Non-coding rRNA-mediated preferential killing in cancer cells is enhanced by suppression of autophagy in non-transformed counterpart

CJ Hwang\(^1\), JR Fields\(^1\) and Y-H Shiao*\(^1\)

Interest to anticancer agents targeting rRNA biogenesis is growing. Cis-non-coding rRNAs, alternative to primary rRNA, have been shown to regulate rRNA biogenesis. We have recently detected bidirectional non-coding rRNAs that carry ribozyme-like properties. Anti-antisense oligonucleotides complementary to antisense non-coding rRNAs markedly stabilized the bidirectional transcripts and induced cell death in mouse lung cells. Here, we demonstrated that the same oligonucleotide killed mouse lung-cancer cells preferentially, compared with non-cancer sister lines, suggesting its potential utility for cancer treatment. A human version of anti-antisense oligonucleotide, complementary to an rDNA intergenic site, mediated apoptosis primarily in cancer cells. Autophagic activation was largely undifferentiated between the anti-antisense and other oligonucleotides and accounted for the undesired cytotoxicity in non-cancer cells. Co-treatment with chloroquine, an autophagy inhibitor, reduced cytotoxicity in the non-cancer cells, but retained the anti-antisense-mediated killings in cancer cells. Furthermore, the anti-antisense oligonucleotide stabilized bidirectional non-coding rRNAs predominantly in human cancer cells and perturbed rRNA biogenesis. Contributions of non-coding rRNAs to cell death were proven by transfection of in–vitro synthesized transcripts. Taken together, cancer/non-cancer cells respond differently to stabilization of non-coding rRNAs, and such differential responses provide a window of opportunity to enhance anticancer efficacy.

Cell Death and Disease (2011) 2, e239; doi:10.1038/cddis.2011.110; published online 8 December 2011

Subject Category: Cancer

Transcription of the rDNA, a tandemly repeated gene and highly conserved across unicellular and multicellular organisms,\(^1\) is coordinated by many proteins, including oncogenes and tumor suppressors.\(^2,3\) Phosphorylation of several transcription factors, such as UBF, TIF-IA, and TIF-IB/SL1, by kinases in EGFR signaling and mTOR-related pathways is essential for the activation of rDNA transcription. Proliferating cells elevate the demand of rRNA production and subsequent ribosomal assembly to support growth and survival. Increase of nucleolar organizer region in size or number, representing actively transcribed rDNA loci, has been a classical biomarker for tumor phenotype.\(^4,5\) The initial 47S transcript undergoes serial steps of rRNA processing to generate mature 18S, 5.8S, and 28S rRNAs, starting with excision of the Leader sequence from the 5′ external transcribed spacer (ETS) at about +650 nucleotide in mice and at the +414 site in humans.\(^6,7\) Many therapeutic drugs targeting rRNA biogenesis, either at transcription or processing step, are common regimens for anticancer treatment.\(^8\) It has been shown that cisplatin suppresses rRNA transcription by displacing UBF and RNA polymerase I to the periphery of the nucleolus\(^9\) and 5′-fluorouracil disrupts rRNA processing.\(^10,11\)

The rRNA transcription is also regulated by nucleolar-remodeling complex, which promotes gene silencing upon binding of an intergenic promoter rRNA of about 100–200 nucleotides in mouse fibroblast cells.\(^12,13\) Non-protein-coding rRNAs, other than rRNA, are ubiquitously expressed in 60–70% of mammalian genomes and frequently regulate the transcription of nearby primary transcripts.\(^14–16\) Of these transcription-rich clusters, over 20% in humans and about 72% in mice are expressed as sense–antisense pairs.\(^17–19\) The levels of sense and antisense non-coding RNA may be concordant, that is, concurrently upregulated or downregulated, or discordant.\(^16,19\) In the rDNA gene, we detected bidirectional cis-non-coding rRNAs (nc-rRNAs) in mouse lung epithelial cells and they followed a feed-forward or concordant mechanism to elevate sense/antisense nc-rRNAs upon introduction of antisense oligonucleotides and to perturb rRNA biogenesis (Supplementary Figure S1a). Oligonucleotides complementary to antisense nc-rRNAs were more potent than those pairing with sense nc-rRNAs to trigger cell death in mouse lung cancer cells (Supplementary Figure S1b). As stabilization of nc-rRNAs is concurrent with perturbation of rRNA biogenesis, this opens an opportunity to explore the potential of targeting nc-rRNAs for anticancer treatment. In this study, we applied antisense strategy to induce preferential cell death in mouse and human lung cancer cells. We determined the efficacy of cancer-selected
cell elimination among antisense oligonucleotides complementary to various regions of sense and antisense nc-rRNAs. Mechanisms of oligonucleotide-mediated cytotoxicity, including apoptosis and autophagy, were examined in human lung cells. Their relevance to preferential killings in cancer cells and reduction of cytotoxicity in non-cancer counterparts are discussed.

Results

Potent cancer cell inhibition by oligonucleotides pairing with antisense nc-rRNAs. Two antisense oligonucleotides, LNA1-S and LNA1-AS (Figure 1a), complementary to mouse antisense and sense nc-rRNAs, respectively, were first applied to test whether stabilization of nc-rRNAs provides any advantage to inhibit lung cancer cells predominantly. The cell viability was compared among treatments and was significantly reduced by LNA1-S, relative to either iFect vehicle control or LNA1-AS, in E10/E9 and C10/A5 pairs of non-cancer/cancer sister lines (Figure 1b). The degree of reduction, calculated by percentage of decrease in formazan intensity, showed that the E9 and A5 cancer lines are more sensitive to LNA1-S-mediated cell inhibition than their non-cancer counterparts, E10 and C10. This presents the evidence that stabilization of nc-rRNAs by targeting antisense nc-rRNAs is a promising strategy for anticancer treatment.

To extend the anticancer potential of nc-rRNA stabilization to humans, the subsequent study focused on human lung cells. Among several oligonucleotides (Figure 1a), the 86S complementary to the antisense nc-rRNA at −86 to −69 upstream from the transcription start site was mostly effective in reducing cell number in H441 and A549 cancer cells (Figure 2a and Supplementary Figure S2). Higher degree of reduction in cell density was appreciable in the 86S treatment, compared with 84AS (Figure 2a). The 86S was therefore chosen for further characterization of mechanisms that lead to its superior cell inhibition in reference to the 84AS or a non-specific scramble oligonucleotide control. The cytotoxic induction by 86S relative to 84AS increased as a function of the dosing concentration (Figure 2b). The same trend was observed using the non-specific oligonucleotide. An approximately 2-fold stronger killing efficiency by the 86S treatment over the 84AS was seen in H441 and A549 cancer lines at 30 and 40 nM. In contrast, similar killing efficiency between 86S and 84AS treatment, also visualized in Figure 2a, was observed in non-cancer HPL regardless of dosing concentration, likely a result of background stress response to oligonucleotides. Undesired cytotoxicity in non-transformed HPL following 86S or 84AS treatment prompted us to determine mechanisms of cell death that may lead to a better treatment strategy for killing human cancer cells predominantly.

Activations of apoptosis and autophagy and their relevance to anticancer efficacy. Small but significant increase of apoptosis was detected 24 h after oligonucleotide treatment, primarily in 86S-treated H441 and
A549 cancer lines, compared with non-specific scramble or 84AS oligonucleotide control (Figure 3). On the contrary, no elevation of apoptosis was detected in HPL non-cancer cells by the 86S treatment at this time point. The second dosing of 40 nM oligonucleotides did not enhance the degree of apoptosis established at 24 h in cancer cells, but the fraction containing fragmented non-apoptotic cells was enlarged in the 86S-treated H441 cells (Figure 4). Other cell death mechanisms might be involved. The significant increase of apoptotic fraction in non-specific oligonucleotide- or 84AS-treated cells, compared with iFect vehicle control, in single (24 h) and double (48 h) treatments suggests a minor degree of cytotoxicity resulting from the presence of oligonucleotide per se. There were not any consensus changes across three phases of cell cycle at 24 and 48 h following the 86S treatment (Supplementary Figures S3 and S4).

Apoptosis was further validated by upregulation of cleaved PARP and/or activated caspase 7, predominantly in 86S-treated cells (Figure 5a). H441 and A549 showed a steady apoptotic induction with increased levels of cleaved PARP and caspase 7 at 48 h. The timing matched to the apoptotic phenotype representing internucleosomal DNA breaks (Figures 3 and 4). In contrast, HPL non-cancer cells followed a very weak apoptotic response to the 86S, evidenced by null
to minimal changes of the internucleosomal DNA breaks (Figures 3 and 4) and cleaved caspase 7 level (Figure 5a) at 24 and 48 h. This suggests that apoptosis is not a major contributor for the undesired cytotoxicity in HPL, shown in Figure 2b. Also, changes of PARP levels in HPL did not correlate with internucleosomal DNA breaks, raising the possibility that PARP is not involved in apoptosis in non-transformed cells. The differential response to apoptosis between cancer and non-cancer cells further supports that cancer-selected cell destruction may be obtained through apoptotic activation by targeting antisense nc-rRNAs.

In addition to apoptosis, conversion of LC3B-I to LC3B-II protein, indicative of autophagy, was observed in 86S-treated cells (Figure 5a). The 84AS and/or non-specific oligonucleotides were also tested in some experiments and their apoptotic inductions were clearly less effective than the 86S treatment; however, the autophagic responses were mostly indistinguishable among oligonucleotides. On the contrary, the timing of autophagic activation relative to vehicle control varied among cells, and therefore the 86S-mediated autophagy was subsequently classified into steady (ocurrence at both 24 and 48 h) and weak response (occurrence at 48 h only). HPL and A549 showed steady response, whereas H441 followed a late or weak autophagic activation. The autophagy was confirmed by the presence of autophagosomes (Figure 5b).

Since autophagy has been implicated in both cell killing and survival, we next determined if co-treatment of oligonucleotide with chloroquine, an autophagy inhibitor, would enhance or suppress overall cytotoxicity. As shown in Figure 6, suppression of oligonucleotide-induced cytotoxicity by chloroquine co-treatment was demonstrated in HPL and A549 cells. The degrees of inhibition, calculated by relative reduction of killing efficiency after chloroquine co-treatment, ranged from about 40% in HPL to over 50% in A549, supporting that 86S and even non-specific oligonucleotide induce cytotoxicity largely by autophagy in these two cells. This is consistent with the steady autophagic response of HPL and A549 cells, measured by the conversion of LC3B-I to LC3B-II protein as described above (Figure 5a). In contrast, killing efficiencies of oligonucleotides in H441 were not reduced by the co-treatment with chloroquine, concordant with its weak response to autophagic activation represented by little changes in LC3B-II/LC3B-I ratio in Figure 5a. This implies that high dosing concentrations of oligonucleotide may
Figure 4  Induction of apoptosis after two transfections of 50 nm oligonucleotides at 0 and 24 h. Comparisons of apoptotic fractions, upper left (UL) and upper right (UR), among duplicate samples from each oligonucleotide treatment. \(^*\)P < 0.05, comparison of 84AS or NS with iFect vehicle control; \(^\dagger\)P < 0.05, \(^\ddagger\)P < 0.01, compared with 84AS; \(^\dagger\dagger\)P < 0.05, \(^\ddagger\ddagger\)P < 0.01, compared with NS. NS, nonspecific scramble oligonucleotide; x-axis, propidium iodide signal for DNA content; y-axis, fluorescein intensity in log scale for apoptosis.

Figure 5  Activation of apoptotic and autophagic pathways by the 86S. (a) Western blot analyses of apoptotic (cleaved PARP and cleaved caspase 7) as well as autophagic (conversion of LC3B-I to LC3B-II) markers in cells 24 or 48 h (2 transfections at 0 and 24 h) after initial transfection of 50 nm oligonucleotides. Apoptotic induction was represented by higher levels of cleaved PARP and/or cleaved caspase 7 relative to the iFect vehicle and/or untreated (UT) controls. (b) The ultrastructural signature of autophagosomes (arrows) was revealed under transmission scanning microscope. Autophagosomes were also detected in controls that did not receive oligonucleotides. Images were taken after two transfections of 50 nm oligonucleotides at 0 h and 24 h (scale bar: 2 \(\mu\)m). NS, nonspecific scramble oligonucleotide.
Stabilization of nc-rRNAs by the 86S oligonucleotide. The roles of nc-rRNAs in the above differential anticancer effects were explored next. A human sense nc-rRNA was reported early\textsuperscript{22} and the presence of an antisense nc-rRNA was confirmed by reverse transcription-polymerase chain reaction (RT-PCR) and sequencing here (nucleotides + 569 to + 887). Search in GenBank Expressed Sequence Tag database also identified many cDNAs showing > 95% homology to the rDNA in antisense orientation. The basal levels of nc-rRNAs and serial rRNA transcripts, that contain the Leader sequence, 5′ETS, 18S, and 28S regions during rRNA biogenesis\textsuperscript{7,23} were measured first to identify transcripts that may differentiate cancer from non-cancer cells. In comparison with non-transformed HPL, the levels of antisense nc-rRNAs and serial transcripts in rRNA biogenesis were in general higher in H441 and A549 cancer cells (Figure 7a), providing evidence that cancer cells acquire elevated levels of these two groups of rRNA species during tumorigenesis. The only exception is that the sense nc-rRNAs were higher in H441 but lower in A549, relative to HPL. Such variability of sense nc-rRNAs among cancer lines was reported before.\textsuperscript{22} Similar expression profiles between sparse and dense growth support that rRNA biogenesis remains steady in cancer and non-cancer cells even at high degree of cell–cell contact.

The transcriptional profiles after treatments with 86S and 84AS oligonucleotides were illustrated in Figure 7b. Stabilizations of both sense and antisense nc-rRNAs were detected, primarily in detached cells. The loss of viability for detached cells was validated by the retention of blue dye in trypan blue exclusion assay. The levels of upregulation for both nc-rRNAs were significantly higher in detached cancer cells induced by the 86S treatment, compared with 84AS. This provides the evidence that human bidirectional nc-rRNAs also follow a feed-forward mechanism, as seen in mouse lung cells, and are preferentially stabilized by the 86S targeting antisense nc-rRNAs. No difference between 86S and 84AS treatment in HPL suggests that increases of nc-rRNAs in non-transformed cells are induced simply by the presence of oligonucleotide per se, which may account for the undesired cytotoxicity (Figures 2b and 6). Strikingly, the levels of sense nc-rRNAs were upregulated 100- to 1000-fold in oligonucleotide-induced detached cancer cells, whereas only about 10-fold accumulation of the transcript were observed in detached HPL cells. It is likely that slight accumulation of nc-rRNAs is detrimental to non-transformed cells, whereas cancer cells tolerate high levels of the transcripts. Elevations of sense and/or antisense nc-rRNAs were sporadically detected in adherent cells and the extent was much lower than detached cells, possibly a result of increased transcripts from dying cells, which were not yet detached.

Concurrent stabilization of nc-rRNAs (Figure 7b) and induction of cell death (Figure 2b) by the 86S and 84AS treatments raised a possibility that sense and antisense nc-rRNAs may function as death mediators. Two lines of evidence support this hypothesis. First, perturbation of the rRNA processing, represented by changes of serial rRNA transcripts containing Leader sequence, 5′ETS, 18S, and 28S regions, was accompanied by upregulation of nc-rRNAs in 86S-and 84AS-induced detached cells (Figure 7b). Interference of rRNA biogenesis is known to trigger cell destruction and is a common strategy in selecting anticancer agents.\textsuperscript{24–26} Second, striking accumulations of nc-rRNAs were observed in detached cells induced by nutrient starvation, namely, no culture media change for over 1 week (Figure 7b). The rRNA biogenesis was also profoundly altered by the starvation treatment.

Induction of cell death directly by in –vitro-synthesized nc-rRNAs. To test the hypothesis that nc-rRNAs can mediate cell death directly, in –vitro-synthesized nc-rRNAs in sense and antisense orientations (Supplementary Table S1) were transfected into human lung cells. Initial trials of eight nc-rRNAs using transfection reagent as a control

---

**Figure 6** Differential suppression of oligonucleotide-induced cytotoxicity by chloroquine (CQ). Cell killing efficiency (KE) was calculated as ratio of averaged cytotoxicity in oligonucleotide-treated cells to the iFect vehicle control from duplicate samples in triplicate assays, measured 48 h after co-treatment of 25 μM chloroquine with single transfection of 40 nM oligonucleotides. # P < 0.05, comparison of NS with iFect vehicle control; ** P < 0.01, compared with NS. Error bars denote ± standard error. UT, untreated; NS, nonspecific scramble oligonucleotide.
detected elevation of cytotoxicity only for 7S (−244 to +203) in the sense direction and two antisense transcripts, 2AS (−1 to −244) and 8AS (+203 to −244). The remaining in vitro-transcribed nc-rRNAs in sense (−244 to −1, −409 to −248, and −709 to −486) and antisense (−248 to −409 and −486 to −709) orientations yielded cytotoxicity similar to the reagent control. The 7S, 2AS, and 8AS transcripts along with a control, 1S (nucleotide −244 to −1), were chosen for further examination under two different dosing conditions (Figure 8). Significant increases of cytotoxicity were readily detectable following transfection of several nc-rRNA transcripts at 1 pmol in HPL and A549. The high degree of cytotoxicity in HPL at 1 pmol recapitulates the susceptibility of HPL to cell detachment in the presence of 10-fold, not 100s to 1000s, accumulation of nc-rRNAs (Figure 7b). Majority of cytotoxic response in H441 began to appear after transfection with 5 pmol of the nc-rRNA transcripts. The requirement of higher dosing concentration to achieve significant cytotoxicity is consistent with the early observation that H441 maintains and tolerates abundant nc-rRNAs during basal

Figure 7 Levels of nc-rRNAs and rRNA processing transcripts before and after oligonucleotide treatments. (a) The basal levels of nc-rRNAs and serial rRNA transcripts representing a signature of rRNA processing were quantified by RT-PCR at two growth conditions in untreated cells (*P<0.05; **P<0.01; compared with HPL from duplicate samples). Error bars denote ± standard error. (b) Heat map of fold changes of nc-rRNAs and serial transcripts during rRNA biogenesis relative to adherent cells from iFect vehicle treatment after two transfections of 50 nM oligonucleotides at 0 and 24 h. The choice of the iFect to obtain fold changes of transcripts avoided the differential effects of the vehicle itself on nc-rRNAs among cells, where upregulation (sense in A549 and antisense in HPL) and downregulation (antisense in A549) were observed (Data not shown). The iFect alone did not alter both nc-rRNAs in H441. Color denotes over 2-fold change and/or P<0.05 (red: upregulation; green: downregulation); * P<0.05, compared with 84AS in corresponding adherent and detached cells.
growth (Figure 7a). Killing efficiencies of both HPL and A549 were enhanced at the 5 pmol dosing concentration as predicted, but the fold increases relative to corresponding treatments at 1 pmol were greatly enhanced in A549 only. The fold changes were 1.3, 1.5, and 1.9 in HPL and 2.5, 3.0, and 2.6 in A549 for 7S, 8AS, and 2AS treatments, respectively. The degrees of changes in killing efficiency from 1 pmol to 5 pmol of nc-rRNAs were relatively small in HPL, implying that further increases of nc-rRNAs would still induce cytotoxicity predominantly in cancer cells. This preferential killing in A549 cancer cells over HPL non-cancer line using in vitro-synthesized nc-rRNA transcripts reminisces the anticancer effects that are induced by stabilization of nc-rRNAs using the 86S oligonucleotide.

**Figure 8** Direct induction of cytotoxicity by in vitro-synthesized nc-rRNAs. Cell killing efficiency (KE) was calculated from averaged cytotoxicity in cells 48 h after single transfection of selected nc-rRNA transcripts at 1 and 5 pmol in 3 ml cell culture in reference to the TransIT transfection (TF) vehicle control from duplicate samples in triplicate assays. *P<0.05, comparison of 1S with TF; **P<0.01, compared with the 1S treatment. Error bars denote ± standard error. UT, untreated

**Discussion**

In this study, we demonstrated that cis-bidirectional nc-rRNAs are potential targets for cancer-specific therapy. The 86S oligonucleotide is more effective than the 84AS in the induction of apoptotic cell death in human lung cancer cells. This recapitulates the killing efficacy of targeting antisense nc-rRNA in mouse lung cancer cells. Although both oligonucleotides stabilized sense and antisense nc-rRNAs, the degree of upregulation was much higher and more steps of rRNA biogenesis were affected in cells committing to multiple death pathways following the 86S treatment, compared with 84AS. This provides evidence that antisense nc-rRNA is a promising target for drug discovery. Its stabilization may trigger massive increase of complementary sense nc-rRNA, similar to types of concordant transcripts described before. Also, cancer cells, relative to non-cancer counterpart, are prone to the stabilization of nc-rRNAs, which are proven to be cell death mediators by transfection of synthetic transcripts, supporting the idea that formulation of drugs to stabilize antisense nc-rRNAs may preferentially eliminate cancer cells. Such regimen is further benefited by co-treatment with chloroquine to reduce 86S-mediated autophagy in HPL non-cancer cells. Although antisense nc-rRNAs are upregulated to a lesser extent than sense transcripts in response to the 86S treatment, detection of cytotoxicity following direct introduction of synthetic antisense nc-rRNA suggests that the 86S exerts dual killing effects by sense and antisense nc-rRNAs.

Previously, an intergenic nc-rRNA at around –43 nucleotide region, also known as promoter rRNA, was detected in mouse and human cells and was shown to form a stem-loop structure essential for the binding of TIP5, a component of nucleolar remodeling complex (NoRC). Disruption of the stem-loop structure or ectopic expression of the nc-rRNA transcript promoted migration of the NoRC from nucleolus to nucleoplasm and upregulated the level of the 47S pre-rRNA that contains the Leader sequence. The 86S in the current study is mapped to the stem-loop core region. The increase of the 47S pre-rRNA following the 86S treatment might be initially interpreted as displacement of NoRC from the nucleolus by excessive sense nc-rRNA and subsequent relief of NoRC-mediated gene silencing, indicative of high demand for rRNA biogenesis. However, such a burst of rRNA transcript level is concomitant with cell death, suggesting that the increase of the 47S rRNA is a ramification of perturbation in the rRNA processing. This assumption is supported by a report finding disruption of rRNA processing in human cancer cells after expression of exogenous sense and antisense nc-rRNAs from the 5'ETS region. The alteration in the rRNA processing is further revealed by the changes of other pre-rRNA intermediates that contain 5'ETS, 18S, and 28S regions. It is likely that level changes of serial pre-rRNA transcripts during processing in detached cancer cells is a result of duplex formation with upregulated antisense nc-rRNAs or competition with excess sense nc-rRNAs. The efficacy of cancer-selected killing by the 86S, better than 84AS, advocates that continuous screening for extensive regions of antisense nc-rRNAs with oligonucleotides may identify additional potent anticancer targets. Besides the strategy to upregulate nc-rRNAs by oligonucleotides, in
theory, it is feasible to utilize the ribozymatic property of nc-rRNAs, demonstrated previously in mouse cells (Personal communication), for discovery of small-molecule ribozyme inhibitors.

Several agents known to interfere with rRNA transcription and processing are first-line anticancer therapy, such as cisplatin for advanced non-small-cell lung carcinomas and 5-fluorouracil for colon cancers. The list of agents inhibiting various stages of rRNA biogenesis continues to grow. The anti-nc-rRNA approach to selectively eliminate cancer cells, described in the current study, provides proof of principle to develop antisense oligonucleotides for anticancer treatment targeting rRNA biogenesis. This strategy may be also applied to other antisense and sense non-coding transcripts, reported to be differentially expressed at genome scale in cancers.

It is reasonable to believe that tyrosine kinase inhibitors, for example, gefitinib and erlotinib, and mTOR antagonists, such as temsirolimus, may also suppress rRNA biogenesis indirectly. The anti-nc-rRNA approach for anticancer treatment targeting rRNA biogenesis. This strategy may be also applied to other antisense and sense non-coding transcripts, reported to be differentially expressed at genome scale in cancers.

Materials and Methods

Cell lines and culture. Non-transformed peripheral lung cells of BALB/c mouse origin, C10 and E10, and corresponding spontaneously transformed A5 and E9 sister lines, respectively, were obtained and cultured, as previously described. HPL1D, a non-transformed line derived from human peripheral lung epithelium, was obtained from Drs. T Takahashi and A Masuda and maintained in a standard medium, as described before, with some modifications to replace fungizone and gentamicin. HPL and H441 cells were seeded in six-well plates at 2 × 10⁵ cells/well and A549 at 1 × 10⁵ cells/well. Cells were transfected with oligonucleotides (Figure 1a and Supplementary Table S1) using RNAiFect kit (Qiagen) according to the manufacturer’s instruction and following the formulation of 1:6 oligonucleotide (μg) to RNAiFect reagent (μl) ratio. The transfection efficiency was determined using the Label IT RNAi Delivery Control (Mirus, Madison, WI, USA) and was comparable among cell lines. For the co-treatment with chloroquine, 50 μl of 1.575 mM chloroquine in 1× phosphate-buffered saline solution was added immediately after transfection, to constitute a 25 μM final chloroquine concentration. In vitro-transcribed nc-rRNAs (1–5 pmol) were transfected in a six-well plate format using TransIT-mRNA Transfection kit (Mirus), according to the manufacturer’s protocol.

Cell viability, cytotoxicity, apoptotic DNA break, and cell cycle analyses. Cell viability was quantified by CellTiter 96 AQ one solution cell proliferation kit (Promega, Madison, WI, USA) converting a tetrazolium compound to formazan for cells cultured in a 96-well plate. The CytoTox 96 Non-Radioactive Cytotoxicity Assay kit (Promega) was used to measure lactate dehydrogenase (LDH) release, indicative of cytotoxicity. In brief, cells were transfected with oligonucleotides in 6-well plates as aforementioned. Three 50 μl aliquots of supernatent before and after direct addition of a 300 μl of Triton X-100 (9% v/v in water) lysis solution were transferred to 96-well plates at designated time points for each of duplicate samples. The intensity of formazan, produced from LDH reaction, was quantified at 492 nm after 30-min incubation. The level of cytotoxicity was expressed as relative density of formazan in supernatent to that of total lysate. Apoptosis as a result of internucleosomal DNA breaks and cell cycle fractions were quantified using the Apo-BrdU kit (Phoenix Flow Systems, San Diego, CA, USA) according to manufacturer’s manual. More than 10,000 stained cells were sorted and counted by flow cytometry. WinMDI 2.8 (http://www.cytop.duke.edu/flowcyt/software.htm) and CyChirei (http://www.facsabi. toxikologie.uni-mainz.de/engl.%20Websites/Downloads-engl.jsp) were used to calculate the fractions of apoptotic cells and 3 phases of cell cycle, respectively.

Transmission electron microscopy. In situ processing technique of cultured cells in 6-well plate for the transmission electron microscopy was described previously. Thin-sections (90 nm) were mounted on 150 mesh grids, stained in uranyl acetate and lead citrate, stabilized by carbon evaporation, and examined with an electron microscope (Hitachi, Tokyo, Japan) at 80 kV. Images were captured with a high resolution CCD camera (AMT, Danvers, MA, USA).

Western blot analysis. Cells seeded in 6-well plates were washed twice with phosphate-buffered saline (pH 7.4) and lysed with 100 μl Cell,ytic MT Mammalian Tissue/Lysis Extraction reagent (Sigma, St. Louis, MO, USA) with inclusion of 1× Complete Mini Protease Inhibitor Cocktail (Roche Applied Science, Mannheim, Germany). Protein concentration was measured using the BCA Protein Assay kit (Thermo Scientific, Wilmington, DE, USA). Proteins were resolved in 4–12% NOVEX NuPAGE Bis-Tris gel (Invitrogen) and electroblotted onto PVDF membranes (Bio-Rad) at 32 V for 1.5 h. Membranes were blocked for 2 h at room temperature with 5% non-fat powder milk in 0.01% Tween 20/phosphate-buffered saline (PBS-T) solution and then probed with specific primary rabbit antibodies against cleaved PARP (1 : 500 dilution), cleaved caspase 7 (1 : 500), LC3B and II (1 : 1000 for beta-Actin). After multiple washes with PBS-T, blots were incubated at room temperature for 1.5 h with donkey anti-rabbit IgG secondary antibody conjugated with horseradish peroxidase (1 : 5000 for PARP, caspase 7, and LC3B; 1 : 1000 for beta-Actin). After washes, blots were developed using a chemiluminescence ECL and ECL Plus kit (Amersham, Piscataway, NJ, USA) for beta-Actin/LC3B and PARP/caspase 7, respectively, and were exposed to X-ray film (Amersham). Beta-Actin served as a loading control.

In vitro transcription. The templates for in vitro transcription were generated by PCR using primer pairs listed in Supplementary Table S1. Transcription was carried out at 37 °C for 18 h using 0.5–1.0 pmol DNA template and MEGAscript T7 kit (Applied Biosystems, Carlsbad, CA, USA) and the transcripts were subsequently purified with MEGAClear kit (Applied Biosystems) according to the manufacturer’s instructions. The sizes and integrity of transcripts were verified in Tris-Urea polyacrylamide gel (Invitrogen).

Statistical analysis. Student’s t-test was applied to determine the differences of cytotoxicity, apoptosis, cell cycle fraction, nc-rRNAs, and serial transcripts.
of rRNA biogenesis among treatments of oligonucleotides or in vitro-transcribed no-RNAs. The differences between comparisons were considered significant if P-values were <0.05.

Conflict of Interest
The authors declare no conflict of interest.

Acknowledgements. We would like to thank Kunio Nagashima and Adam Hamed at Image Analysis Laboratory, SAIC-Frederick for transmission electron microscopy, Robin Steward at the Laboratory of Molecular Technology, SAIC-Frederick for deoxy ISO DNA sequencing, and Kathleen Noer and Gutily Mohammadi at Center for Cancer Research-Frederick Flow Cytometry Core for the flow analysis. We also thank Udo Rudloff for critical reading of the manuscript. This research was supported in part by the Intramural Research Program of the NIH, National Cancer Institute, Center for Cancer Research.

1. Long EO, Dawid IB. Repeated genes in eukaryotes. Ann Rev Biochem 1980; 49: 727–764.
2. Ruggiero D, Pandolfi PP. Does the ribosome translate cancer? Nat Rev Cancer 2003; 3: 179–192.
3. White RJ. RNA polymerases I and III, non-coding RNAs and cancer. Trends Genet 2008; 24: 622–629.
4. Derenzeni M, Trese D, Pession A, Montanaro L, Sirri V, Ochs RL. Nucleolar function and size in cancer cells. Am J Pathol 1998; 152: 1291–1297.
5. Montanaro L, Trese D, Derenzeni M. Nucleolus, ribosomes, and cancer. Am J Pathol 2008; 173: 301–310.
6. Gureyev JT. Characterization of mouse 45S ribosomal RNA subspecies suggests that the first processing cleavage occurs 600±100 nucleotides from the 5’ end and the second 500–550 nucleotides from the 3’ end of a 13.9 kb precursor. Nucleic Acids Res 1985; 13: 4905–4919.
7. Eichler DC, Craig N. Processing of eukaryotic ribosomal RNA. Prog Nucleic Acid Res 1994; 49: 197–239.
8. Drygin D, Rice WG, Grummt I. The RNA polymerase I transcription machinery: an emerging target for the treatment of cancer. Annu Rev Pharmacol Toxicol 2010; 50: 131–158.
9. Jordan P, Camo-Fonsca M. Cisplatin inhibits synthesis of ribosomal RNA in vivo. Nucleic Acids Res 1998; 26: 2831–2836.
10. Cohen MB, Glazer RL. Cytotoxicity and the inhibition of ribosomal RNA processing in human colon carcinoma cells. Mol Pharmauc 1985; 27: 308–313.
11. Ghoshal K, Jacob ST. Specific inhibition of pre-ribosomal RNA processing in extracts from the lymphosarcoma cells treated with 5-fluorouracil. Cancer Res 1994; 54: 632–636.
12. Mayer C, Schmitz KM, Li J, Grummt I, Santoro R. Intergenic transcripts regulate the epigenetic state of rRNA genes. Mol Cell 2006; 22: 351–361.
13. Mayer C, Neubert M, Grummt I. The structure of NoRC-associated RNA is crucial for targeting the chromatin remodeling complex Norc1 to the nucleolus. EMBO Rep 2008; 9: 774–780.
14. Okazaki Y, Furuno M, Kasukawa T, Adachi J, Bono H, Kondo S et al. Analysis of the mouse transcriptome based on functional annotation of 60,770 full-length cDNAs. Nature 2002; 418: 565–573.
15. Ota T, Suzuki Y, Nishikawa T, Otsuki T, Sugiyama T, Ike R et al. Complete sequencing and characterization of 21,243 full-length human cDNAs. Nat Genet 2004; 36: 40–45.
16. Faghihi MA, Wahlstedt C. Regulatory roles of natural antisense transcripts. Nat Rev Mol Cell Biol 2005; 10: 637–643.
17. Yelin R, Dahrari D, Sirek R, Leviyen EY, Goldstein O, Shoshan A et al. Widespread occurrence of antisense transcription in the human genome. Nat Biotechnol 2003; 21: 379–386.
18. Chen J, Sun M, Kent WJ, Huang X, Xie H, Wang W et al. Over 20% of human transcripts might form sense-antisense pairs. Nucleic Acids Res 2004; 32: 4812–4820.

Supplementary Information accompanies the paper on Cell Death and Disease website (http://www.nature.com/cddisis)