Nucleolar Localization of Human Methionyl–tRNA Synthetase and Its Role in Ribosomal RNA Synthesis

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Abstract. Human aminoacyl–tRNA synthetases (ARSs) are normally located in cytoplasm and are involved in protein synthesis. In the present work, we found that human methionyl–tRNA synthetase (MRS) was translocated to nucleoli in proliferative cells, but disappeared in quiescent cells. The nucleolar localization of MRS was triggered by various growth factors such as insulin, PDGF, and EGF. The presence of MRS in nucleoli depended on the integrity of RNA and the activity of RNA polymerase I in the nucleolus. The ribosomal RNA synthesis was specifically decreased by the treatment of anti-MRS antibody as determined by nuclear run-on assay and immunostaining with anti-Br antibody after incorporating Br-UTP into nascent RNA. Thus, human MRS plays a role in the biogenesis of rRNA in nucleoli, while it is catalytically involved in protein synthesis in cytoplasm.

Key words: methionyl–tRNA synthetase • nucleoli • growth signal • ribosomal RNA synthesis • RNA polymerase I

Introduction

Aminoacyl–tRNA synthetases (ARSs) are enzymes decoding genetic information into amino acids. Although these enzymes normally execute their catalytic activities for protein synthesis, recent reports suggest that they are not simple enzymes, and they can play novel regulatory functions in various processes (Martinis et al., 1999). Mammalian tryptophanyl–tRNA synthetase is induced by interferon (Kisselev et al., 1993) and the same enzyme of Drosophila melanogaster is under the control of the homeotic gene, Scr, and highly expressed in salivary gland during development (Seshaih and A ndrew, 1999). Mitochondrial tyrosyl–tRNA synthetase of Neurospora crassa (Akins and Lambowitz, 1987) and leucyl–tRNA synthetase of Saccharomyces cerevisiae (Labouesse, 1990) are involved in the splicing process. Human tyrosyl–tRNA synthetase is converted to two distinct proapoptotic cytokines (Wakasugi and Schimmel, 1999) and human arginyl–tRNA synthetase (RRS) also sequesters the precursor of a proapoptotic cytokine (Park et al., 1999). Thus, we anticipated the unveiling of more diverse functions from these enzymes.

To gain an insight into the novel functions of mammalian ARSs, we investigated cellular localizations of different human ARSs using their specific antibodies. Among the tested ARSs, methionyl–tRNA synthetase (MRS) was uniquely localized in the nucleolus. Although the presence of MRS in nucleoli was previously reported (Dang et al., 1983), the functional reason for the nucleolar localization of MRS is not understood. Here, we investigated the translational control and functional significance of nucleolar MRS.

Human cytoplasmic MRS consists of 900 amino acids (Lage and Dietel, 1996) and is one of the components for the multi-tRNA synthetase complex (M isselev and W olfson, 1994; Y ang, 1996). The core domain is homologous to the corresponding enzymes from prokaryotes (Fig. 1 A). However, it contains the unique NH$_2$-terminal extension of 267 amino acids that is not essential for catalytic activity (data not shown), but is involved in protein–protein interaction (Rho et al., 1999). Similarly, the NH$_2$-terminal extension of yeast MRS is also responsible for the interaction with a nuclear pore–associated protein, A rclp (Simos et al., 1996). A nother motif of ~40 amino acids (Fig. 1A , gray box) is present in the COOH-terminal region (Q847-K897) that is homologous to the motifs present in other ARSs and involved in protein–protein and protein–nucleic acid interactions (Rho et al., 1996, 1998). In addition, putative nuclear
Localization signals (Schimmel and Wang, 1999) are found in the COOH-terminal region as four consecutive lysines from K897 to K900 and PWKRIKG from P724 to G730 (Fig. 1A, bars), implying that MRS may be translocated to the nucleus. Here, we report that human MRS is translocated into nucleoli by various cell proliferation signals and is involved in rRNA synthesis.

Materials and Methods

Cell Culture

HeLa, Chang, HepG-2, COS-1, and human foreskin fibroblast were grown to subconfluency on 5 × 5-mm glass coverslips in 35-mm petri dishes in DME supplemented with 10% FBS (GIBCO BRL). Confluent cells were prepared by growing 5 × 10^5 cells on coverslips in DME/10% FBS for 4–6 d without changing the medium. Quiescent cells were also prepared by serum starvation for 5–7 d in DME. RNA polymerase I and RNA polymerase II were inhibited by the addition of cisplatin (10 μg/ml for 9 h; Jordan and Carmo-Fonseca, 1998) and α-amanitin (2 μg/ml for 16 h; Kedinger et al., 1970; Lindel et al., 1970), respectively. The inhibition of RNA polymerase I and II was achieved by the treatment of α-amanitin (2 μg/ml for 16 h) and actinomycin D (0.2 μg/ml for 16 h; Perry, 1963).

Antibody Preparation

The cDNA encoding the full-length human cytoplasmic MRS was isolated by PCR from pM184 (Rho et al., 1999) as a template using two specific primers. The PCR product was cleaved with EcoRI and HindIII designed into each of the primers, and inserted into the same site of pET28a (Novagen). The resulting plasmid was introduced into Escherichia coli BL21 (DE3) and induced with IPTG. Since the recombinant MRS was insoluble, it was purified as a denatured polypeptide using nickel-affinity chromatography following the manufacturer’s instruction (Invitrogen). The native peptide of EPRS (bifunctional glutamyl-prolyl–tRNA synthetase) from D677 to E884, the native NH2-terminal 236 aa of human QRS (glutaminyl–tRNA synthetase) and the native NH2-terminal 72 aa of human RRS were also expressed as His-tagged proteins. All of these polypeptides were also purified using His tag. Each purified polypeptide was used to raise specific polyclonal rabbit antibodies as described previously (Park et al., 1999). The IgG from each antiserum was purified by protein A affinity chromatography according to the manufacturer’s protocol (BioRad), and the antibody specificity was confirmed by immunoblotting.

Confocal Immunofluorescence Microscopy

Cellular localizations of different ARSs were investigated using confocal immunofluorescence microscopy (μ Radiance, BioRad). The cells were cultured to ~70% confluence on 5 × 5-mm coverslips and then washed with cold PBS, fixed with 10% formaldehyde for 10 min at room temperature, washed with PBS, and then incubated in PBS containing 100 μM NH4Cl for 10 min at room temperature. A 1× washing the fixed cells with PBS, the cells were permeabilized with 0.1% Triton-X 100 in PBS for 5 min at room temperature, washed with PBS, and blocked with 0.2% BSA in PBS for 1 h. The cells were then incubated with polyclonal rabbit antibodies raised against different ARSs in PBS containing 0.2% BSA for 2 h at 37°C. A 1× washing the primary antibody, the FITC-conjugated secondary antibody was added and ARSs were detected using a confocal laser scanning microscope. Nucleolin was stained, as described above, using antinucleolin primary (Santa Cruz) and rhodamine-conjugated secondary antibodies.

Immunoblotting

The nuclear extract and postnuclear supernatant were prepared from the
cultured HeLa cells as described previously (Neufeld and White, 1997). MRS, Y1, lamin B, and heat shock protein 90 (Hsp90) were reacted with their corresponding antibodies (anti-Y1 and -lamin B antibodies from Santa Cruz; anti-Hsp90 antibody from Transduction Laboratories) and detected using the ECL system (A mershham Pharmacia Biotech).

**Gel Filtration of ARSs**

The subconfluent HeLa cells (three 150-mm dishes) were lysed in 1 ml of 25 mM HEPES, pH 7.4, 150 mM NaCl, 10% glycerol, and 0.5% Triton X-100. After the lysate was centrifuged at 25,000 g for 30 min, the supernatant was filtered through a 0.22-μm membrane filter and concentrated using the Viva Spin (VIVASCIENCE) to the protein concentration of 20 mg/ml. The concentrated protein extract was then loaded into Superdex 200 HR (exclusion limit of 1,300 kD) using AKTA-FPLC (Amersham Pharmacia Biotech) and eluted at the flow rate of 0.25 ml/min. The eluted proteins in each fraction were analyzed by immunoblotting with anti-MRS, anti-p43 (Park et al., 1999), and anti-EPRS rabbit antibodies.

**Nuclear Run-on Assay**

Synthesis of rRNA was monitored by nuclear run-on assay as described previously (Giraudo et al., 1998). The nuclei isolated from HeLa cells were mixed with the indicated amounts of rabbit IgG or anti-MRS antibody at room temperature for 10 min. Transcription was initiated by addition of the nuclei to the reaction mixture in the presence of α-[32P]UTP (3,000 Ci/mmol, New England Nuclear). The synthesized radioactive 18S and 28S rRNAs were quantified by hybridization to their respective cDNAs that were immobilized on Hybond membrane (Amersham Pharmacia Biotech.) using Bio-Dot apparatus (BioRad). The amount of the hybridized radioactive transcripts were quantified by phosphor image analyzer (FLA 3000, Fuji). The cDNA encoding β-actin was used as an internal control.

**Immunostaining of Nascent rRNA with Anti-BrdU Antibody**

To monitor the rRNA synthesis, Br-UTP was incorporated into nascent rRNA and detected by confocal immunofluorescence microscopy using an antibody specific to Br (Sigma Chemical Co.). HeLa cells cultured on coverglass were briefly washed twice with a PBS buffer (pH 7.4), and with permeabilization buffer (20 mM Tris-HCl, pH 7.4, 5 mM MgCl2, 0.5 mM EGTA, 0.5 mM PMSF). The cells were then permeabilized with the same buffer containing 0.05% Triton X-100 for 5 min at room temperature and washed with the permeabilization buffer without Triton X-100. The permeabilized cells were used for nuclear run-on assay. The synthesis of rRNA was initiated by adding 50 mM Tris-HCl, pH 7.4, 100 mM KCl, 5 mM MgCl2, 0.5 mM EGTA, 25 U/ml RNasin, 1 mM PMSF, 0.5 mM of ATP, CTP, and GTP, and 0.2 mM Br-U TP (Sigma Chemical Co.) in the presence of different antibodies (5 μg/ml each of anti-MRS, anti-QRS, and mock rabbit IgG), or in the presence of α-amanitin (1 μg/ml) or cisplatin (10 μg/ml). The reactions were continued for 30 min at 37°C. The cells were then washed with a PBS buffer containing 25 U/ml RNasin at room temperature. The cells were then fixed with PBS containing 10% fresh formaldehyde, 25 U/ml RNasin, and 0.1% BSA for 20 min at room temperature and washed twice with 0.1% BSA in PBS for 5 min. The cells were then treated with PBS containing 0.1% Triton X-100 and 0.1% BSA for 10 min at room temperature and washed twice with PBS containing 0.1% BSA for 5 min. An anti-Br antibody (1:100) was then reacted in PBS containing 0.5% BSA and 2% CAS for 2 h at 37°C, and washed three times with PBS containing 0.1% BSA for 5 min. Then, the FITC-conjugated secondary antibody was reacted in PBS containing 0.5% BSA and 2% CAS for 1 h at 37°C. A after washing the cells, the incorporated Br-U TP was monitored by confocal immunofluorescence microscopy. The nuclei were stained with propidium iodide (10 μg/ml) as described previously (A ndreassen et al., 1998).

**A. Cellular distribution of different ARSs**

| MRS | QRS | EPRS | RRS |
|-----|-----|------|-----|

**B. Nucleolar localization of MRS**

Fibroblast HeLa HepG2 COS-1

Figure 2. Nucleolar localization of MRS. A, Distributions of four different ARSs in Chang cells were determined by immunostaining using confocal laser scanning microscopy. B, top, Cellular localization of MRS was monitored in different cell lines. Human foreskin fibroblast, HeLa, HepG2, and COS-1 cells were cultivated and stained with anti-MRS as described above. Bar, 10 μm. Bottom, Localization of MRS and nucleolin in Chang cells was determined using anti-MRS rabbit antibody and mouse monoclonal anti-nucleolin antibody (Santa Cruz) as described in Materials and Methods.
Results

Nucleolar Localization of MRS

To investigate cellular distribution of different ARSs, polyclonal antibodies that were specific to MRS, EPRS, RRS, and QRS were prepared. Their antigenic specificity was determined by immunoblotting of the proteins extracted from HeLa cells. All of the antibodies showed specificity to their antigens (Fig. 1 B). Using these antibodies, cellular distribution of four different ARSs was investigated by immunostaining. Although all of these enzymes were detected in both the nucleus as well as the cytoplasm, relative partitions between the two cellular locations and staining patterns were idiosyncratic (Fig. 2 A). Among them, MRS was uniquely stained at nucleoli, implying its novel function at this site. Removal of anti-MRS IgG from anti-MRS antiserum decreased nucleolar and cytoplasmic staining and mock rabbit IgG or preimmune serum did not give any specific MRS signal (data not shown).

The nucleolar localization of MRS was further investigated in the different cells by confocal immunofluorescence microscopy using an anti-MRS antibody. The nucleolar MRS was detected in all of the human foreskin fibroblast, HeLa, HepG2, and COS-1 cells, indicating that the nucleolar localization of MRS is universal in mammalian cells (Fig. 2 B). The nucleolar localization of MRS was confirmed by coinmunostaining of MRS with nucleolin that was used as a marker for nucleoli. MRS was exactly colocalized with nucleolin, confirming its nucleolar localization (Fig. 2 B).

The presence of MRS at nucleoli was then investigated by immunoblotting of nuclear and postnuclear fractions of HeLa cells. The transcription factor, YY1 (Shi et al., 1991), and nuclear structural protein, lamin B (Moir et al., 1995), and Hsp90 (Koyasu et al., 1986) were used as nuclear and cytoplasmic markers, respectively. The marker proteins were found in the expected fractions, indicating that the nuclei were well isolated. MRS was found both in nucleus as well as in postnuclear supernatant (Fig. 3, top). Since MRS is a component of the multi-ARS complex, the presence of MRS in nucleus implies that at least some portion of MRS should exist as a different form. This possibility was investigated by size exclusion chromatography (exclusion limit of 1,300 kD) of the proteins extracted from the HeLa cells. The multi-ARS complex would be eluted in the void volume from this column because its approximate molecular weight is 1,500 kD. The proteins eluted from the column were resolved by gel electrophoresis. MRS and two other complex-components, EPRS (Fett and Nippers, 1991) and p43 (Quevillon et al., 1997), were detected by immunoblotting with their respective antibodies. The majority of the three proteins were coeluted in the void volume as expected. However, a significant amount of MRS was also detected in the following fractions in which EPRS and p43 were barely detected (Fig. 3, bottom). This result implies that MRS may be loosely associated with the multi-ARS complex or that a portion of MRS may exist unassociated from the multi-ARS complex.

MRS Is Translocated to Nucleoli by Cell Proliferation Signal

We then investigated the condition in which MRS is translocated to nucleoli by cell proliferation signal. Proliferation signal from the multi-ARS complex or that a portion of MRS may exist unassociated with their respective antibodies. The majority of the proteins extracted from HeLa cells were fractionated by size exclusion chromatography (exclusion limit of 1,300 kD). The eluted proteins in each fraction were analyzed by immunoblotting with anti-MRS, -p43 (Park et al., 1999), and -EPRS rabbit antibodies.

Figure 3. Determination of nuclear and free form MRS. Top, MRS was detected by immunoblotting in the whole cell lysate (WCL), nuclear extract (NCL), and postnuclear supernatant (PNS). Cell fractionation was performed as described previously (Neufeld and White, 1997). YY1, lamin B, and Hsp90 were used as nuclear and cytoplasmic markers, respectively. Bottom, The proteins extracted from HeLa cells were fractionated by size exclusion chromatography. The eluted proteins in each fraction were analyzed by immunoblotting with anti-MRS, -p43 (Park et al., 1999), and -EPRS rabbit antibodies.
Fonseca, 1998) that inhibited RNA polymerase I. However, it was not affected by the treatment of the RNA polymerase II inhibitor, α-amanitin (Kedinger et al., 1970; Lindel et al., 1970; Fig. 5, bottom). These results suggest that the nucleolar MRS is related to rRNA synthesis.

**Anti-MRS Antibody Blocks rRNA Synthesis**

The effect of MRS on rRNA synthesis was then investigated by a nuclear run-on assay and immunostaining of nascent rRNA. The HeLa cell nuclei were isolated and the transcription was carried out in the presence of different amounts of anti-MRS antibody. The synthesized RNA's were isolated and hybridized with 18S and 28S rDNAs, as well as cDNA for β-actin that was used for internal control. The synthesis of two rRNA's was decreased by the addition of anti-MRS antibody, in a dose-dependent manner, to 20% of the control in which no antibody was added (Fig. 6). In contrast, mock rabbit IgG did not affect the synthesis of 18S and 28S rRNA's and the amounts of

![Figure 4](image_url) **Figure 4.** MRS is translocated to nucleolus upon a mitogenic signal. Top, Cellular localization of MRS was monitored in Chang cells as described in Fig. 2. The nucleolar MRS was apparent in subconfluent (∼70% confluence) cells, but disappeared in confluent (100% confluence) or 5-d serum-starved cells. Bottom, 10% serum or EGF (40 ng/ml), PDGF (40 ng/ml), and insulin (100 µg/ml) were added to 7-d serum-starved cells and the cells were observed 24 h after the treatment. Bar, 10 µm.

![Figure 5](image_url) **Figure 5.** Nucleolar localization of MRS depends on rRNA synthesis. Top, HeLa cells grown on glass cover-slips were permeabilized in 0.1% Triton X-100 for 5 min at room temperature. The permeabilized cells were incubated with DNase I (0.1 mg/ml) or RNase A (0.1 mg/ml) for 1 h at 37°C and fixed in 10% formaldehyde for 20 min. Digestion of nuclear DNA was confirmed by DAPI staining (data not shown). Bottom, HeLa cells were treated with α-amanitin (2 µg/ml), and α-amanitin (2 µg/ml) + actinomycin D (0.2 µg/ml) for 16 h to inhibit RNA polymerase II and RNA polymerase I + II, respectively. The specific inhibition of RNA polymerase I was performed by incubating HeLa cells with cisplatin (10 µg/ml) for 9 h. The cellular localization of MRS was monitored by immunostaining as described above. Bar, 10 µm.
β-actin RNA were unchanged by the treatment of anti-MRS antibody. These results indicate that anti-MRS antibody specifically affected the synthesis of rRNA.

We also monitored the rRNA and mRNA synthesis by the immunostaining of Br-UTP incorporated to the nascent RNA. Ribosomal RNA synthesis, executed by RNA polymerase I, was shown as nucleolar foci, whereas synthesis of mRNA by RNA polymerase II was detected as nucleoplasmic foci. The amounts of the hybridized transcripts were quantified by phosphor image analyzer. The radioactive intensities of 18S, 28S, and β-actin blots without IgG were taken as 100% and the relative intensities of other blots were shown by percentage. White bars stand for the values of the intensities of 18S and 28S treated with the indicated amounts of mock rabbit IgG. Black, gray, and lined bars represent the relative intensities of the 18S, 28S, and β-actin blots treated with the indicated amounts of anti-MRS antibody, respectively. Similar results were obtained from three independent experiments.

Discussion

Several mammalian ARSs form a macromolecular protein complex (M irande, 1991; Kisselev and Wolfson, 1994; Y ang, 1996). However, the presence of free forms has been reported in a few different ARSs that are the components for the complex (M irande et al., 1983; Vellekamp et al., 1983, 1985). Here, the four complex-forming ARSs showed different patterns in immunostaining (Fig. 2 A). These results imply that at least some portions of the complex-forming ARSs are distributed differently in a cell, while most of them are present within the macromolecular protein complex.

Human MRS is also a component of this multi-ARS complex. However, MRS dissociated from the multi-ARS complex may exist based on the elution profile from size exclusion chromatography and its presence in the nucleus (Fig. 3). MRS in nuclear fraction was also eluted from the gel filtration column as a macromolecular complex (data not shown), implying that it is also associated with other nuclear factors or structure. This observation is consistent with its localization in nucleoli. Human MRS contains peptide extensions attached to the NH₂- and COOH-terminal ends of the core catalytic domain (Fig. 1A). Interestingly, these two peptide appendages are involved in protein–protein interactions (Rho et al., 1999) and phosphorylation sites for casein kinase II are heavily clustered in the NH₂-terminal appendix (data not shown). Since casein kinase II is involved in the growth control and regulation of rRNA synthesis (V olt et al., 1995; H annan et al., 1998), phosphorylation of MRS at these sites may be related to the nucleolar localization of MRS. However, we failed to determine the peptide region responsible for the nucleolar localization of MRS, because various forms of recombinant MRS were not well expressed and gave a cytotoxic effect in transfected cells (data not shown).

The nucleolus is the nuclear site in which rRNA biogenesis takes place. Interestingly, it was reported previously that nucleoli show an independent protein synthesis capability (L amkin et al., 1973) and contain some components of protein synthesis, such as the elongation factor (R ao et al., 1998). Electron microscopic analysis also suggested the presence of a few ARSs in nucleoli (Popenko et al., 1994). However, the condition and physiological meaning for the presence of these factors in nucleolus has not been understood. Here, we report that the cytoplasmic MRS is translocated into nucleolus by various cellular proliferation signals (Fig. 4) and is involved in rRNA synthesis (Figs. 6 and 7). Interestingly, 5S rRNA is associated with MRS and enhances its activity in cytoplasm (O gata et al., 1991). MRS may be also bound to 5S rRNA in nucleolus and regulate the biogenesis of ribosome.

The ribosome biogenesis of prokaryotes is subjected to stringent control mediated by unusual nucleotides in response to uncharged tRNA (C ozzzone, 1980). Amino acid starvation drops the nucleolar rRNA synthesis as well as the cytoplasmic protein synthesis in euukaryotic cells. This suggests the presence of a mechanism sensing the cellular level of the amino acid. Although eukaryotes do not seem to employ unusual nucleotide messengers analogous to those of prokaryotes for this communication (P ollard et al., 1980), many protein factors have recently been found in nucleolus and are involved in rRNA synthesis. For in-

![Image](image_url)
nucleolar rRNA synthesis was blocked with anti-MRS antibody, but not with mock IgG or anti-QRS antibody. The stained foci disappeared with the treatment of cisplatin that inhibits RNA polymerase I, but not with the treatment of α-amanitin that inhibits RNA polymerase II. This confirms that the stained foci resulted from the nucleolar rRNA synthesis. Nucleoplasmic Br-staining is not shown here due to the short exposure.

de, a zinc finger protein, ZPR1, is associated with the cytoplasmic domain of EGF receptor in quiescent cells (Galcheva-Gargova et al., 1996), but translocated to nucleolus to activate rRNA synthesis by a cell proliferation signal (Galcheva-Gargova et al., 1998). In reverse, a tumor suppressor, retinoblastoma protein, is translocated to nucleolus to inhibit rRNA synthesis during cellular differentiation (Cavanaugh et al., 1995). In this sense, MRS is an ideal molecule, coordinating rRNA synthesis in the nucleolus and protein synthesis in cytoplasm.

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