METHOD

Rapid purification of RNAs using fast performance liquid chromatography (FPLC)

INSIL KIM,1 SEAN A. MCKENNA,1 ELISABETTA VIANI PUGLISI,1 and JOSEPH D. PUGLISI1,2
1Department of Structural Biology, School of Medicine, Stanford University, Stanford, California 94305, USA
2Stanford Magnetic Resonance Laboratory, School of Medicine, Stanford University, Stanford, California 94305, USA

ABSTRACT

We present here an improved RNA purification method using fast performance liquid chromatography (FPLC) size-exclusion chromatography in place of denaturing polyacrylamide gel electrophoresis (PAGE). The method allows preparation of milligram quantities of pure RNA in a single day. As RNA oligonucleotides behave differently from globular proteins in the size-exclusion column, we present standard curves for RNA oligonucleotides of different lengths on both the Superdex 75 column and the Superdex 200 size-exclusion column. Using this approach, we can separate monomer from multimeric RNA species, purify the desired RNA product from hammerhead ribozyme reactions, and isolate refolded RNA that has aggregated after long-term storage. This methodology allows simple and rapid purification of RNA oligonucleotides for structural and biophysical studies.

Keywords: RNA; FPLC; purification; NMR; X-ray crystallography

INTRODUCTION

RNA oligonucleotides are typically prepared by in vitro transcription from a DNA template with T7 RNA polymerase (Milligan et al. 1987; Puglisi and Wyatt 1995). After transcription, denaturing polyacrylamide gel electrophoresis (PAGE) is the most commonly used method to separate the desired RNA product from transcription byproducts (Wyatt et al. 1991). While this approach has been successful, it has several disadvantages (Cheong et al. 2004; Kieft and Batey 2004; Lukavsky and Puglisi 2004). First, this lengthy protocol requires multiple days to purify a single RNA sample. According to this method, after denaturing PAGE, the desired RNA is eluted from a gel matrix, concentrated, equilibrated in the desired buffer, and refolded. Water-soluble acrylamide oligomers, which are contaminants from PAGE, are difficult to remove from RNA samples (Lukavsky and Puglisi 2004). Moreover, since RNA is precipitated after the transcription reaction and purified under denaturing conditions, larger RNAs irreversibly aggregate and subsequently up to 90% of the RNA can be lost (Lukavsky and Puglisi 2004).

To overcome these disadvantages in PAGE purification, we previously reported a method using size-exclusion chromatography without requiring the denaturation of RNA (Lukavsky and Puglisi 2004). This approach produced acrylamide-free RNA samples that are suitable for structural studies. However, the protocol was not ideal. First, we found that concentration after phenol/chloroform extraction is not only a time-consuming step, but can also result in RNA aggregation and sample loss. Second, the open column used in the previous method is time consuming and low resolution. Here, we use size-exclusion chromatography columns in the FPLC system, which allows rapid purification of homogeneous RNA samples. Milligram quantities of pure RNA are isolated from transcription and ready for structural studies in <1 d. Another advantage of size-exclusion columns in the FPLC system is their high resolution. The previous size-exclusion method employed open-column size-exclusion chromatography in which the matrix was chosen to separate between plasmid DNA and all other RNA species (Lukavsky and Puglisi 2004). By choosing appropriate size-exclusion columns, we demonstrate high-resolution separation of RNA oligonucleotides of variable sizes. The approach outlined should have broad application in the structural and biophysical investigations of systems involving RNA components.
RESULTS AND DISCUSSION

High-throughput RNA sample preparation

We describe a simple and rapid RNA purification scheme using an FPLC system. Milligram quantities of pure RNA were prepared from transcription in <1 d. As a template for in vitro transcription, we employed plasmid DNA derived from pUC119, which is prepared using the QIAGEN Plasmid Maxi kit (QIAGEN, Inc.). After restriction enzyme (BstZ17I or BsaI) treatment, the linearized plasmid template is used for in vitro transcription using T7 polymerase under standard conditions reported previously (Lukavsky and Puglisi 2004). After 2–3 h of incubation at 37°C, pyrophosphate precipitate is pelleted (3000 g 3 10 min) and the reaction quenched by adding ethylenediaminetetraacetic acid (EDTA). An equal volume of phenol/chloroform is added to the supernatant, mixed, and centrifuged (3000 g 3 10 min). In this process, the restriction enzyme, which is carried over with the linearized plasmid DNA, and the T7 RNA polymerase are removed from the reaction mixture, as only the aqueous phase is retained.

After three phenol/chloroform extractions, the aqueous phase (upper) is directly loaded on a desalting column (10DG column, Bio-Rad) to remove phenol and a significant amount of remaining NTPs. The desalting column is equilibrated with 20 mL of 10 mM phosphate buffer (pH 6.5) and 100 mM NaCl. After application to a desalting column, the eluted transcription reaction is directly loaded on a size-exclusion column without requiring any other treatments (i.e., precipitation or concentration). A summary of the methodology is presented in Figure 1.

An example application of this method is shown for a 60 nucleotide (nt) RNA molecule (HIV TAR). A 5 mL transcription reaction was performed, and the phenol/chloroform-extracted RNA was passed over a desalting column. The desalted RNA fraction was collected and loaded on a Superdex 200 (26/60) column directly. Since we observed that concentration using centrifugation devices accelerates aggregation, we avoid concentrating samples at this stage. The Superdex 200 size-exclusion column was equilibrated with 10 mM phosphate buffer (pH 6.5) and 100 mM NaCl. Chromatography was performed at 3 mL/min, collecting 5 mL fractions. A typical elution profile shows three major peaks: the first contains the plasmid DNA, the second contains the desired RNA, and the third peak contains NTPs and small compounds in the transcription reaction that were not completely excluded by the desalting column (Fig. 2). The fractions containing the desired RNA were combined and concentrated using a Vivaspin 15R concentration device (Vivascience, Inc.). The molecular cutoffs of the centrifugal devices are based on globular proteins, and we observed that the molecular cutoff of RNA, which often adopts elongated conformations, is smaller than that recommended by the vendor.

In order to highlight the usefulness of the method for structural biology studies, the imino proton region of a one-dimensional (1D) NMR spectrum of the HIV TAR RNA purified by this method is shown in Figure 3. The new purification protocol yields highly pure RNA without any contamination.

FPLC size-exclusion column selection (Superdex 200 or Superdex 75)

For globular proteins, the Superdex 200 is used to separate proteins of molecular weight between 10 and 600 kDa and the Superdex 75 is effective for proteins between 3 and 70 kDa. However, as shown in Figure 2, HIV TAR RNA (19.8 kDa) is eluted at 220 mL in Superdex 200 (26/60), which corresponds to a molecular weight of 111 kDa for a globular protein. Roughly a sixfold difference in apparent molecular weight between RNA and protein was observed. We therefore determined standard elution curves for RNA oligonucleotides of various lengths on the Superdex 200 and the Superdex 75 size-exclusion columns (Fig. 4). For the Superdex 200 (10/30), 225-, 177-, 134-, 109-, and 60-nt RNAs, which correspond to the molecular weights of 74.3, 58.4, 44.2, 35.9, and 19.8 kDa, respectively, were used for calibration. These RNAs are the derivatives of HCV IRES, interferon-γ mRNA, and HIV TAR (Ben-Asouli et al. 2002; Lukavsky et al. 2003; Kim et al. 2006). For the protein standards, thyroglobulin, ferritin, and catalase, which correspond to the molecular weights of 670, 440, and 240 kDa, respectively, were used for calibration. The Superdex 200 (10/30) column was equilibrated with 10 mM...
phosphate buffer (pH 6.5) and 100 mM NaCl. Chromatography was performed at 0.4 mL/min. The template plasmid is eluted in the void volume of the column at 8.0 mL. The 225-nt RNA (74.3 kDa) is eluted at 10.9 mL. The ferritin protein (440 kDa) also eluted at 10.9 mL. Likewise, the 177-, the 134-, the 109-, and 60-nt RNA are eluted at 11.20, 12.12, 12.79, and 14.62 mL, respectively. This result implies that RNAs behave approximately six times larger than equivalent globular proteins. NTPs and other small compounds are eluted at 17.5 mL.

For the Superdex 75 (10/30) column, 68-, 57-, 43-, 34-, and 14-nt RNA oligonucleotides were used for calibration, which correspond to molecular weights of 22.4, 18.8, 14.1, 11.2, and 4.6 kDa, respectively. These are also derivatives of HCV IRES, interferon-γ mRNA, and HIV TAR RNA. For the protein standards, aldolase, ovalbumin, and chymotrypsinogen were used for calibration, which correspond to the molecular weights of 158, 43, and 25 kDa, respectively. As shown in Figure 4, RNAs behave about four to five times larger than globular proteins by size-exclusion chromatography.

Taken together, these observations demonstrate that the behavior of RNA greatly differs from that of proteins under native conditions on size-exclusion columns, as RNAs are often not globular but form more extended structures.

Separation of desired RNA product from cis-acting ribozyme

T7 RNA polymerase tends to add extra nucleotides to the 3′ end of synthesized RNA oligonucleotides (Milligan et al. 1987; Draper et al. 1988; Pleiss et al. 1998), often yielding RNA oligonucleotides with heterogeneous ends. As a single RNA species is typically required for high-resolution structural studies of RNAs or RNA-protein complexes, this behavior is problematic. By using cis-acting ribozyme at the 3′ ends of the RNA, this problem is readily overcome (Price et al. 1995; Ferre-D’Amare and Doudna 1996). In addition, proper usage of cis-acting ribozyme allows for efficient segmental labeling of RNA oligonucleotides, as we have demonstrated before (Kim et al. 2002). However, the ribozyme fragment must be removed during purification.

In order to test the purification of a desired RNA product from transcript and ribozyme treatment, we constructed a plasmid containing the T7 promoter, a
A template sequence of HCV IRES fragment (225 nt) and a 5′-cis-acting hammerhead ribozyme (44 nt) (Kim et al. 2002). During the transcription reaction, hammerhead cleavage occurs simultaneously and is completed after 2–4 h of incubation at 37°C. After T7 transcription and ribozyme reaction, the pyrophosphate precipitate is pelleted (3000 × g for 10 min) and the reaction quenched by adding EDTA to chelate Mg2+. Subsequently, the reaction is treated as discussed previously, with phenol/chloroform extraction, application to a desalting column, and size-exclusion chromatography. The size-exclusion elution profile contains four major peaks, the first one containing the plasmid DNA followed by (in order) the desired RNA product, the hammerhead ribozyme, and the NTPs and small compounds in the transcription reaction that were not completely excluded from the desalting column (Fig. 5). The fractions containing the desired RNA are combined and concentrated as described previously. To purify the desired RNA product from the transcript and ribozyme, the RNA product should differ from transcript and ribozyme in length such that efficient separation of the RNAs by size-exclusion chromatography can be achieved.

Separation of monomeric from multimeric RNA

Many RNAs can multimerize by intermolecular formation of either Watson–Crick or noncanonical base pairs (Brion and Westhof 1997; Brunel et al. 2002). After transcription, we have observed that RNAs are often mixtures of monomer and multimers (dimer, trimer, etc.), with these equilibria depending on the RNA sequence (data not shown). In the current methodology, the ability to purify monomer away from all other RNA species is achieved through the use of FPLC in combination with large-scale size-exclusion columns. For high-resolution structural studies, it is often imperative to isolate monomeric RNA, as the inclusion of higher-order RNA complexes can lead to significant confusion in the interpretation of experimental results. Unfortunately, the RNA purification method by denaturing PAGE will not distinguish monomeric from multimeric oligonucleotides.

An example application is shown for CpGAIT, a 29-nt RNA molecule (Sampath et al. 2003) that exists in a...
monomer/dimer equilibrium after T7 transcription. After phenol/chloroform extraction, the RNA-containing species were separated into distinct peaks using the Superdex 75 (26/60) size-exclusion column (Fig. 6), and were characterized based on both denaturing and native gels (data not shown). The standard fractions corresponding to plasmid and NTPs were observed, as well as two peaks corresponding to monomeric and dimerized CpGAIT. Exclusively monomeric RNA can be purified away from all other components in a rapid and large-scale fashion. Furthermore, the purified dimer RNA can be subjected to a refolding process if desired and converted into monomer CpGAIT. Therefore, this methodology represents a significant improvement with respect to the identification and removal of higher-order complexes.

Formation of RNA aggregates during storage

After synthesis and purification, RNA samples are often stored in aqueous buffers at either 4°C or −20°C. Many experimenters use these stored RNAs without concern for their molecular state. This is a particular concern for longer biological RNAs that may adopt multiple unimolecular and higher-order conformations. In Figure 7, we show an FPLC trace for a 69-mer RNA derived from the HIV genome. NMR studies have indicated a flexible ensemble of conformers for this RNA in solution. Upon storage at −20°C for >1 d, higher-order RNA structures are observed using FPLC. These include dimeric and higher-order species. Heating and rapid cooling of the RNA leads to recovery of the monomeric species. More severe aggregation problems were observed with RNAs >100 nt (data not shown). Thus, gel filtration is an essential step for preparation of conformationally homogeneous RNAs after storage.

Conclusion

Purification of RNA samples is essential for biochemical investigation. The common method using PAGE is time consuming and contaminates samples with water-soluble acrylamide oligomers (Cheong et al. 2004; Kieft and Batey 2004; Lukavsky and Puglisi 2004). In this article, using the Superdex 75 or Superdex 200 column in the FPLC system, we have demonstrated fast performance and high-resolution purification of RNA. Milligram quantities of pure RNA can be isolated from transcription in <1 d, and monomeric and higher-order conformations of RNA are readily separated. The speed and relative ease of this approach are advantages over prior methods and should greatly aid biochemical and biophysical investigations of RNA.

MATERIAL AND METHODS

Large-scale plasmid preparation

The plasmid DNA pUC119 is used as a template for in vitro transcription of RNA. A template DNA fragment was designed containing a T7 promoter, a desired nucleotide sequence, and the restriction site (BstZ17I or BsaI) to run off the transcription reaction. A template DNA fragment is subcloned into pUC119 between the HindIII and EcoRI sites and transformed to Escherichia coli DH5α. A 500 mL LB culture usually yields 1 mg of DNA plasmid using QIAfilter plasmid Maxi Kits (QIAGEN, Inc.).

Transcription

After restriction enzyme (BstZ17I or BsaI) treatment, the linearized plasmid template is used for in vitro transcription using T7
polymerase. After 2 or 3 h of incubation at 37°C, pyrophosphate precipitate is pelleted (3000g × 10 min) and the reaction quenched by adding EDTA to chelate Mg²⁺. An equal volume of phenol/chloroform is added to the supernatant and centrifuged (3000g × 10 min). In this process, the restriction enzyme, which is carried over with the linearized plasmid DNA, and the T7 RNA polymerase are removed from the reaction mixture. After 2 or 3 phenol/chloroform extractions, the upper aqueous phase is directly loaded on the desalting column (10DG column, Bio-Rad), which is equilibrated with 20 mL of 10 mM phosphate buffer (pH 6.5) and 100 mM NaCl. A 3 mL sample is loaded to the desalting column. After the entire sample is allowed to enter the column the first 2 mL of effluent are discarded. Five milliliters of desalting column. After the entire sample is allowed to enter the column (pH 6.5) and 100 mM NaCl. The 1D imino proton spectrum was collected in 90% H₂O/10% ²H₂O at 298 K.

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