Commentary. In the article “Lignification of plant cell walls: Impact of genetic manipulation” by Hans-Joachim G. Jung and Weiting Ni, which appeared in number 22, October 27, 1998, of Proc. Natl. Acad. Sci. USA (95, 12742–12743), the authors request that the following corrections be noted. It was accidentally stated that the studies by Kajita et al. (1) and Lee et al. (2) dealt with cinnamoyl-CoA reductase modified plants when in fact they concerned 4-coumarate:coenzyme A ligase (4CL) transgenic plants. Lignin concentration was reduced by down-regulation of 4CL activity in both studies (1, 2). In a subsequent article, Kajita et al. (3) reported a negligible decrease in lignin concentration and a decreased syringyl-to-guaiacyl lignin ratio for anti-sense suppressed 4CL transgenic tobacco line. Kajita et al. (1) rather than Kajita et al. (3) was inadvertently cited when this later report was contrasted with the large decreases in lignin concentration and an increased syringyl-to-guaiacyl lignin ratio for anti-sense suppressed 4CL Arabidopsis transgenics (2). The authors apologize for the confusion these errors have created for readers of their Commentary and to the authors of the cited work for misrepresenting their research.


Biochemistry. In the article “Requirement of GM2 ganglioside activator for phospholipase D activation” by Shun-ichi Nakamura, Toshihiro Akisue, Hitoshi Jinai, Tomohiro Hitomi, Sukumar Sakar, Noriko Miwa, Taro Okada, Kimihisa Yoshida, Shun’ichi Kuroda, Ushio Kikkawa, and Yasutomi Nishizuka, which appeared in number 21, October 13, 1998, of Proc. Natl. Acad. Sci. USA (95, 12742–12743), the authors request that the following corrections be noted. It was accidentally stated that the position of Figs. 3 and 4 were transposed. The correct figures and their legends are reproduced below.

Cell Biology. In the article “Impairing follicle-stimulating hormone (FSH) signaling in vivo: Targeted disruption of the FSH receptor leads to aberrant gametogenesis and hormonal imbalance” by Andree Dierich, M. Ram Sairam, Lucia Monaco, Gian Maria Fimia, Anne Ganssmuller, Marianne LeMeur and Paolo Sassone-Corsi, which appeared in number 23, November 10, 1998, of Proc. Natl. Acad. Sci. USA (95, 13612–13617), the authors request that the following correction be noted: In Fig. 2 appearing on page 13614, the genotype identification for testicular histology in panels C and D were shown reversed. The correct identification is –/– for panel C and +/- for panel D. The fifth sentence of the figure legend should read as follows: “Histological sections at lower (E) and higher (D) magnification of the seminiferous tubuli from a wild-type and mutant (F and C) mouse.”

Biochemistry. In the article “Efficient construction of a large nonimmune phage antibody library: The production of high-affinity human single-chain antibodies to protein antigens” by Michael D. Sheets, Peter Amersdorfer, Ricardo Finnem, Peter Sargent, Ericka Lindqvist, Robert Schier, Grete Hemingsen, Cindy Wong, John C. Gerhart, and James D. Marks, which appeared in number 11, May 26, 1998, of Proc. Natl. Acad. Sci. USA (95, 6157–6162), the following correction should be noted. The fifth author’s name was spelled incorrectly. The correct spelling is Ericka Lindquist. In addition, her department affiliation is also incorrect. Ericka Lindquist’s affiliation should be “Program in Infectious Diseases, School of Public Health, University of California, Berkeley, CA 94720.”
Requirement of \(G_M2\) ganglioside activator for phospholipase D activation

(ADP ribosylation factor phosphatidylcholine)

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ABSTRACT Sequence analysis of a heat-stable protein necessary for the activation of ADP ribosylation factor-dependent phospholipase D (PLD) reveals that this protein has a structure highly homologous to the previously known \(G_M2\) ganglioside activator whose deficiency results in the AB-variant of \(G_M2\) gangliosidosis. The heat-stable activator protein indeed has the capacity to enhance enzymatic conversion of \(G_M2\) to \(G_M3\) ganglioside that is catalyzed by \(\beta\)-hexosaminidase A. Inversely, \(G_M2\) ganglioside activator purified separately from tissues as described earlier [Conzelmann, E. & Sandhoff, K. (1987) Methods Enzymol. 138, 792–815] stimulates ADP ribosylation factor-dependent PLD in a dose-dependent manner. At higher concentrations of ammonium sulfate, the PLD activator protein apparently substitutes for protein kinase C and phosphatidylinositol 4,5-bisphosphate, both of which are known as effective stimulators of the PLD reaction. The mechanism of action of the heat-stable PLD activator protein remains unknown.

Phospholipase D (PLD) in mammalian tissues is activated by a wide variety of external signals (for review, see ref. 1). The enzyme catalyzes the hydrolysis of phosphatidylcholine (PtdCho) to generate lipid mediators for the control of a broad range of physiological and pathological processes including intracellular protein trafficking, vesicular transport, membrane ruffling, cell motility, mitogenesis, oncogenesis, and inflammation (for review, see ref. 2). Biochemical studies have shown that there are at least two types of PLD in mammalian tissues. One is activated by small G protein such as ADP ribosylation factor (ARF) (3, 4), and the other is insensitive to ARF (5). One form of the latter type purified from pig lung is activated by small G protein and does not contain ARF but is activated by PtdIns-4,5-P2 (6). A shorter splice variant PLD1b, which lacks 38 amino acids, shows similar enzymatic properties (9). PLD1 also is activated by protein kinase C (PKC) (10). This activation is enhanced by phorbol ester in the presence of ATP. On the other hand, PLD2 with a molecular mass of 106 kDa shows ≈50% homology to PLD1 and does not require ARF but is activated by PtdIns-4,5-P2 but not by PKC (8).

Several protein factors are shown to be involved in the activation of ARF-dependent PLD. An enzyme obtained from hematopoietic cell lines, which resembles PLD1, is activated by coexistence of a 50-kDa soluble protein and ARF (11, 12).

In earlier reports from this laboratory, cytosol from the rat kidney was essential for the activation of ARF-dependent PLD purified partially from the same tissue (13). The kidney PLD is an ARF-dependent enzyme resembling PLD1. The active components in the cytosol consist of at least three protein factors; ARF, RhoA, and a heat-stable protein (13). The heat-stable protein shows a molecular size of 23 kDa upon SDS/PAGE (14). Sequence analysis now reveals that this protein is highly homologous to previously known \(G_M2\) ganglioside activator. Deficiency of this activator results in the AB-variant of \(G_M2\) gangliosidosis (for review, see ref. 15).

MATERIALS AND METHODS

Materials. 1,2-Di[\(1^{14}\)C]palmitoyl-sn-glycero-3-phosphocholine (\([^{14}\)C]PtdCho, 115 mCi/mmol) and 1-[\(1^{14}\)C]palmitoyl-2-lyso-sn-glycero-3-phosphocholine (\([^{14}\)C]lysoPtdCho, 57.0 mCi/mmol) were purchased from DuPont/NEN. Phosphatidylethanolamine, a standard for TLC, was from Avanti Polar-Lipids. Plasmalogen-rich phosphatidylethanolamine (PtdEtn) (60% plasmalogen) was purchased from Serdary Research Laboratories (Englewood Cliffs, NJ). Guanosine 5′-O-(3-thiotriphosphate) (GTP-γ-S) was from Boehringer Mannheim. Glass-backed silica gel 60 was purchased from Merck. Phorbol 12-myristate 13-acetate (PMA) was a product of LC services (Woburn, MA). Other chemicals were of analytical grade.

Preparation of PLD. PLD was purified 200-fold as described (13) except that the rat kidney was used instead of the bovine kidney as enzyme source and that hydroxyapatite column chromatography was omitted from the purification procedure.

Preparation of ARF. ARF was purified from the bovine brain as described (16). The final preparation of ARF after HiLoad 16/60 Superdex 75 (Pharmacia; 1.6 × 60 cm) was ≈87% pure by silver stain. This preparation was free of heat-stable PLD activator.

Preparation of \(\beta\)-Hexosaminidase A. \(\beta\)-Hexosaminidase A was prepared as described (17) except that the rat liver (100 g) was used instead of the human liver as enzyme source.

Abbreviations: PLD, Phospholipase D; PtdCho, phosphatidylcholine; ARF, ADP ribosylation factor; PtdIns-4,5-P2, phosphatidylinositol 4,5-bisphosphate; PKC, protein kinase C; \([^{14}\)C]PtdCho, 1,2-di[\(1^{14}\)C]palmitoyl-sn-glycero-3-phosphocholine; \([^{14}\)C]lysoPtdCho, 1-[\(1^{14}\)C]palmitoyl-2-lyso-sn-glycero-3-phosphocholine; PtdEtn, phosphatidylethanolamine; GTP-γ-S, guanosine 5′-O-(3-thiotriphosphate); PMA, phorbol 12-myristate 13-acetate.

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β-Hexosaminidase A was separated from β-hexosaminidase B by DEAE-Sephadex A-50 column (1.6 × 5 cm; 10 ml) (17).

**Preparation of G_m2 Ganglioside Activator.** G_m2 ganglioside activator was purified from the rat kidney by the method originally described for the human kidney (18). Purification procedure included heat treatment of the kidney cytosol, adjustment to pH 3.0 with trichloroacetic acid, and column chromatography on DEAE-cellulose, Octyl-Sepharose, and Superdex 200 columns (Pharmacia).

**Gel Filtration of PLD Activator.** The purified heat-stable PLD activator (100 μg) was applied to a HiLoad 16/60 Superdex 200 column (Pharmacia; 1.6 × 60 cm), which had been equilibrated with a buffer (50 mM Hepes-NaOH at pH 7.4, 1 mM MgCl₂, 1 mM EGTA, and 1 μg/ml leupeptin), and eluted from the column with the same buffer at a flow rate of 1 ml/min by using an FPLC system (Pharmacia). Fractions (5 ml each) were collected and assayed for the ability to activate PLD and also to enhance the enzymatic conversion of G_m2 to G_m3 ganglioside catalyzed by β-hexosaminidase A.

**PLD Assay.** PLD activity was routinely assayed by measuring the formation of [14C]phosphatidylethanol from [14C]PtdCho in the presence of ethanol. The reaction mixture (100 μl) contained 100 ng of PLD partially purified from the rat kidney, 200 nM ARF, 100 μM GTP-γ-S, 1.6 M ammonium sulfate, 60 μM [14C]PtdCho (4,583 dpm/nmol), 70 μM PtdEtn, 2% ethanol, 1 mM MgCl₂, and 20 mM Hepes-NaOH at pH 7.4, and each activator fraction to be assayed. [14C]PtdCho and PtdEtn were separately dried, dispersed in distilled water by sonication, and added to the reaction mixture. The detailed conditions were described elsewhere (19).

Where indicated, PLD activity was alternatively determined with PtdIns-4,5-P₂-containing mixed lipid vesicle as substrate under the condition described by Brown et al. (3). The reaction mixture (100 μl) contained 100 ng of PLD purified partially from the rat kidney, 200 mM ARF, 100 μM GTP-γ-S, 5 μM [14C]PtdCho (55,000 dpm/nmol), 80 μM PtdEtn, 7 μM PtdIns-4,5-P₂, 2% ethanol, 1 mM MgCl₂, 20 mM Hepes-NaOH at pH 7.4, and G_m2 ganglioside activator or PKC to be assayed. Lipids were mixed first in chloroform, dried, dispersed in distilled water by sonication, and added to the reaction mixture. Where indicated, PtdIns-4,5-P₂ was omitted from the lipid vesicles, and 1.6 M ammonium sulfate was added to the reaction mixture.

**Assay of Hydrolysis of G_m2 Ganglioside.** Hydrolysis of G_m2 ganglioside by β-hexosaminidase A was assayed in the presence of heat-stable PLD activator or G_m2 ganglioside activator under the condition specified (20). The reaction mixture (100 μl) contained 100 nM G_m2 ganglioside, 0.5 unit of β-hexosaminidase A, 25 μg of ovalbumin, 10 mM acetate buffer at pH 4.6, and either the heat-stable PLD activator or G_m2 ganglioside activator as specified. After incubation, G_m2 and G_m3 gangliosides were extracted and separated by TLC (20). The TLC plate was sprayed with resorcinol, and intensity of the bands was quantitated by a digital image processing program, Image (National Institutes of Health) by using a Macintosh Computer.

**Other Procedure.** Conventional PKC (mixture of PKCα, PKCβI, PKCβII, and PKCγ) was purified from rat brain as described (21). Protein was determined by the method of Bradford (22).

**RESULTS**

**Amino Acid Sequence of PLD Activator.** The heat-stable PLD activator was subjected to gel filtration analysis on a Superdex 200 column. The protein appeared as a symmetric peak with a molecular mass of ~30 kDa, and its elution profile coincided with that of the PLD-stimulating activity (Fig. 1A). This protein showed a single band with 23 kDa upon SDS/PAGE (Fig. 1B).

**FIG. 1.** Gel filtration analysis of PLD activator. (A) Purified PLD activator was loaded on a Superdex 200 column. Each fraction was assayed for PLD activation in the presence of 200 nM ARF. (B) Aliquots of the samples from Superdex 200 column were subjected to SDS/PAGE on a 12.5% gel followed by silver stain. The positions of molecular mass markers are indicated in kilodaltons.

The activator protein (3 μg) was reduced with DTT in the presence of guanidine-HCl and S-pyridylethylated with 4-vinylpyridine as described (23). The protein was collected by 10% trichloroacetic acid precipitation and then subjected to SDS/PAGE. The band corresponding to the 23-kDa protein was treated overnight with 1 μg of lysylendopeptidase (24). The resulting peptide fragments were applied to a C18 column equipped to a Capillary HPLC Microblotter System (Perkin–Elmer Applied Biosystem, model 173A), which had been equilibrated with 0.1% trifluoroacetic acid. The column was washed with the same buffer at a flow rate of 5 μl/min. Peptides were separated with a 70% linear concentration gradient (0–60%) of acetonitrile in 0.1% trifluoroacetic acid. Several peptides were obtained (Fig. 2 Inset). The seven peptides indicated were each subjected to a Protein Sequencer Procise (Perkin–Elmer Applied Biosystems, model 492). Total 76 amino acids were sequenced from these seven peptides by automated Edman degradation analysis. The sequence shows 90% homology with the primary structure of the mouse G_m2 ganglioside activator (ref. 25; Fig. 2). Peptide F, whose sequence was related to that of peptide E, also was recovered with the amount of 1:5 ratio, suggesting that the sample of the original activator isolated is a mixture of G_m2 ganglioside activator isoforms. By comparison, the sequence homology between human and mouse G_m2 ganglioside activators was 74%. After Superdex 200 column (Fig. 1), an aliquot of the heat-stable PLD activator was subjected directly to the sequence analysis. The only amino acid sequence detected was
GGFSWDNCDEGKDPAV, which corresponded to the amino-terminal sequence of G M2 ganglioside activator, verifying the purity of the preparation.

**PLD Activator as G M2 Ganglioside Activator.** The purified PLD activator, after Superdex 200 column, was assayed for its ability to enhance enzymatic conversion of GM2 to G M3 ganglioside that is catalyzed by β-hexosaminidase A. The elution profile to stimulate PLD and that to enhance enzymatic conversion of GM2 to GM3 ganglioside support the notion that a single protein possesses the two separate functions (Fig. 3). The properties of the PLD activator, including thermal stability and abundance in the kidney, were all consistent with those of GM2 ganglioside activator described (for review, see ref. 15).

**GM2 Ganglioside Activator as PLD Activator.** In the next experiment, GM2 ganglioside activator was separately purified from the rat kidney cytosol by the method described (18). The final preparation of the GM2 ganglioside activator was >95% pure as judged by silver stain using SDS/PAGE. In the both presence and absence of ARF, PLD alone was practically inactive, but exhibited its enzymatic activity almost linearly with increasing amounts of GM2 ganglioside activator added, and reached a plateau (Fig. 4A). GM2 ganglioside activator alone showed no effect unless ARF was added. The specific activity of GM2 ganglioside activator was comparable with the heat-stable activator. The reaction of PLD in the presence of both ARF and GM2 ganglioside activator proceeded linearly with time for 40 min (Fig. 4B). Under the given conditions, ≈40% of total PtdCho was converted to phosphatidylethanol after a 60-min incubation.

**PtdIns-4,5-P2 and PKC for PLD Activation.** PLD1 has been shown to require PtdIns-4,5-P2 for enzymatic activity (3). The enzyme also is activated by PKC, and this activation depends on phorbol ester but occurs in the absence of ATP (10). Under the condition comparable with that made for these observations (9), PLD from the kidney exhibited considerable activity in the simultaneous addition of PKC and PtdIns-4,5-P2 (Fig. 5). It is worth noting, however, that the enzymatic activity thus obtained was enhanced further by the addition of GM2 ganglioside activator and that the activity disappeared in the presence of higher concentrations of ammonium sulfate. Further, in the presence of ammonium sulfate, the enzyme was fully active when GM2 ganglioside activator was added irrespective of the presence or absence of PtdIns-4,5-P2 and PKC. ARF alone was very weak to activate the enzyme (Figs. 4 and 5). In this experiment, PKC was added together with PMA, but ATP was omitted from the reaction mixture.

**DISCUSSION**

Several protein factors are known as activators of mammalian PLD1 enzyme, that include small G protein such as ARF (3,
GM2 to G M3 ganglioside catalyzed by PLD activator; F and 200 nM ARF; E, with 200 nM ARF alone. 

heat-stable PLD activator. (■ activator and 200 nM ARF; ■, with 200 nM ARF.

4) and RhoA (26), PKC (9, 10), unidentified cytosolic proteins with 50 kDa (11, 12) and 23 kDa (14). 
PtdIns-4,5-P_2 (3) and PtdEtn (27) also are involved in the reaction, although these lipids do not serve as the substrates. The present study has shown that the cytosolic 23-kDa protein is indistinguishable from the previously known GM2 ganglioside activator. Deficiency of this protein causes the accumulation of a lethal quantity of GM2 ganglioside in the lysosomes, that is Tay-Sachs disease (for review, see ref. 15). The kidney PLD enzyme used in this study is ARF-dependent and shows properties similar to PLD1 (13). The enzymatic activity obtained in the presence of both PKC and PtdIns-4,5-P_2 disappears by the addition of higher concentrations of ammonium sulfate (Fig. 5), which presumably counteracts the proper association of PLD with PKC and PtdIns-4,5-P_2. Yet, in the presence of this salt, the enzyme is fully activated by the addition of both ARF and the heat-stable activator protein irrespective of the presence or absence of PKC and PtdIns-4,5-P_2. Apparently, this activator protein overcomes the inhibitory action of this salt, and it substitutes for PKC and PtdIns-4,5-P_2. ARF is always essential for PLD activity, and PtdEtn is necessary as cosubstrate as described earlier (27). In the reaction of β-hexosaminidase A, GM2 ganglioside activator is proposed to be associated with this lipid substrate rather than acts on the enzyme (for review, see ref. 15). However, no evidence has been available thus far, suggesting that this activator protein is bound to PtdCho, which is the substrate of PLD. It is proposed that PKC activates PLD1 by protein–protein interaction through its regulatory region rather than by protein phosphorylation through its catalytic region (9, 10). Contrarily, several reports have suggested direct involvement of protein phosphorylation in this PKC action (28, 29). Obviously, such a higher concentration of ammonium sulfate is unphysiological, and thus the biological significance of this heat-stable activator protein in the PLD reaction remains to be explored. Equally, the mechanism of PtdIns-4,5-P_2 action in the PLD reaction needs to be clarified further.

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