2-Selenouridine Triphosphate Synthesis and Se-RNA Transcription

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2-Selenouridine triphosphate synthesis and Se-RNA transcription

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ABSTRACT
2-Selenouridine (SeU) is one of the naturally occurring modifications of Se-tRNAs (SeU-RNA) at the wobble position of the anticodon loop. Its role in the RNA-RNA interaction, especially during the mRNA decoding, is elusive. To assist the research exploration, herein we report the enzymatic synthesis of the SeU-RNA via 2-selenouridine triphosphate (SeUTP) synthesis and RNA transcription. Moreover, we have demonstrated that the synthesized SeUTP is stable and recognizable by T7 RNA polymerase. Under the optimized conditions, the transcription yield of SeU-RNA can reach up to 85% of the corresponding native RNA. Furthermore, the transcribed SeU-hammerhead ribozyme has the similar activity as the corresponding native, which suggests usefulness of SeU-RNAs in function and structure studies of noncoding RNAs, including the Se-tRNAs.

Keywords: 2-selenouridine triphosphate; RNA transcription; RNA structure and function; hammerhead ribozyme; RNA crystallography

INTRODUCTION
RNA is involved in numerous biological processes, such as genetic storage, transcription, translation, and regulation (Watson 1963; Serganov and Patel 2007). Moreover, RNA can fold into well-defined three-dimensional structures to interact with proteins and catalyze biochemical reactions (Ponting et al. 2009; Schmeing and Ramakrishnan 2009). The appreciation for the uniqueness of RNAs, especially non-coding RNAs for their structure and function diversities, has increased extensively in the past decade. However, the functional understanding of these complicated macromolecules is often limited. The functional understanding of many natural modifications of the RNAs is even less. Thus, studying RNA natural modifications has become a very active research area in order to better understand biophysical and chemical properties of RNAs (such as tRNA and rRNA). So far, >100 RNA modifications have been discovered in nature (Dunin-Horkawicz et al. 2006), and many of them are frequently discovered in tRNA. 2-Selenouridine (2-SeU or 8-U) is one of naturally occurring nucleosides and exists at the wobble position of the anticodon loop in various bacterial tRNAs (Escherichia coli, Methanococcus vanni-elii, Clostridium sticklandii, etc.) (Ching et al. 1985; Dunin-Horkawicz et al. 2006). This Se-modification might play a critical role in the mRNA decoding process. It was hypothesized that the 2-Se-modification may enhance the accuracy and efficiency of protein translation (Lim and Curran 2001; Sun et al. 2012).

Moreover, another advantage of selenium modification in nucleic acid research is its assistance in addressing phase issue in X-ray crystallography via multiwavelength anomalous dispersion (MAD) or single-wavelength anomalous dispersion (SAD). Heavy atoms, such as selenium (Se) and bromine (Br), are suitable as anomalous scattering centers, which have been extensively applied in protein and nucleic acid crystallography. Encouraged by the successful selenium-assisted MAD phasing (Hendrickson et al. 1990; Ferre-D’Amare et al. 1998), we have pioneered and established nucleic acid X-ray crystallography with selenium derivatization (Carrasco et al. 2001; Caton-Williams and Huang 2008a; Sheng and Huang 2010; Lin et al. 2011b). Among the synthesized Se-derivatives, 2-selenouridine is stable and the only one found in nature so far. Furthermore, the single oxygen atom substitution with selenium at the exo-2 position doesn’t interfere with the hydrogen bonding in the Watson-Crick U/A base pair, thereby preserving the base-pairing function and structure (Sun et al. 2012). Therefore, the 2-selenouridine synthesis and its incorporation into RNAs may largely facilitate both structure and function investigations.

Generally, there are two strategies to synthesize the Se-derivatized RNAs: solid-phase synthesis, and transcription. The first method offers the site-specific incorporation of the Se-nucleoside. However, it is limited to relatively short RNAs (up to 50 nt) for large-scale synthesis. In addition, it requires multiple steps in deprotection and purification. The 2-selenouridine chemical incorporation into RNAs has been...
achieved via solid-phase synthesis (Sun et al. 2012). Our biophysical studies have shown that the 2-Se-modification discriminates against a U/G mispair (wobble pair), while preserving the native U/A pair. This result indicates that SeU can largely improve the RNA base-pairing specificity and the RNA–RNA interaction fidelity. This result has encouraged us to incorporate the Se-modification into RNA by in vitro transcription, in order to further investigate the function and structure of the SeU-RNAs, such as the SeU-containing tRNAs. This enzymatic method can allow synthesis of longer RNAs (>50 nt) in a large quantity (multiple milligrams). Multiple selenium atoms can also be conveniently incorporated into RNA under the mild conditions. As a matter of fact, the transcription strategy with T7 RNA polymerase is favored by most molecular and structural biologists. Herein, we report the first synthesis of 2-selenouridine triphosphate (SeUTP) and the enzymatic incorporation of SeUTP into noncoding RNAs. The active and mutant hammerhead ribozymes (Fig. 1) were successfully transcribed and examined with SeUTP. The transcribed SeU-hammerhead ribozyme is active, suggesting that the SeU-RNAs are useful in both function and structure studies of noncoding RNAs.

RESULTS AND DISCUSSION

In order to minimize by-product formation, the Se-nucleobase modifications are normally protected during chemical synthesis (Salon et al. 2007, 2008; Caton-Williams and Huang 2008b; Hassan et al. 2010; Sun et al. 2012). Since the 2-seleno-modification on uridine is naturally occurring, we decided to directly explore its compatibility with chemical synthesis. We were pleasantly surprised that 2-seleno-uridine, without protection, can be directly converted to the corresponding triphosphate. Thus, the synthesis (Scheme 1) of SeUTP (3) started from deprotection of the 5′-DMTr group of the Se-uridine derivative 1 (Sun et al. 2012) under an acidic condition. Then, 2-Se-uridine (2) was converted to SeUTP (3) via a one-pot synthesis: sequential treatments with phosphorus oxychloride (POCl₃), pyrophosphate, and bicarbonate (Yoshikawa et al. 1967; Caton-Williams and Huang 2008b).

After the synthesis, crude SeUTP (3) was precipitated from the reaction mixture and then purified by reverse-phase high-performance liquid chromatography (RP-HPLC). Purified SeUTP was characterized by MS, HPLC, UV, ¹H-, ¹³C-, and ³¹P-NMR. The profiles of HPLC and UV analyses are shown in Figure 2. Some characterization spectra are included in Supplemental Material (Supplemental Figs. S1–S4). To examine the SeUTP compatibility with RNA polymerase in transcription, the linearized plasmid templates for the wild-type hammerhead-ribozyme (WHR) and the crippled mutant hammerhead-ribozyme (MHR) were used (Fig. 1A) for the SeU-RNA transcription. As expected, SeUTP was recognized by T7 RNA polymerase (Fig. 3). Moreover, the mutant SeU-ribozyme (69-nt; containing 15 selenium atoms) was prepared via RNA transcription, and the integrity of the SeU-ribozyme (SeU-MHR) was confirmed by MS analysis (Fig. 3C). Under standard transcription conditions, the time-course transcriptions of both the native and SeU-modified mutant ribozymes were carried out (Fig. 3A). It was exciting to observe the incorporation of 15 selenium atoms into a RNA molecule by RNA transcription. To increase the SeU-RNA transcription yield, condition optimizations were carried out, such as Mg²⁺ concentration and buffer pH...
After optimization, the SeUTP transcription can reach up to 85% yield of the corresponding native RNA (Fig. 3B). This result suggested that SeUTP does not cause significant disruption of the polymerase catalysis. Detailed experimental conditions are included in Materials and Methods.

To examine whether the SeU-hammerhead ribozyme (Fig. 1A) can cleave itself, we transcribed both the native and Se-modified wild-type self-cleaving ribozyme. The transcription reactions (Supplemental Fig. S6B) were carried out in the standard transcription buffer (containing 6 mM MgCl₂). It was observed that the native ribozyme self-cleaved completely during transcription, while over 80% of the Se-modified ribozyme self-cleaved. When we used the transcription buffer containing 10 mM Mg²⁺, the SeU-ribozyme was self-cleaved completely. To further examine the SeU-ribozyme catalysis, a non-self-cleaving hammerhead ribozyme (35 nt in length) (Fig. 1B) was designed and transcribed with a double-stranded DNA (dsDNA) template (55 nt; dsDNA is more efficient in the SeRNA transcription than single-stranded DNA [ssDNA]). After transcription, the catalytic activity of the SeU-ribozyme was investigated by cleaving the 5'-32P-labeled RNA substrate (20 nt) (Fig. 1B). The Se-ribozyme was examined and compared with the corresponding native ribozyme (Fig. 5). Our results indicated that the SeU-ribozyme is active, and its activity is similar to the native one.

MATERIALS AND METHODS

The synthesis of 2-selenouridine triphosphate was performed under argon, all solvents were redistilled, and all reagents were dried under reduced pressure prior to use. All native NTPs, the transcription buffer, and T7 RNA polymerase used in our transcription experiments were purchased from Epicentre. The templates of the wild-type and mutant hammerhead ribozymes were from the linearized plasmids (Lin et al. 2011a).
To 5′-DMTr-2-Se-uridine (Scheme 1, step 1, 305 mg, 0.5 mmol) (Sun et al. 2012) dissolved in dichloromethane (5 mL), trifluoroacetic acid (11 mg) was added. The solution was heated at 40°C for 30 min, followed by adding methanol (0.2 mL). The reaction was stirred vigorously for another 1 h to obtain a light yellow precipitate product (Scheme 1, step 2). The precipitate was recovered by centrifugation or filtration; the yield of step 2 was almost quantitative. 2-Se-uridine (Scheme 1, step 2, 20 mg) was weighed and dried in a standard procedures from the manufacturer, Epicentre (AmpliScribe T7-Flash Transcription Kit). ATP, UTP, GTP, and TTP (0.5 mM each T7 pol.) was purified by HPLC (Ultimate XB-C18, 250 mm × 4.6 mm, 5 μm) with a gradient from 100% buffer A (20 mM triethylammonium acetate in water) to 40% buffer B (20 mM triethylammonium acetate in 50% acetonitrile and 50% water) for 15 min. The HPLC and UV profiles are shown in Figure 2. The retention times of the native UTP and SeUTP were 11.2 and 14.1 min, respectively.

**HPLC and UV analyses of SeUTP**

The maximal UV absorbance of native uridine triphosphate is 260 nm, while that of the SeUTP-triphosphate is 307 nm. In the HPLC analysis, both the native and selenium-modified UTPs were monitored under two wavelengths (260 and 307 nm). The synthesized SeUTP was purified by HPLC (Ultimate XB-C18, 250 mm × 21.2 mm, 10 μm) with a gradient of 100% buffer A (20 mM triethylammonium acetate in water) to 25% buffer B (20 mM triethylammonium acetate in 50% acetonitrile and 50% water) for 20 min. The HPLC analysis was performed (Ultimate XB-C18, 250 mm × 4.6 mm, 5 μm) with a gradient from 100% buffer A (20 mM triethylammonium acetate in water) to 40% buffer B (20 mM triethylammonium acetate in 50% acetonitrile and 50% water) for 15 min. The HPLC and UV profiles are shown in Figure 2. The retention times of the native UTP and SeUTP were 11.2 and 14.1 min, respectively.

**Transcription of the RNAs with native NTPs**

The native RNAs were transcribed with the transcription protocol (final concentration) in RNA polymerase buffer (40 mM Tris-HCl, 6 mM MgCl₂, 2 mM spermidine, pH 7.9), DTT (10 mM), ATP, UTP, CTP, and GTP (0.5 mM each T7 pol.), DNA template (non-self-cleaving hammerhead ribozyme: 1 μM dsDNA template [55 nt]; self-cleaving hammerhead ribozyme [mutant and wild-type]: 50 ng/μl linearized plasmid), T7 RNA polymerase (2 units/μL, Epicentre), and RNase-free water to adjust to the final volume (e.g., 20 μL). The transcription reaction was incubated for 1 h at 37°C.

**Transcription and analysis of the Se-RNAs**

The transcription experiment was carried out by following the standard procedures from the manufacturer, Epicentre (AmpliScribe T7-Flash Transcription Kit). α-32P-ATP was used as the radioactive labeling material for transcription experiments. Each transcription reaction (5 μL) contained ATP, CTP, GTP, and UTP or SeUTP (0.5 mM each), linearized plasmid DNA template (50 ng/μL), DTT (10 mM), transcription buffer (1×) for T7 RNA polymerase, T7 RNA polymerase (10–20 units), and RNase-free water. In the time-course experiments, a gel-loading dye (5 μL) containing 100 mM EDTA was used to quench the reaction at each time point, followed by denaturing PAGE (15% gel) analysis and autoradiography. The translated material for transcription experiments. Each transcription reaction (5 μL) contained ATP, CTP, GTP, and UTP or SeUTP (0.5 mM each), linearized plasmid DNA template (50 ng/μL), DTT (10 mM), transcription buffer (1×) for T7 RNA polymerase, T7 RNA polymerase (10–20 units), and RNase-free water. In the time-course experiments, a gel-loading dye (5 μL) containing 100 mM EDTA was used to quench the reaction at each time point, followed by denaturing PAGE (15% gel) analysis and autoradiography. The translated material for transcription experiments. Each transcription reaction (5 μL) contained ATP, CTP, GTP, and UTP or SeUTP (0.5 mM each), linearized plasmid DNA template (50 ng/μL), DTT (10 mM), transcription buffer (1×) for T7 RNA polymerase, T7 RNA polymerase (10–20 units), and RNase-free water. In the time-course experiments, a gel-loading dye (5 μL) containing 100 mM EDTA was used to quench the reaction at each time point, followed by denaturing PAGE (15% gel) analysis and autoradiography. The translated material for transcription experiments. Each transcription reaction (5 μL) contained ATP, CTP, GTP, and UTP or SeUTP (0.5 mM each), linearized plasmid DNA template (50 ng/μL), DTT (10 mM), transcription buffer (1×) for T7 RNA polymerase, T7 RNA polymerase (10–20 units), and RNase-free water. In the time-course experiments, a gel-loading dye (5 μL) containing 100 mM EDTA was used to quench the reaction at each time point, followed by denaturing PAGE (15% gel) analysis and autoradiography. The translated material for transcription experiments. Each transcription reaction (5 μL) contained ATP, CTP, GTP, and UTP or SeUTP (0.5 mM each), linearized plasmid DNA template (50 ng/μL), DTT (10 mM), transcription buffer (1×) for T7 RNA polymerase, T7 RNA polymerase (10–20 units), and RNase-free water. In the time-course experiments, a gel-loading dye (5 μL) containing 100 mM EDTA was used to quench the reaction at each time point, followed by denaturing PAGE (15% gel) analysis and autoradiography.
RNAs were WHR and MHR (Fig. 1). The transcribed mutant Se-hammerhead ribozyme with incorporated 15 selenium atoms was confirmed by MALDI-TOF MS analysis (Fig. 3C).

The Se-RNAs were transcribed with the transcription protocol (final concentration) in RNA polymerase buffer (40 mM Tris-HCl, 12 mM MgCl₂, 2 mM spermidine, pH 7.5), DTT (10 mM), ATP, SeUTP, CTP, and GTP (0.5 mM each NTP), DNA template [non-self-cleaving hammerhead ribozyme: 1 μM dsDNA template (55 nt); self-cleaving hammerhead ribozyme (mutant and wild-type): 50 ng/μL linearized plasmid], T7 RNA polymerase (4 units/μL). The cleavages of the RNA substrate by the native and Se-modified ribozymes (y-axis) were normalized via comparison to the substrate cleavage by the native ribozyme at 150 min (defined as 1.0).

Catalytic activity analysis of the Se-RNAs

The non-self-cleaving hammerhead ribozyme (5‘-GGCA-ACCUGAUGAGCCGAAGGCCGAAACGUACA-3′) (Fig. 1B) for the catalytic experiments was transcribed following the standard procedures described above. The DNA template used for this transcription was a 55-nt dsDNA (5′-TGTAGCCTTCGGCGCTTGATCATCAGG TGTCTATATGAGTCGTATACGC-3′ and its complementary sequence). After the transcription, the native and Se-modified ribozymes were purified and adjusted to the same concentration (monitored by UV). The RNA substrate (20 ng, 5′-ACCUGUAACGGCUUGCCUAA-3′) (Fig. 1B) chemically synthesized by solid-phase synthesis was kinased with γ-32P-ATP at the 5′ end for the ribozyme digestion. The digestion was performed in the buffer (10 mM Tris-HCl, 10 mM MgCl₂, pH 7.6) and with 5′-32P-labeled RNA substrate (final concentration: 50 μM) at 27°C. Aliquots (10 μL each) were taken at the time intervals (0, 5, 10, 30, 90, and 150 min), and each was mixed with EDTA (5 μL, 50 mM) dissolved in a saturated urea solution (aqueous) to quench the digestion. The 5′-labeled RNA substrate was digested to the 9-nt fragment and the 5′-32P-RNA fragment (11 nt). The 32P-labeled RNA allowed monitoring the substrate digestion via gel electrophoresis and autoradiography. The time-course results of the ribozyme digestion are shown in Figure 5 and Supplemental Fig. S6A.

Optimization of SeU-RNA transcription

To maximize the transcription yield, condition optimizations have been performed. The linearized plasmid of the mutant hammerhead ribozyme (Fig. 1A) was used as the template, which incorporates 15 SeUs into the ribozyme. The transcription buffers with various pH values were first examined, since the acidity of the imino group (3-NH) of U is higher than that of the native U (Sun et al. 2012). The pH values of the transcription buffer (40 mM Tris base or sodium phosphate, 6 mM MgCl₂, 2 mM spermidine, and 10 mM DTT) were adjusted. The Se-RNA transcription was examined under eight pH values (pH 5.5, 6.0, 6.5, 7.0, 7.5, 8.0, 8.5, and 9.0) and indicated that pH 7.5 was optimal for the Se-RNA transcription (Fig. 4). The pH of the standard transcription buffer is 7.9. Mg²⁺ concentration in the transcription buffer was also examined by varying it from 0 to 12 mM. As the increased MgCl₂ concentration yielded higher transcription yield (Supplemental Fig. S5A), 12 mM MgCl₂ was chosen for the Se-RNA transcription. Other components, such as spermidine (from 0.5 to 1.5 mM), were also examined for the transcription optimization. However, we found that increases of the concentrations of these components slightly decreased the transcription yield (Supplemental Fig. S5B). Moreover, a higher quantity of T7 RNA polymerase can increase the Se-RNA transcription yield (Supplemental Fig. S5C). Finally, after combining these optimized conditions (Table 1), we could increase the yield of the SeU-RNA transcription up to 85% of the corresponding native RNA (Fig. 3B), and these conditions have been used to transcribe various SeU-RNAs.

Thermostability of the SeU-RNA

To examine the thermostability of the SeU-RNA, we designed a short Se-RNA (trimer: 5′-USeU-3′) for this study. This Se-RNA was chemically synthesized by solid-phase synthesis and purified (Sun et al. 2012). We heated the Se-RNA continually at 70°C for a few hours and monitored it by HPLC at both 260 and 307 nm, since the 2-seleulmination modification has a unique UV-absorption at 307 nm, while the native nucleotides absorb strongly at 260 nm. The HPLC analysis was performed (Ultimate XB-C18, 250 mm × 4.6 mm, 5 μm) with a gradient from 100% buffer A (20 mM...
triethylammonium acetate in water) to 40% buffer B (20 mM triethylammonium acetate in 50% acetonitrile and 50% water) for 15 min. No significant decomposition was observed over 4-h heating at 70°C (Fig. 6), indicating that this Se-modification is relatively stable.

SUPPLEMENTAL MATERIAL

Supplemental material is available for this article.

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FIGURE 6. Thermostability study of 5\textsuperscript{46}U-RNA. 5’-\textsuperscript{46}U\textsubscript{5}U-3’ was heated at 70°C for several hours. HPLC was monitored at both 260 and 307 nm (retention time: 10.9 min).
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