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Coenzyme A levels influence protein acetylation, CoAlation and 4'-phosphopantetheinylation: Expanding the impact of a metabolic nexus molecule

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ABSTRACT

Coenzyme A (CoA) is a key molecule in cellular metabolism including the tricarboxylic acid cycle, fatty acid synthesis, amino acid synthesis and lipid metabolism. Moreover, CoA is required for biological processes like protein post-translational modifications (PTMs) including acylation. CoA levels affect the amount of histone acetylation and thereby modulate gene expression. A direct influence of CoA levels on other PTMs, like CoAlation and 4'-phosphopantetheinylation has been relatively less addressed and will be discussed here. Increased CoA levels are associated with increased CoAlation, whereas decreased 4'-phosphopantetheinylation is observed under circumstances of decreased CoA levels. We discuss how these two PTMs can positively or negatively influence target proteins depending on CoA levels. This review highlights the impact of CoA levels on post-translational modifications, their counteractive interplay and the far-reaching consequences thereof.

1. Introduction

Coenzyme A (CoA) is essential for numerous metabolic reactions in eukaryotic and prokaryotic cells. CoA is involved in approximately 9% of all metabolic pathways including the tricarboxylic acid cycle, fatty acid regulation, amino acid synthesis and lipid metabolism [1,2]. In eukaryotes, the de novo biosynthesis of CoA occurs via five consecutive enzymatic steps starting with the substrate vitamin B5 (pantothenic acid/pantothenate). The five enzymatic steps are carried out by pantothenate kinase (PANK), phosphopantothenoylcysteine synthetase (PPCS), phosphopantothenoylcysteine decarboxylase (PPCDC), phosphopantetheine adenylyltransferase (PPAT) and dephospho-CoA kinase (DPCK) respectively (Fig. 1) [1]. In various organisms, including Drosophila melanogaster, mice and humans, PPAT and DPCK enzymatic activities are performed by a single bifunctional protein, CoA synthase (COASY) [3]. It was demonstrated that PANK is a rate limiting enzyme in the CoA biosynthesis pathway [4,5]. In eukaryotic cells, the canonical CoA biosynthesis pathway starts with the uptake of extracellular vitamin B5 via the sodium dependent multivitamin transporter [6]. In addition to this route, in Drosophila melanogaster S2 cells, the CoA precursor 4'phosphopantetheine can be taken up by the cells as well and converted into CoA [7]. It remains to be investigated whether this also occurs in mammalian cells. Indications that this may occur came from a recent publication, demonstrating that metabolic defects induced by the absence of PANK2 in the mouse brain are restored by orally administered 4'-phosphopantetheine, but not by pantetheine or vitamin B5 [8]. Although the mechanism of cellular uptake of 4'-phosphopantetheine remains to be investigated, these publications reveal that eukaryotic cells may be able to synthesize CoA directly, following the uptake of 4'phosphopantetheine.

The existence of alternative ways for cells to generate sufficient levels of CoA underscores the importance of CoA for cellular survival and its role in health and disease. The CoA biosynthesis pathway, compartmentalization of CoA and the essential influence of CoA and acetyl-CoA on metabolism and protein acetylation have been the focus of extensive excellent reviews [1,9–18]. What has received relatively little attention is the newly emerging understanding of the influence of CoA levels on other post-translational modifications of proteins (PTMs). This review discusses how the PTMs CoAlation and 4'-phosphopante-theinylation can be affected by enhanced or disturbed CoA *de novo* biosynthesis (Fig. 2).

2. CoA levels influence protein acetylation

PTMs refer to all covalent modifications that modify protein function. Acetylation is, in addition to phosphorylation and glycosylation, one of the most well-known forms of PTM [24]. Histones were the first acetylated proteins identified over 60 years ago [25,26] and now,

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Fig. 1. CoA biosynthesis pathway. In most eukaryotic cells, *de novo* CoA biosynthesis starts by taking up extracellular pantothenic acid, which is then converted by the action of five consecutive enzymes, PANK, PPCS, PPCDC, PPAT and DPCK, forming 4'-phosphopantothenic acid, 4'-phosphopantethenoylcysteine, 4'-phosphopantetheine and CoA respectively. PPAT and DPCK enzyme activities are combined in COASY in *Drosophila melanogaster*, mice and humans. In *Drosophila melanogaster* S2 cells, 4'-phosphopantetheine can be taken up as well and converted to CoA, only requiring the last enzyme COASY.

profiling of the acetylome in numerous cell types and organisms has revealed a broad spectrum of additional proteins that undergo acetylation [27], indicating that protein acetylation-modifying factors broadly influence cellular processes. Levels of protein acetylation on a lysine residue are mediated by lysine acetylating and deacetylating enzymes. The donor source for the acetyl group is acetyl-CoA, and events that affect acetylation of CoA also affect acetylation of proteins [28]. It is therefore to be expected that levels of CoA also influence levels of protein acetylation. Consistent with this, is the demonstration in C. elegans, Drosophila S2 cells and mammalian cells that a decrease in the cellular CoA content is associated with decreased histone and tubulin acetylation [19]. Such changes are restored upon replenishing CoA in these cells. Furthermore, an association between impaired CoA homeostasis and decreased tubulin acetylation was also demonstrated in mouse liver [29] and in Xenopus laevis embryos, changed levels of acetyl-CoA correlate with changed levels of acetylation of many proteins including histones H3 and H2B [30].

Other lines of research show that the ratio of acetyl-CoA to CoA critically influences overall histone acetylation levels. Most lysine acetyltransferases (KATs) including GCN5, PCAF and p300/CBP, bind acetyl-CoA and CoA with similar affinity, thus the activity of KATs might be more regulated by the ratio of acetyl CoA/CoA than by the concentration of the individual compounds [31]. Consistently, upon reducing the ratio of acetyl-CoA when culturing LN229 cell nuclei *in vitro* together with defined acetyl-CoA and CoA concentrations, histone

acetylation decreased, suggesting that the ratio of acetyl-CoA/CoA in the nucleus affects the levels of histone acetylation [32]. In addition, in a CoA synthase (COASY)-knockdown cell line, the acetylation levels of the majority of acetylated peptides were unchanged compare to control, in line with an unchanged ratio of acetyl-CoA/CoA, even though the individual levels of acetyl-CoA and CoA were decreased. This further underscores that the acetyl-CoA/CoA ratio may be a more relevant determinant of protein acetylation [33]. However, in the COASY knockdown cell line, a small set of hyperacetylated proteins, like CBP and TPX2, was found, leading to an increased activity and subsequently causing a prolongation of mitosis [33]. Those data indicate that decreased levels of acetyl-CoA and CoA may affect acetylation of a small set of proteins, whereas the ratio of acetyl-CoA/CoA influences overall levels of protein acetylation. Cellular compartmentalization of acetyl-CoA/CoA may help to explain the opposing observations of hyper versus hypo acetylation of specific proteins under CoA reducing circumstances and the consequences of decreased CoA levels versus those resulting from a changed ratio of acetyl-CoA to CoA. To further increase the complexity, an increase in KAT activity may lead to a local increase of free CoA and thereby the acetyl-CoA/CoA ratio and levels of protein acetylation are influenced. Taken together, these observations indicate the presence of a multifaceted impact of CoA levels on acetylation.

In addition to acetylation, forms of non-acetyl protein acylation also occur, including succinylation, butyrylation, malonylation, palmitoylation, propionylation, crotonylation and others [34-39]. Among these acylations, recent studies profiling the malonylome, succinylome and butyrylome have revealed the protein landscape influenced by these PTMs [40-42]. The corresponding acyl-CoAs are the likely source of these modifications, further underscoring the influence of CoA levels on cellular processes. Consistently, palmitoylation of the Transferrin receptor is reduced in fibroblasts from patients carrying mutations in PANK2, one of the rate-limiting enzymes in the de novo biosynthesis pathway of CoA. Adding a surplus of CoA to the culture medium restores the level of Transferrin receptor palmitoylation in these cells [43], suggesting a direct link between CoA availability and palmitoylation. Though various types of CoA acylation (succinyl-CoA, butyryl-CoA, etc.) can form PTMs, different CoA-acylation forms have various abilities to impact the efficiency of making such PTMs. KATs can catalyze propionylation and butyrylation of histones, but experiments indicate that KATs catalyze the transfer of acetyl groups more efficiently than propionyl and butyryl groups, suggesting that histone acetylation might occur more rapidly than propionylation and butyrylation [31]. However, malonyl-CoA and succinyl-CoA were observed to cause acylation of BSA more easily than acetyl-CoA [44,45]. UV spectroscopy analysis further confirmed that a malonyl thioester was cleaved much faster than its acetyl analogue [44]. Apart from such enzyme-mediated protein acylation, nonenzymatic protein acylation can also happen due to the interaction of acyl-CoA thioesters with nucleophilic cysteine and lysine residues. This process depends on the local acyl-CoA and nucleophile concentrations, and also the chemical potential energy of acyl-CoA/CoA couples that drive slower nonenzymatic side reactions [46]. These PTMs, which may be related to CoA/acylated-CoA levels, show various cellular functions (Supplementary Table 1). Additional evidence for a link between decreased levels of CoA and altered levels of these PTMs and cellular functional changes remain to be investigated. Altogether, it is likely that levels of CoA influence the protein acylation landscape and thereby a multitude of cellular processes.

3. CoA levels influence protein CoAlation

A recently identified PTM form that is directly influenced by CoA levels is CoAlation [21,23,47–50]. CoA contains a thiol group that can interact with other thiol groups to form a disulphide or mixed disulphide, such as CoA-cysteine and CoA-glutathione. Protein CoAlation was demonstrated by using a CoA-specific antibody to immunoprecipitate, purify and identify CoAlated proteins with mass spectrometry [51].

With this approach over 500 CoAlated proteins have been identified under various cellular conditions, including proteins involved in metabolism, protein synthesis, muscle development and stress responses [21,48,49]. CoAlation is a reversible PTM that is induced by fasting and oxidative stress [49]. Under PANK1_β (CoA de novo biosynthesis enzyme) overexpressing conditions, the level of CoAlation increased compared to the parental HEK293 cells, suggesting that levels of CoAlation correlate with levels of CoA [49]. Moreover, in vitro pre-incubation of CoA with recombinant protein Aurora A showed Aurora A CoAlation, and subsequent microinjection of CoA in mouse oocytes caused abnormal spindle formation and chromosome misalignment [23]. Since Aurora A is a key enzyme involved in mitosis, these data suggest that high levels of CoA might cause Aurora A CoAlation, which could subsequently impair mitosis. In addition, CoAlation was found to preferentially target metabolic enzymes under oxidative stress conditions, including CK (creatine kinase), GAPDH (glyceraldehyde 3-phosphate dehydrogenase), IDH2 (isocitrate dehydrogenase) and PDK2 (pyruvate dehydrogenase kinase), decreasing their activities [49]. It remains to be determined whether the level of CoAlation is decreased under conditions of decreased CoA.

Interestingly, CoAlation seems to be related to stress. In exponentially growing cells, protein CoAlation is present at a low level, but CoAlation levels can be dramatically induced under oxidative or metabolic stress. When the conditions of oxidative stress were decreased, CoAlation disappeared, which suggests that protein CoAlation is an antioxidant defense [23,49,52]. When HEK293/PANK1ß cells were under glucose and pyruvate deprivation conditions, the basal level of protein CoAlation was increased, which is consistent with the observation of induction of protein CoAlation in rat liver cells after fasting. The induction of protein CoAlation under metabolic stress can be relieved by recovering physiological metabolic conditions [49]. In summary, protein CoAlation is a newly identified and potentially influential posttranslational modification. It affects various cellular processes including protection from overoxidation, regulatory interactions between CoA and target proteins, modulation of activity of metabolic enzymes including CK, GAPDH, IDH2 and PDK2, subcellular localization of



Fig. 2. CoA level-related protein post-translational modifications. Decreased levels of CoA reduce protein acetylation, especially histone acetylation, resulting in changes of transcriptional programs [14,19]. The covalent attachment of 4'-phosphopantetheine to specific serine residues in specific proteins, such as acyl carrier proteins (ACPs), is called 4'-phosphopantetheinylation. In *Drosophila*, impaired *de novo* CoA biosynthesis result in the reduction of the availability of 4'-phosphopantetheinylation of ACPs (holo-ACPs). Decreased holo-mitochondrial ACP explains similarities of four genetically distinct CoA-linked neurodegenerative diseases [20]. In contrast, increased levels of CoA boost the covalent binding of the CoA thiol group with cysteine in proteins, forming protein CoAlation, which improves cellular antioxidant defense and leads to decreased activity of CoAlated enzymes [21–23].

CoAlated proteins and changes in conformation of proteins [21]. Since CoAlation has mainly been detected under oxidative or metabolic stress conditions, it will be important to investigate CoAlation under normal physiological conditions as well. Another intriguing question is what the (long-term) consequences are for cells and organisms when CoAlation is disturbed.

4. CoA levels influence 4'-phosphopantetheinylation of mtACP

Protein 4'-phosphopantetheinylation (hereafter referred to as 4PPTylation) is also a type of post-translational modification, directly linked to CoA levels, that affects cellular functions. 4'-Phosphopantetheine is a precursor and moiety of CoA (Fig. 1). Despite the discovery of 4PPTylation over 50 years ago, currently, only a few proteins [53], including 10-formyltetrahydrofolate dehydrogenase, nonribosomal peptide synthase, polyketide synthase [54,55], cytosolic fatty acid synthase and mitochondrial acyl carrier protein (mtACP) [56] have been shown to be 4PPTylated. The 4PPTylome has not been studied and most likely more proteins remain to be identified. The influence of 4PPTylation therefore remains largely unknown. Among the known 4PPTylated proteins, acyl carrier protein (ACP) has been widely investigated. Acyl carrier protein (ACP) was first found as a protein involved in mitochondrial fatty acid synthesis [57]. ACP was later identified as an essential subunit in many enzymes or protein complexes and it plays a key role in mitochondrial functions including lipoic acid metabolism, RNA processing, mito-ribosome function, assembly of respiratory chain complexes and iron-sulphur cluster biosynthesis [58]. A 4'-phosphopantetheine group is covalently attached to a conserved serine site in ACP and thus forms active 4PPTylated mtACP (also referred to as holomtACP) [59,60]. Holo-ACP synthetase catalyzes the transfer of 4'phosphopantetheine from CoA (note: free 4'-phosphopantetheine cannot serve as a direct source for 4PPTylation) to apo-ACP in E. coli according to the following reaction [61,62]:

$CoA + apo-ACP \xrightarrow{Mg^{++}} holo-ACP + 3', 53'$ -adenosine diphosphate

Since 4PPTylation of apo-ACP requires CoA, CoA levels may influence protein 4'-phosphopantetheinylation. Indeed, the turnover of the 4'-phosphopantetheine prosthetic group is dependent on the concentration of CoA in *E. coli*, and the highest rate of turnover was found at the lowest CoA concentration [63]. More recent research also indicated that 4PPTylation levels of mtACP (holo-ACP) are related to CoA levels in *Drosophila melanogaster*. When *Drosophila* S2 cells were treated with HoPan (an inhibitor of CoA biosynthesis), CoA levels decreased [7] and this resulted in reduced levels of holo-mtACP. Holo-mtACP levels recovered when S2 cells were supplemented with CoA [20].

New insights in four neurodegenerative diseases arose recently from the finding that 4PPTylation of mACP is influenced by CoA levels in Drosophila cells. Key to these insights is the knowledge that holo-mtACP is required for lipoylation of and thereby activation of PDH. Indeed, disrupted CoA metabolism led to decreased levels of holo-mtACP, decreased levels of lipoylated PDH-E2 and decreased PDH-E2 enzyme activity in Drosophila S2 cells [20]. This CoA-mtACP-PDH pathway now links the four neurodegenerative diseases: PKAN, CoPAN, MePAN and PDH-E2 deficiency [20]. PKAN and CoPAN are neurodegenerative diseases caused by mutations in two genes of the CoA biosynthesis pathway. Pantothenate kinase-associated neurodegeneration (PKAN) is caused by mutations in the PANK2 gene, and CoA synthase proteinassociated neurodegeneration (COPAN) is caused by mutations in the COASY gene. MePAN, also a neurodegenerative disorder, is caused by mutations in the gene encoding mitochondrial enoyl-[acyl-carrierprotein] reductase (MECR) [64]. The substrate of MECR is trans-2enoyl-holo-mtACP. MECR is required for the formation acyl-holo-mtACP and finally for the formation of lipoic acid. Lipoylation of PDH-E2 is indeed altered in MePAN patients [64]. Finally, PDH-E2-deficiency is a disease associated with mutations causing impairment of the pyruvate

dehydrogenase complex, PDH-E2. PKAN, CoPAN, MePAN and PDH-E2deficiency patients show overlapping clinical and neuroradiographic features, including abnormalities in the globus pallidus [20,64–67]. The CoA-mtACP-PDH axis in which 4PPTylation plays a central role, as demonstrated by Lambrechts et al. in S2 cells, together with the similar phenotypes of the diseases, suggest that the common element in these four diseases is an impairment of PDH activity. In a CoA metabolismdeficient (*Pank2* knock-out) mouse model, abnormalities in the globus pallidus region (GP) were observed together with a decreased activity of PDH and complex I [8]. However, levels of 4PPTylated mtACP (holomtACP) were not investigated.

Interestingly, the disruption of CoA metabolism by PANK2 depletion in mammalian HEK293 and SH-SY5Y cells also caused a reduction of holo-mtACP [20]. However, total CoA levels in HEK293 and SH-SY5Y cells transfected with PANK2 RNAi constructs were not changed, which is in contrast with the findings in S2 cells. These contradictory findings may be due to the variability in distribution of CoA in different subcellular compartments. It is possible that in mitochondria, where human PANK2 is located in the intermembrane space [68], local CoA and holo-mtACP are decreased. Such local compartmental changes could influence 4PPTvlation more than absolute levels of total cellular CoA [20]. CoA and acetyl-CoA are found within the mitochondria, cytosol, nucleus, peroxisomes and ER [69-71]. Total intracellular CoA levels fluctuate in response to metabolic changes [9,10], but the accuracy of measuring metabolites in individual subcellular fractions hampers conclusive interpretations. Because of technical issues regarding purity, efficiency, reproducibility, variability and recovery of different subcellular extractions, one should be careful to directly compare measurements of CoA/Acetyl-CoA between compartments [72]. It is also possible that acetyl-CoA and CoA levels differ between cellular locations that are not limited by a membrane. No techniques currently exist able to measure in situ, local concentrations of acetyl-CoA and CoA. Only after these technical limitations have been solved, an accurate determination of sub-compartmentalization of CoA/acetyl-CoA levels can occur. Subsequently, this information will help to better understand the flow of CoA and acetyl-CoA and how they affect several downstream processes in various tissues.

Taken together, disruption of CoA metabolism causes the reduction of 4PPTylated mitochondrial acyl carrier protein (holo-mtACP), and result in abnormal phenotypes in both *Drosophila* and mouse models of PKAN. These observations highlight the influence of CoA levels on 4PPTylation of mtACP and its importance in cellular metabolism in health and disease. However, whether the findings in *Drosophila*, possessing one pantothenate kinase gene, can also be translated to higher eukaryotes including humans, possessing four pantothenate kinase genes, remain to be investigated. In addition, a causative link between CoA levels and 4PPTylation in eukaryotes other than *Drosophila* also remains to be experimentally proven.

5. Future perspectives

In this review, we discuss how changes in CoA levels influence three different types of post-translational modifications including protein acetylation, CoAlation and 4'-phosphopantetheinylation, all with broad consequences for cellular physiology and underscoring the importance of CoA homeostasis. CoA is recognized as a nexus molecule in intermediary metabolism, serving as a central regulator linking these reactions to more global regulation of cell function. Yet many unanswered questions remain as we gain understanding of a new level of interdependencies and complexities of CoA-influenced processes. Of special interest is the example of the effect of CoA levels on PDH-E2 activity. As explained here, decreased CoA levels are, *via* decreased 4PPTylation, associated with decreased lipoylation of PDH-E2 and thereby decreased activity of PDH-E2. In contrast, an increase in CoA is associated with increased CoAlation of PDK causing decreased activity. PDK is an inhibitor of PDH-E2; thus an increase in CoA levels lead to an increase in

PDH-E2 activity. Ultimately, the activity of PDH-E2 can be positively influenced by increased CoA levels *via* increased CoAlation and negatively by decreased CoA levels *via* decreased 4PPTylation. It will be of interest to identify other enzymes with activities that can be positively and negatively influenced by CoA levels and to determine if CoAdependent post translational modifications are involved. For a deeper understanding of the complexity it will be essential to be able to determine subcellular levels of CoA and specific post-translational modifications. How we accomplish this will be a challenge for the future.

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CRediT authorship contribution statement

Yi Yu: Writing, Reviewing and Editing; Isabele Fattori Moretti: Writing, Reviewing; Nicola A. Grzeschik: Writing, Reviewing, Visualization; Ody C.M. Sibon: Writing, Reviewing and Editing, Supervision; Hein Schepers: Writing, Reviewing and Editing, Supervision.

Declaration of competing interest

The authors declare the following financial interests/personal relationships which may be considered as potential competing interests: Dr. Sibon is a co-inventor on 3 patent applications: (EP 2 868 662 A1, P-201400452, 1933-P48EP/15). Dr. Sibon serves as non-compensated executive for two not-for-profit organizations: the Stichting Lepelaar and the Spoonbill foundation.

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