## Molecular cloning and structural analysis of genes from Zea mays (L.) coding for members of the ras-related ypt gene family

(small guanine nucleotide binding protein/ras gene superfamily)

KLAUS PALME\*, THOMAS DIEFENTHAL\*, MARTIN VINGRON†, CHRIS SANDER†, AND JEFF SCHELL\*

\*Max-Planck-Institut für Züchtungsforschung, D-5000 Köln 30, Federal Republic of Germany; and †European Molecular Biology Laboratory, Postfach 102209, D-6900 Heidelberg, Federal Republic of Germany

Contributed by Jeff Schell, September 16, 1991

**ABSTRACT** We have isolated, cloned, and characterized two cDNAs from Zea mays (L.), denoted yptm1 and yptm2, encoding proteins related to the ypt protein family. Amino acid similarity scores with YPT1 from yeast and ypt from mouse are in the range of 70% for vptm1 and 74% for vptm2, respectively, whereas similarities with p21 ras and other ras-related proteins are <40%. Most amino acid residues showing identity are clustered in the GTP/GDP binding domain. In addition, two cysteine residues close to the C-terminal ends, known to be palmitoylated and necessary for membrane binding in all eukaryotic ras-related proteins that have been characterized so far, are conserved in the maize genes as well. Northern blot hybridization analysis of poly(A)+ mRNA from etiolated maize coleoptiles revealed single mRNA species of approximately the same size as the isolated cDNAs. The gene for yptm1 is expressed at very low levels in maize coleoptiles and tissue culture cells. The gene for yptm2 is expressed at higher levels and is differentially represented in RNAs isolated from various organs of maize plants, with its highest level in leaves and flowers. The structural similarity of the genes identified suggests that they could be involved in the control of secretory processes.

It has been established in animal and fungal cells that guanine nucleotide-binding regulatory (G) proteins exchange bound GDP for GTP thus initiating interaction with the corresponding effector protein(s). The intrinsic GTPase activity then hydrolyzes the bound GTP to GDP to return the complex to the resting state (1). All ras or ras-related genes code for small G proteins with molecular weights of 20,000-28,000 that share structural and biochemical homologies (2). In the past few years >20 small G proteins, including members encoded by the ras oncogene family, have been characterized by molecular cloning, but little is known of their specific functions (3-10). Although the products of the ras oncogenes are implicated in regulation of mammalian cell proliferation, other members of this family may have roles in vectorial membrane traffic (11). It was found that GTP analogs affect intra-Golgi vesicle transport and regulated secretion (12, 13). Moreover, mutations in the yeast genes encoding the ypt1 and SEC4 protein disrupt membrane traffic within the Golgi complex (14, 15) or from the Golgi complex to the plasma membrane (6, 16). The YPT1 gene product, besides sharing GTP-binding and activation properties with other members of the ras gene family, plays an essential role in both mitotic and meiotic stages of the Saccharomyces cerevisiae life cycle (17). Expression of YPT1 mutant alleles results in a dominant lethal phenotype with defects in microtubule organization and function (18). Recent evidence suggests that the YPT1 gene product could be involved in Ca<sup>2+</sup> metabolism (14). Analysis of cell-free protein transport has further indicated

The publication costs of this article were defrayed in part by page charge payment. This article must therefore be hereby marked "advertisement" in accordance with 18 U.S.C. §1734 solely to indicate this fact.

that the ypt1 protein is an essential component for the vesicular transport of proteins from the endoplasmic reticulum to the Golgi complex (19).

Due to their fundamental role in controlling basic cellular functions, it is expected that members of this ubiquitous gene family may also play an important role in plant growth control. In the present study we have searched for ras-related genes in plants. We describe the structural and molecular analysis of cDNA clones<sup>‡</sup> from Zea mays encoding ras-related proteins that are highly homologous to the ypt proteins.

## MATERIALS AND METHODS

Construction and Oligonucleotide Screening of a Maize cDNA Library. A maize coleoptile-specific cDNA library was prepared and screened with oligonucleotides as described (20, 21). The oligonucleotide sequences were 5'-ATTTTA-GATACTGCTGGTCAAGAGGAGTAT-3' (oligonucleotide 1) and 5'-GTTCCTATGGTTGTTGGTAACAAATGT-GATCTT-3' (oligonucleotide 2). Oligonucleotides were synthesized on an Applied Biosystems synthesizer, deprotected, and purified by gel electrophoresis. End-labeling was done with T4 DNA polynucleotide kinase and  $[\gamma^{-32}P]ATP$ . Each probe  $(1 \times 10^6 \text{ cpm/ml}, 1 \text{ pmol/ml})$  was added to two sets of double replicate filters and hybridized for 24 h at 42°C, respectively. Melting temperature  $[4 \times (G + C) + 2 \times (A + C)]$ T)] was calculated (22) and was 46°C and 60°C for oligonucleotides 1 and 2, respectively. Each filter was washed in a solution containing 900 mM NaCl, 90 mM sodium citrate, and 0.5% Nonidet P-40 for three 5-min periods at room temperature, for two 10-min periods at 37°C, and for one 15-min period at each of the following temperatures: 42°C, 45°C, 48°C, and 51°C. The filters were subjected to autoradiography for 24 h. Phage from plaques yielding positive autoradiographic signals with both probes were cloned.

DNA Sequence Analysis. DNA sequence analysis was carried out by the dideoxynucleotide chain-termination method. The sequence alignment was calculated using software described by Vingron and Argos (23) and improved manually. From the distance data, a phylogenetic tree was calculated using the program KITSCH from the PHYLIP package by Felsenstein (24). The tree shown is based on the one calculated by the program but redrawn by hand to scale to the correct homologies between the sequences.

RNA Blot Hybridization. RNA was fractionated in formaldehyde-containing agarose gels, transferred to nylon (Hybond N, Amersham), hybridized for 48 h at 42°C to a nick-translated denatured probe ( $10^8 \text{ cpm}/\mu\text{g}$ ), and analyzed by standard procedures (21).

Abbreviations: G protein, guanine nucleotide-binding regulatory protein; nt, nucleotide(s).

<sup>‡</sup>The sequences reported in this paper have been deposited in the GenBank data base (accession nos. X63277 for yptm1 and X63278 for yptm2).

## **RESULTS**

Identification of YPT1-Related cDNA Clones. Direct sequence comparison of members of the ras gene superfamily by multiple alignments of amino acids of the GTP-binding domain is presented in Fig. 1. Comparison of amino acid sequences representing the GTP-binding domain point to three consensus elements, GxxxxGKSsxl, DTAGQE, and

lxgNKxDL, where lowercase letters are variable amino acids. The first two domains are involved in binding the phosphate moiety of the GTP, whereas elements located further downstream are involved in determining guanine nucleotide specificity (25, 26). To identify related members of the *ras* gene superfamily in plants, oligonucleotides corresponding to amino acids 62–70 and 114–125 were used to screen 200,000 phages. We isolated several clones giving

```
-----KLVVVGAGGVGKSALTIQLIQNHFVDEYDPTIEDSYR-KQVVIDGETCLLD
                                                                       Ha-ras
      -----KIVVVGGGGVGKSALTIQFIQSYFVDEYDPTIEDSYR-KQVVIDDKVSILD
                                                                       RAS1
      -----KLVVVGGGGVGKSALTIQLTQSHFVDEYDPTIEDSYR-KQVVIDDEVSILD
                                                                       RAS2
      -----KLVIVGDGACGKTCLLIVFSKDQFPEVYVPTVFENY-VADIEVDGKQVELA
                                                                       rho
     -----KLLLIGDSGVGKSCLLLRFADDTYTESYISTIGVDFKIRTIELDGKTIKLQ
-----KYIIIGDTGVGKSCLLLQFTDKRFQPVHDLTMGVEFGARMITIDGKQIKLQ
                                                                       rabl
                                                                       rab2
      -----KILIIGNSSVGKTSFLFRYADDSFTPAFVSTVGIDFKVKTIYRNDKRIKLQ
                                                                        rab3a
      -----KLLIIGNSSVGKTSFLLRYADDTFTPAFVSTVGIDFKVKTVYRHEKRVKLO
                                                                       rab3b
      -----KFLVIGNAGTGKSCLLHQFIEKKFKDDSNHTIGVEFGQKIINVGGKYVKLQ
                                                                       rab4
      -----KLVLLGESAVGKSSLVLRFVKGQFHEFQESTIGAAFLTQTVCLDDTTVKFE
                                                                       rab5
      -----KLVFLGEQSVGKTSLITRFMYDSFDNTYQATIGIDFLSKTMYLEDRTVRLQ
                                                                       rab6
      -----KILLIGDSGVGKSCLLVRFVEDKFNPSFITTIGIDFKIKTVDINGKKVKLQ
                                                                       SEC4
      -----KLLLIGNSGVGKSCLLLRFSDDTYTNDYISTIGVDFKIKTVELDGKTVKLQ
                                                                        YPT1
      MSNEFDYLFKLLLIGDSSVGKSCFLLRFADDSYVDSYISTIGVDFKIRTVEVEGKTVKLQ
                                                                       YPTm1
      MNPEYDYLFKLLLIGDSGVGKSCLLLRFADDSYLDSYISTIGVDFKIRTVEQDGKTIKLQ
      ILDTAGQEEYSAMRDQYMRTGEGFLCVFAINNTKSFEDIHQ-YREQIKRVKDSDDVPMVL
                                                                       Ha-ras
51
51
                                                                       RAS1
      ILDTAGQEEYSAMREQYMRTGEGFLLVYSVTSRNSFDELLS-YYQQIQRVKDSDYIPVVV
      ILDTAGQEEYSAMREQYMRNGEGFLLVYSITSKSSLDELMT-YYQQILRVKDTDYVPIVV
                                                                        RAS2
51
52
      LWDTAGOEDYDRLRPLSYPDTDVILMCFSIDSPDSLENIPEKWTPEVRHF--CPNVPIIL
      IWDTAGQERFRTITSSYYRGAHGIIVVYDVTDQESFNNVKQ-WLQEIDRYA-SENVNKLL
52
52
52
52
52
      IWDTAGQESFRSITRSYYRGAAGALLVYDITRRDTFNHLTT-WLEDARQHS-NSNMVIML
                                                                        rab2
      IWDTAGQERYRTITTAYYRGAMGFILMYDITNEESFNAVQD-WSTQIKTYS-WDNAQVLL
                                                                        rab3a
      IWDTAGQERYRTITTAYYRGAMGFILMYDITNEESFNAVQD-WATQIKTYS-WDNAQVIL
                                                                        rab3b
      IWDTAGQERFRSVTTSYYRGAAGALLVYDITSRETYNALTN-WLTDARMLA-SQNIVIIL
                                                                        rab4
      IWDTAGQEGYHSLAPMYYRGAQAAIVVYDITNEESFARAKN-WVKELQRQA-SPNIVIAL
                                                                        rab5
52
      LWDTAGQERFRSLIPSYIRDSTVAVVVYDITNVNSFQQTTK-WIDDVRTER-GSDVIIML
                                                                        rab6
      LWDTAGQERFRTITTAYYRGAMGIILVYDVTDERTFTNIKQ-WFKTVNEHA-NDEAQLLL
52
                                                                        SEC4
52
      IWDTAGQERFRTITSSYYRGSHGIIIVYDVTDQESFNGVKM-WLQEIDRYA-TSTVLKLL
                                                                        YPT1
      {\tt IWDTAGQERFRTITSSYYRGAHGIIIVYDITDMESFNNVKQ-WLDEIDRYAN-DSVRKLL}
                                                                        YPTm1
61
      {\tt IWDTAGQERFRTITSSYYRGAHGIIIVYDVTDQESFNNVKQ-WLNEIDRYAS-DNVNKLL}
                                                                        YPTm2
      VGNKCDLAART-----VESRQAQDLARSYGI-PYIETSAKTRQGVEDAFYTL
110
                                                                        Ha-ras
      VGNKLDLENERQ-----VSYEDGLRLAKQLNA-PFLETSAKQAINVDEAFYSL
VGNKSDLENEKQ-----VSYQDGLNMAKQMNA-PFLETSAKQAINVEEAFYTL
                                                                        RAS1
110
                                                                        RAS2
110
      VGNKKDLRNDESTKRELMKMKQEPVRPEDGRAMAEKINAYSYLECSAKTKEGVRDVFETA
109
                                                                        rho
      VGNKCDLTTKKV------VDYTTAKEPADSLGI-PFLETSAKNEKNVEQSFMTM
IGNKSDLESRRE-----VKKEEGEAPAREHGL-IFMETSAKTASNVERAFINT
110
                                                                        rabl
110
                                                                        rab2
      VGNKCDMEDERV------VSSERGRQLADHLGF-EFFEASAKDNINVKQTFERL
                                                                        rab3a
110
      VGNKCDMEEERV-------VPTEKGQLLAEQLGF-DFFEASAKENISVRQAFERL
                                                                        rab3b
110
      CGNKKDLDADRE------VTFLEASRFAQENEL-MFLETSALTGENVEEAFMQC
                                                                        rab4
110
      SGNKADLANKRA------VDFQEAQSYADDNSL-LFMETSAKTSMNVNEIFMAI
                                                                        rab5
110
      VGNKTDLADKRQ------VSIEEGERKAKELNV-MFIETSAKAGYNVKQLFRRV
110
                                                                        rab6
      VGNKSD-METRV------VTADQGEALAKELGI-PFIESSAKNDDNVNEIFFTL
110
                                                                        SEC4
      VGNKCDLKDKRV------VEYDVAKEFADANKM-PFLETSALDSTNVEDAFLTM
110
      VGNKCDLAENRA------VDTSVAQAYAQEVGI-PFLETSAKESINVEEAFLAM
VGNKSDLTANKV-----VATETAKAFADEMGI-PFMETSAKNATNVQQAFMAM
                                                                        YPTm1
                                                                        YPTm2
      VREIROHKLRKLNPPDESG------PGCMSCKCVLS-----
156
      IRFYSLIRLVRDDGGKYNSMNRQLDNTNE..(107aa)..RKESSGGCCIIC-----
                                                                        RAS1
157
      ARFYTLARLVRDEGGKYNKTLTENDNSKQ..(120aa)..SKSGSGGCCIIS-----
      rab3a
                                                                        rab3b
157
      ARKILNKIESGELDPERMGSGIQYGDAALRQLRSPRRTQAPSAQE--CGC-----AKKLPKNEPQNPGANSARGGGVDLTEPTQPTRNQ------CCSN-----AAALPGMESTQDRSREDMIDIKLEKPQEQPVSE------GGCSC------
157
                                                                        rab4
157
                                                                        rab5
157
                                                                        rab6
      AKLIOEKIDSNKLVGVGNGKEGNISINSGSGNSSK-----SNCC-----
156
                                                                        SEC4
      ARO I KOSMSOONLNETTOKKEDKGNVNLKGOSLTNT-----GGGCC-----
157
                                                                        YPT1
      SAAIKKSKAGSQAALERKPSNVVQMKGRPIQQEQQK-----SSRCCST-----
                                                                        YPTm1
166
       AASIKDRMASQPAAANARPATVQIR-GQPVNQKTS-----CCSS-----
166
```

Fig. 1. Alignment of the amino acid sequences of the conserved GTP-binding domain of ras protein family. All amino acid sequences are from the PIR or the EMBL sequence data bases. Dashes indicate gaps introduced for optimal alignment.

strong hybridization signals. Two of the clones isolated were chosen for further analysis. Both clones, yptml and yptm2, were subcloned into pUC118 and the DNA sequences of both strands were determined. The DNA sequences for both clones are shown in Figs. 2 and 3. Nucleotide sequence analysis of yptm1 gave a 624-nucleotide (nt) open reading frame, with 192 nt in the 5' flanking and 144 nt in the 3' untranslated region followed by 33 adenosine residues (Fig. 2). Similarly, nucleotide sequence analysis of yptm2 demonstrated a 609-nt open reading frame, with 116 nt in the 5' flanking and 257 nt in the 3' untranslated region followed by 82 adenosine residues (Fig. 3). The 5' untranslated regions are G + C-rich (55% and 63%, respectively). The nucleotide sequences surrounding the initiator codons, 5'-cGcaATGa and 5'-cGacATGa, respectively, are consistent with the proposed eukaryotic translation initiation consensus sequence of (A/G)NNATGg, where lowercase letters are variable amino acids (27, 28).

Translation terminates at a TGA (position 817, Fig. 2) or TAA (position 727, Fig. 3) stop codon. No canonical poly-

adenylylation signals typical for other eukaryotic genes are present in either cDNA (29, 30). The motifs AAAAT, located 39 and 133 nt upstream of the poly(A) tail, are possible polyadenylylation signals. Several sequence motifs have been shown to play a role in the 3' processing of mRNA (31, 32). The consensus sequence YGTGTTYY (where Y is a pyrimidine) is located ≈30 base pairs 3' of the polyadenylylation signal in most mammalian genes analyzed (33). Good homology to this consensus sequence is found for yptm1 87 base pairs (TGTGTTTC, position 876) and for yptm2 23 base pairs (TGTGAACA, position 961) downstream from the polyadenylylation signal.

Structural Analysis of the yptm-Encoded Proteins. The amino acid sequences deduced from the DNA sequences for yptm1 and yptm2 are shown in Figs. 2 and 3. Assignment of the initiator codons is based on DNA sequence analysis. The predicted proteins have 208 and 203 amino acids and calculated molecular weights of 23,283 and 22,475, respectively. We compared the derived amino acid sequences with se-

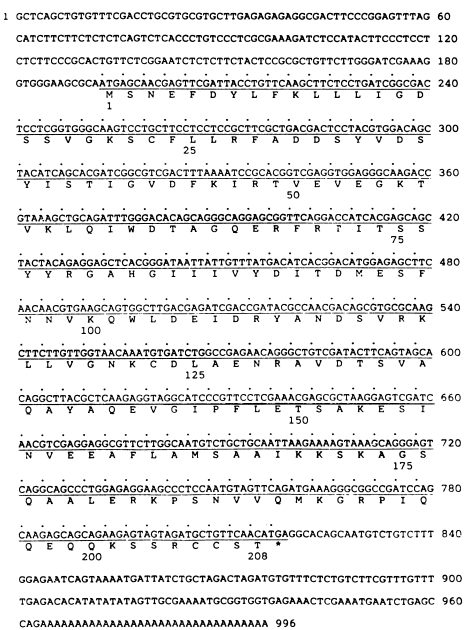


Fig. 2. Nucleotide and deduced amino acid sequence of a cDNA encoding yptm1 protein. Numbering of the nucleotides progresses positively in the  $5' \rightarrow 3'$  direction beginning with nt 1 of the initiator codon.

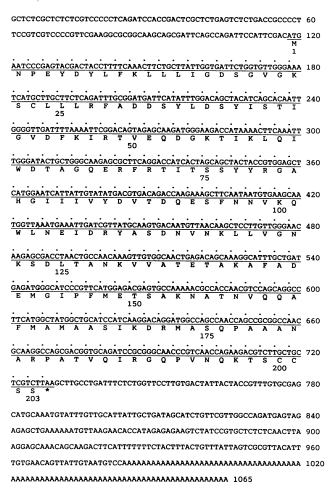


Fig. 3. Nucleotide and deduced amino acid sequence of a cDNA encoding yptm2 protein. Numbering of the nucleotides progresses positively in the  $5' \rightarrow 3'$  direction beginning with nt 1 of the initiator codon.

quences of other ras-related protein sequences deposited in data banks. A comparative analysis showed an unexpectedly strong similarity of the maize ypt proteins to the ras gene superfamily. The yptm2 protein shows a higher level of similarity to the mouse ypt protein or rab1 protein than to the yeast protein. Further inspection of the homologous region at the protein sequence level confirmed that the homology is confined mainly to the GTP-binding domain. At the C terminus, the predicted ypt proteins contain two cysteine residues followed by hydroxylated amino acids. A similar sequence, containing two cysteine residues, one of which is needed for posttranslational lipid binding and subsequent membrane anchoring, is found at the C terminus of all members of the ras gene superfamily (34, 35). To obtain an evolutionary relationship of ras-related proteins, a multiple sequence alignment was prepared based on distance measurements (23). The tree in Fig. 4 shows that rab1 from rat, ypt from yeast, and ypt from maize form a group of proteins within the ras family distinct from other rab or ras proteins. As can be seen from the alignment in Fig. 1, the regions that are likely to be involved in GDP/GTP binding are highly conserved whereas in other positions conservation only within particular subgroupings can be seen. Thus alignment within this tree is according to function and not to species grouping.

Northern Blot Analysis. By using yptm1 cDNA as a probe, a weak 1-kilobase signal was detected in poly(A)<sup>+</sup> mRNA from maize coleoptiles (data not shown). A predominant transcript of  $\approx 1$  kilobase was detected in Northern blot analysis of poly(A)<sup>+</sup> mRNA from coleoptiles using yptm2

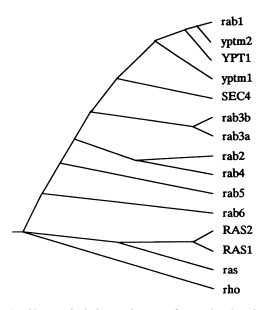


Fig. 4. Unrooted phylogenetic tree of ras-related amino acid sequences.

cDNA as a probe (Fig. 5). To analyze organ-specific expression of the gene for yptm2, total RNAs from various maize organs were separated on an agarose gel, blotted onto nitrocellulose, and hybridized to a <sup>32</sup>P-labeled yptm2 probe. Levels of yptm2 transcripts were quantified by laser scanning densitometry and normalized to the relative level of hybridization to 18S rRNA. The level of expression is low in stems. Intermediate levels of expression are observed in roots, leaves, callus suspension culture cells, and coleoptiles, from 3.7- to 6.2-fold in comparison to stems. We find highest expression of the gene for yptm2 in flower tissues (10-fold in comparison to stems). We also analyzed the expression of ypt-related genes in other plant species. However, as shown in Fig. 5 (lanes f, g, and h), we were not able to detect hybridizing signals with yptm2 cDNA as a probe in total RNA from various other plants.

## **DISCUSSION**

Here we have used an oligonucleotide strategy to search for members of the *ras* superfamily in plants. We were able to

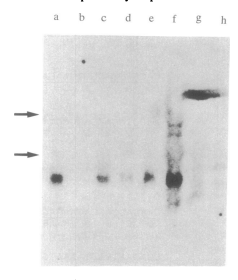


Fig. 5. Organ-specific expression of yptm2 mRNA. Total RNA was fractionated in a formaldehyde gel, transferred to nitrocellulose, and hybridized with a <sup>32</sup>P-labeled yptm2 cDNA. Lanes: a, tassel; b, stem; c, root; d, coleoptile; e, callus suspension culture; f, Arabidopsis thaliana; g, Nicotiana tabacum; h, Datura innoxia.

isolate several cDNAs from a maize coleoptile-specific cDNA library coding for members of the ras-related ypt protein family. Similarity to the Ha-ras protein is 35% and restricted to amino acids in the GTP-binding domain. The homologies are localized in four main blocks corresponding to amino acid residues 18-21, 62-68, 121-124, and 151-153 (using the numbering of the yptm proteins as a reference), which have all been implicated in the binding of GDP/GTP (26). The spacing of regions of highest identity and hydrophobicity demonstrate strong functional similarity for GTP binding and hydrolysis within this family. Amino acids 37–48, which are probably located within the flexible effector loop, are thought to mediate interactions with corresponding effector proteins (26). It is therefore not surprising that precisely these amino acids are very different in yptm proteins from those of ras proteins. The sequence from amino acid 166 to the C terminus diverges considerably within the whole ras family. The only amino acids strictly conserved within this domain are Cys-206/201 and Cys-200/201. In all ras and ras-related proteins, these cysteines are needed for palmitic acid binding and subsequent membrane anchoring. The conservation of these amino acids in the cDNAs isolated here argues that the corresponding plant proteins are localized in the membrane. In contrast to the ras proteins, no basic amino acids are found immediately on the N-terminal side of the cysteine. Moreover, downstream we find, in contrast to other members of the ras family, mostly hydrophilic amino acids. These differences may indicate variations in the specificity of plant acylases catalyzing the addition of the membrane

The identification of a series of ypt-related genes from plants raises the question of their function. In eukaryotes ypt proteins seem to serve apparently basic cellular functions and it is now becoming clear that, at specific steps in the secretory pathway, G proteins regulate the traffic of proteins in membrane vesicles. Genes encoding GTP-binding proteins involved in regulation of transport between early compartments of the secretory pathway have been cloned from yeast and from mammalian cells (36, 37). Members of this rapidly growing gene family of small GTP-binding proteins may be key elements in the regulation of vesicular transport and delivery of proteins to the cell surface.

The identification of ras-related proteins in maize, which apparently have homologues in other plants as well, is of particular interest. Most plant tissues respond to growth stimuli, such as auxins, by cell elongation. Active expansion of membranes requires regulated secretion and correct targeting of secretory vesicles. Thus, in plant cells, GTP hydrolysis could trigger the vectorial flow of membrane material through a metabolic cascade to the surface. This may explain why we find members of the ypt gene family expressed in elongating maize coleoptile cells.

We thank Sybil Schwonke for her skilled technical assistance and Dr. I. Moore for critical reading of the manuscript.

- Bourne, H. R., Sanders, D. A. & McCormick, F. (1991) Nature (London) 349, 117-127.
- Hall, A. (1990) Science 249, 625-640.

- Gallwitz, D., Donath, C. & Sander, C. (1983) Nature (London) 306, 704-707.
- Madaule, P., Axel, R. & Myers, A. M. (1987) Proc. Natl. Acad. Sci. USA 84, 779-783.
- Chardin, P. & Tavitian, A. (1986) EMBO J. 5, 2203-2208.
- Salminen, A. & Novick, P. J. (1987) Cell 49, 527-538.
- Touchot, N., Chardin, P. & Tavitian, A. (1987) Proc. Natl. Acad. Sci. USA 84, 8210-8214.
- Lowe, D. G., Capon, D. J., Delwart, E., Sakaguchi, A. Y., Naylor, S. L. & Goeddel, D. V. (1987) Cell 48, 137-146.
- Sewell, J. L. & Kahn, R. A. (1988) Proc. Natl. Acad. Sci. USA 85, 4620-4624.
- Haubruck, H., Disela, C., Wagner, P. & Gallwitz, D. (1987) EMBO J. 6, 4049-4053.
- Balch, W. E. (1990) Trends Biochem. Sci. 15, 473-477.
- Bourne, H. R. (1988) Cell 53, 669-671.
- Melancon, P., Glick, B. S., Malhotra, V., Weidman, P. J., Serafini, T., Gleason, M. L., Orci, L. & Rothman, J. E. (1987) Cell 51, 1053-1062.
- Schmitt, H. D., Puzicha, M. & Gallwitz, D. (1988) Cell 53, 635-637
- Segev, N., Mulholland, J. & Botstein, D. (1988) Cell 52, 15. 915-924.
- Walworth, N. C., Goud, B., Kabcenell, A. K. & Novick, P. J. (1989) EMBO J. 8, 1685-1693.
- Segev, N. & Botstein, D. (1987) Mol. Cell. Biol. 7, 2367-2377.
- Schmitt, H. D., Wagner, P., Pfaff, E. & Gallwitz, D. (1986) Cell 47, 401-412
- Baker, D., Wuestehube, L., Scheckman, R., Botstein, D. & Segev, N. (1990) Proc. Natl. Acad. Sci. USA 87, 355-359.
- Glover, D., ed. (1985) DNA Cloning (IRL, Oxford). Maniatis, T., Fritsch, E. F. & Sambrook, J. (1982) Molecular Cloning: A Laboratory Manual (Cold Spring Harbor Lab., Cold Spring Harbor, NY).
- Suggs, H. V., Hirose, T., Miyake, K., Kawashima, E. H., Johnson, M. J. & Wallace, R. B. (1981) ICN-UCLA Symp. Mol. Cell. Biol. 23, 682-693.
- Vingron, M. & Argos, P. (1989) Comput. Appl. Biosci. 5, 115-121.
- Felsenstein, J. (1985) Evolution 39, 783-791.
- De Vos, A. M., Tong, L., Milburn, M. V., Matias, P. D., Jancarik, J., Noguchi, S., Nishimura, S., Miura, K., Ohtsuka, E. & Kim, K. H. (1988) Science 239, 888-893.
- Pai, E. F., Kabsch, W., Krengel, U., Holmes, K. C., John, J. & Wittinghofer, A. (1989) Nature (London) 341, 209-214.
- Kozak, M. (1989) J. Cell Biol. 108, 229-241.
- Lütcke, H. A., Chew, K. C., Mickel, F. S., Moss, K. A., Kern, H. F. & Scheele, G. A. (1987) EMBO J. 6, 43-48.
- Dean, C., Tamaki, S., Dunsmuir, P., Favreau, M., Katayama, C., Dooner, H. & Bedbrook, J. (1986) Nucleic Acids Res. 14, 2229-2240.
- Joshi, C. P. (1987) Nucleic Acids Res. 15, 9627-9640.
- Proudfoot, N. J. & Whitelaw, E. (1987) in Frontiers in Molecular Biology: Transcription and Splicing, eds. Glover, D. M. & Hames, B. D. (IRL, Oxford), pp. 97-129.
- Ingelbrecht, I. L. W., Herman, L. M. F., Dekeyser, R. A., Van Montagu, M. C. & Depicker, A. G. (1989) Plant Cell 1, 671-680.
- McLauchlan, J., Gaffney, D., Whitton, J. L. & Clements, J. B. (1985) Nucleic Acids Res. 13, 1347-1368.
- Willumsen, B. M., Norris, K., Papageorge, A. G., Hubbert, N. L. & Lowy, D. R. (1984) EMBO J. 3, 2581-2585.
- Molenaar, C. M. T., Prange, R. & Gallwitz, D. (1988) EMBO J. 7, 971–976.
- Santos, E. & Nebreda, A. R. (1989) FASEB J. 3, 2151-2162.
- Zahraoui, A., Touchot, N., Chardin, P. & Tavitian, A. (1989) J. Biol. Chem. 264, 12394-12401.