THE p60-S6K1 ISOFORM OF RIBOSOMAL PROTEIN S6 KINASE 1 IS A PRODUCT OF ALTERNATIVE mRNA TRANSLATION

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Ribosomal protein S6 kinase 1 (S6K1) is a well-known downstream effector of mTORC1 (mechanistic target of rapamycin complex 1) participating primarily in the regulation of cell growth and metabolism. Deregulation of mTOR/S6K1 signaling can promote numerous human pathologies, including cancer, neurodegeneration, cardiovascular disease, and metabolic disorders. As existing data suggest, the S6K1 gene encodes several protein isoforms, including p85-S6K1, p70-S6K1, and p60-S6K1. The two of these isoforms, p85-S6K1 and p70-S6K1, were extensively studied to date. The origin and functional significance of the p60-S6K1 isoform remains a mystery, however, it was suggested that the isoform could be a product of alternative S6K1 mRNA translation. Herein we report the generation of HEK-293 cells exclusively expressing p60-S6K1 as a result of CRISPR/Cas9-mediated inactivation of p85/p70-S6K1 translation. Moreover, the generated modified cells displayed the elevated level of p60-S6K1 expression compared to that in wild-type HEK-293 cells. Our data confirm an assumption that p60-S6K1 is alternatively translated, most probably, from the common for both p70- and p85-S6K1 mRNA transcript and reveal a link between p60-S6K1 expression and such cellular processes as cell proliferation and motility. In addition, our findings indicate that the p60-S6K1 isoform of S6K1 may undergo a mode of regulation distinct from p70- and p85-S6K1 due to the absence of mTOR-regulated p60-S6K1 phosphorylation at T389 that is important for S6K1 activation.

Keywords: CRISPR/Cas9 technology, p60-S6 kinase 1 (p60-S6K1) isoform, Akt/mTOR/S6K1 signaling pathway.

Diverse environmental signals from growth factors, hormones, energy and nutrients converge at mammalian target of rapamycin complex 1 (mTORC1) signaling enabling cells to promote anabolic and inhibit catabolic processes [1, 2]. Aberrant mTORC1-dependent signaling can contribute to cancer, diabetes, obesity and several other pathological states [3].

A number of downstream effects of mTORC1, including protein biosynthesis, cell growth, proliferation and survival [4, 5] are mediated via ribosomal protein S6 kinase 1 (S6K1), a well-studied mTORC1 substrate.

The S6K1 gene (RPS6KBI) was shown to encode two well-known S6K1 isoforms, p85-S6K1 and p70-S6K1, that differ only by the presence of the N-terminal 23 a.a. extension in p85-S6K1 due to the use of alternative (the first and second ATG) translational start sites [6]. Recently, it has been discovered that the splicing factor SF2/ASF promotes the expression of the oncogenic and the only known S6K1 splice variant termed p31-S6K1 or S6K1-isoform 2 that is truncated from the C-terminus [7]. A mechanism underlying oncogenic properties of p31-S6K1 is unclear but it seems to be kinase-independent, since the kinase domain of the given isoform is severely truncated.
According to the existing data, the usage of an alternative translation start site (the third ATG) may yield the expression of the additional putative S6K1 isoform [8], p60-S6K1, which is 30 a.a. and 53 a.a. shorter than p70-S6K1 and p85-S6K1, respectively, and has an intact kinase domain. However, there is no direct evidence supporting this assumption and one cannot exclude the possibility of p60-S6K1 generation via limited proteolytic degradation of the N-terminal kinase region.

Numerous experimental data suggest the implication of the S6K1 N-terminal domain in the regulation of kinase activity [reviewed in 2]. Thus, deletion of N-terminal 53 amino acids in p85-S6K1 (as in the p60-S6K1 isoform) inhibits S6K1 activity and phosphorylation similar to the action of mTOR inhibitor rapamycin [9]. Surprisingly, deletion of additional 24 a.a. (Δ77 a.a.) has rescued the negative effect of 53 a.a. deletion causing generation of a constitutively active S6K1 form [10]. Later on, a short sequence at the N-terminus of p70-S6K1 was identified to be important for mTOR-mediated S6K1 activation and was termed the TOS (mTOR signaling) motif. Deletion of the TOS motif or its mutagenic inactivation abolishes S6K1 kinase activity indicating the critical regulatory function of the N-terminus [11]. In addition, we have demonstrated that CK2-mediated phosphorylation of Ser-17 located within the p70-S6K1 N-terminus is implicated in the regulation of kinase nuclear export that could be a mode of its nuclear substrates regulation [12].

The above data indicate that the p60-S6K1 isoform should have regulation and functional activity in a cell different from that of p70-S6K1 and p85-S6K1, however, this aspect of p60-S6K1 is poorly investigated.

To date, several studies indicate an important role for S6K1 in cancer initiation and progression. Overexpression of the S6K1 gene and deregulated S6K1 signaling have been found in different malignancies, including breast, prostate, thyroid, brain and gastric cancers [13-21] that according to multiple studies correlate with poor prognosis of disease [14, 20-22]. However, implication of different S6K1 isoforms in carcinogenesis is poorly understood.

The aim of the present study was, firstly, to verify the hypothesis that p60-S6K1 is alternatively translated from the common for p70-S6K1 and p85-S6K1 template and, secondly, to evaluate the functional activity of the p60-S6K1 isoform in a cell.

For this purpose we applied the CRISPR/Cas9 technology to generate HEK-293 cell lines with disrupted expression of p70-S6K1 and p85-S6K1 isoforms by editing the DNA sequence between the second and hypothetical third alternative start of p60-S6K1 isoform translation. According to our data, inactivation of translation start sites for p70-S6K1 and p85-S6K1, located upstream to the third ATG, completely suppressed the expression of these isoforms, but had no inhibitory effect on expression of p60-S6K1 supporting an existing assumption that p60-S6K1 is alternatively translated from the third ATG.

Upon the generation of p85-/p70-/p60-HEK-293 cells, we elucidated proliferative and migratory properties of the cells comparing them with the p85/p70/p60-HEK-293 cells generated previously [23]. A state of S6K1 substrates phosphorylation has also been assessed. A phenotype of HEK-293 expressing only p60-S6K1 isoform confers both proliferative and migratory advantage over the S6K1 null cells, despite a decreased rate of cell proliferation and migration compared to the parental HEK-293 cells. The given cell model and the data obtained in this study can facilitate unraveling the p60-S6K1 cellular regulation and function.

Materials and Methods

CRISPR gRNA design and cloning. To create anti-S6K1 gRNA the http://crispr.mit.edu/ [24] web server was utilized. This gRNA design web tool allows for an off-target search over the genome and calculates quality scores for each gRNA selected. The higher score certain gRNA has the more specifically it binds to a target. The pair of oligonucleotides for targeting the region of the S6K1 gene with a highest score is:

Top gRNA strand: 5’-CACCCTCCTCAGAGCCCGCGGTCTCTC-3’
Bottom gRNA strand: 5’-AAACGGAGGACGGGGCTCTGAGGAC-3’

Underlined sequences correspond to the target S6K1 gene sequence, and unstressed sequences correspond to the Esp3I restriction site. The complementary oligonucleotide duplexes were ligated with the pSpCas9(BB)-2A-Puro (PX459) V2.0 cloning vector driven by the U6 promoter. This vector is dedicated for gRNA cloning and encodes the human Cas9 endonuclease necessary for gRNA functioning. The Esp3I restriction site was used for anti-S6K1 gRNA insertion. PCR amplification and DNA sequencing (Applied Biosystems™ 3130 DNA Analyzer) were used to prove that gRNA was inserted
into the indicated vector. The U6 primer was applied as a forward one (5′-GAGGGCCTATTTCCCAT-3′) and the oligonucleotide conforming to the bottom gRNA strand was employed as a reverse primer (5′-AAACGAGGACGCAGGCTCTGAGGAC-3′).

Cell culture and transfection. The human embryonic kidney cell line HEK-293 was maintained in Dulbecco’s Modified Eagle’s Medium (DMEM) supplemented with 10% fetal bovine serum (FBS) [GIBCO], 100 units/ml penicillin, and 100 μg/ml streptomycin, and cultured at 37 °C (5% CO2). Derivatives of HEK-293 cells expressing Cas9 and anti-S6K1 gRNA were developed by transfecting the HEK-293 cells [jetPEI transfection reagent (Polyplus-transfection ® SA)] for 24 h with the pSpCas9(BB)-2A-Puro (PX459) V2.0 vector encoding corresponding gRNA. Selection was carried out by the incubation of the transfected HEK-293 cells in puromycin containing media (4 μg/ml) for 24 h. Single clones were isolated for further analysis.

Antibodies. Rabbit polyclonal antibodies and mouse monoclonal antibodies (C3/10) specific to the C-terminal region of S6K1 (amino acids 453-525 in p85-S6K1) were generated as described in [25, 26]. The β-actin antibody was obtained from Sigma-Aldrich. Anti-phospho-p70-S6K1 (Thr389), anti-phospho-pS6 (Ser240/244), anti-phospho-eEF-2K (Ser366), and anti-phospho-rpS6 (Ser235/236), anti-phospho-p70-S6K1 (Thr389), anti-phospho-p54-S6K1 (Thr546), and anti-phospho-p52-S6K1 (Thr525 in p85-S6K1) were generated as described in [25, 26]. The β-actin antibody was obtained from Sigma-Aldrich. Anti-phospho-p70-S6K1 (Thr389), anti-phospho-pS6 (Ser240/244), anti-phospho-eEF-2K (Ser366), and anti-phospho-rpS6 (Ser235/236) were obtained from Cell Signaling Technology. Anti-mouse and anti-rabbit antibodies coupled to horseradish peroxidase were from Jackson ImmunoResearch.

Cell lysis, immunoprecipitation and Western blotting. Cells were lysed on ice for 30 min in 20 mM Tris-HCl, pH 7.5, 150 mM NaCl, 1 mM EDTA, 1 mM EGTA, 0.5% Triton X-100, supplemented with a Complete EDTA-free protease inhibitor cocktail tablet (Roche) and phosphatase inhibitors (Sigma-Aldrich). Lysates were cleared by centrifugation at 12,000 g for 15 min at 4 °C. Bradford assay was used to calculate protein concentrations. The precipitating antibodies were incubated with protein-A-agarose (Pierce) for 1 h followed by incubation with a cell lysate overnight at 4 °C with shaking. Immunoprecipitates were washed three times with a lysis buffer before the immune complexes were eluted from beads with Laemml loading buffer. Whole-cell lysates and immunoprecipitates were resolved by 10% SDS-PAGE. The proteins were transferred to a polyvinylidene difluoride (PVDF) membrane. The membrane was blocked with 5% low-fat milk in TBS-T (50 mM Tris-HCl, pH 7.4, 150 mM NaCl and 0.1% Tween-20) and after that it was incubated with the indicated primary antibodies used according to the manufacturer’s instructions. Antibodies against the S6K1 C-terminus were diluted 1 : 2500 (0.4 μg/ml) and incubated overnight at 4 °C. Afterwards, secondary antibodies linked to horseradish peroxidase were diluted 1 : 10000 and incubated at room temperature for 1 hour. The proteins of interest were revealed using enhanced chemiluminescence reagent (GE Healthcare).

Immunofluorescence. The wild-type HEK-293 and p85-/p70-/p60−HEK-293 cells were plated on glass coverslips in a 24-well plate and incubated to reach 70-80% confluence. Afterwards, cells were fixed with 10% neutral buffered formalin for 15 min and then washed three times with PBS. After fixation, 0.2% Triton X-100 in 100 μl of PBS was added to permeabilize the cells. To block non-specific staining the coverslips were incubated in a blocking solution (10% fetal calf serum in PBS) for 30 min at 37 °C. The cells were immunostained with rabbit polyclonal antibodies against the S6K1 C-terminus [25] diluted 1 : 100 in PBS overnight at 4 °C, followed by incubation with secondary FITC-labeled anti-rabbit antibodies (Jackson ImmunoResearch) diluted 1 : 400 in PBS. Thereafter, the samples were mounted on microscope slides using the Mowiol medium (Sigma) containing 2.5% DABCO (Sigma) and 0.5 mg/ml of DAPI (Pierce). Images were taken using the Zeiss LSM 510 META microscope (Germany).

In vitro scratch assay. In the present study, the wtHEK-293, p85−/p70−/p60−HEK-293 and p85−/p70−/p60−HEK-293 cells were plated (5×10⁴) in 6-well plates and grown to 90% confluence. The confluent monolayer was scraped with 200 μl sterile pipette tips to leave a scratch. After the scratch was created, digitized images of a wound closure were taken at 0 h and 24 h with the Leica DM 1000 microscope and a digital camera. The average distance (μm) between the wound edges was determined at each time point using the ImageJ software. The scratch wound assay was performed in triplicates for each cell line used in this study.

MTT assay. For the MTT assay, the wtHEK-293, p85−/p70−/p60−HEK-293 and p85−/p70−/p60−HEK-293 cells were plated in a 96-well culture plate at 5×10⁴ per well in 100 μl of DMEM supplemented with 10% FBS. The cells were incubated
for 24, 48 and 72 h. At each time point, 20 µl of 5 mg/ml MTT (thiazolyl blue tetrazolium bromide, Sigma-Aldrich) were added to corresponding wells. After incubation with MTT for 3 h and removing the medium, 150 µl of DMSO were added to each well and formazan crystals were dissolved. Absorbance was recorded with the microtiter plate reader, BioTek ELx800 (BioTek Instruments, Inc.), at 570 nm. The MTT assay was carried out thrice for each cell line tested in the study.

Statistical analysis. All quantitative results are expressed as the means ± standard deviation (SD). All experiments were performed independently and at least three times. The Student’s t-test was used to compare mean values between experimental groups. \( P < 0.05, \ P < 0.01, \) and \( P < 0.005 \) were considered to be significant.

Results and Discussion

It is generally recognized that the S6K1 gene encodes two protein isoforms (p70-S6K1 and p85-S6K1) generated by alternate ATG start site utilization. In addition, a smaller and the only known splice variant of S6K1 has been reported, p31-S6K1, that is required for cellular transformation induced by the splicing factor SF2/ASF (splicing factor 2/alternative splicing factor) [7].

According to the existing data, utilization of the third alternative ATG start site in S6K1 mRNA transcript may trigger translation of additional, truncated from the N-terminus, kinase isoform – p60-S6K1 [8]. Based on this assumption we hypothesized that disruption of a region located between the second and the hypothetical third ATG by the CRISPR/Cas9 editing tool may lead to the silencing of p85/p70-S6K1 protein expression enabling the cells to express solely p60-S6K1 [8]. Thus, we designed gRNA oligonucleotides (20 nt in size) corresponding to 106-125 nt in the p85-S6K1 mRNA coding sequence for targeting the S6K1 gene region in human embryonic kidney cells HEK-293 (Fig. 1). The DNA vector encoding both the Cas9 endonuclease and the constructed gRNA oligonucleotide was incorporated into HEK-293. After selection individual clones were analyzed by Western blot and immunofluorescent staining using anti-S6K1 antibodies specific to its C-terminal region. According to the data of Western blot analysis, CRISPR/Cas9-mediated gene editing induced the generation of a set of clones with a different S6K1 forms expression profile (Fig. 2, A). However, among six clones selected for analysis only two (clone 2 and 6) met our expectations, since they express exclusively the p60-S6K1 isoform. The specificity of p60-S6K1 expression was further confirmed by an immunoprecipitation assay. As shown in Fig. 2, B, alternative anti-S6K1 mAbs (C3/10) specific to the S6K1 C-terminus [26] precipitated p60-S6K1 from the CRISPR/Cas9-modified p85/p70/p60 HEK-293 cells, determining the specific recognition of the p60-S6K1 isoform in the samples verified in Western blot by rabbit polyclonal anti-S6K1 antibodies.

Thus, it is the first experimental evidence confirming that the p60-S6K1 isoform is a product of alternative mRNA translation initiated from the third ATG. In favor of a given conclusion are our previous data demonstrating that inactivation of all three S6K1 translation starts, including the hypothetical third one, abolished expression of all S6K1 isoforms [23].

![Fig. 1. A scheme of p85-S6K1 mRNA coding for p85/p70/p60-S6K1 isoforms with indication of alternative translation starts and a region that is targeted by gRNA in the S6K1 gene. Specifically designed gRNA is directed to the indicated DNA sequence to shut off the expression of both p85-S6K1 and p70-S6K1 isoforms. A black box corresponds to the approximate location of the gRNA target site (34-53 nt downstream of the second ATG site)](image-url)
Apart from the clones with detected p60-S6K1 expression, clone 5 did not display the expression of any S6K1 isoform, though clone 1 expressed the 80 kDa immunoreactive protein that does not correspond to any of known S6K1 forms. The possible explanation of such an effect of CRISPR/Cas9-mediated targeting of the S6K1 gene can be linked to the nature of the gRNA/Cas9 action. The CRISPR/Cas9 system causes the error-prone NHEJ repair to fix double-strand breaks in DNA. The indicated DNA repair system generally introduces either insertions or deletions, thus leading to unpredictable changes in a target sequence. In our case, it is possible that CRISPR/Cas9 induced specific mutagenic changes affecting in some cases the third ATG and, as a consequence, the disruption of p60-S6K1 expression or affecting mRNA splicing causing the expression of unusual 80 kDa S6K1 forms.

Another interesting observation was that the disruption of p85/p70-S6K1 expression (Fig. 2, A, clone 2 and 6) or even their down-regulation (Fig. 2, A, clone 3) correlated with the up-regulation of p60-S6K1 expression.

The generated HEK-293 clones with the disrupted expression of the S6K1 isoforms was further analyzed by the immunofluorescent detection of S6K1 with anti-S6K1 Abs. In overall, the data of immunofluorescent analysis (Fig. 3) confirmed the data of Western blot, but the detectable level of the p60-S6K1 protein in the p85/p70/p60-HEK-293 cells was even higher than the total level of S6K1 forms in the parental HEK-293 cells that differs from the data of Western blot (Fig. 2, A). It seems that truncation of the N-terminal region may enhance recognition of the S6K1 C-terminal domain by specific antibodies which is consistent with existing data suggesting interaction of both N- and C-terminal regions in the inactive S6K1 form [2].

Recently, we have reported the characterization of S6K1-deficient HEK-293 cells (p85-/p70-/p60-HEK-293) generated by the application of the CRISPR/Cas9 technology. We have demonstrated that the disruption of S6K1 isoforms expression had no effect on phosphorylation of S6K1 substrates analyzed [23] that could be explained by the presence of the intact highly homologous S6K2 kinase that shares with S6K1 most substrates. At the same time, we found the significant inhibition of p85-/p70-/p60-HEK-293 proliferation and migration as well as the inhibition of Akt activity, based on the extent of Ser-473 phosphorylation, known to be critical for cell survival and cell proliferation [23]. In the present study, analysis of p85/p70/p60-HEK-293 cells revealed that selective silencing of only p85- and p70-S6K1 isoforms has very similar effect. We did not find any effect on phosphorylation of some known S6K1 substrates (Fig. 4) and detected the inhibition of cell proliferation and motility (Fig. 5, 6) that accompanied reduction of Akt Ser-473 phosphorylation (Fig. 4). It should be noted that the extent of Akt phosphorylation was significantly reduced only at the condition of serum depletion. However in case
Fig. 3. Immunofluorescent analysis of p85^-/p70^-/p60^+ HEK-293 generated by CRISPR/Cas9 (clone 6) with the antibody specific to the S6K1 C-terminus. Wild-type HEK-293 and the generated clone 5 (knockout of the p85-, p70- and p60-S6K1 protein expression) were used as a positive and negative control, respectively. Confocal images of the HEK-293 and p85^-/p70^-/p60^+ HEK-293 cells are shown.
Akt phosphorylation was downregulated not only in serum-starved cells, but at regular cell growth conditions as well [23]. It is a well-known fact that the activation of p85/p70-S6K1 requires mTORC1-mediated phosphorylation of Thr-412/389 [27-29]. Our investigation of Thr-412/389 phosphorylation in p60-S6K1 with the phospho-Thr412/389 specific antibodies revealed that this site remains inactive in p60-S6K1 under different growth conditions in p85-/p70-/p60+HEK-293 (Fig. 4). These data suggest that the p60-S6K1 isoform functions differently from the p85/p70-S6K1 main isoforms and could be a subject of a different regulation mode that doesn’t require mTORC1-dependent phosphorylation. Nevertheless, the ability of p60-S6K1 to be phosphorylated at Thr-412/389 in MCF-7 cells had been highlighted in [8]. Indeed, the latter result seems to be somewhat surprising taking into account that hypothetical p60-S6K1 mRNA lacks an N-terminal region where the binding site for Raptor (TOS motif) is located.

The Raptor protein was determined to be a critical scaffold protein functioning within mTORC1 [30]. In contrast to [8], our observation of the absence of Thr-412/389 phosphorylation in p60-S6K1 is completely consistent with the idea that the p60-S6K1 isoform cannot be phosphorylated at this site by mTORC1 due to a lack of the TOS motif.

However, one cannot exclude other possibilities of p60-S6K1 control via phosphorylation of the Thr-412/389 site through the signaling pathway(s) distinct from mTORC1 signaling. Yet results reported in [8, 31] could underscore the presence of a linkage between other kinase/kinases activity and p60-S6K1 phosphorylation at Thr-412/389 in a breast carcinoma cell line MCF-7 [8]. Despite this, the HEK-293 cells used in our study seem not to involve such the kinase/kinases in p60-S6K1 regulation. In case of p60-S6K1 expressed in the p85-/p70-/p60+HEK-293 cells, the absence of the Thr-412/389 phosphorylation event arises intriguing questions about p60-S6K1 activity. If the activity of the major S6K1 isoforms is dependent on mTORC1-mediated phosphorylation, this could explain the lack of Thr-412/389 phosphorylation in p60-S6K1.

Fig. 4. Alterations in Akt/mTOR signaling under the condition of CRISPR/Cas9-mediated p85- and p70-S6K1 knockout in HEK-293 cells. The wild-type HEK-293 and p85-/p70-/p60-HEK-293 cells were starved in serum-depleted DMEM for 24 h and then restimulated with 20% FBS for 1 hour. Cell lysates were analyzed by Western blotting using the indicated antibodies. Blotting with antibodies against β-actin was used to confirm equal loading of proteins. rpS6, ribosomal protein S6; PKB, protein kinase-B; S6K1, ribosomal protein S6 kinase 1; eEF-2K, eukaryotic elongation factor 2 kinase; FBS, fetal bovine serum.

Fig. 5. HEK-293 cells expressing the sole p60-S6K1 isoform exhibit a partially restored ability to proliferate compared to S6K1 knockout HEK-293. Growth rates of the p85-/p70-/p60-HEK-293, p85-/p70-/p60-HEK-293 and wtHEK-293 cells were defined by the MTT assay. The data are the mean ± SD for three independent experiments. *P < 0.005, p85-/p70-/p60-HEK-293 versus wtHEK-293 and p85-/p70-/p60-HEK-293 versus p85-/p70-/p60-HEK-293; **P < 0.05, p85-/p70-/p60-HEK-293 versus wtHEK-293.
Fig. 6. Analysis of cell migration by the in vitro scratch assay reveals a partial recovery of the migration rate of HEK-293 expressing sole p60-S6K1 compared to S6K1 knockout HEK-293 cells. The confluent HEK-293, p85/p70/p60 HEK-293 and p85/p70/p60 HEK-293 cells were scratch-wounded and images were acquired 0 and 24 h post-scratching. A – Bars represent the mean ± S.D. for three independent experiments. * (P < 0.05) and ** (P < 0.01) versus wtHEK-293. B – Representative images of a wound closure from three independent experiments are presented.

phosphorylation, does the novel p60-S6K1 isoform possess kinase activity and what substrates are? Are well-known phosphorylation events common for p85- and p70-S6K1 activation except for Thr-412/389 phosphorylation sufficient for the control of p60-S6K1 function? What other mechanisms of p60-S6K1 regulation could exist? These questions will be addressed in future studies.

S6K1 has been linked to the regulation of cell proliferation [32] and motility [33, 34] that are affected in transformed cells and numerous data suggest implication of S6K1 in carcinogenesis [13-21]. To evaluate the contribution of the S6K1 isoforms to the modulation of cell proliferation and cell motility we have compared the p85-/p70-/p60-HEK-293, p85-/p70-/p60 HEK-293 and parental HEK-293 cells.
As it has already been mentioned disruption of all three S6K1 isoforms (p85, p70, and p60) or only two of them (p85, p70) had an inhibitory effect on cell proliferation and motility, however, the extent of inhibition was not so profound in case of sustained p60-S6K1 expression. It seems that expression of p60-S6K1 to some extent rescues cells from inhibitory effect caused by the disruption of p85- and/or p70-S6K1 expression.

In summary, we provide direct evidence supporting an assumption that expression of the p60-S6K1 isoform is determined by mRNA translation initiated from the third alternate ATG. In addition, we conclude that a phenotype of p85-/p70-/p60+HEK-293 has proved to be distinct from that of the S6K1 knockout cells suggesting different functional activity of p60-S6K1 in a cell. Future research on discerning p60-S6K1 regulation and function within a cell is required and will enable us to understand a role for p60-S6K1 in cellular physiology, as well as its impact on carcinogenesis.

Acknowledgements. This work was supported in part by a fellowship from Boehringer Ingelheim Fonds (BIF) awarded to Igor Zaiets.
кодирует несколько протеиновых изоформ киназы, включая p85-S6K1, p70-S6K1 и p60-S6K1. На сегодняшний день наиболее изученными являются изоформы p85-S6K1 и p70-S6K1. Происхождение и функциональное значение изоформы p60-S6K1 остаётся непонятным, однако существует предположение, что она может быть продуктом альтернативной трансплации мРНК. В работе представлены данные о создании клеточной линии HEK-293, в которой экспрессируется исключительно p60-S6K1 вследствие CRISPR/Cas9-опосредованной инактивации трансляции протеинов p85- и p70-S6K1. Более того, в созревших клетках выявлен повышенный уровень экспрессии p60-S6K1 по сравнению с исходными клетками HEK-293. Наши результаты подтверждают предположение, что изоформа p60-S6K1 образовывается в результате альтернативной трансляции, скорее всего, общего для p85- и p70-S6K1 транскрипта и раскрывают связь между экспрессией p60-S6K1 и такими процессами, как клеточная proliferация и подвижность. Кроме того, наши данные указывают на то, что изоформа p60-S6K1 может подвергаться отдельному способу регулирования по сравнению с p70-S6K1, о чём свидетельствует отсутствие mTOR-зависимого фосфорилирования p60-S6K1 по Т389, важного для активации киназы.

Ключевые слова: CRISPR/Cas9-технология, p60-S6 киназа 1 (p60-S6K1), Akt/mTOR/S6K1-сигнальный путь.

References


Received 13.04.2018