal in enhancing rRNA transcription and proliferation.

**DISCUSSION**

Ras is one of the most frequently mutated oncogenes in human cancers. Three members of the Ras family, sharing 85% primary sequence identity, have been found to be activated in human cancers: H-Ras, N-Ras, and K-Ras (25). Up to 30% of human lung cancers harbor K-Ras mutations and, in pancreatic cancers, the K-Ras mutation rate is >90% (25). Ras signaling is a complex...
network whose downstream components play multiple roles in cell growth and cell proliferation. In its active, GTP-bound state, Ras is able to activate two major oncogenic signaling cascades: Raf/MEK/ERK and PI3K/AKT pathway (28). Aside from its role in promoting cell proliferation and cell survival, cell invasiveness and enhanced production of angiogenic factors, Ras activation also causes a significant elevation in the production of rRNA and increases in mRNA translation. Ras enhanced ribosome RNA synthesis are due to a variety of contributions from several Ras downstream effectors such as ERK (38–40), cyclin D1 (41, 42), and mTOR (43, 44), all of which can promote RNA Pol I transcription through different mechanisms. However, these enhancements and gains were observed in cells lacking an intact ARF/p53 pathway. The canonical roles of the ARF tumor suppressor reside in its ability to sense activated Ras alleles and prevent downstream cellular processes normally augmented by oncogenic Ras. Thus, it seems that proteins central to these processes must be under the control of both ARF and Ras regulators. Identifying these key players was our focus.

In this report, we identified a new downstream target of Ras, the DHX33 DEAH-box RNA helicase. DHX33 plays as an important role in promoting rRNA synthesis and ribosome biogenesis (30). In cells that maintain an intact Arf locus, oncogenic RasV12 overexpression resulted in a significant reduction in DHX33 protein expression without any lowering of DHX33 mRNA. The timing of DHX33 downregulation coincided with the classical induction of ARF expression by oncogenic RasV12 alleles. This negative regulation was not observed in cells lacking p53, arguing that the attenuation of DHX33 protein expression relied on the canonical ARF/p53 tumor suppressor pathway. Our results support the notion that other than cell cycle regulation, a p53-dependent role of ARF might also reside in inhibiting ribosome biogenesis. The ARF tumor suppressor has been found to inhibit rRNA synthesis (14, 24) through its ability to prevent UBF phosphorylation (24) and by translocating TTF-I, a RNA polymerase I termination factor, from the nucleolus into the nucleus (31). Nonetheless, in the absence of Arf, RasV12 was quite capable of dramatically increasing DHX33 protein expression, squarely placing DHX33 in the nexus...
of ARF and Ras regulation. In contrast to ARF, RasV12 considerably shifted existing DHX33 mRNAs onto translating polysomes. Thus, we have identified a new route through which ARF inhibits rRNA synthesis.

We have provided evidence that elevated expression of DHX33 is critical for Ras V12-induced cellular transformation. Importantly, our experiments utilized shRNAs that target and reduce DHX33 expression back to just above baseline. As such, we are not entirely removing DHX33 from these cells. Reduction of DHX33 in RasV12-expressing Arf-null cells resulted in a return of 47S rRNA and mRNA translation back to levels normally seen in Arf-null cells. These cells no longer grow in soft agar and do not form tumors in immunocompromised mice. Much of the focus on ARF tumor biology has been on its ability to respond to oncogenic signals, such as those emanating from RasV12, to induce a potent p53-dependent cell cycle arrest. Even more recently, a significant amount of interest has also shifted to ARF’s ability to directly inhibit ribosome biogenesis independent of p53. Our new find-

FIG 8 DHX33 induction is required for RasV12-initiated tumor formation. (A) Arf-null ear fibroblasts from 2-month-old mice were infected with retroviruses encoding pBABE-empty vector (EV) or pBABE-RasV12. Cells were then infected with lentiviruses encoding shScrambled (shSCR), shLuciferase (shLuc), or shDHX33. Whole-cell lysates were extracted and analyzed by Western blotting with Ras, DHX33, and tubulin antibodies. (B) A total of 5 × 10^5 infected cells were plated onto soft agar 60-mm plates in triplicate to measure anchorage-independent cell growth after 14 days. Quantitation of the colony numbers is presented from three representative fields under ×4 magnification. Error bars represent the standard deviation calculated from three different fields of colonies on triplicate plates. (C) Arf-null NIH 3T3 cells were infected with retroviruses encoding RasV12, followed by infection with lentiviruses encoding shScrambled (shSCR) or shDHX33. Whole-cell lysates were subjected to Western blot analysis with Ras, DHX33, and tubulin antibodies. (D) A total of 3 × 10^6 infected NIH 3T3 cells were subjected to cytosolic ribosome profiles. (E) The upper panel shows NIH 3T3 cells infected with retroviruses encoding RasV12 that were then infected with shSCR or shDHX33 lentiviruses. A total of 10^6 infected NIH 3T3 cells were injected into the flanks of nude mice. Tumor formation was visualized and photographed after 14 days. For the lower panel, mice were sacrificed at day 14 postinjection, and tumors were excised and photographed.
ings herein imply that ARF and Ras are in a constant struggle for
downstream target activation/inactivation. Normal cells harboring
activated \textit{Ras} alleles in the face of wild-type ARF are unable to
gain access to critical downstream targets, such as DHX33, to fully
activate critical processes required for tumorigenesis. ARF effect-
ively eliminates these proteins by removing their mRNAs from
actively translating polyribosomes. When \textit{Arf} is lost, Ras gains
access to these targets, and active translation of them ensues. How
ARF might selectively repress mRNA translation remains to be
investigated, but our findings with DHX33 are reminiscent of
ARF’s regulation of VEGFA translation (45).

Enhanced ribosome biogenesis is tightly correlated with en-
hanced cell proliferation in human cancers. Targeting RNA Pol I
transcription has been regarded as a potential treatment for cancer

\begin{figure}
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\caption{DHX33 is overexpressed in Ras-mutated cancer cell lines and is required for their efficient growth and proliferative properties. (A) A panel of K-Ras mutated or wild-type cancer cell lines (mutation status is shown at the bottom) were screened for total DHX33 protein expression, p14ARF status is also shown at the bottom. (B) 47S rRNA was measured by RT-PCR and normalized to total RNA levels. Error bars represent the standard deviation from three separate experiments. *, \(P < 0.001\). (C) A total of 5 \times 10^6 cells were plated onto six-well culture plates. Cell numbers were counted daily and graphed. The doubling time was calculated based on growth curves and is shown in the table. (D) The indicated cell lines were infected with lentiviruses encoding shScrambled (shSCR) or shDHX33. Whole-cell lysates were extracted 4 days postinfection and subjected to Western blot analysis with antibodies recognizing DHX33 and tubulin. (E) shSCR or shDHX33-infected cells (10^4) from indicated cancer cell lines were plated onto 100-mm culture dishes. The cells were fixed 10 or 20 days later with 100% methanol and incubated with Giemsa stain for 1 h. Stained colonies were air dried and photographed. (F) shSCR or shDHX33-infected cells (10^4) from Miapaca-2 and A549 cancer cells were subjected to cell cycle analysis by flow cytometry after propidium iodide staining.}
\end{figure}
patients (3). Several drugs, including actinomycin D, cisplatin, 5-fluorouracil, and camptothecin have been shown to inhibit RNA Pol I transcription (46–48). However, this field is still at the early stage and will require more selective targets in order to develop efficient therapeutic drugs that preferentially inhibit tumor growth while sparing normal cells. Recently, selective drugs that target rRNA synthesis have been developed, shedding a more positive light onto our ability to target RNA synthesis as a way of treating cancers. One of the latest compounds, CX-3543, is a small molecule nucleolus-targeting agent that selectively disrupts nucleolin/rDNA G-quadruplex complexes in the nucleolus (49). CX-3543 inhibited Pol I transcription and induced apoptosis in cancer cells and is currently in phase II clinical trials. Another compound, CX-5461, selectively inhibits Pol I-driven relative to Pol II-driven transcription, DNA replication, and mRNA transcription (50). CX-5461 inhibits the initiation stage of rRNA synthesis and induces both senescence and autophagy through a p53-independent process in solid tumor cell lines. Although more work needs to be done in order to develop efficient and more specific drugs to target RNA Pol I transcription as a way for cancer treatment, the validity of the approach itself has proven fruitful.

It seems uncertain whether other consensus targets of both ARF and oncogenic RasV12 exist. However, given the pleiotropic effects of ARF and Ras, identification of other common proteins seems likely. In fact, given the large number of RNA helicases in the DEAD/DEAH-box family, DHX33 may signal the first of many dually regulated helicases. Given our findings that the helicase activity of DHX33 was required for the RasV12-driven phenotype, generating novel compounds that inhibit its helicase activity seems likely. In fact, given the large number of RNA helicases in the DEAD/DEAH-box family, DHX33 may signal the first of many dually regulated helicases. Given our findings that the helicase activity of DHX33 was required for the RasV12-driven phenotype, generating novel compounds that inhibit its helicase activity seems likely. In fact, given the large number of RNA helicases in the DEAD/DEAH-box family, DHX33 may signal the first of many dually regulated helicases.

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