## Biochemical and Structural Characterization of the *Arabidopsis* Bifunctional Enzyme Dethiobiotin Synthetase–Diaminopelargonic Acid Aminotransferase: Evidence for Substrate Channeling in Biotin Synthesis

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Diaminopelargonic acid aminotransferase (DAPA-AT) and dethiobiotin synthetase (DTBS) catalyze the antepenultimate and the penultimate steps, respectively, of biotin synthesis. Whereas DAPA-AT and DTBS are encoded by distinct genes in bacteria, in biotin-synthesizing eukaryotes (plants and most fungi), both activities are carried out by a single enzyme encoded by a bifunctional gene originating from the fusion of prokaryotic monofunctional ancestor genes. In few angiosperms, including *Arabidopsis thaliana*, this chimeric gene (named *BIO3-BIO1*) also produces a bicistronic transcript potentially encoding separate monofunctional proteins that can be produced following an alternative splicing mechanism. The functional significance of the occurrence of a bifunctional enzyme in biotin synthesis pathway in eukaryotes and the relative implication of each of the potential enzyme forms (bifunctional versus monofunctional) in the plant biotin pathway are unknown. In this study, we demonstrate that the BIO3-BIO1 fusion protein is the sole protein form produced by the *BIO3-BIO1* locus in *Arabidopsis*. The enzyme catalyzes both DAPA-AT and DTBS reactions in vitro and is targeted to mitochondria in vivo. Our biochemical and kinetic characterizations of the pure recombinant enzyme show that in the course of the reaction, the DAPA intermediate is directly transferred from the DAPA-AT active site to the DTBS active site. Analysis of several structures of the enzyme crystallized in complex with and without its ligands reveals key structural elements involved for acquisition of bifunctionality and brings, together with mutagenesis experiments, additional evidences for substrate channeling.

### INTRODUCTION

Biotin (vitamin B8) is an essential covalently linked cofactor to some carboxylases, decarboxylases, and transcarboxylases that deal with crucial metabolic processes, such as fatty acid and carbohydrate metabolism (Knowles, 1989; Alban et al., 2000). In mammals, biotin is also known to regulate gene expression through different mechanisms, including histone biotinylation (Zempleni, 2005; Beckett, 2007). Despite its essential functions, de novo synthesis of this vitamin is restricted to bacteria, a few fungi, and plants. Thus, inhibition of the biotin synthesis pathway key enzymes is an attractive approach for antibiotic, fungicide, and herbicide development (Alban et al.,

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2000: Marquet, 2010), All animals, including humans, cannot synthesize biotin as part of their normal metabolism and therefore rely on the supply of biotin from exogenous sources. The biotin biosynthetic pathway has been extensively studied in bacteria, especially Escherichia coli (reviewed in Streit and Entcheva, 2003, among others). In all known microbes, biotin is synthesized from pimeloyl-CoA through four enzymatic steps comprising 7-keto-8-aminopelargonic acid (KAPA) synthase, 7,8-diaminopelargonic acid aminotransferase (DAPA-AT), dethiobiotin synthetase (DTBS), and biotin synthase encoded by the bioF, bioA, bioD, and bioB genes, respectively. Enzymes encoded by these genes in E. coli, Bacillus sphaericus, and more recently in Bacillus subtilis or Mycobacterium tuberculosis have been characterized biochemically, and/or structurally, and their reaction mechanisms elucidated (Schneider and Lindqvist, 2001; Berkovitch et al., 2004; Dey et al., 2010). In plants, the biosynthetic pathway is similar to the one described for bacteria (Schneider et al., 1989; Shellhammer and Meinke, 1990; Baldet et al., 1993; Patton et al., 1996, 1998). However, if the biotin biosynthetic genes have been identified and genetically characterized in the model plant Arabidopsis thaliana, to date only

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the first and the last enzymes of the plant pathway, namely, KAPA synthase and biotin synthase, have been characterized at the biochemical level (Picciocchi et al., 2001, 2003; Pinon et al., 2005). The intimate biochemistry of plant DAPA-AT and DTBS reactions remains unknown.

In bacteria, DAPA-AT is a homodimeric pyridoxal 5'-phosphate (PLP)-dependent aminotransferase that catalyzes the conversion of KAPA to DAPA (Figure 1A). In most bacteria, the enzyme uses S-adenosyl-L-Met (AdoMet) as amino group donor, an unusual feature among aminotransferases (Izumi et al., 1975; Breen et al., 2003; Mann and Ploux, 2006). DTBS is a homodimeric enzyme that catalyzes the formation of the ureido ring of dethiobiotin (DTB) from DAPA and CO<sub>2</sub> in the presence of ATP and divalent metal ions (Figure 1A) (Alexeev et al., 1995; Huang et al., 1995). In Arabidopsis, genetic studies revealed that DAPA-AT (BIO1; BioA ortholog) and DTBS (BIO3; BioD ortholog) are encoded by adjacent genes, defining a single genetic locus, and are expressed in both single and chimeric BIO3-BIO1 transcripts through alternative splicing events (Muralla et al., 2008). One of the fused transcripts is monocistronic and encodes a bifunctional fusion protein capable of complementing the orthologous auxotrophs of E. coli (bioD and bioA). The second transcript includes 10 additional nucleotides that introduce a premature stop codon. As a conseguence, this splice variant is bicistronic, with distinct but overlapping reading frames. This bicistronic transcript is potentially capable of producing separate BIO3 and BIO1 proteins (Figure 1B). Other plants and most fungi also present a chimeric BIO3-BIO1 homolog gene (Hall and Dietrich, 2007; Muralla et al., 2008; Magliano et al., 2011).

In this study, we present evidence that the BIO3-BIO1 fusion protein is the major, if not the exclusive, protein form produced by the BIO3-BIO1 locus in Arabidopsis. The enzyme catalyzes both DAPA-AT (BIO1) and DTBS (BIO3) reactions and is targeted to mitochondria. Biochemical and kinetic properties of the pure recombinant mature enzyme were investigated. Our data are consistent with the hypothesis that DAPA, the product of DAPA-AT domain, is not released in the bulk solvent in the course of the reaction but is directly transferred to the DTBS domain for its subsequent conversion to DTB. Finally, the three-dimensional structure of the enzyme was solved to 2.5-Å resolution, in its holoform (i.e., in the presence of bound-PLP) as well as in complex with various substrates and/or products of the enzyme-catalyzed reactions. The structural analyses show key elements involved in acquisition of bifunctionality. These structures and accompanying kinetic data combined with mutagenesis experiments form the basis for mechanistic considerations, in particular with respect to DAPA intermediate transit between both catalytic sites of the bifunctional enzyme.

## RESULTS

## Expression of the *BIO3-BIO1* Locus in *Arabidopsis* at the Protein Level and Subcellular Distribution

DAPA-AT and DTBS activities in plants are catalyzed by a bifunctional fusion protein, including both reaction domains, BIO1 and BIO3. In addition, the *Arabidopsis BIO3-BIO1* gene fusion also produces a bicistronic mRNA transcript with the potential to encode separate proteins (Muralla et al., 2008): a long BIO3 form exhibiting a C-terminal extension and an N-terminal truncated BIO1 form lacking conserved amino acid residues essential for activity in bacteria (Figure 1B). To clarify the situation in the plant cell at the protein level, we performed an immunological characterization of BIO3-BIO1 gene products in Arabidopsis cell extracts using polyclonal antibodies raised against the fulllength recombinant BIO3-BIO1 protein overproduced in E. coli (see Supplemental Figure 1 online). Immunoblot analysis of protein samples from whole plants (aboveground organs from 35-d-old plants) or cultured cells identified a single polypeptide of  $\sim$ 90 kD (i.e., the expected size for the BIO3-BIO1 fusion protein) (Figure 2A). No band of lower molecular mass was specifically detected using BIO3-BIO1 antibodies (the faint signal bands of low molecular mass are nonspecific background from the preimmune).

This contrasts with the situation at the mRNA level, where the bicistronic splice variant is somewhat more abundant than the monocistronic version in most parts of the plant (Muralla et al., 2008) (see Supplemental Figure 2 online). To investigate the distribution of the BIO3-BIO1 protein in plant cells, intact chloroplasts and mitochondria from Arabidopsis leaves were purified on Percoll density gradients, thus providing organelles devoid of contamination from the other compartments. Also, a cytosolic-enriched fraction was prepared. Soluble proteins from purified chloroplasts (stroma), mitochondria (matrix), and the cytosolic-enriched fraction were then analyzed by immunoblots. Purified antibody detected the BIO3-BIO1 protein exclusively in the mitochondrial matrix fraction (Figure 2B). As a complementary approach, subcellular localization of BIO3-BIO1 was also investigated by fusing its full-length sequence upstream to the green fluorescent protein (GFP) marker. As shown in Figure 2C, the transient expression of BIO3-BIO1-GFP in Arabidopsis protoplasts resulted in a punctuate pattern of green fluorescence, similar to the one observed with GFP fused to the transit peptide of an authentic mitochondrial protein. Collectively, these results strongly suggest that in Arabidopsis, the BIO3-BIO1 locus produces a unique BIO3-BIO1 fusion protein form that takes place in mitochondria.

## Production of the Recombinant Mature BIO3-BIO1 Protein (mBIO3-BIO1)

To further characterize the plant DAPA-AT/DTBS reactions, BIO3-BIO1 was overproduced in *E. coli* as a truncated protein, hereafter designated as mBIO3-BIO1 (for mature BIO3-BIO1) with an N-terminal 22-residue deletion. This sequence was predicted to correspond to the mitochondrial targeting peptide by all the major prediction programs available on the Web and gathered on the *Arabidopsis* subcellular database (Heazlewood et al., 2007). Since the cloning procedure added an N-terminal hexa-His tag, the recombinant mature protein was purified to near homogeneity by standard nickel-affinity chromatography (Figure 3A). The apparent molecular mass of the native protein, as estimated by gel filtration chromatography on a Superdex S200 column, was found to be  $\sim$ 200 kD, consistent with a homodimeric structure of the recombinant enzyme in solution

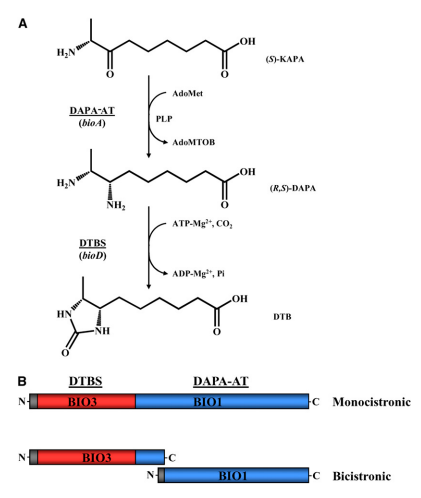


Figure 1. The Two-Step Synthesis of DTB from KAPA.

(A) In bacteria, DTB is synthesized from KAPA by two distinct enzymes encoded by *bioA* and *bioD* genes, respectively, catalyzing successively DAPA-AT and DTBS reactions. AdoMTOB, S-adenosyl-2-oxo-4-methylthiobutyric acid.

(B) Representation of BIO3 (BioD ortholog) and BIO1 (BioA ortholog) domains of BIO3-BIO1 fusion protein and of separate BIO3 and BIO1 proteins, putatively encoded by *Arabidopsis* monocistronic and bicistronic BIO3-BIO1 transcripts, respectively.

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(Figure 3B). Typically, 10 to 15 mg of pure enzyme was obtained per liter of cell culture using this two-step purification procedure. UV-visible spectroscopy analysis of the recombinant enzyme revealed an absorption spectrum characteristic of the internal aldimine of PLP-dependent enzymes with an absorbance maximum (band) at 415 nm and a shoulder at 335 nm (Figure 3C).

## Biochemical Characterization of DAPA-AT and DTBS Reactions

To investigate the overall DAPA-AT + DTBS or DTBS alone activities catalyzed by mBIO3-BIO1 in vitro, we developed a radiochemical assay that monitors the formation of acid stable [<sup>14</sup>C]-DTB from acid-labile <sup>14</sup>CO<sub>2</sub> in the presence of appropriate substrates and cofactors. This assay, adapted from a previously published procedure designed for measuring *E. coli* DTBS activity (Gibson et al., 1995), was sensitive and more reliable in our hands than published bioassays (paper disc plate method) and spectrophotometric assays for bacterial DTBS (Krell and Eisenberg, 1970; Gibson et al., 1995; Yang et al., 1997). Identification of [<sup>14</sup>C]-DTB as a reaction product of the in vitro assays was confirmed by high-performance thin layer chromatography (HP TLC) analysis on a silica gel plate, by comparison of its retention time value with that of authentic [<sup>3</sup>H]-DTB sample (Figure 3D). Plant DAPA-AT activity was measured using a discontinuous assay in which DAPA synthesized by mBIO3-BIO1 was subsequently titrated using the *E. coli* monofunctional DTBS (see Methods).

When DAPA-AT + DTBS activity was assayed under optimized conditions, pure recombinant *Arabidopsis* enzyme exhibited a specific activity of 25 to 35 nmol  $H^{14}CO_3^{-}$  incorporated  $h^{-1}$  mg<sup>-1</sup> protein, confirming that mBIO3-BIO1 is actually a bifunctional enzyme catalyzing both DAPA-AT and DTBS reactions (Figure 4A). Recorded reaction rates were low but

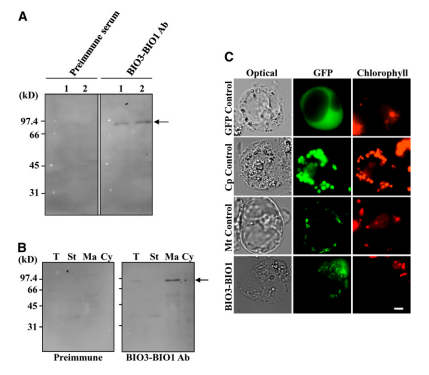


Figure 2. Analysis of the Expression of the Arabidopsis BIO3-BIO1 Locus at the Protein Level and Subcellular Distribution.

(A) Immunological detection of *BIO3-BIO1* gene products in *Arabidopsis*. Soluble proteins (40 μg) from aboveground organs from *Arabidopsis* plants (lane 1) and *Arabidopsis* cultured cells (lane 2) were separated by SDS-PAGE and analyzed by immunoblotting using affinity-purified antibodies raised against recombinant BIO3-BIO1 (BIO3-BIO1 Ab) or preimmune serum (for negative control).

(B) Immunolocalization of BIO3-BIO1 in *Arabidopsis*. Soluble proteins (50 µg) from total plant extracts (T), chloroplast stroma (St), mitochondrial matrix (Ma), and cytosolic enriched fraction (Cy) were separated by SDS-PAGE and analyzed by immunoblotting as in (A). Arrows in (A) and (B) point to the BIO3-BIO1 polypeptide band.

(C) Expression of BIO3-BIO1 fused to GFP in *Arabidopsis* protoplasts. Constructs encoding GFP alone (GFP Control), GFP fused to the C terminus of the small subunit of ribulose-1,5-bis-phosphate carboxylase/oxygenase transit peptide (a chloroplastic marker) from *Arabidopsis* (Cp Control), GFP fused to the C terminus of the dihydropterin pyrophosphokinase-dihydropteroate synthase transit peptide (a mitochondrial marker) from pea (*Pisum sativum*; Mt Control), and GFP fused to the C terminus of full-length BIO3-BIO1 (BIO3-BIO1) were introduced into *Arabidopsis* protoplasts. Images are optical photomicrographs (Optical), GFP fluorescence (GFP; green pseudocolor), and chlorophyll fluorescence (Chlorophyll; red pseudocolor). Bar = 10 µm.

consistent with estimated in vivo rates of biotin production and requirement in living organisms (Alexeev et al., 1998; Alban et al., 2000; Farrar et al., 2010; Lin et al., 2010). The reaction was strictly dependent upon the presence of (S)-KAPA, AdoMet, ATP-Mg, and NaHCO<sub>2</sub> in the reaction medium. In most bacteria, DAPA-AT uses AdoMet as an amino group donor. B. subtilis uses L-Lys, another unusual amino donor for the reaction (Van Arsdell et al., 2005). Besides AdoMet, however, no other substrate was found to act as an amino donor for Arabidopsis mBIO3-BIO1, among a variety of potential substrates, including L-Asp, L-Glu, L-Met, L-Lys, S-adenosylhomocysteine, and 5'-deoxy-5'-methylthioadenosine. Interestingly, the overall DAPA-AT + DTBS activity catalyzed by mBIO3-BIO1 was linear for at least 60 min under optimized conditions, and no lag in DTB production was observed (Figure 4A). This suggested a direct transfer of the DAPA intermediate between the DAPA-AT and the DTBS domains (Figure 4A).

When *Arabidopsis* mBIO3-BIO1 was assayed for DTBS reaction, a specific activity of 700 to 1200 nmol [ $^{14}C$ ]-DTB formed h<sup>-1</sup> mg<sup>-1</sup> protein was measured, a value roughly 25 to 30 times higher than the one recorded for the overall DAPA-AT + DTBS reaction (Figure 4B). This suggested that the overall reaction is rate limited by the DAPA-AT activity. Interestingly, it also demonstrated that in vitro the enzyme can use external DAPA for the synthesis of DTB (i.e., that the BIO3 active site is accessible to the outside).

When DAPA-AT was assayed alone, the initial velocity of the reaction was roughly similar to that of the overall reaction, but surprisingly, DAPA formation rapidly reached a plateau (Figure 4A). At best, ~0.7 to 0.8 nmol DAPA could be synthesized per nmol enzyme monomer in the reaction mixture. A similar kinetic behavior was observed when ATP-Mg, one of the DTBS substrates, was added to the standard DAPA-AT assay. Noteworthy, in the presence of NaHCO<sub>3</sub> in lieu of ATP-Mg in the reaction medium, DAPA-AT activity was found to be linear with time, with a resulting specific activity close to that for the overall DAPA-AT + DTBS reaction (Figure 4B). Thus, in the absence of NaHCO<sub>3</sub>, one of the substrates of the DTBS domain, the DAPA-AT reaction was (sub)stoichiometric, suggesting that the DAPA-AT domain is blocked in a noncatalytically competent form. Interestingly,

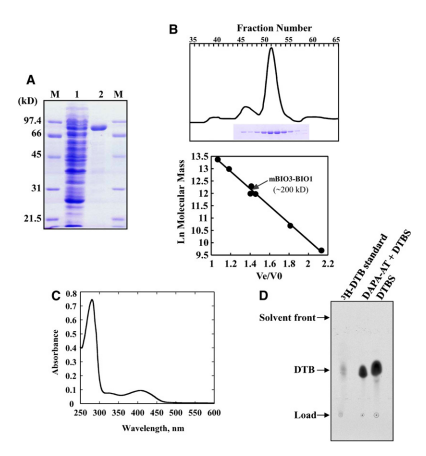


Figure 3. Physicochemical and Biochemical Properties of Recombinant Arabidopsis mBIO3-BIO1.

(A) Documentation of mBI03-BI01 purification on nickel-nitrilotriacetic acid-agarose resin. Polypeptides were separated by SDS-PAGE and stained with Coomassie blue. Lane 1, soluble proteins (25 µg) from *E. coli* Rosetta cells producing mBI03-BI01; lane 2, proteins eluted from the column (10 µg); lanes M, molecular mass markers.

(B) Purification and molecular mass estimation of native mBIO3-BIO1 by gel filtration. Purified protein was resolved by chromatography onto a Superdex 200 HiLoad column ( $2.6 \times 60$  cm; GE Healthcare). Eluted fractions (3 mL) were analyzed by SDS-PAGE (top panel). Standards proteins for column calibration (bottom panel) were as follows: thyroglobulin (669 kD), ferritin (443 kD), catalase (232 kD),  $\gamma$ -globulin (158 kD), aldolase (158 kD), ovalbumin (43 kD), and myoglobulin (17 kD). Ve, elution volume; V0, void volume.

(C) Spectroscopy analysis of purified recombinant mBIO3-BIO1. Absorption spectrum was recorded at 30°C in 100 mM HEPES-KOH, pH 7.5, in the presence of 20 µM pure enzyme.

(D) HP TLC detection of DTB produced by recombinant mBIO3-BIO1. The overall (DAPA-AT + DTBS) reaction and DTBS reaction were performed by measuring the formation of acid-stable [ $^{14}$ C]-DTB from acid-labile H $^{14}$ CO<sub>3</sub> as described in Methods. Assays contained 0.2 nmol (DAPA-AT + DTBS) reaction) or 0.1 nmol (DTBS reaction) enzyme, and reaction mixtures were incubated for 120 or 20 min, respectively. [ $^{14}$ C]-DTB produced was separated by TLC and detected by phosphor imaging analysis (Picciocchi et al., 2001). Migration was compared with that of authentic [ $^{3}$ H]-DTB standard (Rf = 0.53).

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preincubation of the enzyme in the presence of exogenous DAPA did not prevent de novo DAPA production nor did it preclude appearance of the plateau, showing that the DAPA-AT active site was not accessible to external DAPA and that production of the stable enzyme-DAPA complex was dependent on catalysis (Figure 4C). Also, addition of ATP-Mg and NaHCO<sub>3</sub> 15 min after the start of the DAPA-AT reaction (i.e., when most of the enzyme is bound up in a noncatalytic enzyme-product complex) resulted in a rapid synthesis of DTB from already produced DAPA, followed by a slower but linear production of DTB, presumably from newly synthesized DAPA (Figure 4C). Collectively, our results suggest that DAPA produced by the

DAPA-AT domain is somehow trapped in the active site. Upon addition of the DTBS substrates, DAPA would be rapidly transferred to the DTBS active site for DTB synthesis, without previous release in solution.

To learn more about *Arabidopsis* DAPA-AT and DTBS reactions, we determined the kinetic parameters for both the overall (DAPA-AT + DTBS) and DTBS activities (Table 1). The most striking feature of these analyses is that the enzyme exhibited a kinetic cooperativity with respect to all tested substrates and for both reactions (Hill coefficient values calculated from Hill plots varied from 1.3 to 2.4 according to the substrate considered). This kinetic behavior has not been described for

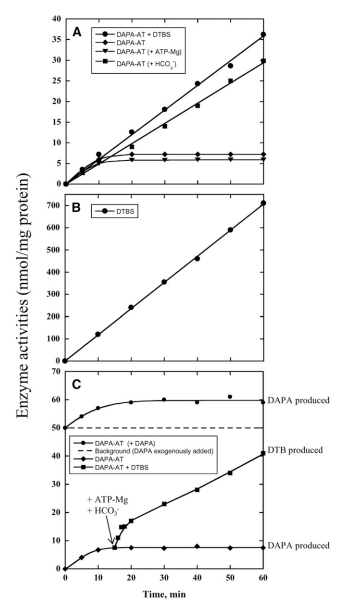


Figure 4. Kinetic Study of *Arabidopsis* mBIO3-BIO1 Catalyzed Reactions.

Activities were assayed under optimized assay conditions as described in Methods. Data are from a representative experiment repeated at least three times.

(A) Time-course formation of DAPA (DAPA-AT reaction; diamonds, triangles, and squares) or DTB (DAPA-AT + DTBS reaction; circles). DAPA-AT was assayed in a reaction medium containing  $20 \ \mu$ M (S)-KAPA, 0.2 mM PLP, 3 mM AdoMet, and 0.4  $\mu$ M enzyme, in the absence (standard assay medium; diamonds) or in the presence of 0.3 mM ATP + 5 mM MgCl<sub>2</sub> (triangles) or 10 mM NaHCO<sub>3</sub> (squares). DAPA-AT + DTBS was assayed in a reaction medium containing 20  $\mu$ M (S)-KAPA, 0.2 mM PLP, 3 mM AdoMet, 0.3 mM ATP, 5 mM MgCl<sub>2</sub>, 10 mM NaHCO<sub>3</sub>, and 0.4  $\mu$ M enzyme. (B) Time-course formation of DTB (DTBS reaction). DTBS was assayed in a reaction medium containing 200  $\mu$ M DAPA, 0.3 mM ATP, 5 mM MgCl<sub>2</sub>, 10 mM NaHCO<sub>3</sub>, and 1  $\mu$ M enzyme.

(C) Effect of exogenous DAPA on DAPA-AT reaction. DAPA-AT was assayed in a reaction medium containing 20  $\mu$ M (S)-KAPA, 0.2 mM PLP,

bacterial monofunctional enzyme counterparts studied so far. It is also interesting to note the ~30 times lower apparent  $k_{0.5}$ value for ATP measured when the overall DAPA-AT + DTBS reaction was assayed compared with that determined for the DTBS reaction alone, without affecting the catalytic efficiency (unaltered  $k_{cat}/k_{0.5}$  value, since  $k_{cat}$  for DTBS reaction is 25-fold higher than for the overall reaction) (Table 1). This probably reflected conformational changes of the ATP binding site upon binding of DAPA-AT substrates or products and suggested interdependence of both DAPA-AT and DTBS domains.

Nevertheless, kinetic parameters ( $k_{0.5}$  and  $k_{cat}$  values for the plant bifunctional enzyme) compared well with those ( $K_m$  and k<sub>cat</sub> values) of bacterial monofunctional enzymes (Krell and Eisenberg, 1970; Gibson et al., 1995; Yang et al., 1997; Eliot et al., 2002; Mann and Ploux, 2006; Dey et al., 2010) (Table 1). A noticeable exception concerned the k<sub>cat</sub> value for plant overall reaction (which reflects DAPA-AT activity) that was found to be one order of magnitude lower than that for bacterial DAPA-AT (Eliot et al., 2002; Mann and Ploux, 2006). Many bacterial DAPA-AT enzymes contain a conserved Tyr residue at the N terminus (Tyr-17 in E. coli) that plays an essential role in binding of Ado-Met or KAPA substrates and for catalytic efficiency (Sandmark et al., 2004; Dey et al., 2010). A Phe residue at this position (Phe-326 in Arabidopsis) is present in all available plant of fungal sequences (see Supplemental Figure 3 online). This difference could possibly account for the low DAPA-AT activity measured with the Arabidopsis enzyme compared with the bacterial enzymes. To test this hypothesis, we constructed a F326Y BIO3-BIO1 mutant. However, in Arabidopsis, Phe-to-Tyr substitution did not impact importantly the enzyme kinetic parameters. In particular,  $k_{cat}$  values were poorly affected (Table 1).

Taken together, our biochemical and kinetic analyses are consistent with a mechanism enabling the enzyme to directly transfer the DAPA intermediate from the DAPA-AT domain active site to the DTBS domain active site without diffusion to the aqueous environment. To address this question and to gain insight into the mechanism of the overall DAPA-AT + DTBS reaction, we undertook a structural characterization of the enzyme holoform (with bound PLP) and the complexes with various substrates and/or products of the enzyme-catalyzed reactions.

## Overall Unliganded Structures of mBIO3-BIO1 from Arabidopsis

The native and the selenomethionine ([SeMet]) enzyme have been crystallized without ligand in two different space groups. In both cases, the structures show a dimer made of two monomers related by a noncrystallographic twofold axis. In the space group C2, one dimer of [SeMet]-mBIO3-BIO1 was present in the asymmetric unit. The C $\alpha$  atoms of the monomers from the [SeMet]-mBIO3-BIO1 can be superimposed with a 0.23 Å root mean square deviation (rmsd) value. In the native form crystallized

3 mM AdoMet, and 0.4  $\mu$ M enzyme, in the absence (standard assay medium; diamonds) or in the presence of 2  $\mu$ M DAPA (circles). After 15 min incubation of the enzyme in the standard DAPA-AT reaction medium, 0.3 mM ATP, 5 mM MgCl<sub>2</sub> and 10 mM NaHCO<sub>3</sub> were added and the time-course formation of DTB was followed (squares).

 Table 1. Kinetic Parameters for the Arabidopsis Wild Type (mBIO3-BIO1) and F326Y Mutant (mBIO3-BIO1 F326Y) DTBS-DAPA-AT: Comparison with

 Those from Bacterial Monofunctional DAPA-AT (bioA) and DTBS (bioD)

Enzyme	Reaction	К <sub>0.5</sub> AdoMet (n <sub>H</sub> ) (mM)	К <sub>о.5</sub> КАРА (л <sub>Н</sub> ) (µМ)	К <sub>0.5</sub> АТР (л <sub>Н</sub> ) (µМ)	К <sub>о.5</sub> DAPA (л <sub>Н</sub> ) (µМ)	k <sub>cat</sub> (min <sup>-1</sup> )	k <sub>cat</sub> /K <sub>0.5</sub> (min <sup>-1</sup> Μ <sup>-1</sup> )
mBIO3-BIO1ª	DAPA-AT + DTBS DTBS	0.63 ± 0.06 (1.34)	4.78 ± 0.6 (1.29)	. ,	18.7 ± 1.5 (1.75)	$\begin{array}{c} 0.072  \pm  0.009 \\ 1.85  \pm  0.05 \end{array}$	$1.28 imes10^5$ (ATP) $0.98 imes10^5$ (ATP)
mBIO3-BIO1 F326Y <sup>a</sup>	DAPA-AT + DTBS	$1.65 \pm 0.19$ (1.23)	4.6 ± 0.6 (1.36)	$0.80 \pm 0.21$ (2.1)		$0.062 \pm 0.005$	$0.77 imes10^5$ (ATP)
	DTBS			29.5 ± 2.7 (1.35)	28.5 ± 3.8 (1.74)	2.16 ± 0.11	$0.73 imes10^5$ (ATP) Organism
bioA <sup>b</sup>	DAPA-AT	0.15 <sup>c</sup>	<2°			0.78	E. coli
bioA <sup>d</sup>	DAPA-AT	0.78 <sup>c</sup>	3.8 <sup>c</sup>			1	M. tuberculosis
bioD <sup>e</sup>	DTBS			5–19°	1.3–3°	2–2.9	E. coli
bioD <sup>f</sup>	DTBS			29 <sup>c</sup>	2 <sup>c</sup>	-	M. tuberculosis

The results are means  $\pm$  sp from three separate determinations, and parameters were estimated by direct fitting of experimental data to the Hill equation by nonlinear regression analysis. Hill coefficient values ( $n_{\mu}$ ) are given in parentheses. Empty cell, not applicable; –, not available.

<sup>a</sup>This work.

<sup>b</sup>Eliot et al. (2002).

<sup>c</sup>K<sub>m</sub> values.

<sup>d</sup>Mann and Ploux (2006).

<sup>e</sup>Krell and Eisenberg (1970), Alexeev et al. (1995), and Gibson et al. (1995).

<sup>†</sup>Dey et al. (2010).

in the space group P1, two dimers were present in the asymmetric unit and the monomers in each dimer well superimposed (0.21 and 0.19 Å rmsd values for the monomer A/B and C/D, respectively). The two dimers of native mBIO3-BIO1 superimposed with a 1.0 Å rmsd value. [SeMet]-mBIO3-BIO1 can be superimposed with 0.48 and 1.04 Å rmsd values with the dimer A/B and C/D of native mBIO3-BIO1, respectively. These results show that the monomer folds in the dimers are roughly identical.

Each monomer can be divided into two domains: the DTBS (BIO3) domain from Ser-6 to Thr-299 (mature enzyme form numbering) and the DAPA-AT (BIO1) domain from Met-300 to Thr-811. A long α-helix Ile283-Val321 (53.3 Å in length) linked the two domains (Figure 5). The catalytic sites of BIO1 and BIO3 are ~48 Å apart. The monomers interact by involving several zones of both domains. In the space group P1, the superimposition of the dimers (A/B onto C/D) showed large differences in the position of the DAPA-AT domain. These differences were also observed when dimer C/D was superimposed onto [Se-Met]-mBIO3-BIO1, mBIO3-BIO1/KAPA, or mBIO3-BIO1/DTB. The average interface area of mBIO3-BIO1 calculated with Protein Interfaces Surfaces Assemblies is 7300 Å<sup>2</sup> (22.4% of the accessible surface) for the most complete dimer (A/B) and 5420 Å<sup>2</sup> (18.4%) for the second dimer (C/D) (Krissinel and Henrick, 2007). The interface area is 7040 Å<sup>2</sup> (22.6%) in [SeMet]-BIO3-BIO1. Because the number of residues observed in the electron density map was different between the dimers in the space group P1 (1525 amino acids in A/B versus 1390 in C/D), the dimer C/D was rebuilt by superimposition with the monomers A and B. In this putative dimer, the interface area observed was 6680 Å<sup>2</sup> (20.6%). Differences in buried surfaces result from flexibility of monomers as also revealed by the differences of rebuilt residues in the structures (see Methods). Observation of the molecular surface at the interface revealed the presence of a small discontinuous groove going from the catalytic site of DAPA-AT to the catalytic site of DTBS. This groove is larger in the dimer C/D.

## Structures of the DTBS and DAPA-AT Domains of the Holoform

The DTBS domain is composed of a nine-stranded  $\beta$ -sheet, with  $\beta$ 5 being antiparallel to the others  $\beta$ -strands, surrounded by three  $\alpha$ -helices on both sides of the  $\beta$ -sheet. Residues His-9 to Pro-11 form the  $\beta$ -strand  $\beta$ 1 involved in the interaction with the DAPA-AT domain through an antiparallel four-stranded β-sheet (see Supplemental Figure 4 online). The two catalytic sites of DTBS are fully accessible to the solvent and are composed of residues from both monomers. The Walker-A motif (Walker et al., 1982) that interacts with phosphates of nucleotide is in the loop between B2 and a1 (Ala22-Thr29). The Walker-B motif (Walker et al., 1982) (Lys177-Glu188) is in the loop connecting  $\alpha 5$  to  $\beta 7$  and in  $\beta 7$ . Lys-177, Ser-178, Glu-179, Lys-180, and Ser-181 were not observed in the electron density whatever the crystal form was. The Walker-A motif stabilized two sulfate ions in one monomer and only one bound to a magnesium ion in the second monomer in mBIO3-BIO1. The magnesium ion was also stabilized by the Thr-29 hydroxyl group from the Walker-A motif, the Asp-66 and Glu-188 carboxylate groups, and one water molecule. Two other sulfate ions were observed in the dimer. They were stabilized by the Ser-195 hydroxyl group of one monomer and the dipole of the  $\alpha$ -helix  $\alpha$ 7 of the second monomer as well as hydrogen bonds with the Gly-223, Gly-224, Ile-225, and Ser-226 NH groups belonging to the  $\alpha$ -helix  $\alpha$ 7.

The DAPA-AT domain can be divided into three subdomains. The first subdomain (Val339-Asp364) is composed of a three-stranded antiparallel  $\beta$ -sheet, which forms a four-stranded  $\beta$ -sheet with  $\beta$ 1 from the DTBS domain. This subdomain is connected to the core of the DAPA-AT domain by two  $\alpha$ -helices. The core (Arg424-Ser665) is formed by a seven-stranded  $\beta$ -sheet ( $\beta$ 14- $\beta$ 23- $\beta$ 22- $\beta$ 21- $\beta$ 20- $\beta$ 15- $\beta$ 16) with the  $\beta$ -strand  $\beta$ 23 being antiparallel to the six others and by an antiparallel three-stranded  $\beta$ -sheet that connects  $\beta$ 16 to  $\beta$ 20 involving the  $\alpha$ -helix  $\alpha$ 14. The C-terminal part of the DAPA-AT

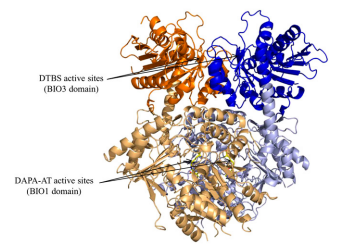


Figure 5. View of the Overall Fold of the Dimer of mBIO3-BIO1 from *Arabidopsis*.

The protein is represented as a ribbon, and the DAPA-AT (BIO1) and DTBS (BIO3) domains from each monomer are colored in light blue/light orange or dark blue/orange. PLP is displayed as a yellow stick at the active sites.

domain (Val732-Thr811) is connected to the core by two  $\alpha$ -helices and is formed by an antiparallel three-stranded  $\beta$ -sheet and two  $\alpha$ -helices. In the DAPA-AT domain, Lys-644 linked to PLP and Val-583 are in disallowed part of the Ramachandran plot but are well defined in the electron density. A large pocket corresponding to the AdoMet and KAPA binding site is observed in the DAPA-AT domain, going from the solvent to the PLP. PLP from one monomer is stabilized by interacting with Asn-430, Gly-431, Ser-432, Tyr-473, His-474, Gly-475, Glu-581, Asp-615, Val-617, Phe-618, Ala-643, Leu-646, and Thr-647 and with His-679, Ser-680, and Tyr-681 of the second monomer.

A large polar crevice of  $\sim$ 12 to 14 Å width at the surface of the enzyme is observed going from the DAPA-AT catalytic binding site to the bottom of the DTBS catalytic site (Figure 6; see Supplemental Figure 5 online).

## Structure of mBIO3-BIO1 Bound to KAPA

Four KAPA molecules were observed in the electron density map of mBIO3-BIO1/KAPA: Two are bound into the DAPA-AT catalytic site and two were found within the catalytic site of DTBS (Figure 7; see Supplemental Figure 6 online). The presence of KAPA into the DTB binding site likely results from its structural similarity with DAPA and the high KAPA concentration (5 mM) used for cocrystallization experiments. Two tartrate ions from the crystallization solution are also observed in the DTBS domain active site. In the DTBS active site, KAPA interacts with Thr-24, Thr-59, Ala-141, Ile-142, Ser-143, Ser-195, and a tartrate ion of one monomer, and Gly-223, Leu-222, Ile-225, and Ser-226 of the second monomer. The tartrate molecule is hydrogen bonded to Thr-24 and Lys-28 from the Walker-A motif and to the Asp-66 carboxylate group, Lys-55 ammonium group, and Gly-191 nitrogen atom. In the DAPA-AT active site, KAPA is stabilized by several interactions with Phe-326, Trp-369, Trp-370, Tyr-473, Ala-587, Arg-775, Leu-777, and PLP from one monomer and with Gly-678, His-679, and Ser-680 from the other monomer. Phe-326 is located in a loop connected to the  $\alpha$ -helix  $\alpha$ 9, which links BIO3 to BIO1.

mBIO3-BIO1/KAPA superimposed onto mBIO3-BIO1 with a 0.62 Å rmsd value calculated on C $\alpha$ . This comparison showed that binding of KAPA leads only to several local conformational changes in the DAPA-AT domain: The Trp-369 side chain rotates to interact with KAPA and the loop connecting  $\alpha$ 13 to  $\beta$ 16 moves toward the binding site and decreases its solvent accessibility (Figure 8). In the DTBS domain, only weak displacements of the main chain are observed (Figure 7B).

### Structure of mBIO3-BIO1 Bound to DTB

Two DTB molecules and two tartrate ions were observed bound to the DTBS domains (Figure 9; see Supplemental Figure 6 online). DTB interacts with Thr-24, Gln-58, Thr-59, Ser-143, Gly-191, Ser-195, Pro-196, and tartrate from one monomer and Leu-222, Gly-223, Gly-224, and Ile-225 from the second monomer. The rmsd values with mBIO3-BIO1 and mBIO3-BIO1/KAPA are 0.48 and 0.55 Å, respectively, when calculated on C $\alpha$ . Conformational changes of the Trp-369 side chain are observed in the DAPA-AT domain as compared with the holoenzyme, although no ligand molecule is bound in the DAPA-AT catalytic site. This observation strongly suggests close communication between both enzyme catalytic sites. Albeit present into the crystallization solution, no ADP was observed in the protein structure resulting from the presence of 0.2 M ammonium tartrate into the crystallization solution that occupied the ATP binding site.

#### Structure Comparisons with Monofunctional Enzymes

The Arabidopsis DAPA-AT domain from one monomer superimposed with rmsd values in a range 1.78 to 2.01 Å, calculated on C $\alpha$ , with DAPA-AT from *M. tuberculosis* (PDB entry: 3BV0) (Dey et al., 2010) and *E. coli* (PDB entry: 1QJ5) (Käck et al.,

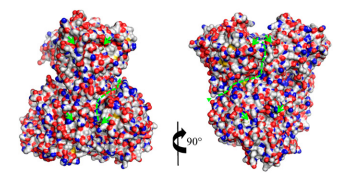


Figure 6. Views of the Crevice at the Surface of mBIO3-BIO1 Going from DAPA-AT (BIO1) to DTBS (BIO3).

The two structures of mBIO3-BIO1 are shown after a rotation of 90° following a vertical axis. KAPA observed in the DAPA-AT and DTBS catalytic sites is drawn as a stick and marked by green arrows. The crevice is highlighted with a dashed line.

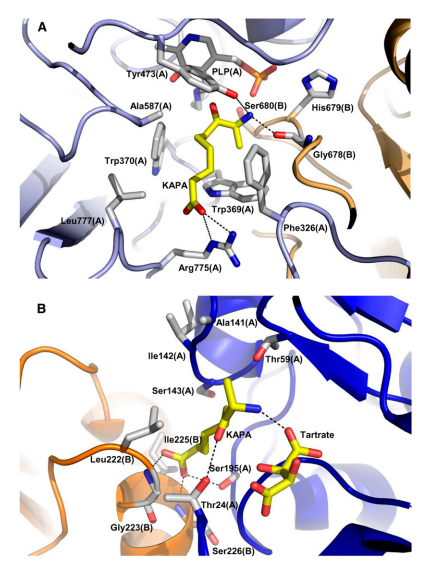


Figure 7. Views of the DAPA-AT (BIO1) and the DTBS (BIO3) Catalytic Sites Bound to KAPA.

(A) The DAPA-AT catalytic site. Each monomer is colored in light blue or light orange. The residues interacting with KAPA and PLP are shown as sticks. (B) The DTBS catalytic site. Each monomer is colored in dark blue or orange. The tartrate ion and the residues interacting with KAPA are shown as sticks.

Hydrogen bonds were calculated by COOT with a distance cutoff of 3.2 Å. They are drawn with dashed lines.

1999), respectively. The main structural difference is the additional three  $\beta$ -stranded  $\beta$ -sheet  $\beta$ 17- $\beta$ 18- $\beta$ 19 in the DAPA-AT domain of the bifunctional enzyme (Figure 10). The *Arabidopsis* DTBS domain superimposed with DTBS of known structures with rmsd values in a range from 1.78 Å (PDB entry: 3OF5; DTBS of *Francisella tularensis*) to 2.47 Å (PDB entry: 3QXX; DTBS of *Helicobacter pylori*). The *Arabidopsis* DTBS domain has three more  $\beta$ -strands,  $\beta$ 1,  $\beta$ 4, and  $\beta$ 5, and the  $\alpha$ -helix  $\alpha$ 3 (Figure 10). These low rmsd values reflect the conservation of the BIO3 and BIO1 folds. The dimers of the monofunctional DTBS and DAPA-AT from bacteria superimposed onto *Arabidopsis* mBIO3-BIO1, and only a few residues are then missing to form the long  $\alpha$ -helix  $\alpha$ 9 connecting BIO3 to BIO1 in mBIO3-BIO1. However, a marked difference between the *Arabidopsis* and prokaryotic structures is the absence of the crevice at the surface of bacterial DAPA-AT and DTBS proteins.

## Mutations Obstructing the External Crevice Affect the Channeling

To test the possibility that the external crevice of the *Arab-idopsis* enzyme could be the route for the DAPA intermediate between both plant enzyme actives sites, S360Y and I793W mutants (mature enzyme numbering) were produced and assayed for the overall DAPA-AT + DTBS activity. These amino acid substitutions are predicted to obstruct the observed external crevice (Figure 11A). In contrast with the wild-type enzyme, mutant variants exhibited progress curves with obvious

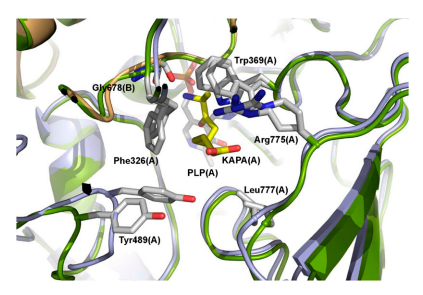


Figure 8. Conformational Changes in the DAPA-AT (BIO1) Catalytic Site upon KAPA Binding.

The unliganded protein is colored in green, and the DAPA-AT bound to KAPA is colored in light blue or light orange. PLP and residues in the active site are drawn as sticks.

lag phases with extrapolated transient times of  $\sim$ 10 to 12 min, respectively (Figure 11B), as expected for nonchanneling controls.

## DISCUSSION

The molecular organization of DAPA-AT and DTBS functional domains as a fusion protein, based on homology searches among eukaryotic and prokaryotic genomes, appears to be a common feature to most biotin prototrophic eukaryotes (flowering plants, mosses, green and red algae, and most ascomycete and basidiomycete fungi with sequenced genomes), whereas in prokaryotes, the presence of separate monofunctional BIO1 and BIO3 orthologs seems to be the rule (Rodionov et al., 2002; Hall and Dietrich, 2007; Muralla et al., 2008; Magliano et al., 2011). This suggests that a fusion event between prokaryotic *bioD* and *bioA* ancestor genes occurred early in the evolution of modern-day eukaryotes. The ability to produce a bicistronic transcript through differential splicing is probably a more recent event because it seems limited to few angiosperms, including *Arabidopsis*, *Brassica*, and rice (*Oryza sativa*; Muralla et al., 2008). Such read-through transcription of

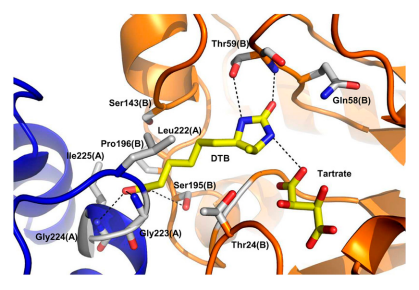


Figure 9. View of the DTBS (BIO3) Catalytic Site Bound to DTB.

Each monomer is colored in dark blue or orange. The tartrate ion, DTB, and the residues interacting with DTB are shown as sticks. Hydrogen bonds were calculated by COOT with a distance cutoff of 3.2 Å. They are drawn with dashed lines.

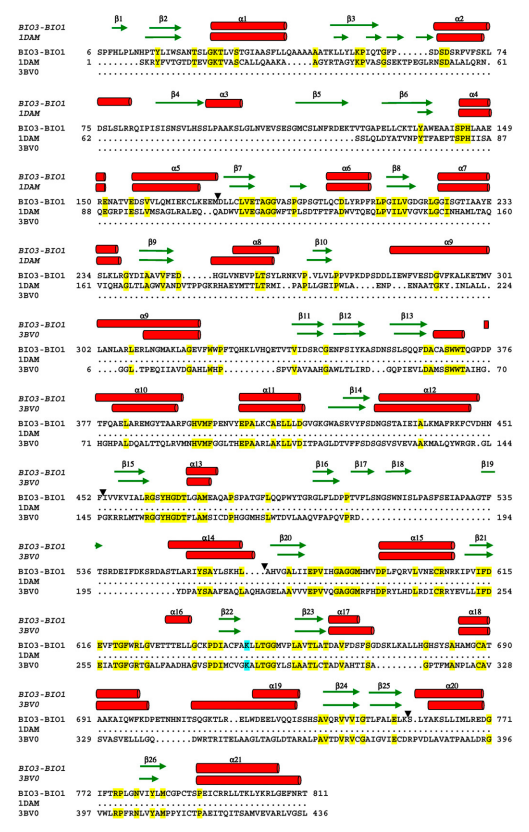


Figure 10. Structure-Based Sequence Alignment between mBIO3-BIO1 and DAPA-AT (bioA) from *M. tuberculosis* (3BV0) (Dey et al., 2010) and DTBS (bioD) from *E. coli* (1DAM) (Käck et al., 1998).

Only residues observed in the density map are reported in the alignment (gaps are indicated by arrowheads). The conserved residues are highlighted in yellow. The Lys covalently linked to PLP is shaded in cyan. The  $\alpha$ -helices and the  $\beta$ -strands are represented by red cylinders and green arrows, respectively.

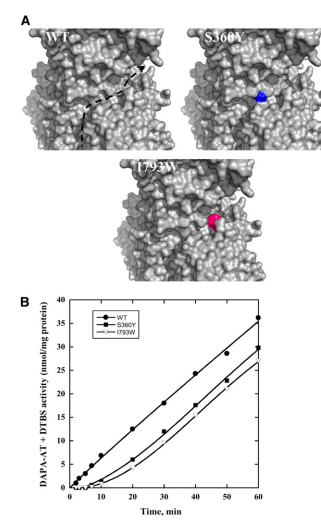


Figure 11. Single Amino Acid Substitutions within the External Crevice of *Arabidopsis* mBIO3-BIO1 Affect Substrate Channeling.

**(A)** Close-up view of the crevice of the wild-type (WT) and mutant enzymes. The crevice is shown with an arrow. Tyr-360 and Trp-793 are colored in blue and pink, respectively.

**(B)** Kinetic study of mBIO3-BIO1 (wild-type, S360Y, and I793W variants) overall catalyzed reaction. DAPA-AT + DTBS activity (time-course formation of DTB) was assayed as in Figure 4A. Data are from a representative experiment repeated three times.

a neighboring tail-to-head tandem genes phenomenon giving rise to mono- and/or bicistronic chimeric mRNA is not uncommon in eukaryotes. For example, computational and experimental analyses estimated that at least 4 to 5% of the tandem gene pairs in the human genome can be eventually transcribed into a single RNA sequence encoding a putative chimeric protein (Parra et al., 2006). Also, up to 155 tandem genes in *Arabidopsis* and rice plants (110 in *Arabidopsis* and 45 in rice) could be transcribed into chimeric mRNA potentially encoding chimeric or separate proteins (Shahmuradov et al., 2010). While the functional significance of the production of both mono- and bicistronic chimeric transcripts remains to be established, it may be a means to increase protein variety in eukaryotic genomes.

Here, we demonstrate through immunodetection using antibodies produced with pure recombinant BIO3-BIO1 protein, that in Arabidopsis, the bifunctional fusion protein is the unique enzyme form found in the plant cells. We cannot exclude the possibility that monofunctional BIO3 and BIO1 protein forms are actually translated from the bicistronic transcript, but if so, could be in amounts too low to be accurately detected by an immunodetection approach. Alternatively, separate BIO3 and BIO1 proteins could be produced but rapidly degraded in the plant cell due to instability. Indeed, owing to the overlapping nature of its reading frames, the bicistronic BIO3-BIO1 transcript is predicted to encode a BIO3 protein form presenting a C-terminal extension corresponding to the N terminus of BIO1 domain and a putative N terminus truncated BIO1 domain form lacking regions conserved among all known DAPA-AT and essential for catalysis, which would be catalytically inactive, if actually produced. Interestingly, our initial attempts to overproduce BIO1 protein alone failed. On the other hand, separate long BIO3 protein could be overproduced in an active form in E. coli but proved to be highly susceptible to proteolytic degradation upon purification (see Supplemental Figure 1 online). Another possibility is that bicistronic BIO3-BIO1 transcript could have a function as a large noncoding RNA. Recent evidence points to a widespread role of these molecules in eukaryotic cells (Costa, 2007).

Production of a bifunctional fusion protein through the nonstop transcription of adjacent tandem genes imply that the destination compartment of the protein expressed by the second tail-to-head gene is determined by transit signals of the first gene. Our data are consistent with Arabidopsis BIO3-BIO1 protein being specifically targeted to mitochondria. This underscores the central role played by plant mitochondria in biotin biosynthesis. Indeed, with the noticeable exception of KAPA synthase, the first committed enzyme of biotin synthesis pathway, all the biotin biosynthetic enzymes are localized within mitochondria in Arabidopsis (Baldet et al., 1997; Picciocchi et al., 2001, 2003; Pinon et al., 2005; Arnal et al., 2006). This also emphasizes the complex compartmentalization of the whole biotin network since biotin-utilizing enzyme are scattered among multiple cell compartments (Alban et al., 2000). With few exceptions, such a situation is also prevalent in many plant vitamin pathways, such as those of folates, ascorbate, pantothenate, niacin, or thiamine (Lunn, 2007; Rébeillé et al., 2007).

In this study, we report the biochemical and structural characterizations of the *Arabidopsis* BIO3-BIO1 protein. The *Arabidopsis* BIO3-BIO1 protein catalyzes both DAPA-AT and DTBS activities. Kinetics of the DAPA-AT reaction alone suggests that under these assay conditions, DAPA produced is sequestered within the DAPA-AT active site, forming a stable enzyme-product complex preventing further catalytic activity. In the presence of CO<sub>2</sub> (a substrate of the DTBS domain), the activity is catalytic, suggesting that DAPA can have access to the DTBS active site, leaving free the DAPA-AT domain for additional turnovers. Finally, the bifunctional enzyme exhibits kinetic cooperativity with respect to all substrates, suggesting conformational changes upon fixation on active sites, consistent with allosteric communication between the two sites. Such conformational changes could favor the transit of DAPA intermediate between both active sites. These changes are particularly well illustrated by the observed kinetics changes relative to ATP when the overall reaction is measured compared with the DTBS reaction alone and by the singular behavior of DAPA intermediate according to the reaction medium conditions. Because DTBS is about 30 times more active than DAPA-AT, it seems likely that the fusion protein arrangement would convey a kinetic improvement owing to proximity effects (reduced soluble phase diffusion) or substrate channeling (direct transfer between catalytic sites).

Several lines of evidence from kinetic and structural studies show that in the course of the overall reaction, DAPA could be directly delivered from the DAPA-AT active site where it is produced to the DTBS active site, without prior diffusion into the bulk solvent. Indeed, when the overall (DAPA-AT + DTBS) reaction is measured in vitro, no delay in the production of DTB, the end product of the whole reaction, is observed (Figure 4). This result contrasts with the expectations for noninteracting DAPA-AT and DTBS activities. In such a situation, the progress curve is predicted to exhibit a lag phase with a transient time  $(\tau)$ equal to the ratio of  $K_{\rm m}$  to  $V_{\rm max}$  for the second enzyme activity (Ovádi et al., 1989). The value of  $\tau$  estimated from the kinetic constants in Table 1 is  $\sim$ 10 min. The absence of such a lag phase in the experimental progress curve for the overall reaction indicates that direct transfer (i.e., channeling) occurs in BIO3-BIO1. Substrate channeling is also supported by the structure analysis of mBIO3-BIO1. The active site for the DAPA-AT domain is a pocket formed by domains of both BIO1 monomers. The active site of each DTBS domain is an open cleft at the interface of both monomers, with more solvent exposed than the DAPA-AT active sites. Both catalytic domains are distant and too far apart (~48 Å) to allow direct transfer of the DAPA intermediate simply by proximity of the two active sites. DAPA-AT and DTBS active sites are connected by a small groove at the interface of the BIO3-BIO1 monomers. However, in all the structures solved, this groove is interrupted and not wide enough to allow the transfer of a DAPA molecule between both active sites. Thus, this route for the intermediate transfer is hardly predictable unless large protein conformational rearrangements.

Because the three-dimensional structure and the dimer interface are similar to those from bacterial enzymes and no channel or putative channel is observed, the surface was also analyzed. Indeed, surface channeling has been proposed for the bifunctional enzyme dihydrofolate reductase-thymidylate synthase (Knighton et al., 1994). The main difference between the bifunctional and the monofunctional enzymes is an  $\sim$ 95 Å long and 12 Å average width crevice at the enzyme surface going from the DAPA-AT catalytic site to that of DTBS in each monomer of mBIO3-BIO1. Inspection of the crevice shows that it consisted of polar amino acids corresponding to nonconserved residues belonging to common secondary structures. This polar crevice could be a candidate for the DAPA transfer from DAPA-AT to DTBS. The fact that mutations blocking the trench conduce to the recording of overall DAPA-AT + DTBS activity progress curves exhibiting lag phases with extrapolated transient times close to the theoretical value of  $\tau$  for noninteracting DAPA-AT and DTBS activities supports this hypothesis.

Interestingly, analysis of the crystal structures also shows that the additional  $\beta$ -strand  $\beta$ -1 and the  $\alpha$ -helix  $\alpha$ -9 are the key structural elements for acquisition of the bifunctionality. Indeed,  $\alpha$ -helix  $\alpha$ -9 covalently connects DAPA-AT to DTBS, and the  $\beta$ -strand  $\beta$ -1 of DTBS interacts with the  $\beta$ -strand  $\beta$ -13 of DAPA-AT to stabilize the interaction between the two domains through an antiparallel four-stranded  $\beta$ -sheet together with  $\beta$ -11 and  $\beta$ -12.

Finally, our work raises the question of the biological significance of intermediate channeling, if any, in biotin metabolism. Substrate channeling has many advantages over the free diffusion within the bulk solvent. First, the transit time from one active site to the next is reduced; therefore, unfavorable equilibria, competing enzymatic transformations, or unwanted side reactions can thus be circumvented (Ushiroyama et al., 1992; Herrmann, 1995). Chemically labile intermediates can be protected from breakdown by the aqueous external environment during the transfer between distant active sites (Rudolph and Stubbe, 1995). Sequestration also provides a means of preventing a potentially toxic intermediate product from being released to the solvent (Manjasetty et al., 2003). Last, substrate channeling can trap metabolites with high diffusivity, thus preventing their loss in the surrounding medium or across the cell membranes (Chittur et al., 2001; Barends et al., 2008). DAPA is not known to be particularly labile or toxic and its only fate is biotin synthesis; thus, there is no apparent practical requirement for the sequestration of this intermediate. Because aminotransferases catalyze reversible reactions and because DAPA-AT is much less active than DTBS in Arabidopsis bifunctional enzyme, DAPA trapping in the BIO1 active site and direct transfer to the BIO3 domain upon binding of the BIO3 substrates could be a means of avoiding having the DAPA-AT reaction proceed in the reverse direction, hence increasing the efficiency of the overall reaction. This could be especially critical owing to the low abundance of biotin intermediates in the plant cells (Alban et al., 2000).

In conclusion, we provide an exhaustive biochemical, kinetic, and structural characterization of a bifunctional enzyme of the biotin synthesis pathway. Because this enzyme form is specific to biotin-synthesizing eukaryotes (Hall and Dietrich, 2007; Muralla et al., 2008; Magliano et al., 2011), this characterization of the *Arabidopsis* enzyme provides a unique working model for ongoing mechanistic studies critical for a better understanding of biotin synthesis and regulation in eukaryotes. Also, elucidation of the structure of a plant bifunctional DTBS–DAPA-AT creates a rich molecular platform that will be useful for the design of new inhibitor families having herbicidal and/or fungal activities.

## METHODS

#### **Plant Materials**

Arabidopsis thaliana (ecotype Columbia) seeds were grown on soil under greenhouse conditions (23°C with a 16-h photoperiod and a light intensity of 100  $\mu$ E m<sup>-2</sup> s<sup>-1</sup>) until harvested for analysis. *Arabidopsis* (ecotype Columbia) cell suspension cultures were grown under continuous white

light (40  $\mu E$  m<sup>-2</sup> s<sup>-1</sup>) at 23°C with rotary agitation at 125 rpm in Gamborg's B5 medium supplemented with 1  $\mu M$  2-naphtalene acetic acid and 1.5% (w/v) Suc.

#### Chemicals

*Rac*-KAPA and DAPA were obtained as described previously (Baldet et al., 1993). (S)-KAPA was a kind gift from Olivier Ploux and Stéphane Mann (Mann et al., 2009). Isopropylthio-β-D-galactoside was from Bioprobe Systems. NaH<sup>14</sup>CO<sub>3</sub> (52 mCi/mmol) was from Perkin-Elmer. His binding resin was from Qiagen. All other biochemicals were obtained from Sigma-Aldrich and were the purest grade available.

### Protein Determination, SDS-PAGE, and Immunoblot Analysis

Proteins were measured either by the Bradford method (Bradford, 1976) using Bio-Rad protein assay reagent, with  $\gamma$ -globulin as a standard, or for pure proteins by measuring the absorbance at 205 nm (Scopes, 1974). Total soluble proteins from Arabidopsis tissues were extracted by grinding powdered samples in 20 mM MOPS-KOH, pH 7.5, 5% (w/v) glycerol, 1 mM DTT, 1 mM EDTA, 1 mM benzamidine-HCl, 5 mM ε-amino caproic acid, and the complete protease inhibitor cocktail from Roche Applied Science. Samples were centrifuged at 130,000g for 20 min at 4°C, and the supernatant was used as a source of soluble proteins. Arabidopsis cell subfractions were purified as described previously (Pinon et al., 2005). Briefly, chloroplast and mitochondria were isolated from Arabidopsis leaves and purified on Percoll gradients. A cytosolic-enriched fraction was prepared from Arabidopsis cultured cells by gentle rupture of isolated protoplasts followed by differential centrifugation steps. Purity of cytosol, chloroplast, and mitochondrial preparations was assessed by measurement of specific subcellular markers and was found to be higher that 95%, in each case (Pinon et al., 2005). Proteins from Arabidopsis cell subfractions were resolved by SDS-PAGE and electroblotted to nitrocellulose membrane. The blots were probed using affinity-purified BIO3-BIO1 antibodies produced in guinea pigs (Charles River Laboratories, France) and horseradish peroxidase-conjugated anti-guinea pig IgGs, and detection was achieved by chemifluorescence using the ECL Plus Western Blotting detection reagents and a Typhoon 9400 scanner (GE Healthcare Europe).

#### **Cloning, Protein Expression, and Purification**

Sequences of primers used in this study are listed in Supplemental Table 1 online. Full-length monocistronic and bicistronic BIO3-BIO1 cDNAs and cDNA encoding mature BIO3-BIO1 protein (mBIO3-BIO1; residues 23 to 833) were obtained by PCR amplification of an RT reaction using total RNAs isolated from Arabidopsis seedlings (RNeasy plant mini kit from Qiagen). Primers were designed on the basis of the predicted sequence of BIO3-BIO1 locus and available cDNA sequences (Muralla et al., 2008) and generated appropriate restriction sites for the cloning of cDNA into expression vectors. The PCR fragments were first cloned into pPCR-Script (Stratagene) for maintenance and sequenced in both strands (Beckman Coulter Genomics). For overexpression, the monocistronic BIO3-BIO1 sequences (full length or devoid of the transit peptide sequence) were inserted between the Ndel and Sacl sites of pET-28b (Novagen). This procedure added an N-terminal hexa-His tag. The F326Y, S360Y, and 1793W variants of mBIO3-BIO1 were constructed using the Quikchange II site-directed mutagenesis kit (Stratagene) and the pET-mBIO3-BIO1 plasmid as a matrix. The resulting pET-BIO3-BIO1 constructs were introduced first into Escherichia coli DH5 $\alpha$  cells and then into the E. coli overexpression host Rosetta 2 (DE3) (Stratagene).

The transformed cell cultures were grown at 37°C in Luria-Bertani medium supplemented with 10  $\mu$ M pyridoxine, 15  $\mu$ M thiamin, and the appropriate antibiotics until  $A_{600}$  was 0.6, at which point isopropylthio-

β-D-galactoside was added to a final concentration of 0.4 mM. Incubations were continued for 40 h at 20°C. Pelleted cells from 2.5-liter cultures were resuspended in 30 mL of buffer A (20 mM Tris-HCl, pH 8, 500 mM NaCl, 100 µM PLP, and EDTA-free complete protease inhibitor cocktail from Roche Applied Science) supplemented with 10 mM imidazole and disrupted by sonication with a Vibra-Cell disrupter (Branson Ultrasonics) The soluble protein extracts were separated from the cell debris by centrifugation at 18,000g for 30 min and applied onto nickelnitrilotriacetic acid-agarose columns (1  $\times$  3 cm) previously equilibrated with the same buffer. The columns were washed with buffer A supplemented with 50 mM imidazole. The recombinant proteins were then eluted with buffer A supplemented with 250 mM imidazole. Fractions containing proteins were pooled and dialyzed against Tris-buffered saline. Purifications were achieved by size exclusion chromatography (Superdex 200 HiLoad, 26/60; GE Healthcare) in buffer B (20 mM Tris-HCl, pH 8, 150 mM NaCl, 10 µM PLP, and 10% [v/v] glycerol). Purified enzymes were concentrated by filtration (Macrosep, 30-kD cutoff; Pall Filtron), aliquoted, and stored at -80°C until use.

The seleno-Met F326Y mutant enzyme ([SeMet]-mBIO3-BIO1) was produced and purified similarly except that the expression was performed in the *E. coli* B834 (DE3) pLys S host strain, and growth was conducted in 1 liter of a minimal medium containing 40 mg of seleno-Met.

The *E. coli bioD* JW0761 strain carrying the pCA24N-bioD plasmid encoding *E. coli* DTBS headed by an N-terminal hexa-His tag was obtained from the National BioResource Project: *E. coli* (National Institute of Genetics). BioD protein was produced and purified in the same way as BIO3-BIO1 proteins, except for the omission of PLP in purification buffers.

#### **GFP Fusion Targeting Analyses**

Primers were designed to amplify the sequence encoding for the complete BIO3-BIO1 protein (see Supplemental Table 1 online) and allowed its cloning in frame to the 5' extremity of the GFP-coding region in the pTH2 vector, at Sall and Ncol sites. The GFP reporter plasmid pTH2 expressed an engineered version of GFP under the control of the cauliflower mosaic virus 35S promoter (Chiu et al., 1996). The transit peptide sequences of the small subunit of ribulose-1,5-bisphosphate carboxylase/oxygenase from Arabidopsis (55 residues, ATS1A gene; GenBank accession number X13611) and dihydropterin pyrophosphokinase-dihydropteroate synthase from pea (Pisum sativum; 28 residues; (Rébeillé et al., 1997)) were used as controls for the targeting of GFP to plastids and mitochondria, respectively. Transient transformation of Arabidopsis protoplasts prepared from a 4-d-old cell suspension culture was achieved using 40 µg of plasmid construct by the polyethylene glycol method, essentially as described (Pinon et al., 2005). Transformed cells were incubated at 23°C for 36 h and analyzed by epifluorescence microscopy using a Zeiss Axioplan2 fluorescence microscope, and images were captured with a digital charge-coupled device camera (Hamamatsu). The filter sets used were Zeiss filter set 13, 488013-0000 (exciter BP 470/20, beamsplitter FT 493, emitter BP 505-530) and Zeiss filter set 15, 488015-0000 (exciter BP 546/12, beamsplitter FT 580, emitter LP 590) for GFP and chlorophyll fluorescence, respectively.

#### **Activity Measurements**

The overall (DAPA-AT + DTBS) reaction was assayed by measuring the formation of acid-stable [<sup>14</sup>C]-DTB from acid-labile H<sup>14</sup>CO<sub>3</sub>. The standard assay consisted of 100 mM EPPS, pH 8.6, 1 mM DTT, 0.1 M NaCl, 0.2 mM PLP, 3 mM AdoMet, 20  $\mu$ M (S)-KAPA, 0.3 mM ATP, 5 mM MgCl<sub>2</sub>, 10 mM NaH<sup>14</sup>CO<sub>3</sub> (1.6 mCi/mmol), and 0.4 nmol/mL pure mBIO3-BIO1. The assays were initiated by the addition of the enzyme. After 2 to 120 min incubation at 37°C, 500  $\mu$ L aliquots of the reaction mixture were mixed with 250  $\mu$ L of 20% (v/v) acetic acid to stop the reaction. The solution was taken to dryness, and the acid-stable radioactivity was quantified in

a liquid scintillation counter. Duplicate assays without KAPA were run as controls.

DAPA-AT activity was assayed by measuring the amount of [<sup>14</sup>C]-DTB formed in a discontinuous coupled reaction using as the coupling enzyme monofunctional DTBS (bioD) from *E. coli*, which catalyzes the synthesis of DTB from DAPA and CO<sub>2</sub> in the presence of ATP-Mg. The standard assay consisted of 100 mM EPPS, pH 8.6, 1 mM DTT, 0.1 M NaCl, 0.2 mM PLP, 3 mM AdoMet, 20  $\mu$ M (S)-KAPA, and 0.2 to 2 nmol/mL pure mBIO3-BIO1. The reaction was performed at 37°C and arrested by heating 500- $\mu$ L aliquots of the mixture for 2 min at 80°C and filtration through a 30-kD cutoff membrane (Nanosep; Pall Filtron) to remove the proteins. Duplicate assays without KAPA or AdoMet were run as controls. The amount of DAPA formed was determined by radiochemical assay after complete conversion to DTB, by incubating the filtrate for another 2 h in the presence of 0.3 mM ATP, 5 mM MgCl<sub>2</sub>, 10 mM NaH<sup>14</sup>CO<sub>3</sub> (1.6 mCi/mmol), and 4 nmol pure bioD. The reaction was stopped by acetic acid treatment and acid-stable radioactivity quantified as described above.

DTBS activity was assayed as described above for the overall (DAPA-AT + DTBS) reaction with the following modifications: The standard assay consisted of 100 mM HEPES, pH 8, 1 mM DTT, 0.1 M NaCl, 0.3 mM ATP, 5 mM MgCl<sub>2</sub>, 200  $\mu$ M DAPA, 10 mM NaH<sup>14</sup>CO<sub>3</sub> (1.6 mCi/mmol), and 1 nmol/mL pure enzyme. The reaction was stopped by mixing 100- $\mu$ L aliquots of the reaction mixture with 250  $\mu$ L of 10% (v/v) acetic acid. Duplicate assays without DAPA were run as controls.

Kinetics of product formation were linear and rates varied linearly with enzyme concentration, demonstrating adherence to steady state conditions. Kinetic data were fitted to the appropriate rate equations by nonlinear regression analyses using the KaleidaGraph program (Abelbeck Software). Production of [<sup>14</sup>C]-DTB as the end product of the reactions was controlled by HP TLC analysis and phosphor imaging detection according to the procedure described by Picciocchi et al. (2001).

### **Crystallization and Data Collection**

Searches for crystallization conditions of native mBIO3-BIO1 were performed using the TECAN robot at the Institut de Biologie Structurale in Grenoble, France. One microliter of protein (10 mg/mL) was mixed with 1 µL crystallization solution. The mutation F326Y (mature protein numbering) was found to improve the recombinant enzyme solubility and stability, with no significant impact in Arabidopsis on the enzyme kinetic parameters (Table 1). Then it was subsequently used for crystallization of [SeMet]-mBIO3-BIO1. The native protein (mBIO3-BIO1) was crystallized in 0.1 M BisTris, pH 5.9, 0.2 M LiSO<sub>4</sub>, and 15% (w/v) polyethylene glycol 3350. [SeMet]-mBIO3-BIO1 F326Y (7.5 mg/mL) was crystallized in the same conditions with the addition of DTT in a 2 to 10 mM range. Extensive searches of crystallization conditions of mBIO3-BIO1/complexes with either 5 mM DAPA, KAPA, DTB, AdoMet, ATP, AMP-PNP, or CO<sub>2</sub>, or various combinations of these ligands, were performed using the highthroughput crystallization facilities of EMBL. Good diffracting crystals were obtained for mBIO3-BIO1/KAPA and mBIO3-BIO1/ADP/DTB using 0.2 M ammonium tartrate, pH 6.2, and 15% (w/v) polyethylene glycol 3350. Data for the native protein, [SeMet]-mBIO3-BIO1, and mBIO3-BIO1/KAPA were collected on the FIP-BM30A beamline (Roth et al., 2002) at the European Synchrotron Radiation Facility (Grenoble, France) using an ADSC Q315r charge-coupled device detector. Data for mBIO3-BIO1/ ADP/DTB were collected on ID14-1 at the European Synchrotron Radiation Facility using an ADSC Q210 charge-coupled device detector. All the data were processed and scaled using XDS (Kabsch, 1993) (see Supplemental Table 2 online).

### Single-Wavelength Anomalous Diffraction Phasing

The phase problem could not be solved using molecular replacement and atomic coordinates of the bacterial enzymes. A three-wavelength multiwavelength anomalous diffraction data set was collected on FIP-BM30A at 2.7-Å resolution, at the peak, inflection, and remote high energy of the Se absorption edge. A total of 32 selenium atoms were found using Phenix (Adams et al., 2002) and the data collected at the peak of the Se absorption edge. The resulting phases were further improved by solvent flattening and density averaging using Phenix allowing for partial automated chain tracing (see Supplemental Table 2 online).

#### Model Building and Refinement

The polyalanine model of [SeMet]-mBIO3-BIO1 was constructed at 2.7-Å resolution, and side chains were added using COOT (Emsley and Cowtan, 2004). The model was refined by restrained refinement and molecular dynamic using Phenix and water molecules were added, also using Phenix. A simulated-annealing 2Fo-Fc omit electron density map was calculated to check the model.

The phase problem for mBIO3-BIO1 was solved by molecular replacement using PHASER (McCoy et al., 2007) and the [SeMet]-mBIO3-BIO1 F326Y atomic coordinates as template. The phase problem for mBIO3-BIO1/ADP/DTB and mBIO3-BIO1/KAPA was solved by molecular replacement using PHASER and the mBIO3-BIO1 atomic coordinates as template. All the structures were refined by restrained refinement and molecular dynamic with Phenix and using non-crystallographic symmetry restrains. In the case of mBIO3-BIO1, the noncrystallographic symmetry was applied only in the dimers. Water molecules were added using Phenix. At the end of the refinement, the occupancies of the ligands (Mg<sup>2+</sup>, SO<sub>4</sub><sup>2-</sup>, DTB, KAPA, and tartrate ion) were also refined. The refinement statistics are given in Supplemental Table 3 online.

In all the models, several residues and side chains were not observed into the electron density maps. Also, the electron density for the six His tag and the preceding five residues at the N terminus was not visible. The structure comparisons with the monofunctional enzymes were performed using the most complete dimer (dimer A/B) of mBIO3-BIO1 (crystals obtained in space group P1). This dimer was also chosen to describe the *Arabidopsis* mBIO3-BIO1 structure. The structure superimposition was performed using PDBeFold (Krissinel and Henrick, 2004) at European Bioinformatics Institute. All the models were checked using PROCHECK (Laskowski et al., 1993) and WHATCHECK (Hooft et al., 1996). All the figures were prepared using Pymol (Delano, 2002).

#### **Accession Numbers**

Sequence data from this article can be found in the GenBank/EMBL or Arabidopsis Genome Initiative databases under the following accession numbers: *BIO3-BIO1* (*At5g57590*; BOF481); monocistronic *BIO3-BIO1* (HQ857557); bicistronic *BIO3-BIO1* (HQ857558); *E. coli BioD* (1DAM; NP\_415299.1); and *Mycobacterium tuberculosis BioA* (3BV0; AAK45886.1). The [SeMet]-mBIO3-BIO1 F326Y, mBIO3-BIO1, mBIO3-BIO1/KAPA, and mBIO3-BIO1/ADP/DTB atomic coordinates and structure factors have been deposited in the Protein Data Bank with the accession code numbers 4A0F, 4A0G, 4A0H, and 4A0R, respectively.

#### Supplemental Data

The following materials are available in the online version of this article.

**Supplemental Figure 1.** Analysis of BIO3-BIO1 and BIO3 Production in *E. coli* Using Polyclonal Antibodies Raised against Recombinant *Arabidopsis* BIO3-BIO1 Protein.

**Supplemental Figure 2.** Expression Analysis of *Arabidopsis* Bicistronic and Monocistronic *BIO3-BIO1* mRNAs by Real-Time PCR.

**Supplemental Figure 3.** Part of an Amino Acid Alignment of DAPA Aminotransferases from a Selection of Bacteria, Plants, and Fungi, Highlighting Residue Tyr-17 (*E. coli* Numbering).

Supplemental Figure 4. Topology of mBIO3-BIO1.

**Supplemental Figure 5.** View of the Crevice at the Surface of mBIO3-BIO1.

Supplemental Figure 6. Simulated Annealing Omit Maps for the mBIO3-BIO1 Ligands.

Supplemental Table 1. Synthetic Oligonucleotides Used in This Study.

Supplemental Table 2. Data Collection.

Supplemental Table 3. Refinement Statistics.

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#### AUTHOR CONTRIBUTIONS

D.C., R.D., and C.A. designed the research. D.C., R.D., V.P., and C.A. performed the research. D.C., R.D., C.M., J.-L.F., and C.A. analyzed data. D.C. and C.A. wrote the article with contribution from R.D.

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