Molecular and Biochemical Analysis of the Plastidic ADP-glucose Transporter (ZmBT1) from Zea mays

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Physiological studies on the Brittle1 maize mutant have provided circumstantial evidence that ZmBT1 (Zea mays Brittle1 protein) is involved in the ADP-Glc transport into maize endosperm plastids, but up to now, no direct ADP-Glc transport mediated by ZmBT1 has ever been shown. The heterologous synthesis of ZmBT1 in Escherichia coli cells leads to the functional integration of ZmBT1 into the bacterial cytoplasmic membrane. ZmBT1 transports ADP-Glc in counterexchange with ADP with apparent affinities of about 850 and 465 μM, respectively. Recently, a complete ferredoxin/thioredoxin system has been identified in cereal amyloplasts and BT1 has been proposed as a potential Trx target protein (Balmer, Y., Vensel, W. H., Cai, N., Manieri, W., Schurmann, P., Hurkman, W. J., and Buchanan, B. B. (2006) Proc. Natl. Acad. Sci. U. S. A. 103, 2988–2993). Interestingly, we revealed that the transport activity of ZmBT1 is reversibly regulated by redox reagents such as diamide and dithiothreitol. The expression of ZmBT1 is restricted to endosperm tissues during starch synthesis, whereas a recently identified BT1 maize homologue, the ZmBT1–2, exhibits a ubiquitous expression pattern in hetero- and autotrophic tissues indicating different physiological roles for both maize BT1 isoforms. BT1 homologues are present in both mono- and dicot-yledonous plants. Phylogenetic analyses classify the BT1 family into two phylogenetically and biochemically distinct groups. The first group comprises BT1 orthologues restricted to cereals where they mediate the ADP-Glc transport into cereal endosperm storage plastids during starch synthesis. The second group occurs in mono- and dicotyledonous plants and is most probably involved in the export of adenine nucleotides synthesized inside plastids.

Cereal crops accumulate starch in seed endosperm plastids as main energy reserve. The pathway of starch synthesis in cereal endosperms is unique and requires enzyme isoforms that are not present in other tissues or non-cereal plants. The ability of heterotrophic plastids to utilize cytosolic precursors to support their biosynthetic and catabolic pathways depends on the presence of specific transporters in the plastid envelope. In cereal endosperms, the ADP-glucose pyrophosphorylase (AGPase), which catalyzes the first committed and rate-limiting step in starch biosynthesis, is mainly localized in the cytosol with a total AGPase activity of about 85–95% (2). Therefore, ADP-glucose (ADP-Glc) is synthesized in the cytosol of cereal endosperms as the main precursor for starch synthesis and has to be subsequently imported into the storage plastids.

Several maize (Zea mays L.) endosperm mutants affected in starch quality or quantity were used to elucidate critical steps in amyloplast starch synthesis. The Waxy gene, which encodes a starch-granule-bound starch synthase involved in amylose synthesis (3) and the Shrunken-2 and Brittle-2 genes, which encode subunits of the AGPase (4–6), were shown to be important for starch synthesis in maize. The Brittle-1 (BT1) maize mutant was identified in 1926 (7, 8) and corresponding endosperm is severely reduced in starch content, which results in kernels with a collapsed angular appearance at maturity. The BT1 protein from Z. mays (ZmBT1) belongs to the mitochondrial carrier family and is located in the amyloplast envelope membrane (9, 10). The absence of ZmBT1 correlates with a 12-fold higher level of ADP-Glc in the cytosol of BT1 mutant endosperm than in wild-type endosperm (11) and BT1 mutant kernels accumulate about 80% less starch than wild-type kernels (12). The incorporation of externally applied ADP-Glc into starch in amyloplasts isolated from BT1 mutant endosperm was reduced to about 25% compared with wild-type amyloplasts (13). These results indicate that ZmBT1 is involved in the transport of ADP-Glc into maize endosperm plastids (14), but up to now, no direct ADP-Glc transport mediated by ZmBT1 has been shown.

Recently, a BT1 homologue from a non-cereal plant, namely potato (StBT1 from Solanum tuberosum), was functionally characterized as a plastidic adenine nucleotide uniporter expressed in sink and source tissues (15). The physiological role of StBT1 is most likely the export of purine nucleotides that are exclusively de novo synthesized in plastids of autotrophic and heterotrophic tissues (15, 16). These novel findings further

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† The on-line version of this article (available at http://www.jbc.org) contains supplemental Table S1 and Figs. S1–S4.

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2 The abbreviations used are: AGPase, ADP-glucose pyrophosphorylase; RT, reverse transcriptase; IPTG, isopropyl β-D-thiogalactopyranoside; EF, elongation factor; Trx, thioredoxin; DTT, dithiothreitol.
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urged for a comprehensive analysis of the biochemical transport features of the maize homologue ZmBT1. We present here direct transport properties of the heterologously expressed ZmBT1. For this we exploited an Escherichia coli expression system that was successfully used to analyze plastidic or mitochondrial carrier proteins (15, 17, 18). We also reviewed the endosperm-specific expression of ZmBT1 and proved the occurrence and expression pattern of further ZmBT1 homologues in maize.

EXPERIMENTAL PROCEDURES

Cultivation of Plants—Maize plants (Z. mays L.) were grown in a greenhouse at 22–26 °C and watered once a day. The ambient light period was extended to 16 h/day with Philips Sont-Agro lights (200 μmol quanta m⁻² s⁻¹). For RNA and genomic DNA isolation, maize tissues were collected and immediately frozen in liquid nitrogen until use.

Southern Blot Analyses—Genomic DNA was isolated from 4 g of frozen leaf tissue as described by Dellaporta et al. (19). About 10 μg of genomic DNA were digested with high concentrated restriction enzymes (BamHI, HindIII, Xbal, and XhoI). DNA fragments were separated in 0.8% agarose gels, denatured, and neutralized according to standard procedures (20). After transfer to nylon membranes (Hybond-N membranes, Amersham) DNA fragments were fixed to the nylon filters by optimal UV cross-link (254 nm, 120 millijoule cm⁻²) in a Spectrolinker. The membranes were prehybridized for 60 min at 68 °C in 6× SSPE buffer (20× SSPE contains 3.6 M NaCl, 0.2 M sodium phosphate, 0.02 M EDTA, pH 7.7) supplemented with 1% SDS, 100 μg/ml herring sperm DNA, and 5× Denhardt’s reagent. Hybridization was performed for 16 h at 68 °C in the same solution plus the complete ZmBT1-cDNA fragment labeled by the random primer method with [α³²P]dCTP using the Ready-To-Go prime kit (Amersham Biosciences). Afterward, the membranes were washed once at 37 °C for 15 min in 7× SSPE, 0.5% SDS and twice at 60 °C for 15 min in 0.1× SSPE, 1% SDS. Blots were visualized by a Cyclon PhosphorImager (Packard).

Quantitative Real-time RT-PCR—Total RNA was prepared from various maize tissues using the RNeasy Plant Mini Kit (Qiagen, Hilden, Germany). To remove any contaminating DNA, the samples were treated with deoxyribonuclease (RNase-free DNase Kit, Qiagen). Quantitative PCR was performed using MyIQ-Cycler (Bio-Rad) and IQ SYBR Green Supermix (Bio-Rad) according to the manufacturer’s instructions with the following cycler conditions: 20 min at 50 °C; 15 min at 95 °C; 55 cycles: 15 s at 95 °C, 25 s at 58 °C, and 40 s at 72 °C. The sequences of the gene-specific oligonucleotides used for real-time RT-PCR are the following: ZmBT1-sense, 5′-CAA-GGCTATGAGGACATTTCCC-3′; ZmBT1-antisense, 5′-TCG-TAGGCGTAG-AAGTTACAGG-3′; ZmBT1–2-sense, 5′-AAC-AGCATATTACCCGTGGA-3′; ZmBT1–2-antisense, 5′-GCTTC-ACG-GACTTACCCGTGGA-3′; EF-1α-sense, 5′-GCTTC-ACG-GACTTACCCGTGGA-3′; and EF-1α-antisense, 5′-TGGCTTG-GAGCTTTGATCA-GTCA-3′. The housekeeping gene in Z. mays encoding the elongation factor EF-1α (AF136829) was used for quantitative normalization. The specificity of the obtained RT-PCR products was controlled on 2% agarose gels.

Plasmid Construct for Heterologous Expression of ZmBT1 in E. coli—DNA manipulations were performed essentially as described by Sambrook et al. (20). The expression plasmid (pET 16b, Novagen, Heidelberg, Germany) encoding the recombinant ZmBT1 protein with an additional N-terminal tag of 10 histidine residues was constructed as follows: the cDNA coding the entire ZmBT1 was generated by PCR from first strand cDNA of maize endosperm tissue using Pfu DNA polymerase (Invitrogen). A sense primer including an NdeI restriction site and an antisense primer were used for the PCR (sense: 5′-GACT-GAcatATGGCGGCGACAATG-3′; the lower case letters indicate the introduced base exchange to create an NdeI restriction site; antisense: 5′-CTACCTTTGCGGCGACAATG-3′). The obtained PCR product was purified (Nucleospin Extract II, Macherey & Nagel, Düren, Germany), subcloned into the EcoRI restriction site of the plasmid pBSK (Stratagene), and checked by sequencing on both strands by chain termination reaction (MWG-Biotech, Ebersberg, Germany). For the construction of the E. coli expression plasmid (encoding His₁₀–ZmBT1), the Ndel/BamHI DNA insert of the pBSK-plasmid was introduced in-frame into the corresponding restriction sites of the isopropyl β-D-thiogalactopyranoside (IPTG)-inducible T7-RNA polymerase bacterial expression vector pET16b (Novagen, Heidelberg, Germany). Transformations of E. coli were carried out according to standard protocols. The nucleotide sequence of the ZmBT1 reported in this paper is available at the EMBL data base (www.ebi.ac.uk/embl/) under the accession number BT016796.

Heterologous Expression of ZmBT1 in E. coli—The E. coli strain Rosetta(DE3) was used for heterologous expression. The cDNA sequence encoding ZmBT1 under control of the T7-promoter was transcribed after IPTG induction of the T7-RNA polymerase (21). E. coli cells harboring the ZmBT1 expression plasmid (or control expression plasmid pET16b) were grown at 37 °C in TB/Amp/Clin medium (TB: 2.5 g/liter KH₂PO₄, 12.5 g/liter NaCl, 15% (v/v) glycerol, pH 7.0). An optical density (A₆₀₀) of 0.5–0.6 was required for the initiation of T7-RNA polymerase expression by addition of IPTG (final concentration, 1 mM). Cells were grown for 1 h after induction and collected by centrifugation for 5 min at 4,000 × g (room temperature, Sorvall RC5B centrifuge, rotor type SS34; Sorvall-Du Pont, Dreieich, Germany). The pellet was resuspended to an A₆₀₀ of 6 using potassium phosphate buffer (50 mM, pH 7.0) (22) and promptly used for uptake experiments.

Membrane integration of the recombinant full-length ZmBT1 in E. coli was confirmed by enrichment of the histidine-tagged protein and Western blot analysis using a ZmBT1 specific antibody (10). E. coli cells (20 ml) harboring the ZmBT1 expression plasmid (or the pET16b control plasmid) were collected 1 h after IPTG induction (controls without IPTG) and transferred to liquid nitrogen to destroy cell intactness. After resuspension in a medium consisting of 10 mM Tris/HCl (pH 7.5), 1 mM EDTA, 0.1 mM Pefabloc, and 15% (v/v) glycerol, cells were further disrupted by ultrasonication (250 W, 3 × 30 s, 4 °C) and the suspension was centrifuged (10 min, 15,800 × g, 4 °C) to remove unbroken cells and inclusion bodies. Membranes extracted in the supernatant were sedimented for 45
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Sequence Analysis of ZmBT1—The ZmBT1 cDNA clone was isolated from the first strand cDNA of maize endosperm as given under “Experimental Procedures.” Six independent PCR products of ZmBT1 have been sequenced and found to exhibit 100% identity excluding failure of the Pfu polymerase. Compared with the open reading frame of the ZmBT1 sequence published by Sullivan et al. (28), our ZmBT1 cDNA sequence differs in the following positions: six additional nucleotides (G at bp position 246, 295, and 411, and GGA at bp position 1257–1259) and two nucleotide changes (GC instead of CG at bp position 338–339). The resultant triple frameshift modified the deduced amino acid sequence ahead of the first predicted transmembrane domain (amino acid position 83–137) and the additional amino acid residue at the C terminus (position 419) led to a substantial higher similarity to other BT1 isoforms (Fig. 1). Interestingly, the ZmBT1 cDNA sequence we determined was recently submitted by Lai et al. as well (2004, direct submission, accession number BT016796, www.ncbi.nlm.nih.gov). The deduced ZmBT1 protein sequence was aligned with putative BT1 homologues from wheat (Triticum aestivum), rice (Oryza sativa), barley (Hordeum vulgare), maize (Z. mays), barrel medic (Medicago truncatula), potato (S. tuberosum), and Arabidopsis thaliana (Fig. 1).

The amino acid sequence of the BT1 homologues consists of three tandem repeats of ~100 residues showing 6 putative transmembrane helices (15). All BT1 homologues possess three conserved mitochondrial energy transfer signatures that are characteristic for membrane proteins belonging to the mitochondrial carrier family (29, 30). Phylogenetic analyses classify BT1 homologues as members of the mitochondrial carrier family forming a monophyletic cluster (31). A comparison of the BT1 isoforms revealed a high similarity of ZmBT1 to TaBT1 (66% identity, 75% similarity), OsBT1–1 (66% identity, 73% similarity), HvNST (65% identity, 74% similarity), and to a lower extent to OsBT1–3 (58% identity, 68% similarity), ZmBT1–2 (58% identity, 68% similarity), OsBT1–2 (48% identity, 60% similarity), MtBT1 (46% identity, 63% similarity), StBT1 (45% identity, 61% similarity), and AtBT1 (44% identity, 61% similarity). Phylogenetic analyses of the BT1 homologues indicate that the BT1 protein family can be divided into two subgroups. The first phylogenetic cluster comprises only BT1 homologues from monocotyledonous plants including ZmBT1 (Fig. 2). The second cluster includes BT1 homologues from both mono- and dicotyledonous plants indicating a putative distinct function compared with the monocotyledonous-specific group (Fig. 2).

Several computer programs (i.e. ChloroP 1.1, PredSL, etc.) predict for all BT1 homologues a plastidic localization due to the N-terminal plastidic transit peptides (32, 33). For ZmBT1, HvNST1, and StBT1, a plastidic localization was experimentally verified by immunological studies, green fluorescent protein fusion, and/or targeting experiments into isolated plastids (9, 15, 34).

Sequence Analysis—Multiple alignments of amino acid sequences from known BT1 homologues available at the EMBL data base (www.ebi.ac.uk/emb/seq) were obtained using Clustal X (25). The phylogenetic tree was created with PhyML (26) after aligning the sequences with Muscle (27).

RESULTS

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mono- and dicotyledonous plants based on the rice and Arabidopsis genomes (35, 36), we assessed BT1 gene copy numbers in maize by Southern blot analyses. We chose restriction enzymes that do cut either ZmBT1 or the recently published ZmBT1-2 homologue (accession number BT016800). The digestion of the genomic maize DNA with the restriction enzymes (XbaI, XhoI, and HindIII) led to two distinct bands on the Southern blot (Fig. 3, lanes 1–3). Only digestion with BamHI showed three bands, which is most probably due to a restriction site inside an intron of one of the two BT1 homologues. These results clearly indicate that two BT1 homologues occur in maize, in contrast to one homologue in dicotyledonous plants (15) (Fig. 2) and three homologues in rice (35).

Heterologous Expression of ZmBT1 in E. coli Cells—We showed previously that the heterologous synthesis of the plastidic StBT1 homologue in E. coli leads to the functional integration of StBT1 into the bacterial cytoplasmic membrane (15). Astonishingly, we identified StBT1 as a plastidic adenine nucleotide uniporter despite the attributed function to BT1 proteins to transport ADP-Glc in countercurrent with ADP and/or AMP (14, 34, 37). However, a high degree of sequence similarity for homologues belonging to the same protein family cannot be taken as proof for the same function (see Ref. 38), specifically when there are multiple homologues per genome indicating paralogy and potential functional differentiation between the homologues. The kinetic properties as well as the substrate specificity should be identified for each putative homologue.

The heterologous expression of ZmBT1 in different E. coli strains was analyzed using a peptide-specific antibody raised against a fusion protein including the 56 amino acid residues in the C-terminal sequence of ZmBT1. Substantial differences in the expression level of carrier...
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Proteins in *E. coli* have been reported (18, 39), probably due to the unfavorable codon usage of several carrier proteins in *E. coli* (39). In fact, a screening of the ZmBT1 cDNA sequence revealed several codons rarely used in *E. coli* so that synthesized ZmBT1 protein in BL21, the most widely used expression host, was hard to detect (data not shown). In marked contrast, the use of *E. coli* Rosetta strains, which carry the pRARE plasmid and thus supply tRNAs for codons rarely used in *E. coli*, allows a high expression of ZmBT1. We confirmed the integration of the heterologously expressed ZmBT1 in the *E. coli* Rosetta cytoplasmic membrane by Western blotting by use of the above-mentioned peptide-specific ZmBT1 antibody (Fig. 4).

We tested ADP-Glc as a putative transport substrate for ZmBT1. Uptake of [14C]ADP-Glc into *E. coli* Rosetta cells harboring ZmBT1 in their membranes was linear with time for at least 15 min. The ADP-Glc uptake was strictly dependent on the membrane integrity and ZmBT1 protein synthesis (supplementary materials Fig. 1). Thus, we analyzed the affinity for ADP-Glc at different substrate concentrations. The results are given in Fig. 5A. Increased exogenous radioactively labeled ADP-Glc induced increased rates of ADP-Glc transport into *E. coli* cells harboring ZmBT1. Lineweaver-Burk analyses revealed an apparent *Km* value for ADP-Glc of 847.8 ± 39.6 μM and a *Vmax* of 191.5 ± 28.7 nmol of ADP-Glc/mg of protein h⁻¹ (Fig. 5A).

To investigate the substrate specificity of ZmBT1, we measured the effect of various non-labeled metabolites on the rate of [14C]ADP-Glc uptake (Table 1). Substantial inhibition of [14C]ADP-Glc import could be observed with non-labeled ADP and ADP-Glc reducing the transport rate below 15 and 24% of the control (without effector), respectively. None of the other metabolic intermediates tested showed any substantial influence on [14C]ADP-Glc uptake, which confirms ADP-Glc and ADP as the main substrates for ZmBT1 (Table 1). Furthermore, we analyzed the affinity of ZmBT1 for ADP (Fig. 5B). Increased exogenous radioactively labeled ADP induced increased rates of ADP transport into *E. coli* cells harboring ZmBT1. Lineweaver-Burk analyses revealed an apparent *Km* value for ADP of 465.2 ± 36.2 μM and a *Vmax* of 6.1 ± 0.7 nmol of ADP/mg of protein h⁻¹ (Fig. 5B). The higher affinity for ADP compared with ADP-Glc is reflected in the competition experiments of ADP-Glc uptake (Table 1). The substantial higher maximal velocity of ADP-Glc uptake seems to be a particular feature of this carrier (Fig. 5). Thus, the resulting relative catalytic efficiency (*Vmax*/*Km*) of ZmBT1 is 18 times higher for ADP-Glc than for ADP.

Biochemical studies on isolated maize endosperm plastids led to the assumption that ADP-Glc is transported across the envelope membranes in counterexchange with ADP or AMP (37). AMP can be excluded as exogenous substrate for ZmBT1 because we could not detect any substantial inhibition of non-labeled AMP on the ADP-Glc uptake or any direct uptake of radioactively labeled [14C]AMP into *E. coli* cells harboring ZmBT1 (Table 1, data not shown). However, we could also exploit the *E. coli* system to investigate the counterexchange properties of the heterologously synthesized ZmBT1. The principle of this approach is to initiate a putative efflux of imported radioactively labeled [14C]ADP-Glc through a high dilution

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**FIGURE 1.** Alignment of the predicted amino acid sequence of ZmBT1 with BT1 homologues. The residues identical or similar among all family members are indicated by black shading and the residues conserved by at least five proteins are shaded in gray shading. Solid black bars underline six putative membrane-spanning regions (H1–H6). Three conserved mitochondrial energy transfer signatures (METS = PX(D/E)L(IV)L(R/K)/H/L(V/M/F/Y)(Q/G/A/V/M)), following each odd membrane-spanning domain are marked by white bars. Conserved cysteine residues are indicated by triangles. Dashes represent gaps introduced to improve the similarity among the proteins. The numbers indicate the amino acid positions. ZmBT1, Brittle1 from Zea mays (NCBI accession number BT016796); TaBT1, Brittle1 from T. aestivum (NCBI accession number BT009587); OsBT1–1, Brittle1–1 from O. sativa (NCBI accession number BAD15863); HvNST1, nucleotide sugar transporter (brittle1) from *H. vulgare* (NCBI accession number AAT12275); OsBT1–1, Brittle1 from O. sativa (NCBI accession number BAD15863); ZmBT1–2, Brittle1–2 from Z. mays (NCBI accession number BT016800); OsBT1–2, Brittle1–2 from *O. sativa* (NCBI accession number AAU44334); MtBT1, Brittle1 from *M. truncatula* (NCBI accession number ABE88494); StBT1, Brittle1 from *S. tuberosum* (NCBI accession number CAA67107); AtBT1, Brittle1 from *A. thaliana* (NCBI accession number CAB79957).

**FIGURE 2.** Phylogeny of the Brittle1 proteins. A phylogeny was created with PhyML (26) after aligning the sequences with Muscle (27). The phylogeny was created with 2 γ-distributed rate parameters and the JTT model for amino acid substitutions. Bootstrap values above 85 out of 100 are indicated. A phylogeny created with Neighbor Joining using the identity matrix and correcting for multiple replacements had an identical topology. The phylogeny indicates a split between the AMP/ADP/ATP carriers and the ADP-glucose carriers, although the exact division between the groups as indicated in the tree is not very strongly supported. The extra sequences are a plant specific group (Arabidopsis and rice) of unknown function that is phylogenetically close to the Brittle1 group. The deoxynucleotide carrier group was used as outgroup because it is phylogenetically closest in a phylogeny of the 500 “best hits” to the Brittle1 group of proteins. At the right, experimentally determined substrate specificity of the carriers are indicated. Accession numbers of the BT1 proteins are given in the legend to Fig. 1 with the exception of TaBT1–2 (NCBI accession number BT009587), which is not a full-length clone.
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During an uptake experiment (15, 22), we disrupted the cells at several time points after preloading with \([14C]ADP-Glc\) and analyzed the cytosolic fraction by thin layer chromatography. Most of the \([14C]ADP-Glc\) (about 90%) was found not to be metabolized by the \(E. coli\) cells over a time span of about 12 min, which suggests the involvement of \([14C]ADP-Glc\) in a putative nucleotide exchange (Fig. 6A).

Fig. 6B shows a typical time course for \([14C]ADP-Glc\) uptake (at 1 \(\mu M\)) into \(E. coli\) cells harboring ZmBT1. Right after the start of the chase with non-labeled ADP (1 \(\mu M\)), a rapid efflux led to a total release of labeled nucleotides of about 65% (7 min after the start of the chase). GTP, used as a control, is known not to be a substrate for ZmBT1 (Table 1) and, therefore, showed no influence on the uptake of radioactively labeled ADP-Glc after the chase. These results clearly reveal that ZmBT1 mediates a counterexchange of ADP-Glc and ADP. Non-labeled ADP-Glc (1 \(\mu M\)) inhibited significantly the

**FIGURE 3.** Genomic Southern analysis of maize DNA. Nuclear DNA of \(Z. mays\) was digested with XbaI (lane 1), Xhol (lane 2), HindIII (lane 3), and BamHI (lane 4), and subjected to Southern analysis. The coding region of ZmBT1 cDNA was radioactively labeled and used as hybridization probe. The sizes of DNA molecular mass standards are indicated.

**FIGURE 4.** Western blot analysis of ZmBT1 heterologously expressed in \(E. coli\) Rosetta cells. \(E. coli\) cells harboring the plasmid encoding ZmBT1 and \(E. coli\) control cells (pET16b without any insert) were induced by application of IPTG for protein synthesis. Details of induction, purification, and Western blotting are given under “Experimental Procedures.” Immunoblotting was carried out with a ZmBT1-specific antiserum. Lane 1, membrane fraction of uninduced \(E. coli\) control cells; lane 2, membrane fraction of IPTG-induced \(E. coli\) control cells; lane 3, membrane fraction of uninduced \(E. coli\) cells harboring the plasmid encoding ZmBT1; lane 4, membrane fraction of IPTG-induced \(E. coli\) cells harboring the plasmid encoding ZmBT1.

**FIGURE 5.** Substrate saturation of ADP-Glc and ADP uptake into intact \(E. coli\) cells. IPTG-induced \(E. coli\) cells harboring the plasmid encoding ZmBT1 were incubated for 8 min with the indicated concentrations of \([14C]ADP-Glc\) (A) or \([\alpha-32P]ADP\) (B). Data are the mean of three independent experiments with four replicates each. Background rates of the control (pET16b without insert) have been subtracted. The insets represent a double reciprocal plot of the data for uptake indicating a \(K_m\) value for ADP-Glc of 847.8 \(\pm\) 39.6 \(\mu M\) with a \(V_{max}\) of 191.5 \(\pm\) 28.7 nmol of ADP-Glc/mg of protein \(h^{-1}\) (A) and a \(K_m\) value for ADP of 465.2 \(\pm\) 36.2 \(\mu M\) with a \(V_{max}\) of 6.1 \(\pm\) 0.7 nmol of ADP/mg of protein \(h^{-1}\) (B).

(chase) with non-labeled substrates at a certain time point during an uptake experiment (15, 22). To analyze whether imported \([14C]ADP-Glc\) is metabolized by \(E. coli\) cells harboring ZmBT1, we disrupted the cells at several time points after preloading with \([14C]ADP-Glc\) and analyzed the cytosolic fraction by thin layer chromatography. Most of the \([14C]ADP-Glc\) (about 90%) was found not to be metabolized by the \(E. coli\) cells over a time span of about 12 min, which suggests the involvement of \([14C]ADP-Glc\) in a putative nucleotide exchange (Fig. 6A).
[14C]ADP-Glc uptake mediated by ZmBT1 but did not initiate any exchange of ADP-Glc (Fig. 6B).

Due to the fact that commercially available ADP-Glc is contaminated by other adenylates, namely AMP, ADP, and ATP (15), we determined the contamination of ADP-Glc by high performance liquid chromatography analysis (1.8% ADP; 0.6% ATP; 0.5% AMP) and carried out the same experiment using the calculated nucleotide contaminations as a non-labeled nucleotide mixture (18 μM ADP, 6 μM ATP, and 5 μM AMP) for the chase during [14C]ADP-Glc uptake. The chase with this nucleotide mixture did not lead to any competitive inhibition of [14C]ADP-Glc uptake, so that the above mentioned competitive inhibition of non-labeled ADP-Glc is not influenced by the contamination (Fig. 6B). These results indicate a selective exchange of ADP-Glc with ADP. To further validate this mode of transport we preloaded E. coli cells harboring ZmBT1 with 500 μM non-labeled exogenous ADP for 5 and 10 min, washed the cells, and performed [14C]ADP-Glc uptake in comparison to not preloaded cells. Indeed, [14C]ADP-Glc uptake into E. coli cells harboring ZmBT1 increased up to 174% compared with the control when the cells were preloaded with the counterexchange substrate ADP (Fig. 6C). To clarify whether endogenous AMP might have an influence on [14C]ADP-Glc uptake into E. coli cells harboring ZmBT1 we preloaded these cells with 1 mM non-labeled exogenous AMP for 5 and 10 min, washed the cells, and performed [14C]ADP-Glc uptake in comparison to non-preloaded cells (supplementary materials Fig. S2). In strong contrast to the experiment with ADP-preloaded cells (Fig. 6C), additional endogenous AMP has no influence on the ZmBT1 mediated ADP-Glc uptake (supplementary materials Fig. S2).

We also determined the influence of several inhibitors on [14C]ADP-Glc uptake (Table 2). The highly specific inhibitors of the mitochondrial ADP/ATP carriers (AACs), bongkrekic acid and carboxyatractylloside (40, 41), showed

| Effector | Rate of transport |
|----------|------------------|
| None     | 100.0            |
| ADP      | 14.5             |
| ADP-Glc* | 23.9             |
| ATP      | 70.8             |
| AMP*     | 82.3             |
| UDP-Glc  | 98.2             |
| UDP-Gal  | 92.6             |
| Adenosine| 91.1             |
| Adenine  | 90.1             |
| GTP      | 101.1            |
| CTP      | 101.6            |
| UTP      | 99.7             |
| IDP      | 95.3             |
| NADP     | 98.9             |
| NADPH    | 89.9             |

* ADP-Glc value is corrected by inhibition (0.5%) of the contamination with ATP, ADP, and AMP.
* AMP value was determined at the presence of 1 mM adenosine (to prevent the cleavage of AMP by E. coli cells (15)) and corrected by the adenosine inhibition of 8.9%.

**TABLE 1**

Effects of various metabolites on [14C]ADP-glucose transport activities of ZmBT1

Metabolic effectors were given at a concentration of 1 mM. [14C]ADP-glucose was present at a concentration of 200 μM. Uptake into IPTG-induced E. coli cells harboring ZmBT1 was carried out for 8 min and stopped by rapid filtration (see “Experimental Procedures”). Data are the mean of three independent experiments with four replicates each. S.E. is less than 8% of the mean values.

**FIGURE 6.** Exchange-mediated efflux studies of intracellular radioactivity. A, E. coli cells harboring ZmBT1 were disrupted after preloading with 1 μM [14C]ADP-Glc for 3 (lane 1) and 12 min (lane 2), and the cytosolic radioactively labeled compounds were separated by thin-layer chromatography. Most of the imported ADP-Glc (90%) was found not to be metabolized. Cells used were harvested in the exponential phase that leads to a negligible glycogen synthase activity. B, E. coli cells harboring ZmBT1 were incubated in the presence of 1 μM [14C]ADP-Glc. After 5 min of [14C]ADP-Glc incubation, unlabeled nucleotides were added (chase) to a final concentration of 1 mM (1000-fold) and a possible induced efflux was monitored for the given time span. Data are the mean of three independent experiments with four replicates each. S.E. values were less than 8% of the mean values. *, the contamination of 1 mM ADP-Glc was determined by high performance liquid chromatography analysis; the corresponding concentrations of 18 μM ADP, 6 μM ATP, and 5 μM AMP were added and monitored as described above for all unlabeled nucleotides. C, E. coli cells harboring ZmBT1 were preloaded with 500 μM non-labeled exogenous ADP for 5 and 10 min. After washing the preloaded cells, [14C]ADP-Glc uptake was measured as above for 8 min in comparison to not ADP preloaded cells (control). Data are the mean ± S.E. of three independent experiments with four replicates each. Lane 1, [14C]ADP-Glc uptake into E. coli control cells harboring ZmBT1 (100% uptake = 1.56 nmol of ADP-Glc/mg of protein*); lane 2, [14C]ADP-Glc uptake into E. coli cells harboring ZmBT1 (5 min preloaded); lane 3, [14C]ADP-Glc uptake into E. coli cells harboring ZmBT1 (10 min preloaded).
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TABLE 2

| Effects of various inhibitors on [14C]ADP-glucose transport activities of ZmBT1 |
|---------------------------------------------------------------|
| [14C]ADP-glucose uptake into IPTG-induced E. coli cells harboring ZmBT1 was measured at a concentration of 200 μM. E. coli cells were preincubated for 10 min with lysozyme (1.25 mg/ml) to allow penetration of the reagents across the outer membrane. Uptake was carried out for 8 min and stopped by rapid filtration (see “Experimental Procedures”). The inhibitors were used in following concentrations: bongkrekic acid (BKA, 10 μM); carboxyatractyloside (CAT, 1 mM); pyridoxal 5′-phosphate (PLP, 2 mM), and mersalyl (100 μM). Data are the mean of three independent experiments with four replicates each. S.E. is less than 9% of the mean values. |
| Effector | Rate of transport % |
|----------|---------------------|
| None     | 100.0               |
| BKA      | 109.5               |
| CAT      | 88.1                |
| PLP      | 91.4                |
| Mersalyl | 11.6                |

Effects of various inhibitors on [14C]ADP-glucose transport activities of ZmBT1 | 

no inhibitory effect at the given concentrations (Table 2). In addition, no considerable inhibition was observed with pyridoxal 5′-phosphate, a potential inhibitor of StBT1 and the plastidic phosphate translocator (15, 42) (Table 2). Interestingly, the sulphydryl reagent mersalyl, which is known to initiate a blockage of thiol groups (43), significantly inhibited the [14C]ADP-Glc transport mediated by ZmBT1 below 12% of the control (Table 2). This indicates that SH-groups are of considerable importance for the activity of the ZmBT1.

The activity of various chloroplast enzymes is known to be regulated by reversible thioldisulfide interchange mediated by the ferredoxin-thioredoxin system (44). Recent studies on wheat starchy endosperm amyloplasts suggest also a thioredoxin (Trx) regulation of proteins in heterotrophic tissues. 42 amyloplast proteins were identified as potential Trx target proteins using a proteomic approach in combination with affinity chromatography and a fluorescent thiol probe (1). Interestingly, BT1, a major envelope protein of wheat amyloplasts, was also recognized as a thioredoxin target protein (1). To determine whether the activity of ZmBT1 is dependent on the redox state, we performed uptake of [14C]ADP-Glc into E. coli Rosetta cells harboring ZmBT1 in the presence of diamide and/or dithiothreitol (DTT). In E. coli, the thioredoxin and glutaredoxin systems are known to keep reduced conditions for cytoplasmic proteins (45). This implicates that in ZmBT1 a disulfide bond formation is prevented under normal E. coli growth conditions. The addition of the membrane-permeant thiol-specific oxidizing agent diamide (diazenedicarboxylic acid bis(N,N′-dimethylamide) (45)) during the uptake experiments causes a strong inhibition of ADP-Glc uptake of E. coli Rosetta cells harboring ZmBT1 (Fig. 7A). At a concentration of 10 mM diamide the ADP-Glc uptake mediated by ZmBT1 is decreased up to 50%. In control experiments we confirmed the specificity of this effect on ZmBT1. The mitochondrial ADP/ATP carrier AAC2 from A. thaliana is not influenced by 10 mM diamide when heterologously synthesized in E. coli (Fig. 7A). Interestingly, the inhibition of ADP-Glc uptake into E. coli Rosetta cells harboring ZmBT1 was shown to be reversible. The decrease of ADP-Glc uptake mediated by ZmBT1 after preincubation with 10 mM diamide was mainly compensated by the addition of 40 mM reduced DTT (Fig. 7B). The addition of 40 mM DTT (without diamide preincubation) does not affect any ADP-Glc uptake into E. coli Rosetta cells harboring ZmBT1, which indicates an optimal activation of ZmBT1 by reduced conditions in these E. coli expression cells (Fig. 7B).
Expression Analyses of ZmBT1 and the BT1 Maize Homologue, ZmBT1–2, in Different Maize Tissues—To obtain further insight into the putative physiological roles of ZmBT1 and the recently identified ZmBT1–2 homologue, we analyzed the accumulation of the corresponding mRNAs in various autotrophic and heterotrophic tissues. Gene expression levels of low abundant proteins can be determined by real-time RT-PCR, which is more specific and sensitive than Northern blot analyses (15, 46). The use of the total RNA mass (or 18 S rRNA) for normalization might be misleading because it consists predominantly of rRNA molecules that are often not representative for the mRNA fraction in different plant tissues (47). A reliable internal control should be similarly expressed in different cell types (48). Therefore, we chose the elongation factor EF-1α, which catalyzes the first step of protein synthesis by binding the aminoaeryl-tRNA to the aminoaeryl site of ribosomes. EF-1α is described to be a suitable housekeeping gene commonly used in plants as an internal control for real-time RT-PCR (48–50).

The real-time RT-PCR quantification shows an exclusive expression of ZmBT1 in maize endosperm tissues (15 days after pollination), which indicates an essential role for ZmBT1 in starch metabolism as reported earlier (10, 14). In strong contrast, the ZmBT1–2 homologue is expressed in heterotrophic and autotrophic tissues indicating a general role in plant metabolism that was also observed for the BT1 homologues of potato and autotrophic tissues indicating a general role in plant metabolism (53). The comparable low affinity of ZmBT1 for ADP-Glc (850 μM) seems to be sufficient for effective transport due to a highly active cytosolic ADP-Glc pyrophosphorylase (AGPase), which maintains high concentrations of ADP-Glc in the cytosol of maize endosperm (51–54).

The expression of ZmBT1 was completely inhibited any further [14C]ADP-Glc uptake. Furthermore, uptake of [14C]ADP-Glc into E. coli cells harboring ZmBT1 was stimulated when cells were preloaded with nonlabeled ADP as a putative counterexchange substrate (Fig. 6C). These results characterize ZmBT1 as a carrier that mediates a strict counterexchange of ADP-Glc with ADP, which is generated during the ADP-Glc-dependent starch synthesis inside plastids (14, 37).

Recently, efflux experiments on isolated wheat amyloplasts revealed that during the ADP-Glc-dependent starch synthesis ADP is the major adenylate exported from intact amyloplasts (53). The components of a complete ferredoxin/thioredoxin system have been recently identified in amyloplasts isolated from wheat starchy endosperm. In cereal endosperm, ferredoxin is reduced not by light, as in chloroplasts, but by the metabolically generated NADPH via ferredoxin NADP reductase (1). Interestingly, the wheat BT1 was revealed as a potential Trx target protein (1). BT1 homologues possess several conserved cysteines, which is a prerequisite for the presence of a regulatory disulfide bond (Fig. 1). The inhibition and the reversible activation of ZmBT1 by diadime and DTT provide the first experimental evidence that the ADP-Glc carrier from cereal endosperm is in fact redox regulated as postulated for the wheat BT1 (Fig. 7). However, the specific cysteine residues that are involved in the redox regulation of ZmBT1 have still to be identified. Therefore, we aim at generating ZmBT1 Cys mutants by site-directed mutagenesis for further functional analyses.

The reducing power (NADPH) for the activation of the ADP-Glc carrier from cereal endosperm is derived from the oxidative pentose phosphate pathway residing in amyloplasts (55). NADPH is generated enzymatically by glucose 6-phosphate and 6-phosphogluconate dehydrogenases after import of Glc-6-P (56, 57). In this context, the high expression of the glucose 6-phosphate/phosphate transporter in developing maize kernels makes sense. Glc-6-P imported by this carrier is used for the oxidative pentose phosphate pathway and not for the starch synthesis (57, 58).

The regulation of BT1 gained prominence in subsequent studies showing that the cytosol, rather than the plastid, is the major site of ADP-Glc formation in cereal endosperm (2). Thus, the regulation of the BT1-mediated ADP-Glc transport into cereal endosperm plastids is of high physiological importance. The transport of sucrose into sink tissues such as endosperm would finally increase the NADPH/NADP ratio, which indicates a high energy status of the plant cells leading to a higher starch synthesis via the Trx-regulated ADP-Glc transport into heterotrophic plastids (1).

The occurrence of BT1 homologues in non-cereal plants such as A. thaliana, S. tuberosum, and M. truncatula was first unexplainable because in these plants ADP-Glc was not imported into the storage plastids for starch synthesis due to the plastidic localization of the AGPase. Recently, a BT1 homologue from a dicotyledonous plant, the StBT1 from potato, was characterized as a novel plastidic adenine nucleotide uniporter used to provide the cytosol and other compartments with adenine nucleotides exclusively synthesized inside plastids (15).

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should possess at least a second BT1 isofrom. Indeed, Southern blot analyses revealed that the maize genome carries two distinct BT1 homologues (Fig. 3) and the second maize BT1 homologue (ZmBT1–2) was recently identified as well. Furthermore, the analysis of the rice genome revealed even three different BT1 homologues (35).

Interestingly, the phylogenetic analyses exhibit a subdivision of the BT1 protein family into two biochemically distinct groups (Fig. 2): the first group comprises BT1 homologues (ZmBT1–2) was recently identified as well. Furthermore, the analysis of the rice genome revealed even three different BT1 homologues (35).

The biochemical classification of the maize BT1 homologues into two subgroups (Fig. 2) is further supported by the functional analyses of OsBT1–2 and OsBT1–3, which are identified as adenine nucleotide transporters that are unable to transport ADP-Glc (supplementary materials Table S1). The recently identified ZmBT1–2 homologue clearly belongs to the group of AMP/ADP/ATP carriers. The bootstrap value for the branch that contains OsBT1–3 and ZmBT1–2 is almost maximal (97/100) giving strong support for the notion that ZmBT1–2 is orthologous to an AMP/ADP/ATP carrier. However, the exact position of these two sequences relative to the other AMP/ADP/ATP carriers and the ADP-glucose carrier group is not well resolved (65/100). Nevertheless, based on the phylogeny and the functional information about the proteins, a single separation of the BT1 proteins into a monophyletic group of AMP/ADP/ATP carriers and a monophyletic group of ADP-glucose carriers appears the most parsimonious and therefore most likely scenario. Given the phylogenetic location of these carriers close to the deoxynucleotide carrier group, it is likely that the ancestor of the BT1 group was a nucleotide carrier rather than an ADP-glucose carrier (Fig. 2).

The biochemical classification of the maize BT1 homologues into two subgroups (Fig. 2) is also supported by the expression analysis. The ZmBT1 ADP-Glc carrier is exclusively expressed in endosperm tissues during starch synthesis, whereas ZmBT1–2 shows a ubiquitous expression pattern in hetero- and autotrophic maize tissues as well, with the highest expression levels in silk and tassel (Fig. 8). The ubiquitous expression of ZmBT1–2 is in line with the proposed function of the recently identified plastidic adenine nucleotide uniporter (StBT1), which is supposed to have the general task to provide the cell with adenine nucleotides exclusively synthesized inside plastids (15). Similar observations were made for the expression patterns of the three BT1 homologues identified in the rice genome. In contrast to the endosperm-specific expression of the BT1 orthologue OsBT1–1, both BT1 paralogues (OsBT1–2 and OsBT1–3) exhibit a wide expression pattern comprising hetero- and autotrophic rice tissues, with the lowest expression level in seeds and roots (35).

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