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Human selenophosphate synthetase 1 has five splice variants with unique interactions, subcellular localizations and expression patterns

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Abstract

Selenophosphate synthetase 1 (SPS1) is an essential cellular gene in higher eukaryotes. Five alternative splice variants of human SPS1 (major type, ΔE2, ΔE8, +E9, +E9a) were identified wherein +E9 and +E9a make the same protein. The major type was localized in both the nuclear and plasma membranes, and the others in the cytoplasm. All variants form homodimers, and in addition, the major type forms a heterodimer with ΔE2, and ΔE8 with +E9. The level of expression of each splice variant was different in various cell lines. The expression of each alternative splice variant was regulated during the cell cycle. The levels of the major type and Δ E8 were gradually increased until G2/M phase and then gradually decreased. ΔE2 expression peaked at mid-S phase and then gradually decreased. However, +E9/+E9a expression decreased gradually after cell cycle arrest. The possible involvement of SPS1 splice variants in cell cycle regulation is discussed.

Keywords

alternative splicing; cell cycle; selenium; selenocysteine; selenophosphate synthetase 1

1. Introduction

Selenium is an essential trace element and provides many health benefits for animals including humans. For example, this element has been known to play a role in cancer prevention, aging retardation, immune augmentation, prevention of cardiovascular disease, muscle development and immune function [1-4]. Most of these various selenium effects are believed to be mediated by selenoproteins which contain the selenium-containing amino acid, selenocysteine (Sec), at their active site(s). Sec, which is the $21st$ amino acid in the

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genetic code, is inserted into a growing peptide during translation in response to the codon, UGA [5-7]. The biosynthesis of Sec is carried out through a unique pathway that is distinct from that of other amino acids. Most amino acids are synthesized prior to being aminoacylated onto their cognate tRNAs; however, Sec is synthesized on its tRNA, designated tRNA[Ser]Sec. In eukaryotes, tRNA[Ser]Sec is initially aminoacylated with serine by seryl-tRNA synthetase, the hydroxyl moiety of serine on seryl-tRNA^{[Ser]Sec} is then replaced by phosphate group to form O-phosphoseryl-tRNA[Ser]Sec that is catalyzed by a specific kinase, phosphoseryl-tRNA^{[Ser]Sec} kinase [8]. Finally, the phosphate group on Ophosphoseryl-tRNA^{[Ser]Sec} is exchanged with activated selenium to form selenocysteyltRNA[Ser]Sec catalyzed by the enzyme, Sec synthase (SecS) [9,10]. The O-phosphoseryltRNA[Ser]Sec intermediate is not synthesized in eubacteria wherein SecS in *E. coli* (SelA) removes the hydroxyl group of serine on seryl-tRNA[Ser]Sec to form aminoacrylyltRNA[Ser]Sec and the resulting aminoacrylyl intermediate in turn accepts activated selenium to form Sec-tRNA^{[Ser]Sec} [11]. Mono-selenophosphate (SeP) is the activated selenium form in both bacteria and eukaryotes [12], and it is synthesized from selenide and ATP. Selenophosphate synthetase (SPS) catalyzes the synthesis of SeP [13]. Two isoforms of SPS were found in higher eukaryotes, SPS1 and SPS2, whereas only one type of SPS (SelD) exists in lower eukaryotes and eubacteria [14]. The sequence homology between SPS1 and SPS2 is different depending on the species. For example, the amino acid sequence homology between human SPS1 and SPS2 is 72% and that between *Drosophila* SPS1 and SPS2 is approximately 45%. One of the major differences between SPS1 and SPS2 is that SPS1 has an arginine at the position corresponding to Sec in SPS2 [15,16]. The sequence homology between *Drosophila* SPS1 and *Drosophila* SPS2 is 45%, while that between human SPS1 and *Drosophila* SPS1 is 71%, indicating that the SPS1 sequence has been well conserved during evolution (NCBI, Build 37.1 for human sequence and BDGP Release 5.12 for *Drosophila*).

Recently, it was shown that SPS1 does not have SeP synthesis activity, but SPS2 catalyzes SeP synthesis. In *in vitro* experiments, SPS2 synthesized SeP from selenide and ATP, but SPS1 did not have this activity [9]. The ablation of SPS2 mRNA in NIH3T3 cells led to the loss of selenoprotein biosynthesis, while the inhibition of SPS1 expression did not affect the biosynthesis of selenoprotein, and the synthesis of selenoproteins in the cells in which SPS2 was knocked down was rescued by expression of SPS2, but not by SPS1 [17]. Although some insects such as the red beetle and silkworm have lost the selenoprotein synthesizing machinery including *SPS2, SPS1* is still encoded in the genome of those organisms suggesting SPS1 is not essential for selenoprotein synthesis [18].

Although SPS1 does not catalyze SeP biosynthesis [12], there are several lines of evidence suggesting that it plays important roles in the cell. Targeted removal of the gene encoding SPS1 (also designated *patufet*) in *Drosophila* causes embryonic lethality suggesting that SPS1 participates in pathways important for cell viability [19]. It was also found that knockout of *SPS1* caused an increase in reactive oxygen species (ROS) and the haploinsufficiency of *SPS1* suppressed the abnormal phenotypes caused by overexpression of genes involved in *Ras*-regulated signaling pathway [20,21]. These latter findings suggest that SPS1 participates in the regulation of oxidative stress and cell growth. In agreement with this predicted function of SPS1, knockdown of *Drosophila SPS1* led to growth inhibition and ROS generation [22]. Interestingly, knockdown of *Drosophila SPS1* caused generation of megamitochondria through the increase of intracellular glutamine levels.

When cDNA of *SPS1* was cloned from human lung adenocarcinoma and transformed into a *SelD* deficient mutant of *E. coli*, the *selD* mutation was not complemented in selenite containing medium, but complemented when the cells were cultured in the medium supplemented with L-Sec [23]. From these findings, it was suggested that SPS1 is involved

in the recycling of Sec. However, the mechanism of Sec recycling and how SPS1 regulates Sec recycling have not been determined. Interestingly, SPS1 was found to interact with soluble liver antigen (SLA) which was recently identified as eukaryotic Sec synthase (SecS) and the binding reaction was enhanced by a protein designated SECp43 [24].

In the present study, we found that *hSPS*1 produces five alternative splice variants and one of them was generated by mis-splicing. These alternative splice variants showed unique characteristics of intermolecular interaction, subcellular localization, specific tissue and cell cycle expression suggesting that each splice variant has a specific function for cell growth.

2. Materials and methods

2.1. Cell culture and DNA transfection

A549, CRL7407, MCF7, Chang liver, HeLa, and 293 cells were grown in Dulbecco's Modified Eagle's Medium supplemented with 10% fetal bovine serum (FBS) (Hyclone) and 1% antibiotic-antimycotic (AA), and BJAB cells in RPMI 1640 medium supplemented with 10% FBS and 1% AA. All cultures were maintained in a humidified atmosphere of 5% $CO₂$ at 37°C. Transient transfections in HeLa cells were carried out using Lipofectamine Plus (Invitrogen) according to the manufacturer's instructions and calcium phosphate method was used in 293 cell transfection.

2.2. Vector constructions

Vectors that express alternative splice variants were constructed in two steps: In the first step, a pool of clones containing all types of splice variants was prepared. In the second step, each splice variant was screened using colony PCR. Vectors that can express Flag- or HAtagged proteins were prepared by PCR using primers that contain the sequences encoding Flag or HA, respectively (see Supplementary Materials and methods for detail).

2.3. Immunostaining and confocal microscopy

HeLa cells transfected with Flag-tagged hSPS1 expression vector were transferred onto a cover glass one day before staining. Immunocytochemistry was carried out as described previously [22] with a slight modification. After the cells were fixed, permeabilized and blocked, anti-Flag antibody (Sigma) diluted in 3% BSA in PBS (1:2,500) was treated for 1 h at room temperature. The cells were then incubated with Alexa 488 anti-mouse IgG (Invitrogen) diluted in 3% BSA in PBS (1:2,000) for 1 h at room temperature. For counterstaining, cells were stained with 50 μg/ml of propidium iodide (PI) (Sigma) for 20 min at room temperature. Cells were observed with a LSM 510 NLO confocal microscope (Carl Zeiss). Excitation was at 488 nm for Alexa 488 and 543 nm for PI.

2.4. Quantitative real-time RT-PCR

To determine the number of hSPS1 mRNA molecules, total RNA was isolated from cells using TRIzol reagent (Invitrogen), first-strand cDNA was synthesized from 3 μg of total RNA with Mo-MuLV reverse transcriptase (Super-Bio), and used for the PCR reaction. cDNAs were amplified with SYBR Green PCR master mix and specific primers using the ABI 7300 real-time PCR system (Applied Biosystems). Vectors which can express the hSPS1 alternative splice variant were serially diluted from $10¹$ to $10⁸$ copy numbers and applied to real-time PCR. Standard curve was obtained by plotting the log of the number of DNA templates versus the measured C_T (cycle-threshold). Using the equation calculated from the standard curve, the copy number of hSPS1 mRNA per nanogram of total RNA was determined.

2.5. Immonoprecipitation and immunoblotting

A vector which can express a Flag-tagged splice variant was cotransfected with a HA-tagged splice variant expression vector in 293 cells in a manner that a vector expressing one type of Flag-tagged splice variant was cotransfected with a vector expresses one of all types of HAtagged splice variants. Cell lysates were coimmunoprecipitated with anti-Flag M2 affinity gel (Sigma) and the precipitates subjected to western blotting using anti-HA antibody or anti-Flag antibody (see Reference [22] and Supplementary Materials and methods for detail).

2.6. Synchronization of cells and flow cytometry

For synchronization, HeLa cells were incubated with 5 μ g/ml of aphidicolin (Sigma) for 24 h. Cells were washed with fresh medium and released from the aphidicolin block for indicated times. Cell cycle profiles were analyzed by flow cytometry with standard PI staining methods. Asynchronized and synchronized cells were harvested and washed once with PBS. Cells were fixed overnight in 70% ethanol at -20°C, washed with PBS twice and then resuspended with PBS containing 200 μg/ml of RNase A. After 10 min incubation at room temperature, cells were further incubated with 100 μg/ml of PI in PBS for 30 min at room temperature. Stained cells were analyzed by using a FACS Calibur flow cytometer and CellQuest Pro software (BD Biosciences).

3. Results

3.1. Identification of alternative splice variants in hSPS1 mRNA

To elucidate the molecular function of hSPS1, its cDNA was amplified using total cDNAs prepared from HeLa cells and cloned into a vector. The sequences of several clones harboring hSPS1 cDNA were determined. Interestingly, we found that one of the clones was a truncated form in which exon 8 was deleted. We, therefore, assumed that there would be alternative splice variants in hSPS1 mRNA. To find possible alternative splice forms, we searched EST clones from the University of California, Santa Cruz Genome Browser [\(http://genome.ucsc.edu](http://genome.ucsc.edu)) in scrutiny. Among a total of 283 EST clones of hSPS1, 277 ESTs were the same as known hSPS1 mRNA and designated major type (MT). Except MT, there were two ESTs (CX787008, DB001684) in which exon 2 was deleted (ΔE2), two ESTs (AI741523, AI742171) where exon 8 was deleted (ΔE8) and additional 2 ESTs (BX385308, $CR604492$) where exon 9 was added to the major type $(+E9)$. These data suggest that there are 4 alternative splice variants of hSPS1 mRNA in human cells.

To confirm that the predicted alternative splice variants are present in human cells, we performed RT-PCR in several steps and surprisingly, detected 5 splice variants instead of the 4 predicted splice variants (see Supplementary Result and Supplementary Fig. S1). The sequence of each splice variant was determined and found to be consistent with the expected sequence (GenBank Accession No. GU954545, GU954546, GU954547, GU954548, and GU954549; Supplementary Fig. S2).

As summarized in Fig. 1, hSPS1 pre-mRNA is alternatively spliced to produce 5 different splice variants. The translation initiation codon is located in exon 2 and the termination codon in exon 10. As expected, ΔE2 and ΔE8 do not have exon 2 and exon 8, respectively. Since the normal initiation codon is located in exon 2, ΔE2 cannot use the original initiation codon. Therefore, $\Delta E2$ must use another initiation codon. In exon 3 (at +561), there is an ATG codon and the surrounding Kozak's consensus sequence. Both forms are void of exon 9 as found in MT. The sequence of exon 9 is very short and there is the translation termination codon (TAA) within exon 9 suggesting that the translated protein from +E9 will be truncated from MT, even though the size of mRNA of +E9 is larger than that of MT.

+E9a has an additional sequence which is part of 5' region of intron 9. Interestingly, the beginning and end of the additional sequence contain splice junction sequences (GT and AG), respectively. In the genomic DNA sequence, there is no gap between exon 9 and the additional sequence. This suggests that +E9a is generated by mis-splicing. The protein sequences of +E9 and +E9a should be the same, because there is a termination codon within exon 9.

3.2. Subcellular localizations of splice variants

Recently, it was reported that hSPS1 makes supramolecular complexes by interacting with SecS both *in vivo* and *in vitro* and this supramolecular complex shuttles between the nucleus and the cytoplasm [24]. Therefore, it is important to determine the subcellular localization of each alternative splice variant to elucidate their functions. The Flag-tagged splice variants were expressed and detected by immunocytochemistry using anti-Flag antibody. As shown in Fig. 2, MT was localized on both plasma and the nuclear membrane, but the other three types were localized in the cytoplasm homogeneously. These results suggest that each alternative splice type plays a unique role by interacting with different proteins or molecules.

3.3. Interactions between splice variants

By analyzing the X-ray crystallography structure of hSPS1, it was suggested recently that hSPS1 could form a homodimer [25]. We therefore examined whether all the splice variants could make dimers (homodimers as well as heterodimers). In addition to Flag-tagged constructs, HA-tagged constructs were also prepared (see Supplementary Materials and methods). After co-transfecting a Flag-tagged construct and a HA-tagged construct into 293 cells, cell lysates were subjected to immunoprecipitation using anti-Flag antibody and then the co-purified products were detected by western blot analysis using anti-Flag or anti-HA antibody. As shown in Fig. 3, MT-Flag interacts with MT-HA and Δ E2-HA, and Δ E2-Flag with MT-HA and ΔE2-HA. Similarly, ΔE8-Flag interacts with ΔE8-HA and +E9-HA and +E9-Flag with ΔE8-HA and +E9-HA. These results suggest that all the splice variants can make homodimers as well as heterodimers between MT and Δ E2, and between Δ E2 and +E9. Although it cannot be excluded that they form multimers such as trimers or tetramers, each splice variant makes specific interactions. It should be noted that MT forms a heterodimer only with Δ E2, and the C-terminal sequences of them are the same suggesting C-terminal sequence is important for heterodimer formation.

3.4. Expression of alternative splice variants during cell cycle progression

It has been reported that *patufet* which is the *Drosophila* homolog of *SPS1* is expressed highly in actively dividing cells and its deletion caused embryonic lethality during fly development [19,26]. More recently it was also reported that knockdown of *Drosophila* SPS1 led to growth inhibition of cells [22]. These reports suggest that the expression of hSPS1 can be controlled during the cell cycle. We examined the expression level of alternative splice variants during the cell cycle. After synchronizing the cells at G1/S phase, the synchronized cells were then released to proceed through the cell cycle by removing aphidicolin. At 6, 12, and 18 h after release, cells were in the mid-S phase, in the G2/M phase, and in the G1/S phase, respectively. However, at 24 h after release, the cells were detected in all phases of the cell cycle (Fig. 4A).

The expression levels of each alternative splice variant were measured by real-time PCR as described in the Materials and methods. As shown in Fig. 4B, the expression of the MT variant was gradually increased until the G2/M phase and then decreased. For example, cells arrested at the G1/S phase by aphidicolin treatment showed that the MT variant expression level was decreased to 56% compared to that in asynchronized cells (*P*<0.05). The level of

MT expression in the cells of the G2/M phase was increased to 160% compared to that of cells arrested at the G1/S phase (*P*<0.05). The levels of MT were similar at 0 h and 18 h when the cells were in G1/S phase. ΔE2 expression was peaked at mid-S phase and then gradually decreased. ΔE8 showed similar expression patterns as MT, but +E9/+E9a levels were decreased gradually as time lapsed suggesting the formation of +E9/+E9a was impaired by synchronization. These results suggest that alternative splicing of hSPS1 mRNA is closely related with cell cycle control.

4. Discussion

Five alternative splice variants of hSPS1 mRNA (MT, Δ E2, Δ E8, +E9 and +E9a) were identified in HeLa cells. Through searching the web browser of the University of California at Santa Cruz, four of these variants could be predicted with only the exception of +E9a. Interestingly, +E9a contains an extra 29 nucleotide extension of +E9. In genomic DNA, there is no gap between the 3' end of +E9 and the 5' end of +E9a. Furthermore, the consensus sequence of intron/exon junction (GT/AG) occurs at the 5' and 3' boundary of the additional sequence of +E9a suggesting +E9a is generated by attaching a part of intron 9 to the exon 9 by mis-splicing (see Fig. 1 and Supplementary Fig. S2).

The average amount of each splice variant was different. Approximately 90% of SPS1 population consists of MT. Δ E2 and Δ E8 comprise 2% of the total SPS1 mRNA, while +E9 and +E9a make up less than 1%.

Although there are 5 alternative splice variants in hSPS1 mRNA, only four proteins are produced, because there is a translational termination codon in exon 9. In terms of protein size, the MT protein is the largest, ΔE2 the second, +E9 /+E9a the third and ΔE8 the smallest.

The hSPS1 splice variants can interact with specific combinations; MT with Δ E2 and Δ E8 with $+E9$. It should be noted that the C-terminal sequences of MT and $\Delta E2$ are the same and the C-terminus of ΔE8 and +E9 are truncated. These results suggest that the C-terminus of SPS1 is important for interaction.

As described in the Introduction, SPS1 participates in Ras mediated cell signaling in *Drosophila* [20,21]. MT is located on both the plasma membrane and nuclear membrane and the other splice variants in the cytoplasm. All splice variants interact in a specific manner. Therefore, it can be speculated that the interactions of SPS1 between these molecules mediate cellular signal transductions. However, interactions of splice variants with other proteins have not been found. Because only MT is localized in the membranes, membrane localizing signals may occur in MT. However, we could not find any known common membrane localizing signals suggesting MT may interact with other membrane localizing protein(s).

Each splice variant was expressed differentially depending on the transformed cell lines derived from various tissues. For example, MT and +E9/+E9a are expressed in highest levels in BJAB cells which were derived from B cells; ΔE2 was expressed in the highest level in 293 cells which were derived from kidney; and ΔE8 was expressed in highest level in MCF7 cells which were derived from breast tissue. It is not clear whether these expression patterns are specific to each tissue or they are changed during transformation. However, if we compare Supplementary Fig. S3 with Supplementary Fig. S4, the expression pattern of splice variants at least in liver and kidney was changed during cell transformation. The expression level of splice variants is also regulated in a specific manner during the cell cycle and this result is consistent with the proposal that SPS1 is required for cell growth and that SPS1 is an essential gene; i.e., when *Drosophila* SPS1 was knocked out, development

was halted at the embryonic stage. Our findings in the present study suggest that alternative splice variants of SPS1 play important roles in cell growth presumably by controlling cell cycle progression through their specific subcellular localization and interaction.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

Acknowledgments

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Abbreviations

Sec selenocysteine Sec_S Sec synthase

Fig. 1. Five types of alternative splice variants of hSPS1 mRNA

The black line on the top represents genomic DNA, on which exons are shown as a bar. Numbers designate exons of each predicted alternative splice variant. The open and filled area in the exons designate the untranslated region and the open reading frame, while open and closed triangles designate the position of translational initiation and termination codons, respectively. Bent lines within cDNA designate a deleted exon in each splice variant. The numbers underneath MT indicate nucleotide positions.

Fig. 2. Subcellular localization of hSPS1 alternative splice variants

Flag-tagged hSPS1 overexpression vectors which can express each alternative splice variant were transfected into HeLa cells. Forty eight hours after transfection, cells were fixed and permeabilized, then hSPS1 was detected by mouse anti-Flag antibody and Alexa 488 antimouse IgG (green). Nucleus was stained with PI (red) and cells were observed under the confocal microscope. Scale bars represent 10 μm.

Fig. 3. Interactions between each hSPS1 splice variant

Flag or HA tagged splice variant expression vectors were transfected in 293 cells in the combinations shown, extracts were prepared, splice variants were immunoprecipitated, electrophoresed and detected by western blotting.

Fig. 4. Expression of hSPS1 alternative splice variants during cell cycle progression

(A) FACS (fluorescent associated cell sorter) analysis of cells after release from synchronization. X axis represents DNA contents and Y axis % of cell number. (B) Quantification of the amount of splice variants during the cell cycle. Real-time RT-PCR was performed using the same primers given in Supplementary Table S2. Experiments were performed in triplicate, and error bars denote the standard deviation from the mean of three experiments. Statistical significance was tested by pairwise Student's *t*-test with Bonferroni correction for multiple testing $(p<0.05)$. Asy represents asynchronized cells.