Stress Induction of Mitochondrial Formate Dehydrogenase in Potato Leaves

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In higher plants formate dehydrogenase (FDH, EC 1.2.1.2.) is a mitochondrial, NAD-dependent enzyme. We previously reported that in potato (Solanum tuberosum L.) FDH expression is high in tubers but low in green leaves. Here we show that in isolated tuber mitochondria FDH is involved in formate-dependent O₂ uptake coupled to ATP synthesis. The effects of various environmental and chemical factors on FDH expression in leaves were tested using the mitochondrial serine hydroxymethyltransferase as a control. The abundance of FDH transcripts is strongly increased under various stresses, whereas serine hydroxymethyltransferase transcripts decline. The application of formate to leaves strongly enhances FDH expression, suggesting that it might be the signal for FDH induction. Our experiments using glycolytic products suggest that glycolysis may play an important role in formate synthesis in leaves in the dark and during hypoxia, and in tubers. Of particular interest is the dramatic accumulation of FDH transcripts after spraying methanol on leaves, as this compound is known to increase the yields of C₃ plants. In addition, although the steady-state levels of FDH transcript increase very quickly in response to stress, protein accumulation is much slower, but can eventually reach the same levels in leaves as in tubers.

FDH catalyzes the oxidation of formate into CO₂ and is a widespread enzyme in bacteria, yeasts, fungi, mammals, and plants. Several types of FDHs have been reported with differing cofactors (NAD or FAD), electron acceptors, substrates, and cellular locations. A wide diversity of FDH types is found in the bacteria, where they are involved in respiration (for review, see Sawers, 1994) and possibly in the maintenance of a reducing environment (Haynes et al., 1995). The NAD-dependent FDHs (EC 1.2.1.2) have been extensively studied in yeast and bacteria over the past decade because the favorable thermodynamics of the reaction, the easy removal of the product (CO₂), and the low cost of the substrate (formate) have made this enzyme a good candidate for industrial NADH regeneration (Allen and Holbrook, 1995). In methanol-utilizing yeasts such as Hansenula polymorpha, Candida boidinii, and Pichia pinus, NAD-dependent FDH plays a crucial role, together with formaldehyde dehydrogenase, in providing NADH to the respiratory chain (Van Dijken et al., 1976). Aside from formate, it is possible that S-formylglutathione might also be a substrate for FDH, as was suggested for yeasts (Van Dijken et al., 1976), pea (Uotila and Koivusalo, 1979), and Pseudomonas sp. 101 (Popov and Lamzin, 1994).

In higher plants only NAD-dependent FDHs have been found, and these are localized in the mitochondrial matrix (Halliwell, 1974; Oliver, 1981; Colas des Francs-Small et al., 1993). Oliver (1981) showed that isolated leaf mitochondria from spinach (and, to a lesser extent, that from beet and tobacco) can oxidize formate with a tight coupling to oxidative phosphorylation. The first plant FDH cDNA to be published encodes a potato (Solanum tuberosum L.) enzyme (Colas des Francs-Small et al., 1993) that is 8 to 10 times more abundant in mitochondria isolated from tubers and dark-grown shoots than in leaf mitochondria (Colas des Francs-Small et al., 1992). Similar variations were found in many other plant species, suggesting drastic changes in formate metabolism between tissues. For example, there is a variable abundance of a polypeptide designated W in pea (Pisum sativum L.) tissues (Humphrey-Smith et al., 1992), which was subsequently identified in our laboratory as FDH.

In this paper we describe the expression of potato mitochondrial FDH in various tissues and under several stresses. The mRNA levels of the mitochondrial SHMT isoform were studied in parallel because this enzyme was previously described to be abundant in pea photosynthetic tissues and scarce in green tissues kept in the dark (Turner et al., 1992, 1993). Like FDH, SHMT is involved in C1 metabolism, which is important for the biosynthesis and degradation of some amino acids (Ser, Gly, and Met), purines, sugars, and organic acids (Cossins, 1980). Our results show that FDH expression is increased by various stresses and that SHMT is either decreased or unchanged. These

Abbreviations: α- and β-ATPase, α- and β-subunits of the mitochondrial ATP synthase, respectively; FDH, formate dehydrogenase; MDH, malate dehydrogenase; SHMT, Ser hydroxymethyltransferase; THF, tetrahydrofolate; TPP, triphenylphosphonium.

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changes in expression were relatively slow and therefore probably result from metabolite changes rather than direct induction or repression by the environmental alterations. As formate would be a good candidate for the final signal in transduction pathways leading to FDH responses, we tested the effects of spraying leaves with metabolites that could be directly or indirectly involved in formate biosynthesis at the level of FDH and SHMT mRNAs.

MATERIALS AND METHODS

Plant Material and Growth Conditions

Leaves, stems, roots, flowers, shoots, and tubers were obtained from potato (Solanum tuberosum L. cv BF15 or Désirée) plants (2n = 4x = 48) grown in 15-cm pots in a greenhouse at 20°C under a 16-h photoperiod. Hypoxia (2–3% O2 [v/v]) was created for 8 to 48 h in an airtight Perspex chamber (made in the lab) flushed with N2 for 2 hours out of 12 h. Cold stress was induced at 4°C for 24 or 48 h. To induce drought stress, plants were dug up carefully and placed for 24 to 48 h on a sheet of Parafilm. These stresses were performed under the same photoperiod as the control plants. Dark treatment was performed at 20°C in a dark chamber for 8 h to 5 d, and dark-to-light shift treatment was performed by transferring plants from the dark chamber (24- or 48-h treatment) back to the greenhouse for an extra 24 h. In the wounding experiment, leaves were lacerated and left for 20 min to 3 d on the plants prior to extraction. Chemical treatments were performed by spraying the plants with various solutions, such as 100 μM ABA, 10 mM formate, acetate, pyruvate, Ser, or sarcosine, 10% ethanol, or 20% methanol. Twenty-four to 48 h after treatment, the leaves were harvested and immediately frozen in liquid N2 for mRNA isolation.

RNA Isolation and Northern Analysis

Total RNAs were isolated from various potato tissues according to either the procedure of Logemann et al. (1987) or a modified method of Sambrook et al. (1989). Plant tissues were ground in liquid N2, and the powder was homogenized in extraction buffer (8 M guanidium hydrochloride, 20 mM Mes, pH 7, 20 mM Na2EDTA, and 50 mM 2-mercaptoethanol) and centrifuged for 20 min at 17,000 rpm (rotor SS-34, model RC5C, Sorvall) at 4°C. RNAs were then purified by an overnight centrifugation through a 4-ml 5.7 M CsCl cushion at 20°C at 32,000 rpm (rotor SW41, model L7-55, Beckman). For each sample, 10 μg of RNA was run on a 3% formaldehyde, 1.5% agarose gel, transferred by blotting onto nylon membranes (Hybond-N+, Amersham), and probed with either an 850-bp potato FDH cDNA (Colas des Francs-Small et al., 1993) or a 1.5-kb pea (Pisum sativum L.) SHMT cDNA (Turner et al., 1992). RNA loading on gels was controlled by EtBr staining under UV light, and transfer efficiency by hybridization with a 2.42-kb wheat mitochondrial 18S rRNA gene fragment (Bonen and Gray, 1980).

Measurement of Formate Oxidation by Isolated Mitochondria

Potato tuber mitochondria were isolated and subsequently purified on Percoll gradients as previously described (Diolez and Moreau, 1985). O2 uptake and membrane potential were measured simultaneously with an O2 electrode (Hansatech, Kings Lynn, UK) and a TPP-sensitive electrode (Diolez and Moreau, 1985). Experiments were carried out at 25°C in a medium containing 1 mL (0.5 mg) of mitochondrial protein in 400 mM mannitol, 10 mM K2PO4 buffer, pH 7.2, 100 mM KCl, 5 mM MgCl2, 1 mg/mL BSA, and 4 μM TPP+ as final concentrations. Respiration was initiated with 15 mM formate, and a limiting amount of ADP (100 μM) was then added to ensure a state 3-to-state 4 transition. Membrane potential was calculated using the equation:

$$\Delta \Psi = 59 \log (v/V) - 59 \log (10^{AE/59})$$

where v is the mitochondrial matrix volume taken as 1 μL/mg protein, V is the volume of the incubation medium, and AE is the electrode potential in mV (Kamo et al., 1979).

Mitochondrial Protein Analysis

Two-dimensional SDS-PAGE of mitochondrial proteins was performed on a Mini Protean II apparatus (Bio-Rad), as described previously (Colas des Francs-Small et al., 1992). The gels were Comassie blue stained and the polypeptides were quantified (Scanalytics MasterScan 1, Computer Signal Processing, Billerica, MA) equipped with two-dimensional gel-analysis software. The polypeptides of interest were identified by immunodetection with various antibodies after transfer onto nitrocellulose sheets (Hybond C, Amersham) using the Mini Trans-Blot cell (Bio-Rad).

RESULTS

FDH and SHMT mRNA Expression in Various Potato Tissues

The levels of FDH and SHMT mRNAs were studied in parallel in various tissues from greenhouse-grown potato plants and are presented in Figure 1. The lowest FDH transcript levels were found in leaves and stamen-free flowers, whereas developing tubers exhibited a very high amount of FDH mRNAs (about 100-fold higher than in leaves). Veins, stems, stolons, stamens, and roots showed intermediate amounts of FDH transcripts: about 16-fold higher in stamens than in leaves (used as the reference for quantitations), 6-fold higher in veins, 8-fold higher in stamens, 6-fold higher in stolons, and 2.5-fold higher in roots. Compared with FDH, SHMT transcript levels were high in...
leaves, one-half as abundant in veins and stolons, and very scarce or even undetectable in other tissues. FDH protein amounts followed mRNA levels in the various tissues tested. The histogram of the quantities of FDH, MDH (matrix protein), and β-ATPase (membrane protein) illustrates the variations in FDH and the relatively constant amounts of MDH and β-ATPase in these tissues (Fig. 2).

Formate Oxidation in Potato Tuber Mitochondria

From simultaneous measurements of O
2 uptake and membrane potential in mitochondria isolated from potato tubers, it could be demonstrated that NAD-linked FDH produced NADH, which was reoxidized by the respiratory chain (Fig. 3). Formate oxidation in state 4 (10.5 ± 0.5 nmol O
2 min
−1 mg
−1 protein) was associated with high values of membrane potential (228 ± 2 mV), whereas the addition of ADP induced a large depolarization (55 ± 5 mV) (Fig. 3). The rates of formate oxidation (32 ± 2 nmol O
2 min
−1 mg
−1 protein) were one-half that of those obtained with malate (64 ± 14 nmol O
2 min
−1 mg
−1 protein). By contrast, the ADP-to-O
2 ratio found with either formate or malate was identical (2.15 ± 0.35), indicating that formate oxidation was coupled to the three proton translocation sites in plant mitochondria (see also Oliver, 1981).

The Expression of FDH in Leaves Is Enhanced under Various Environmental Stresses

Hypoxia

Hypoxia led to a strong FDH mRNA induction. Although no effect was observed after 8 h of hypoxia, the FDH mRNA level increased 8-fold after 16 h (Fig. 4), and remained high after 24 and 48 h of hypoxia (data not shown).

Chilling and Drought

Chilling and drought also considerably increased FDH mRNA (respectively, 10- and 24-fold higher than the control) after 24 h of treatment (Fig. 4). Chilling, however, induced a slower response than drought, as the transcript abundance continued to increase for another 24 h.

Figure 1. Northern analysis of RNA isolated from various potato tissues and probed with either potato FDH cDNA or pea SHMT cDNA. Transcript sizes are, respectively, 1.4 and 1.5 kb. Ten micrograms of RNA was loaded onto each lane, except for developing tubers, the lanes of which were loaded with only 5 μg of RNA. Lf, Leaves; Ve, veins; Ste, stems; Sto, stolons; Rt, roots; sfF, stamen-free flowers; Sta, stamens; and dT, developing tubers. The autoradiographs were scanned using the leaf sample as the reference for both probes, and the figures were corrected according to the RNA loadings.

Figure 2. Relative contents of FDH, MDH, and β-ATPase proteins in potato mitochondria isolated from different tissues, as estimated by quantitation of the corresponding spots on Coomassie blue-stained two-dimensional gels, as described in “Materials and Methods.” The relative abundance of each spot was expressed as a percentage of the total number of protein spots detected on the gels. The figures are an average of five quantitations performed on five gels resulting from five distinct mitochondrial preparations.

Figure 3. Simultaneous recording of respiratory rate and membrane potential in purified mitochondria from potato tuber oxidizing formate. Experimental conditions are described in “Materials and Methods.” Mitochondria (0.5 mg protein/mL) were incubated in the presence of 4 μM TPP− and 1 mM NAD+. Concentrations used were: formate, 15 mM; ADP, 100 μM; and carbonyl cyanide m-chlorophenylhydrazone, 1 μM. PM, Purified mitochondria.

Stress Induction of Mitochondrial Formate Dehydrogenase
Dark and Dark-to-Light Transition Effects

A dramatic increase of FDH mRNA was observed after 16 h of dark treatment (Fig. 4), which continued to increase up to 24 h (17- and 20-fold higher than the control, respectively), and started to decrease after 3 d (down to 10-fold higher than the control). Dark induction of FDH mRNAs was almost completely reversible by placing the plants under a 16-h photoperiod for another 24 h (Fig. 4, lanes D24-L24 and D48-L24).

Wounding

Unlike the previous stress treatments, wounding led to a very rapid FDH mRNA response, as the transcripts were very abundant only 20 min after wounding. They accumulated (13-fold higher than the control) for up to 1 h (Fig. 4), remained high for 24 h, and then decreased between 24 h and 3 d (data not shown).

SHMT and FDH Leaf mRNAs Show Opposite Responses to Various Environmental Stresses

SHMT mRNA levels were unchanged after chilling or wounding (Fig. 4). On the contrary, they were undetectable after 8 h of hypoxia, 24 h of drought, or 24 h of dark treatment. SHMT mRNAs reappeared when plants kept for 24 to 48 h in the dark were placed back under a 16-h photoperiod for another 24 h. In summary, SHMT and FDH mRNAs showed opposite responses (except for chilling and wounding), demonstrating that we are dealing with specific regulations. In regard to dark and hypoxia stresses, however, the kinetics of the increase or decrease were different, i.e. more rapid and drastic for SHMT transcripts.

The Expression of FDH and SHMT mRNAs in Leaves Are Altered Differently by External Addition of Various Metabolites

Formate

Spraying plants with 10 mM formate (the substrate for FDH) increased FDH mRNAs 13-fold after 24 h, whereas the amount of SHMT mRNAs was unchanged (Fig. 5).

ABA

ABA, a compound involved in the response to several stresses such as cold, hypoxia, drought, and wounding, was tested for its ability to regulate FDH or SHMT mRNA levels. Twenty-four hours after spraying 100 mM ABA, the amount of FDH mRNAs was multiplied by five (Fig. 5), and the amount of SHMT mRNAs decreased.

Ser and Sarcosine

Ser and sarcosine, two products derived from Gly and methylene THF by the action of SHMT and sarcosine dehydrogenase, respectively, induced a 2.5- to 3-fold decrease of SHMT mRNA levels in leaves, whereas the amount of FDH mRNA was increased (about 6-fold higher in both cases) (Fig. 5).

Pyruvate, Acetate, and Ethanol

Because glycolysis is known to be induced by dark and hypoxia, some products of glycolysis (pyruvate, acetate, and ethanol) were tested. Northern analyses showed an increase of FDH expression level 24 h after spraying 10 mM acetate (5.5-fold) or 10 mM pyruvate (3-fold), and a greater increase (15-fold) 24 h after 10% ethanol treatment (Fig. 5). On the contrary, SHMT mRNAs decreased after spraying.
acetate or pyruvate (3- and 5-fold, respectively), and disappeared after ethanol treatment (Fig. 5).

Methanol

Methanol led to a dramatic increase (30-fold higher than the control) of FDH mRNAs (Fig. 5) and a 3-fold decrease of SHMT mRNAs (Fig. 5).

Two-Dimensional Analysis of Mitochondrial Polypeptides

Figure 6, A and B, shows the polypeptide patterns obtained with mitochondria isolated from tubers and leaves from young, actively growing plants (summer plants). The relative FDH content was 10-fold higher in tubers than in leaves, whereas MDH and α- and β-ATPase were roughly constant in both tissues (Fig. 2). However, it was evident that in mitochondria isolated from the leaves of old, slowly growing plants (winter plants, mostly grown under artificial light), the relative FDH content observed was similar to that found in tubers (Fig. 6C). The two-dimensional mitochondrial protein patterns of tubers from summer and winter plants were similar (not shown).

DISCUSSION

Differential Expression of FDH in Various Potato Tissues

Our results show that under normal growth conditions, FDH transcript and protein contents are high in nonphotosynthetic tissues (stems, stolons, stamens, tubers, and dark-grown sprouts) and scarce in leaf mitochondria. An exception to this rule is the roots, where FDH content is only slightly higher than in leaves. The fact that formate can be excreted by roots, as reported for oats and maize under poorly oxygenated conditions (Davison, 1951), might explain this discrepancy.

The highest FDH expression levels were found in developing tubers, as opposed to dormant tubers, where no FDH transcripts were detected (Colas des Francs-Small et al., 1993). FDH activity, however, remains high in dormant tubers, sprouting tubers, and sprouts. Why do nonphotosynthetic tissues need more FDH than green tissues? What does this particularly high FDH activity in potato tubers mean? In some plant tissues, such as tubers and sprouts, FDH might be involved in cell respiration when the TCA cycle is not fully operative. Norton (1963) suggested that in fresh potato tuber slices, the bulk of respiration is mediated by systems other than the TCA cycle, because only 10% of the O₂ uptake is inhibited by malonate, a potent inhibitor of succinate dehydrogenase. In addition, it was shown that during the aging of tuber slices, malonate-sensitive respiration increases greatly, whereas malonate-insensitive respiration remains roughly constant.

In isolated mitochondria a formate-dependent O₂ uptake coupled with ATP synthesis was described by Oliver (1981) and found to vary greatly between tissues and plant species (it was very high in spinach leaves). Our results clearly demonstrate that formate is a good respiratory substrate for isolated potato tuber mitochondria and is consistent with the high level of FDH protein found in tuber (Fig. 2) and other nongreen tissues (Colas des Francs-Small et al., 1993). Furthermore, formate is able to strongly inhibit the oxidation of single substrates such as malate or succinate (not shown).

FDH Response to Stress

Some environmental signals to which tubers are naturally submitted (e.g. dark and hypoxia) and a few other stresses were investigated for their ability to enhance FDH expression in potato leaves, where it is low under normal conditions. The mRNA level of the mitochondrial isoform
of SHMT, a key enzyme of C1 metabolism, was studied in parallel as a control. Our results show that FDH mRNA expression is strongly increased by dark, hypoxia, wounding, cold, and drought. We performed western analyses and activity tests for most of these stresses and observed that FDH quantity and activity follow the mRNA increase 1 to 3 d after the stress treatment. In all cases, the final activity was 1.5- to 1.7-fold higher than the control (not shown).

Under all of these stresses, SHMT was regulated in an opposite manner, except during chilling, when the mRNA level remains as high as in control leaves. The disappearance of SHMT mRNAs is faster than the increase of FDH mRNAs. Only wounding led to a very rapid response of FDH mRNAs (20 min). The relatively slow responses to all the other stresses tested here (8 h on average) probably reflect regulation of FDH expression by changes in concentration of certain intracellular metabolites due to the activation of specific metabolic pathways; for example, dark, hypoxia, and chilling all induce glycolysis. The fact that the response of FDH transcripts to ABA is minor compared with their induction by most stresses suggests that ABA-independent transduction pathways are probably involved in the FDH response to stress.

An important observation reported here is that FDH can accumulate to high levels in leaves of continuously stressed plants, e.g. after 2 to 3 months of poor growth conditions in small pots. We hypothesize that in such stressed tissues, formate respiration might complement classic respiration.

FDH Response to Metabolites

To elucidate the transduction pathways involved in the FDH plant response to stress, the effects of spraying leaves with various metabolites were studied. The effects of the metabolites should be considered to be qualitative rather than quantitative because of the uncertainties concerning the amount of metabolite actually entering the plant cells. The observation that formate application greatly enhances FDH expression in potato leaves led us to focus on formate biosynthesis. Under normal growth conditions, formate can arise from various pathways (Fig. 7). The photorespiratory origin of formate in leaves was the first to be described (Tolbert et al., 1949), and in a recent review, Fall and Benson (1996) suggested rapid formate biosynthesis via methanol metabolism (reactions 2, 3, and 4), with the methanol arising from cell wall degradation or synthesis. Finally, glycolytic products may generate glyoxylate from malate (via malate synthase, reaction 15) or from isocitrate (via isocitrate lyase, reaction 16). This glyoxylate will give rise to formate in peroxisomes. In stressed plants formate biosynthesis could result from the enhancement of any of these pathways or from newly induced ones. Three metabolic pathways thought to be involved in response to stress were studied here.

The Ser and Sarcosine Degradation Pathway

Our results show that spraying Ser and sarcosine increases FDH transcription in potato leaves. The catabolism of sarcosine and Ser to formate was first described in yeast and mammals (Barlow and Appling, 1988), but whether plants use sarcosine is not clear. When photorespiration occurs, the mitochondrial SHMT isozyme catalyzes the synthesis of Ser, whereas the thermodynamic equilibrium of this reaction is normally in favor of Ser degradation (Rebeille et al., 1994). In nongreen tissues Ser is thought to be cleaved by other isozymes of SHMT to yield Gly, 5,10-methylen THF, and, subsequently, formate (Schirch,
1984). Furthermore, Ser catabolism was shown to occur in grapevine leaves under water stress (Morchiladze, 1969). We suggest that under drought, Ser might be partly metabolized into formate (reactions 5, 6, 7, 8, 9, and 10) and possibly account for a part of the FDH induction in potato leaves. The advantages of such a degradation pathway under stress would be to produce ATP (reaction 10) and to provide NADH to the respiratory chain (reaction 1). It would be interesting to study this Ser catabolism in more detail in tissues in which FDH expression is high.

Glycolysis

Dark and hypoxia are known to induce glycolysis, which leads to the production of pyruvate, acetate, and, in the case of hypoxia, ethanol. Spraying acetate and pyruvate increased FDH transcript levels, but less than dark or hypoxia. This suggests that these compounds could be involved in the FDH response to dark or hypoxia, but the concentrations used might not be sufficient to reflect these stresses. On the other hand, 10% ethanol (2 mM) led to a higher FDH increase and to total disappearance of SHMT transcripts, close to what was observed under dark and hypoxic stresses. Acetate, pyruvate, and ethanol could act either directly as formate precursors, or indirectly by induction of the enzymes involved in formate biosynthesis. These results suggest that glycolysis may play an important role in the FDH response to dark and hypoxia. The important glycolytic flux that occurs in nonphotosynthetic tissues (germinating tubers, sprouts, germinating pollen, stems, and leaves in the dark) might to a certain extent account for the high FDH levels in such tissues. However, the pathway from glycolytic products to formate remains obscure. We suggest two possibilities: (a) via malate and isocitrate or (b) directly from pyruvate to formate via a pyruvate formate lyase (reaction 17), which has been described in both cytosol and mitochondria; reactions 5 and 13 are both plastidial and mitochondrial; and reaction 17 has only been described in microorganisms and mitochondria from unicellular algae.

The Metabolism of Methanol

Both cell wall growth and degradation lead to methanol production in plants (Fall and Benson, 1996). Methanol
catabolism to formate is particularly interesting, because methanol-treated plants exhibited very high levels of FDH mRNA. The lack of response of SHMT after methanol treatment indicates that FDH induction is specific. Moreover, recent papers demonstrate that the application of methanol to leaves enhances the yields in a wide range of C₃ plants ( unlike C₄ plants) (Nonomura and Benson, 1992; Devlin and al., 1994; Rowe et al., 1994). These authors suggest that an increase in available CO₂ might contribute to this response. Cossins (1964) showed that methanol is rapidly metabolized into CO₂ in leaves, but although formaldehyde dehydrogenase and FDH exist in plants, methanol oxidase has so far only been found in microorganisms. Furthermore, their role in this metabolic pathway is not yet established (Fall and Benson, 1996). Formaldehyde dehydrogenase, in particular, could be partly or entirely overshadowed by other pathways, as formaldehyde can also give rise to formate directly or via methylene THF (Doman and Romanova 1962; Cossins, 1964). The dramatic induction of FDH after methanol treatment strongly favors the hypothesis of its participation in methanol oxidation to CO₂. Mutants deficient in one or more enzymes of this latter pathway would be very helpful for studying the role of these enzymes in C₃ plant growth.

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