Swinging arms in multifunctional enzymes and the specificity of post-translational modification
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Introduction
Covalently attached prosthetic groups serve as swinging arms in several different and well-characterized multifunctional enzymes. Notable among these are the lipoyl-lysine residues in 2-oxo acid dehydrogenase multienzyme complexes [1–3] and the glycine cleavage system [4–6], the biotinyl-lysine residues in various ATP-dependent carboxylases [7,8] and the phosphopantetheinyl-serine residues in fatty acid [9] and polyketide [10] synthases. In all these instances, the swinging arm is used to ferry substrate between the active sites that function successively in a multistep catalytic reaction. Several explanations have been proposed for the advantages supposedly conferred on a multifunctional protein by virtue of these swinging arms [11,12]: an enhancement of catalytic efficiency, the potential for substrate channelling, and an opportunity to protect an otherwise unstable catalytic intermediate (the ‘hot potato hypothesis’).

Another important feature is the mechanism by which the target side chain in the parent protein is selected for post-translational modification. In each case it has become apparent that there is an enzyme, a ligase or transferase, that catalyses the formation of an amide bond with the amino group of a lysine residue (lipoylation or biotinylation) or phosphodiester bond with the hydroxyl group of a serine residue (pantetheinylation) as a means of attaching the prosthetic group ([13–16] and references therein). However, only recently has it become clear, at least in the case of lipoylation [17] and biotinylation [18], how these ligases recognize their target protein and identify the particular side chain for modification.

We review here our current knowledge of the lipoyl-lysine and biotinyl-lysine swinging arms in multienzyme complexes. Parallels between lipoic acid and biotin have long been drawn and it has now become apparent that there is an underlying structural similarity in the proteins that become lipoylated or biotinylated.

Moreover, the similarity extends to the molecular basis of the post-translational modifications. There are no obvious structural features in common with the enzymes that utilize pantetheinic acid as a swinging arm, but it is likely that similar considerations will apply.

The lipoyl domain and the biotinyl domain
In lipoylated and biotinylated proteins, the prosthetic group is covalently attached to the N\(^\text{\theta}\)-amino group of a lysine residue in an independently folded domain of about 80 amino acid residues. In 2-oxo acid dehydrogenase complexes, the lipoyl domain is found at the N-terminus of the dihydrolipoyl acyltransferase (E2) component, as a single entity or repeated up to three times in tandem array, depending on the origin of the complex [2,3]. Its structure has been determined by NMR spectroscopy [19–23] and takes the form of a flattened \(\beta\)-barrel composed of two four-stranded \(\beta\)-sheets. The N- and C-terminal residues lie close together in space in one sheet and the lipoyl-lysine residue is located in an exposed \(\beta\)-turn in the other sheet (Figure 1). A similar structure has been determined for the H-protein of the glycine cleavage system, except that the 80 amino acid residue lipoyl domain has N- and C-terminal extensions of approximately 20 residues each [24].

The biotinyl domain in biotin-dependent carboxylases shows only vestigial amino acid sequence similarity to the lipoyl domain but, as predicted [25], has an homologous three-dimensional structure (Figure 1). As determined by X-ray crystallography [26] and NMR spectroscopy ([27] E. L. Roberts, N.-C. Shu, M. J. Howard, R. W. Broadhurst, A. Chapman-Smith, J. C. Wallace, T. W. Morris, J. E. Cronan and R. N. Perham, unpublished work), it too is a flattened \(\beta\)-barrel, comprising two four-stranded \(\beta\)-sheets, with the N- and C-terminal residues close together in space in one sheet and the biotinyl-lysine residue located in a highly conserved Met-Lys-Met sequence in an exposed \(\beta\)-turn in the other sheet. In both instances, therefore, the modified lysine residue is well placed to act as a
Importance of the lipooyl and biotinyl domains

On the face of it, there is nothing in the chemical mechanism of the oxidative decarboxylation of 2-oxo acids [28] that would require the lipooyl group to be attached to a protein domain in the E2 component. However, cleavage of the lipooyl domain from the remainder of the E2 chain by limited proteolysis, while leaving the residual multienzyme complex and the E1 (a thiamin diphosphate-dependent decarboxylase) and E3 (dihydrolipooyl dehydrogenase) components intact and fully active, causes the overall catalytic activity of the pyruvate dehydrogenase complex of *Escherichia coli* to fall virtually to undetectable levels [29,30]. Similar results are obtained when the overall complex is reconstituted from the truncated C2 chain and free lipooyl domain [31]. This presumably illustrates the rate enhancement that is achieved by covalent attachment of the lipooyl domain to the complex, with the consequent increase in its local concentration.

More subtly, perhaps, attachment of the lipooyl group to the lipooyl domain is required for the reductive acylation of the dithiolane ring by the E1 component of 2-oxo acid dehydrogenase complexes. Thus free lipoic acid is not a substrate for E1, although it is for E2 and E3 [32]. However, the lipooyl group attached to a lipooyl domain becomes a highly effective substrate, as evidenced by the value of $k_{cat}/K_m$ being raised by a factor of $10^4$ for the lipooyl domain of the *E. coli* pyruvate dehydrogenase complex [33]. Moreover, the lipooyl domain confers on its pendant lipooyl group specificity for reductive acylation only by the E1 component of its parent 2-oxo acid dehydrogenase complex [33]. E1 catalyses the first committed step in the oxidative decarboxylation of 2-oxo acids and, given that the lipooyl group is common to a number of enzymes, the specificity of the protein–protein interaction between E1 and its cognate lipooyl domain thereby provides an elegant molecular basis for substrate channelling in these complexes [2].

For it to serve as a substrate, attachment of the biotin to its domain is likewise essential for all but two biotin–dependent carboxylases: acetyl-CoA carboxylase and $\beta$-methylcrotonyl-CoA carboxylase [34]. Again, it is likely that the biotinyl domain is not merely an anchor for the swinging arm but participates actively in the multienzyme system in which it is found.

Interactions of the lipooyl domain

Detailed studies have been made of the interactions of the lipooyl domain with the E1 component of the pyruvate dehydrogenase complex of *Bacillus stearothermophilus*. In particular, it has been found that replacement of the amino acid residues flanking the lipooyl-lysine residue in the $\beta$-turn (Figure 1) can have a major deleterious effect on the reductive acetylation by E1 in the presence of pyruvate [17]. Likewise, deletion of part of the prominent surface loop between the first and second $\beta$-strands which lies close in space to the lipooyl-lysine $\beta$-turn (Figure 1) virtually abolishes reductive acetylation, and this
cannot be restored by reconstituting the loop with amino acid sequences taken from other lipoyl domains [35].

The idea that the E1 component recognizes its cognate lipoyl domain, at least in part, by interactions with residues in this surface loop and the β-turn is reinforced by NMR data [35]. On the basis of chemical shift changes noted in the heteronuclear multiple quantum coherence (HMQC) spectrum of the apo form of the B. stearothermophilus lipoyl domain in the presence of its E1 component, it appears that significant and specific contact is made between the two proteins over this region, as indicated in Figure 2. The lipoylated form binds too tightly to E1 for comparable experiments to be possible [35] but insufficiently tightly [36] for it to be possible to co-crystallize a complex. The recognition of the lipoyl domain by E1 by transient contact and the presentation of the pendant lipoyl group for reductive acylation is doubtless more complicated than these experiments have thus far been able to reveal; further analysis may have to await the determination of a three-dimensional structure of the E1 component to complement that of the lipoyl domain.

The specificity of post-translational modification

The ligases that catalyse the post-translational modification of the lipoyl and biotinyl domains have much in common. The mechanism of lipoylation is a two-step process: activation of the carboxyl group of lipoic acid by reaction with ATP to form lipoyl-5'S-AMP and elimination of pyrophosphate, followed by transfer of the lipoyl group to the target lysine residue of the lipoyl domain and release of AMP. In E. coli two genes (iplA and iplB) have been postulated to encode independent pathways for the attachment of the lipoyl group to the apo-protein [14]. The iplA gene has been cloned and over-expressed [37]; the product, lipoyl protein ligase (LPL), utilizes ω-lipoic acid as its preferred substrate but can use octanoic acid and α-lipoic acid [38], corresponding to the B form of the enzyme described previously [39]. The R-enantiomer of lipoic acid is required by the E1 component in the pyruvate and 2-oxoglutarate dehydrogenase complexes [40]. In contrast, the product of the iplB gene relies on an endogenous source of lipoic acid supplied by the iplA-dependent biosynthetic route [14] and is likely to be responsible for most of the aberrant octanoylation of lipoyl domains that occurs under conditions of lipoic acid deficiency [41,42]. The bovine lipoyl protein ligase, for which a cDNA has been cloned and expressed in E. coli [43], shows about 35% sequence identity with the E. coli LPL. Curiously, however, it cannot catalyse the initial activation of the lipoic acid to form lipoyl-AMP, for which another enzyme appears to be required [15]. A detailed comparison of the bovine and E. coli enzymes may thus help to delineate the two active sites.

In E. coli biotinyl protein ligase, BPL, is encoded by the birA gene [13] and its X-ray crystal structure has been solved [44]. The reaction mechanism resembles that of LPL: formation of biotinyl-5'-AMP from biotin and ATP, followed by transfer of the biotinyl group to the target lysine residue of the biotinyl domain and release of AMP. Interestingly, BPL also acts as a repressor to regulate the biosynthesis of biotin by the biotin operon, with biotin-5'-AMP as the co-repressor [45].

Perhaps the most notable observation to emerge from recent work on LPL and BPL is the molecular basis of their selection of the correct target residue [17,18]. Despite the structural similarity between the lipoyl and biotinyl domains, the ligases are highly specific for their respective partners. In the lipoyl domain the lysine residue is found in a moderately conserved Asp-Lys-Ala sequence, and in the biotinyl domain the lysine residue resides in a strongly conserved Met-Lys-Met sequence. Site-directed mutagenesis experiments on the biotinyl domain,
in which the Met-Lys-Met motif was systematically varied, have demonstrated that BPL is relatively insensitive to changes in the amino acid sequence flanking the lysine residue but is unable to function if the lysine residue is shifted one residue to the N- or C-terminal side of its normal position in the exposed β-turn in the apo-domain (Figure 3). Importation of the Asp-Lys-Ala motif into the biotinyl domain renders it incapable of biotinylation in vivo but the domain then becomes a poor substrate for lipoylation and aberrant octanoylation [18].

Similar results have been obtained with the post-translational modification of the lipoyl domain. Again, the target lysine residue must be correctly positioned in the β-turn if it is to become lipoylated and the flanking sequence turns out to be relatively unimportant. Likewise, replacing the Asp-Lys-Ala motif with the Met-Lys-Met sequence widely conserved in biotinyl domains is insufficient to designate the domain for biotinylation [17]. Thus, unlike many enzymes responsible for post-translational modification, e.g. phosphokinases, LPL and BPL require a folded domain to act on; and the target lysine residue must be presented in the exposed

β-turn in the particular domain in which it normally resides.

Envoi

The structural analysis of the lipoyl and biotinyl domains has uncovered some further intriguing features that are clearly important to the enzymic function of their parent multi-enzyme complexes. Although attaching the lipoyl group to a lipoyl domain has a profound effect on the value of $k_{cat}/K_m$ for the reductive acylation of the pendant dithiolane ring (see above), the lipoyl-lysine residue is effectively free to rotate on the surface of the protein, as befits a classical swinging arm [2]. In contrast, the biotinyly-lysine residue on the surface of the biotinyl domain of the acetyl-CoA carboxylase of E. coli is clearly localized by interaction with the protein in the structure deduced from X-ray crystallography [26] and solution NMR spectroscopy (E. L. Roberts, N.-C. Shu, M. J. Howard, R. W. Broadhurst, A. Chapman-Smith, J. C. Wallace, T. W. Morris, J. E. Cronan and R. N. Perham, unpublished work), though there is no evidence of similar biotin–protein interactions in the 1.3 S subunit of Propionibacterium shermanii transcarboxylase [46]. It is not immediately obvious what purpose, if any, is served by the prior localization of the biotinyllysine residue in a holo-domain. The lipoyllysine residue in the H-protein of the glycine cleavage system of pea is also localized [24], but switches to a new position when charged with substrate, such that the aminomethylated derivative, which would otherwise be unstable, is sequestered in a surface cavity of the domain unique to the H-protein [47]. In this instance, the swinging arm is fulfilling the expectation of the ‘hot potato hypothesis’ [12].

Swinging arms occupy a special place in the study of multi-step catalysis by multi-enzyme systems. The recent spate of new knowledge about the structure of the proteins that harbour them, and the enzymes that attach them, has put them in a new light while leaving plenty of interesting dark corners still to be explored.

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