UDC 577.218+616-006

OVEREXPRESSION/KNOCKDOWN OF ADAPTOR PROTEIN RUK/CIN85 IN HUMAN LUNG ADENOCARCINOMA A549 CELLS RESULTS IN OPPOSITE CHANGES BETWEEN MMP-2/MMP-9 EXPRESSION LEVELS/ACTIVITIES AND CELL'S INVASION

M. I. BEKALA^{1,2}, D. S. GERASCHENKO¹, O. V. KHUDIAKOVA¹, T. D. SKATERNA¹

¹Palladin Institute of Biochemistry of NAS of Ukraine, Kyiv ²Taras Shevchenko National University, Kyiv, Ukraine

E-mail: markbekala2@gmail.com

Received 2023/03/16 Revised 2023/04/14 Accepted 2023/04/28

To acquire the ability to metastasize, cancer cells undergo molecular reprogramming in the course of epithelial-mesenchymal transition resulting in an increased motility and invasiveness [1]. Cells with elongated spindle-like mesenchymal phenotype require ECM-degrading enzymes, mainly MMP-2 and MMP-9, to generate the path for migration. In our previous works we demonstrated that overexpression of adaptor protein Ruk/CIN85 in breast cancer cells was associated with their aggressive metastatic behavior [2, 3]. In this study we aimed to investigate the changes in MMPs expression and activity as well as invasiveness of human lung adenocarcinoma A549 cells with up-/down-regulation of Ruk/CIN85.

Methods. We used A549 cells with stable overexpression (subline RukUp) and knockdown of Ruk/CIN85 (subline RukDown), as well as corresponding vector control sublines Mock and Scr.

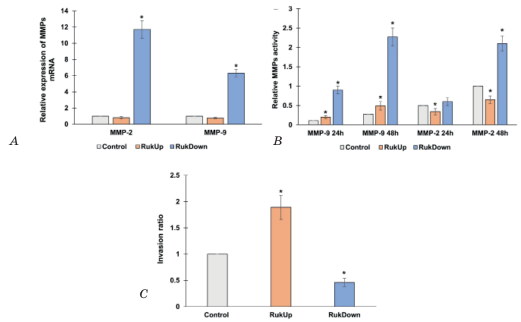


Fig. Up-/down-regulation of Ruk/CIN85 in human lung adenocarcinoma A549 cells lead to opposite changes between MMP2/MMP9 genes expression (*A*); their enzymatic activities (*B*) and invasion ratio of cell sublines (*C*):

 $M \pm m$, n = 3; * — P < 0.05 compared to the corresponding control

Cells were cultured in the complete DMEM medium under standard conditions. mRNA expression levels were estimated by RT^2 -PCR, enzymatic activity was assessed using gelatin zymography. Invasiveness of cancer cells was studied using Boyden chambers coated with Matrigel.

Results. Analysis of mRNA expression of MMPs in RukUp and RukDown cells revealed that MMP-2 and MMP-9 were preferentially overexpressed in RukDown cells, while RukUp subline did not exhibit significant difference compared with corresponding control (Fig. A). These findings were confirmed and complemented by study of enzyme activities. As can be seen from Fig. B, the gelatinolytic activities of both MMP-2 and MMP-9 were dramatically increased in RukDown subline, compared to respective control (Fig. B). Surprisingly, we revealed that MMPs regulation was inversely correlated with invasion potential of Ruk/CIN85 up/down A549 cells. In particular, it was established that invasiveness of RukUp cells was 2 times higher in comparison with respective control subline. Alternatively, invasion ratio was significantly decreased in RukDown cells (0.5 times) in comparison with control (Fig. C).

Discussion. In this study we found that expression level of Ruk/CIN85 in A549 cells is strongly associated with opposite changes between their invasiveness and MMPs expression/activities. It is known that the role of MMPs in carcinogenesis remains ambiguous. MMPs are involved in tumor progression. In particular, MMP's associated degradation of ECM components modulates cancer cells motility as well as leads to activation of proangiogenic factors in various cancerous tissues. On the other hand, MMP-2/MMP-9 take part in digestion of plasminogen resulting in generation of angiostatins (kringle-containing fragments of plasminogen) that could function as inhibitors of angiogenesis and tumor growth *in vitro* and *in vivo* [4].

Conclusions. According to the data received, it is possible to suggest that up-regulation of adaptor protein Ruk/CIN85 in A549 cells can lead to the very aggressive MMP-independent mode of migration that rely on cycles of expansion and contraction of the cell body mediated by the cortically localized actin and myosin [5].

Key words: Lung Adenocarcinoma; Motility; Invasion; Epithelial-Mesenchymal Transition; MMPs; Adaptor Protein Ruk/CIN85.

Author's contribution. Bekala M. I. has performed estimation of mRNA expression levels by RT²-PCR as well as was involved in zymography assay, data analysis and thesis writing. Geraschenko D. S. has received sublines of A549 cells with stable expression and knockdown of Ruk/CIN85. Khudiakova O. V. worked on the RNA extraction. Skaterna T. D. has performed zymography and Boyden chamber assays, as well as has curated research planning, data analysis and thesis writing.

Acknowledgments. We express our gratitude to Professor Liudmyla Drobot for scientific guidance and support of our work.

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