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Stabilisation of the Fatty Acid Decarboxylase from *Chlorella variabilis* by Caprylic Acid

Yinqi Wu,[a] Caroline E. Paul,[a] and Frank Hollmann*[a]

The fatty acid photodecarboxylase from *Chlorella variabilis* NC64 A (CvFAP) catalyses the light-dependent decarboxylation of fatty acids. Photoinactivation of CvFAP still represents one of the major limitations of this interesting enzyme en route to practical application. In this study we demonstrate that the photostability of CvFAP can easily be improved by the administration of medium-chain length carboxylic acids such as caprylic acid indicating that the best way of maintaining CvFAP stability is ‘to keep the enzyme busy’. The recently discovered fatty acid photodecarboxylase from *Chlorella variabilis* NC64 A (CvFAP) catalyses the light-driven decarboxylation of fatty acids into their corresponding (C1-shortened) alkanes.[3] CvFAP-catalysed transformations may play a role in the synthesis of fuel alkanes[3] or value-added fine chemicals.[5] Next to DNA photolyase[6] and protochlorophyllide oxidoreductase,[5] the flavin adenine dinucleotide (FAD)-containing CvFAP so far represents the only known example of a photoenzyme.

Upon illumination, the photoexcited FAD cofactor (‘FAD*) inside the enzyme abstracts a single electron from the active site-bound carboxylate substrate resulting in the FAD semiquinone radical (FAD*-) and carboxyl radical. The latter rapidly decarboxylates yielding a short-lived alkyl radical (R*) which abstracts a H-atom from conserved cysteine or asparagine active site residues. The catalytic cycle is closed by an electron transfer from FAD* to the amino acid radical (Scheme 1).[1b,6]

Similar to other flavin-containing enzymes,[7] CvFAP is prone to photochemical inactivation.[8] Scrutton and co-workers therefore suggested keeping CvFAP as much as possible under dark (or red light) conditions to minimise its inactivation.[8] The same authors also reasoned that the photoinactivation may originate from a CvFAP malfunction upon photoactivation in the absence of a carboxylate substrate. In this situation, the photoexcited, high redox-potential flavin is assumed to oxidise nearby active site amino acid residues leading to irreversible inactivation of the enzyme.

We therefore set out to investigate whether the light-dependent inactivation of CvFAP may simply be alleviated by incubating the enzyme with carboxylic acids.

Prior to investigating the effect of irradiation on the stability of CvFAP, we first tested the thermal stability of the enzyme to rule out possible effect of thermal inactivation on the photoinactivation experiments. CvFAP rapidly lost its catalytic activity upon incubation at temperatures higher than 30 °C (t<sub>1/2</sub> (40 °C) ~ 2.7 h; t<sub>1/2</sub> (50 °C) < 1 h; after 24 h incubation at the temperatures, the catalytic activity was completely lost), whereas at 30 °C, more than 90% of its initial activity was retained for at least 22 h (Figure 1 and Figure S2). We therefore continued our investigations under conditions at 30 °C.

Next, we compared the photochemical inactivation of CvFAP as purified enzyme and as crude cell extract preparation (Figure 2). In previous studies we had already observed a significant difference in CvFAP performance in purified form and as crude cell extract.[2] Under blue light illumination, the crude cell extract preparation of CvFAP was significantly more stable exhibiting a half-life time of approx. 19 h. Under the same conditions, the purified enzyme was almost completely
inactivated within 2 h ($t_{1/2}$ ~ 1 h). Under dark conditions, both enzyme preparations exhibited comparable robustness.

It is worth mentioning here that the inactivation of the purified enzyme also depended on the wavelength of the light applied during incubation. Blue light ($\lambda_{max} = 450$ nm) had the most pronounced inactivating effect, followed by green light ($\lambda_{max} = 550$ nm) whereas red light ($\lambda_{max} = 650$ nm) slightly influenced the stability of purified CVFAP (Figure 3). This corresponds well with the UV/Vis spectrum of CVFAP-bound FAD and supports the assumption of photocexcited FAD being the main cause of photoinactivation.

The strikingly higher photostability of CVFAP in crude cell extracts (Figure 2) may be attributed to the presence of E. coli borne carboxylic acids in these preparations. We therefore examined the influence of various carboxylic acids on the photostability of purified CVFAP (Figure 4).

Alkanes (as reaction products) did not exceed a significant stabilising effect on illuminated CVFAP, whereas several fatty acids stabilised the illuminated enzyme. This indicates that the photostability of CVFAP is linked to its decarboxylation reaction. Interestingly, caprylic acid had the most pronounced stabilising effect. This was somewhat unexpected as according to the previously determined substrate scope of CVFAP, caprylic acid should be a much poorer substrate compared to C$_{16}$-C$_{18}$ carboxylic acids. Currently, we are lacking a plausible explanation for why octadecanoic/eicosanoic acid were less efficient than caprylic acid, possibly the concentration of free octadecanoic/eicosanoic acid (more hydrophobic and hence exhibiting a lower critical micelle concentration) was lower than the one of caprylic acid.

Figure 1. Thermal stability of CvFAP under dark conditions. (A) Residual activity of CvFAP after 15 min incubation at the temperature indicated; (B) Time course of CVFAP activity upon incubation at 30 °C (●), 40 °C (▲), 50 °C (○). Incubation conditions: [CVFAP] = 18 μM, buffer: 100 mM Tris-HCl (pH 8.5), protected from light. Activity assay conditions: [palmitic acid] = 13 mM, [DMSO] = 30 vol%, buffer: 100 mM Tris-HCl (pH 8.5), [CVFAP] = 3–6 μM, light intensity of blue light = 14.5 μE L$^{-1}$ s$^{-1}$, T = 37 °C, reaction time = 30 min. Data represent the mean ± SD of two independent experiments.

Figure 2. Comparison of photochemical inactivation of CVFAP as purified enzyme (●) and crude cell extract preparation (○). Incubation conditions: [CVFAP] = 18 μM, buffer: 100 mM Tris-HCl (pH 8.5), light intensity of blue light = 14.5 μE L$^{-1}$ s$^{-1}$, T = 30 °C. Activity assay conditions: [palmitic acid] = 13 mM, [DMSO] = 30 vol%, buffer: 100 mM Tris-HCl (pH 8.5), [CVFAP] = 3–6 μM, light intensity of blue light = 14.5 μE L$^{-1}$ s$^{-1}$, T = 37 °C, reaction time = 30 min. Data represent the mean ± SD of two independent experiments.

Figure 3. Residual activity of purified CvFAP illuminated under different wavelengths of LEDs. Incubation conditions: [CVFAP] = 18 μM, buffer: 100 mM Tris-HCl (pH 8.5), protected from light or illuminated by different colour of light, T = 30 °C, pre-illumination time = 2 h. Activity assay conditions: [palmitic acid] = 13 mM, [DMSO] = 30 vol%, buffer: 100 mM Tris-HCl (pH 8.5), [CVFAP] = 3–6 μM, light intensity of blue light = 14.5 μE L$^{-1}$ s$^{-1}$, T = 37 °C, reaction time = 30 min. Data represent the mean ± SD of two independent experiments.

Figure 4. Influence of additive on the photostability of purified CvFAP. Incubation condition: [CVFAP] = 18 μM, buffer: 100 mM Tris-HCl (pH 8.5), [DMSO] = 5 vol%, light intensity of blue light = 14.5 μE L$^{-1}$ s$^{-1}$, T = 30 °C, pre-illumination time = 2 h, (additive) = 10 mM. Activity assay conditions: [palmitic acid] = 13 mM, [DMSO] = 30 vol%, buffer: 100 mM Tris-HCl (pH 8.5), [CVFAP] = 3–6 μM, light intensity of blue light = 14.5 μE L$^{-1}$ s$^{-1}$, T = 37 °C, reaction time = 30 min. Data represent the mean ± SD of two independent experiments.
We further investigated the concentration-dependency of the stabilising effect of caprylic acid (Figure 5). Increasing the concentration of caprylic acid up to 10 mM steadily increased the stabilising effect, which may well be explained with increasing saturation of the active site of CvFAP. Further increase of the caprylic acid concentration apparently gradually decreased the stabilising effect. Most likely, this, however, is an artefact from the activity assay based on the accumulation of pentadecane. At high caprylic acid concentrations, its decarboxylation competes with the decarboxylation of palmitic acid and thereby reduce the pentadecane formation rate. In fact, caprylic acid could be converted during the pre-illumination process catalysed by CvFAP. After 24 h pre-illumination, little caprylic acid was detected and CvFAP exhibited little residual activity, compared with 46% residual activity of that incubated in dark for 24 h (Figure 2 and Figure S3).

Finally, we confirmed the stabilising effect of caprylic acid on purified, pre-illuminated CvFAP used in semi-preparative transformation of palmitic acid. Figure 6 impressively demonstrates the positive effect of simple caprylic acid in the pre-illumination reaction. While in the absence of caprylic acid, very low concentration of pentadecane was detected, the product accumulation rate of the caprylic acid protected CvFAP was comparable to activities previously observed (without pre-illumination).

Photoinactivation of CvFAP remains a major limitation to preparative application of this promising catalyst. In accordance with the CvFAP inactivation mechanism proposed by Scrutton and co-workers, our results indicate that photoexcitation of CvFAP in the absence of a convertible substrate (carboxylic acid) represents the main cause for CvFAP photoinactivation. In this situation, the photoexcited flavin oxidises active-site borne amino acids, which in light of the recent mechanistic studies by Beisson and co-workers, may be the catalytically active cysteine or arginine (Scheme 2).

In the case of cysteine, it may be argued that the thiyl radical (lacking suitable reaction partners) may react with further amino acids such as tyrosine 466. It is also conceivable that arginine 451 undergoes deguanidination. These inactivation reactions may be expected to be dependent on the protonation stage of the amino acids and that pH-dependent inactivation experiments may shed a light on this. Moreover,

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digestion of the photoinactivated enzyme and mass-spectroscopic analysis of the fragments will shed a light on the inactivation mechanism and possibly serve as guiding principle for future CvFAP engineering to increase its photorobustness. Evidently, the generally observed ‘simple’ photobleaching of the flavin prosthetic group may also considerably contribute to the photoinactivation of CvFAP.

In the present study we have shown that the photostability issue of CvFAP, at least under illumination conditions, can significantly be alleviated by the addition of caprylic acid (and possibly some other carboxylic acids not tested yet). In line with previous suggestions by Scrutton and co-workers, these results point towards ‘keeping CvFAP catalytically busy’ as the most promising strategy to minimise photoinactivation of the enzyme.

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Conflict of Interest

The authors declare no conflict of interest.

Keywords: caprylic acid · fatty acids · fatty acid photodecarboxylase · photochemical inactivation · photostability

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Keep me busy: The photoactivated fatty acid decarboxylase from *Chlorella variabilis* NC64 A (CvFAP) is a promising catalyst to generate next generation biodiesel from (waste) fatty acids. CvFAP, however is also plagued by pronounced photoinactivation, which can be alleviated by keeping it busy, i.e. converting carboxylic acids such as caprylic acid.

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