

Proline rich regions of coenzyme A synthase and interact with SH3 domains of signaling proteins *in vitro*

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Coenzyme A-synthases and (CoASy and CoASy) contain proline rich regions which may bring them into complexes with SH3-domain containing proteins. To test whether CoASy isoforms can bind to SH3 domains we performed in vitro pull down experiments. It was found that CoASy N-terminal extension, which is especially abundant in prolines, can interact specifically and directly with SH3 domains of tyrosine kinases Fyn and CSK, phospholipase C, NADPH oxidase activator 1 – p67phox, and cytoskeleton protein spectrin. Furthermore, C-terminal SH3 domain of p67phox can also interact with SH3 binding site that resides on the shared part of CoASy and CoASy. These data demonstrated that CoA Synthases could be involved in complexes with signaling proteins in living cells which may regulate enzymatic activities of CoA Synthases or vice versa CoA Synthase may modulate some steps in signal transduction in the cell in currently unknown way.

Key words: CoA Synthase, SH3 domain, proline rich regions, signaling proteins.

Introduction. Coenzyme A (CoA) is ubiquitous cofactor essential for living, involved in numerous enzymatic reactions in cell. Reduced CoA and its acyl- and acetyl-derivatives play a key role in the energetic, carbohydrate, and lipid cell metabolism [1], regulation of transcription via histone acetylation [2], proteins acylation [3], and oxidative stress protection via cysteamine biosynthesis [1]. Both quantity and form of available CoA and its derivatives depend on cell type and cell compartment, and vary in different physiological circumstances [1].

It has been shown that the level of intracellular CoA is regulated by various extracellular stimuli including insulin, glucose, fatty acids, pyruvate, glucagon, and

glucocorticoids [1, 6–8]. The rate of CoA biosynthesis changes with fasting, re-feeding, and several pathological conditions, such as diabetes, Reye syndrome, and cancer [1, 6–8].

However, very little is known about the pathways leading to such regulation. Up-to-date, the mechanisms of regulation are known only for the first enzymatic step in the pathway of CoA biosynthesis – phosphorylation of pantothenate by pantothenate kinase (PanK). There are at least four *PANK* genes in human genome (*hPANK1*, *hPANK2*, *hPANK3*, and *hPANK4*) and at least two alternative splicing isoforms for *PanK1* and *PanK2*. *PanKs* share the same catalytical core but are differently regulated by reduced CoA and its thioesters such as acetyl-CoA and acyl-CoA [4, 9–11]. Differential transcription of

PANKs mRNA in response to different stimuli such as hypolipidemic agents, glucose, and lipids was also shown to modulate intracellular CoA level [10].

4'-phosphopantetheine is one of the major pantothenate metabolites in cell along with CoA and pantothenate itself. It means that CoA Synthase (CoASy), which utilizes 4'-phosphopantetheine to generate dephosphoCoA and CoA, may be the second rate limiting enzyme in the pathway of CoA biosynthesis [4]. Nevertheless, no physiological regulators of CoASy have been reported so far. Recently two isoforms of CoA Synthase – CoASy₁ and CoASy₂ – have been cloned [13, 15]. We have demonstrated that CoASy₁ associates specifically with ribosomal S6 kinase – one of the key regulators of cell size and growth. The association was observed between native and transiently overexpressed proteins *in vivo* [15]. New interactions with signaling molecules could uncover novel mechanisms of CoA biosynthesis regulation.

The distinctive feature of CoASy₁ is a proline-rich 29 amino acids (29 aa) long N-terminal extension (Fig. 1, A). According to bioinformatic analysis this proline-rich sequence may bring CoASy₁ into different signaling complexes with Src homology 3 (SH3) domain containing proteins in cell. SH3 domains are small modular domains about 60 amino acids in size. They function as adapters, mediating protein-protein interactions, and are common domains for proteins building signaling networks and for some cytoskeleton proteins. SH3 domains are found in a wide range of functionally distinct signaling proteins including adapter proteins, protein kinases, lipid kinases, GTPases, GAPs, GEFs, lipases, protein phosphatases, etc. These domains bind with moderate affinity and selectivity to proline-rich ligands and play critical roles in numerous biological processes ranging from regulation of enzymes by intramolecular interactions, increasing the local concentration or altering the subcellular localization of components of signaling pathways, and mediating the assembly of large multiprotein complexes [5].

In this paper, we show that full-length CoA Synthase₁ and N-terminal 29 aa extension of CoASy₁ isoform can directly interact with SH3 domains of a number of signaling proteins *in vitro*.

Material and Methods. *Bioinformatics.* Scansite program was used for bioinformatical identification of protein motives, which could potentially bind to domains such as SH2 and/or SH3. Scansite search is based on the matrix of selectivity values for amino acids at each position relative to an orienting residue as determined by the oriented peptide library technique [11].

Plasmid Construction and Expression studies. The human EST clone 5187222, corresponding to the full length CoASy₁, was received from I.M.A.G.E. consortium and verified by sequencing. The coding sequence corresponding to the first N-terminal 29 aa of CoASy₁ was amplified by PCR and cloned into *pET42a* vector («Novagen», USA) in frame with GST and 6HIS (29 aa CoASy₁ -*pET42a*). 29 aa CoASy₁ -*pET42a* were expressed in *Escherichia coli* BL21 DE3 cells and recombinant protein was then purified using Ni-NTA agarose according to Ni-NTA Spin Handbook manual («Qiagen», USA).

Pull Down Experiments. In a pull-down assay, 1 µg of 6His-GST, 6His-GST-29 aa CoASy₁ fusion protein and His-CoASy₁ protein were incubated with Ni-NTA («Qiagen») for 2 h at 4 °C in a buffer for native condition (50 mM NaH₂PO₄, pH 8.0, 300 mM NaCl). The beads were then washed with the binding buffer to remove unbound proteins and incubated at 4 °C for 3 h with 1 µg of purified GST-SH3 domains [16]. Non-specific interactions were removed by washing with binding buffer and bound proteins were eluted by boiling in Laemmli sample buffer.

Immunoblot Analysis. Proteins were separated by SDS-PAGE and transferred onto polyvinylidene difluoride membrane. After that the membrane was incubated for 1 h in blocking solution (5 % non-skim dry milk in TBS/Tween 0.1 %). After blocking the membranes were cut on three pieces according to molecular weight marker to visualize 62 kDa full length CoASy₁-6His protein with CoASy₁ specific antibodies, SH3 domains with molecular weights ranging from 55 to 30 kDa with anti-GST antibodies or 28–25 kDa respectively 6His-GST-29 aa CoASy₁, and 6His-GST proteins with anti-6His antibodies. After washing with TBS, 0.1 % Tween, the membrane was incubated for 1 h with secondary horseradish peroxidase-conjugated anti-mouse antibody («Prome-

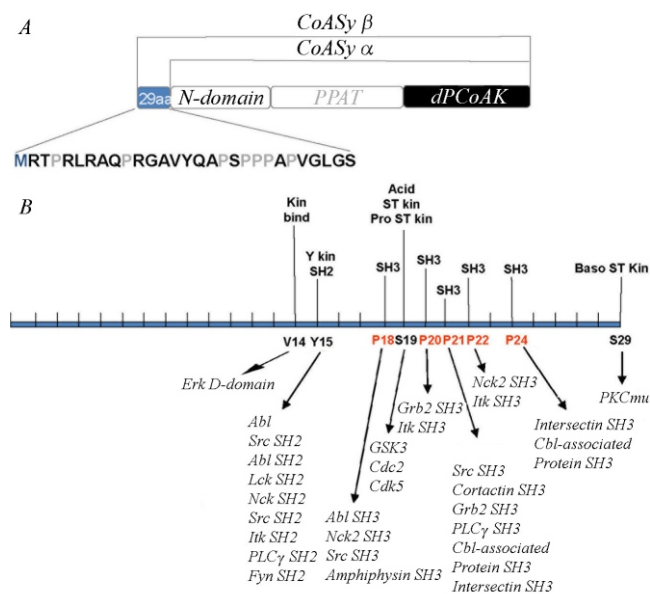


Fig. 1. Schematic representation of mammalian CoA Synthases: *A* – domain organization of mammalian CoA Synthases and ; dPCoAK – dephosphoCoA kinase domain; PPAT – phosphopantetheine adenylyltransferase domain; N-domain – N-terminal domains; amino acids are shown in the one letter code; *B* – bioinformatic prediction of interactive sites on CoASy N-terminal 29 aa extension. For prediction Scansite program was used (<http://scansite.mit.edu>)

ga)). The antigen-antibody complexes were detected using the ECL system («Amersham Biosciences»).

Results and Discussion. To elucidate specific function of CoASy isoform, which has 29 aa proline rich extension at the N-terminus, we employed bioinformatical approach. Using a medium stringency Scansite [12] search, in both isoforms we revealed potential binding sites for SH2/SH3 domains of different proteins. Moreover, recognition motives specific for SH3 domains of Crk, Abl, Nck2, PLC, Itk proteins were identified only in 29 aa N-terminal extension of the CoASy, whereas SH3 domains of cortactin, Grb2, Src, amphiphysin, Cbl-associated protein and intersectin had predictive binding sites residing on the CoASy specific extension as well as on the part of molecule, shared by and isoforms (Fig. 1, *B*). These predictions are an important indication of the possibility of CoASy involvement in complexes with signaling proteins in living cells.

At the next stage we tested interactions of full length CoASy and CoASy specific 29 aa extension

with SH3 domains of different intracellular proteins. For this purpose, we cloned cDNA coding for 29 aa fragment in frame with 5' sequence of GST-6His tag in *pET42a* vector («Novagen»). Fusion with GST-6His tags allowed 29 aa peptide to be expressed in bacteria and purified on Ni-NTA agarose («Qiagen») (Fig. 2, *B*). Purified protein was tested in Western blotting with rabbit polyclonal antibodies generated against the synthetic peptide composed from the first N-terminal 16 aa of CoASy [13] (Fig. 1, *C*).

To examine direct interactions of 29 aa CoASy peptide and full-length CoASy with SH3 domains, we used a panel of 13 SH3 domains originated from various mammalian proteins in pull down experiments. This panel was generated as SH3-GST fusion proteins expressed in *E. coli*. GST-6His-29 aa CoASy purified from bacteria and full-length 6His-CoASy which was purified earlier in our laboratory using baculovirus system [13] were immobilized on Ni-NTA agarose. SH3-GST domains were added to the resin coated with the bait proteins, incubated for 2 h in binding buffer and

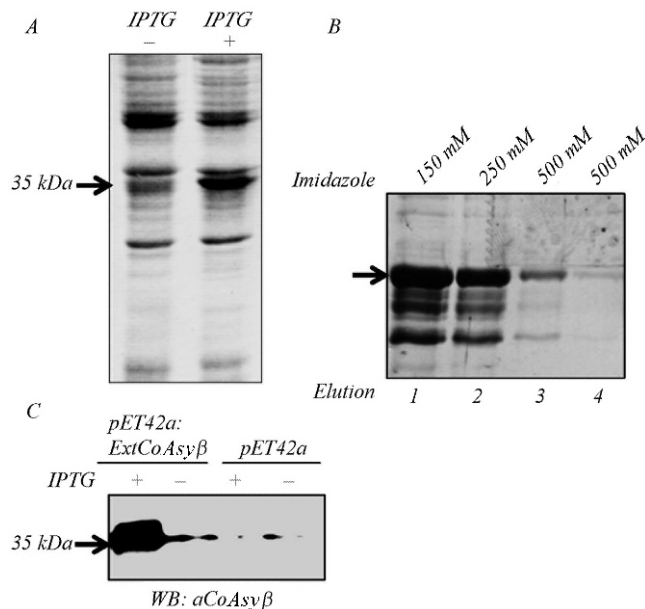


Fig. 2. Expression in *E. coli* and purification of GST-6His fused N-terminal 29 aa extension of CoASy: *A* – PAGE of total lysates of BL-21 cells transformed with His-GST-29 aa CoASy *pET42a* before or after induction with 100 μ M IPTG; Coomassie blue staining; *B* – PAGE of His-GST-29 aa CoASy fractions eluted from Ni-NTA agarose with different imidazole concentrations; Coomassie blue staining; *C* – Western blot analysis of purified 6His-GST-29 aa CoASy protein with rabbit polyclonal antibodies to N-terminal peptide of CoASy

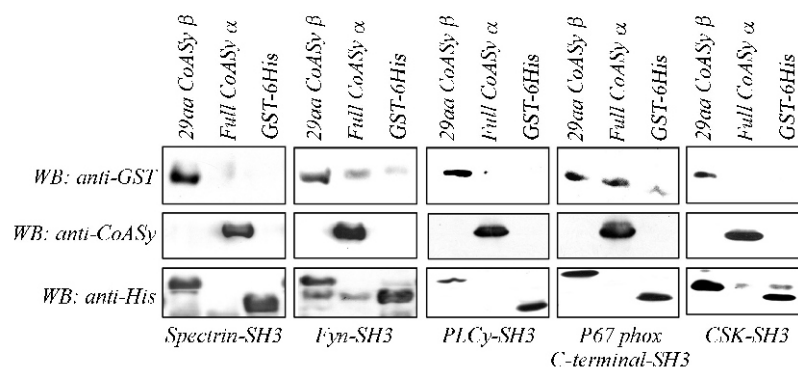


Fig. 3. *In vitro* binding of CoASy and 29 aa N-terminal extension of CoASy with SH3 domains of different proteins. The top panel, visualized with anti-GST antibodies shows binding of GST fused SH3 domains with immobilized 29 aa N-terminal extension of CoASy or full-length CoASy or with control 6His-GST protein. The middle panel, visualized with anti-CoASy antibodies, shows immobilized on Ni-NTA full-length 6His-CoASy protein. The low panel, visualized with anti-6His antibodies shows, immobilized on Ni-NTA 6His-GST-29 aa N-terminal extension of CoASy and 6His-GST protein

Summary of SH3 domains tested in the pull down experiment as direct interacting partners of CoASy N-terminal extension and full size CoASy protein

Tested SH3 domains	Binding to N-terminal extension of CoASy	Binding to CoASy	Protein function
Ruk1(SH3A)	-	-	Adapter protein involved in regulating diverse signal transduction pathways
Ruk1(SH3B)	-	-	-
Ruk1(SH3C)	-	-	-
Grb2	-	-	Adapter protein mediating coupling of activated growth factor receptors with proline rich regions containing ligands
Spectrin	+	-	One of the key components of membrane skeleton involved in membrane trafficking
c-Src	-	-	Non-receptor tyrosine protein kinase of Src family
p67phox C-terminal	+	+	NADPH oxidase activator 1
Fyn	+	-	Non-receptor tyrosine protein kinase of Src family
Crk	-	-	Adapter protein recruiting cytoplasmic proteins to the vicinity of tyrosine kinase
Fgr	-	-	Non receptor tyrosine kinase of Src family
PLC	+	-	Phospholipase C ; Hydrolyzes phosphatidylinositol 4,5-biphosphate (PIP2) in response to various extracellular stimuli to produce two second messengers, diacylglycerol and inositol 1,4,5-triphosphate
GAP(Ras)	-	-	GTPase activating protein of Ras
Csk	+	-	Non-receptor tyrosine protein kinase. Negatively regulates Fyn and other non receptor Src family tyrosine kinases

washed. As a control, non-specific binding of SH3-GST domains to Ni-NTA agarose along with immobilized GST-6His was tested for each SH3 domain. Interaction of SH3-GST domains with immobilized proteins was analysed by Western blot with GST specific antibodies.

Specific binding of different SH3 domains with N-terminal 29 aa extension of CoAsy and/or full-length CoAsy was detected (Fig. 3). Table summarizes results obtained in the pull down experiment. We showed direct specific *in vitro* interactions of N-terminal extension CoAsy with SH3 domains of Src family non-receptor tyrosine kinases Fyn and Csk, phospholipase C, NADPH oxidase activator 1 – p67phox, and structural membrane skeleton protein spectrin. Using GST-fused C-terminal SH3 domain of p67phox we showed that CoAsy can also directly interact with SH3 domains, and, contrary to N-terminal 29 aa proline rich sequence, has several separate proline rich motives. These data indicate that p67phox protein may have several binding sites on CoAsy molecule, one of them located on N-terminus of CoAsy and others – within the shared part of CoAsy and .

Noteworthy, in this experiment we confirmed interaction predicted by Scansite program for PLC SH3 domain. However, we failed to detect interactions between CoAsy and SH3 domains of tyrosine kinases Src, Crk, and adapter protein Grb2 predicted by ScanSite.

Although this type *in vitro* interaction experiments has to be validated for *in vivo* systems in co-immunoprecipitation experiments, we consider this approach to be suitable for primary screening of protein interactions. Our data indicate definitely that CoA Synthase isoforms can potentially form complexes with SH3 containing proteins involved in signal transduction in cell. Furthermore, CoAsy isoform has an additional N-terminal proline rich peptide. As it was demonstrated here, this proline rich peptide substantially extends the range of SH3 domains which can bind directly to CoAsy. Whether these interactions interfere with regulation of the rate of Coenzyme A biosynthesis, intracellular localization of CoAsy or with other intracellular functions remains to be elucidated.

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Збагачені проліном області коензим А-синтаз та взаємодіють з SH3 доменами сигнальних білків *in vitro*

Резюме

Кофермент А-синтази та містять збагачені проліном ділянки, які можуть сприяти їхньому комплексоутворенню з білками, що містять SH3 домени. Для перевірки цієї гіпотези ми провели експерименти in vitro по зв'язуванню повно-розмірного білка КоА-синтази та специфічного для КоА-синтази N-кінцевого пептиду із SH3 доменами сигнальних білків. Виявилось, що специфічний для КоА-синтази N-кінцевий пептид, багатий на пролін, специфічно взаємодіє з SH3 доменами тирозинових протеїнкіназ Фун і ССК, фосфоліпази С, активатора NADPH оксидази 1 – р67рhox і цитоскелетного білка спектрину. Крім того, SH3 домен р67рhox зв'язується із збагаченою проліном послідовністю, спільною для обох ізоформ. Отримані результати дозволяють припустити, що КоА-синтази перебувають у комплексах з сигнальними білками і, таким чином, відбувається регуляція їхньої ферментативної активності або ж КоА-синтази невідомим на даний час способом модулюють певні етапи передавання сигналів у клітині.

Ключові слова: КоА синтаза, SH3 домен, області збагачені проліном, сигнальні білки.

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Обогащенные пролином области кофермент А-синтаз и взаимодействуют с SH3 доменами сигнальных белков в условиях *in vitro*

Резюме

Кофермент А-синтазы и содержат богатые пролином области, возможно, способствующие их комплексообразованию с белками, содержащими SH3 домены. Для проверки этой гипотезы мы провели эксперименты по связыванию in vitro полно-размерного белка КоА-синтазы и специфического для КоА-синтазы N-концевого пептида с SH3 доменами сигнальных белков. Оказалось, что пролин-богатый N-концевой пептид КоА-синтазы специфически взаимодействует с SH3 доменами тирозиновых протеинкиназ Фун и ССК, фосфоліпазы С, активатора NADPH оксидазы 1 – р67рhox и белка цитоскелета спектрина. Кроме этого, SH3 домен р67рhox взаимодействует с обогащенной пролином последовательностью, общей для обеих изоформ. Полученные результаты позволяют предположить, что КоА-синтазы и образуют комплексы с сигнальными белками и, таким образом, регулируется ферментативная активность КоА-синтаз либо же КоА-синтазы неизвестным на данный момент способом модулируют определенные этапы передачи сигналов в клетке.

Ключевые слова: КоА-синтаза, SH3 домен, области, обогащенные пролином, сигнальные белки.

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