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Assays of dioxins and dioxin-like compounds in actually contaminated soils using transgenic tobacco plants carrying a recombinant mouse aryl hydrocarbon receptor-mediated β -glucuronidase reporter gene expression system

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 β -glucuronidase reporter gene expression system E'HERSOP EUN⁴ C'UNESE OK IS IM^{4,5} and HIPEO OHIKAWA ^{12,6} TR

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The transgenic robacco plant XD4V-26 carrying the recombinant mouse anythydrocarbon receptor XD4V-mediated \$\text{B}\$ iglucuronidase (GUS) reporter gene expression system was used for assay of dioxins and dioxin-like compounds consisting of polychlorodibenzo-p-dioxins polychlorinated dibenzofurans, and coplanar polychlorinated biphedyls (Co-PCBs) in actually contaminated soils. The transgenic tobacco plant XD4V-26 showed a significant dose-dependent induced GUS activity when cultured on MS medium containing PCB/26 [toxic equivalency factor (PEF) = 0.1]. In contrast PCB 169 and PCB 180, which have 0.03 of TEF and unassigned TEF values, respectively, did not significantly induce GUS activity under the same conditions as with PCB 126. When the tobacco plants were cultivated for up to 5 weeks on actually contaminated spils with dioxins and dioxin-like compounds collected from the periphery of an incinerator used for disposal of life and industrial wastes, GUS activity in the leaves was dose-dependently increased. The plants clearly detected 360 pg-TEQ g⁻¹ of dioxins and dioxin-like compounds in this assay. There was a positive correlation between GUS activity and TEQ value of dioxins and dioxin-like compounds in the plants. This assay does not require any extraction

and purification processes for the actually contaminated soil samples.

Keywords: Biochemical assay, dioxin, aryl hydrocarbon receptor, transgenic plants, polychlorinated biphenyl. TRACTED

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Polychlorinated dibenze p choxins (PCDDs), polychlorin nated dibenzofurans (PCDFs) and coplanar polychlo-ribated biphenyls (Co-PCBs), the so-called dioxins and dioxin-like compounds, are highly lipophilic and persistent in the environment. These compounds widely contaminated the environment. These were deposited in sediments in aquatic environments, and then highly accumulated at the tops of food chains, including humans. This contamination mainly/art occurred at the sites of incinerators of

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Hie and industrial wastes. Therefore, it is important to continuously monitor dioxins and dioxin-like compounds in the periphery of incinerators from the standpoint of risk assessment and management. RCDDs, RCDFs, and Co-PCBs each consist of a number of congeners with different numbers and positions of chlorines attached on two ben-zene rings and are represented by a toxic equivalency factor (TEF), which was updated by the World Health Organization in 2005. A The THE is a relative toxicity value based on the value 1 of the most toxic 2,3,7,8-tetrachlorodibenzo-n-dioxin (2,3,4,8-TCDD). A mixture of these congeners is represented by the total toxic equivalency (TEQ), which is the sum of concentration of each of the congeners multiplied by its TEF.

High-resolution gas chromatography/high-resolution mass spectrometry (HRGC/HRMS) is used to identify and quantify residues of PCDDs, PCDFs and Co-PCBs in

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environmental samples. This method is sensitive and accurate for measuring extremely low amounts of the residues in a variety of environmental samples. However, extraction and purification of these compounds from samples are imperative and result in a high cost of analysis. In contrast, blochemical assays based on prolecular mechanisms of modes of actions of chemicals are suitable for rapid screening of a number of different kinds of samples and are advantageous for assessing the toxicity of these residues in mammals. Most of the biochemical assays estimate toxicity levels of dioxins and dioxin-like compounds in recombinant mammalian hepatoma cells expressing anyl/hydrocarbon receptors (AlfRs), since AhRs are primarily responsible for toxicity in mammals towards these compounds. [2,3] However, like instrumental analyses, these methods require several steps for extraction and purification of the samples.

The transgenic tobacco plants carrying a gene encoding the recombinant AhR, XDV, consisting of the ligandbinding domain of mouse AhR, the DNA-binding domain of bacterial repressor profein LexA, and the transactivation domain of the virus VP16 as well as β -glucuronidase (GUS) reporter gene were genetically engineered. These transgenic tobacco plants showed a significantly increased GUS activity when treated with the AhR ligands such as indigo. β -naphthoflavone, and 3-methylcholanthrene (MC). [4] The transgenic tobacco plants seemed to be useful for a biochemical assay of dioxins and dioxin-like compounds toxic to mammals. The assay did not need any extraction and purification of chemicals, since the plants took up these chemicals in medium through their developed roots by passive diffusion.

In this study, the transgenic tobacco plant XII/4V-26 was examined for practical assays of PCDDs, PCDFs, and Co-PCBs in actually contaminated soils collected from the periphery of an incinerator used for disposal of life and industrial wastes, since the transgenic plants showed low background activity and dose- and time-dependent GUS arrivity induced in response of METRACTED

RETRACTED RETRACTED Materials and Methods RETRACTED RETRACTED

The compounds 13',4,4,5-Pentachlorobipheny (PCBI26:) TEF = 0.1), 3,3',4,4',5,5'-hexachlorobiphenyl (PCB169; TRF = 0.03), and 2.2', 3,4,4',5,5'-heptachlorobiphenyl (PCB180; TEF value was not assigned) were purchased from Accustandard Inc. (New Heaven, CTAUSA). They were each dissolved in dimethyl sulfoxide (DMSO) for application to Murashige and Skoog (MS medium. The final concentration of DMSO was 0.1 % in MS medium The compounds R4-Methyl-umbelliferyl-11-D-glucuronide (4MUG) for a substrate of GUS and 4-methyl-umbelliferone (4MU) as a product of GUS reaction were purchased from Nacalai Tesque (Kyoto, Japan).

RETRACTED RETRACTED were transformed by the use of Agrobacterium tumefa-cions carrying the plasmid pGPXXD4VGUS as previously described. The resultant transgenic tobacco strain XD4V-26 carries the gene encoding recombinant AhR, which consists of the DNA-binding domain of bacterial LexA [amino acids (AA) Lto 2027, the ligand-binding do main of mouse AhR (a.a. 83 to 494), and the transactivation domain of virus VP16 (a.a. 413 to 490), as well as the gene encoding the reporter GUS. The transgenic tobacco plant KD4V-16 was a eptically and separately incubated in MS medium containing each of the PCB congeners in a growth ckamber at 23 C under 16-1 Right 18-R dark cycle con ditions or grown on the soils contaminated with PCDDs, PCDFs and Co-PCHs ma closed greenhouse undernatural day-length light conditions.

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Weathered contaminated soils with PCDDs, PCDFs, and Co-PCBs (\$100 pg-TEQ g) were collected from the pe riphery of an incinerator used for disposal of life and in-dustrial wastes in Japan. Table V shows concentrations of PCDD, PCDF, and Co-PCB congeners in the collected soils. The soils were diluted with uncontanimated soils (1.1) pg-TEQ g^{-1}) purchased from Suntory Flowers Ltd. (Tokyo, Japan) prior to assays. The uncontaininated soils were also

used as the control soils. RETRACTED RETRACTED

Fluorometric GUS assay of transgenic tobacco plants E

Axillary buds of the transgenic tobacco plants were cultured on MS medium containing 3.0% (w/v) sucrose, 0.7 % (w/y) again and 0.1 to 1000 ng mL P6B126, P6B169, or P6B180. After two weeks, either the second or third leaf from the top of the plant, or both, were used for fluorometric GUS assay as previously described. [3] Sol uble fractions were prepared from leaves and incubated with 4MUG. After stopping GUS reaction, fluorescence at 360 nM excitation and 450 nM emission was measured with a microplate reader (MTP-100F; CORONA, Kat suta, Japan). GUS activity was determined from a standard curve of the fluorescence of AMU, and data were presented

as means ± standard deviation (SD).

RAxillar bods were also aceptically cultured or MS medium for about a month. Approximately 10 cm high plants were individually transferred into 1/5000-acre pots filled with the soils contaminated with dioxins and dioxinlike compounds or the uncontaminated soils. The surface of the soils was covered with the uncontaminated soils or aluminum foil. The tobacco plants were grown in a closed greenhouse for several weeks, and then either the second or third leaf from the top of the plants, or both, were assayed for GUS activity as described above.

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Transgenic tobacco plant for dioxin assay RETRACTED RETRA Table 1. Concentrations of PCDD, PCDF, and Co-PCB congeners in the contaminated soils containing 5100 pg-TEQ grant Co-PCB, and Co-PCB, and Co-PCB. PCODSTRA(1005.06 346.84 22.71 1,3,6,8-TetraCDD R.3,7,9 Tetra CDD 2,3,7,8-Tetra CDD D,23,7,8-PentaCDD 1,2,3,4,7,8-HexaCDD 123,67,8-HexaCDD EI 1,2,3,4,6,7,8-HeptaCDD K23,4,5,6,7,8-OctaCDD R23222 10 T 79735.53 Total PCDDs A PROFSTRA(1,2,7,8-TetraCDF 186.37 2,3,7,8-TetraCDF 1,2,3,7,8-PentaCDF **A**11.29 193.18 (11.T3EI 35.80 1,2,3,6,7,8 HexaCDF 1,2,3,7,8,9 HexaCDF 2.3.4.6.7.8-HexaCDF R.2.3.4.6.7.8-HeptaCDF RETR48611.88 A 486. 18 E $99.\overline{0}1$ 1,2,3,4,7,8,9-HeptaCDF 9901.31 P.213,415,67,840ctaCDF Total PCDFs RA003' R83064.51TE AC432ED CRPCBSTRACTED
3,3',4,4'-TetraCB(#77) 3,4,4',5-Tetra GB(#81) 3,3,4,4',5 Penta GB(#126) 33',4,4',5,5'-HexaCB(#169) 33,3',4,4' Renta CB(#105) 0.00003 2,3,4,4',5-PentaCB(#114) 2,3,4,4',5-PentaCB(#118) 2',3,4,4',5-PentaCB(#123) 0.00003 941.48 129.42 0.**0**3 E 0.00003 $0.\overline{00003}$ 3,3',4,4' \$-HexaCB(#156) 3,3',4,4',5'-HexaCB(#157) $R_{0.00003}$ 2,3,4,4,5,5'-HexaCB(#167) 2,3,3',4,4',5,5'-HeptaCB(#189) 2 2′ 3,3′ ,4,4′,5 HeptaCB(#170) R2,3,44R5 HeptaCB(#180) RET 5238,69 940,58 T Total Co-PCBs 7945.89 TRITETRA RETRACTED RETRACHED HRGC/HRMS analysis of PCDDs, PCDFs, and RETRACTED RETRACTED congeners in the transgenic tobacco plants as well as dif-RETRACTED ferences between uncontaminated and contaminated soils. Co-PCBs in tobacco plants Pearson's correlation coefficient was used for analysis of As previously described, PCDDs, PCDFs, and Co-PCBs were extracted from the aerial parts of the tobacco correlation A RETR ED plants. [6] Congeners of PCDDs, PCDFs, and Co-PCBs were each quantified by HRGO/HRMS (HP6899) Micromass Autospec-Ultima, Micromass Ltd., Manchester, UK) RETRACTED RETRACTE
6US activity in the transgenic tobacco plant XD4V-26 der the same conditions as previously reported.[7] aultured on MS medium containing PGB congeners RFTRAC Statistical analysis The transgenic tobacco plant XD4V-26 was cultured on MS medium Achtaining each of PCB 26/(TEF = 0,1) Statistical analysis used Student's t-test for GUS activity in the transgenic tobacco plants in the response to PCB PCB169 (TEF = 0.03), and PCB180 (not assigned TEF RETRAC RETRAC

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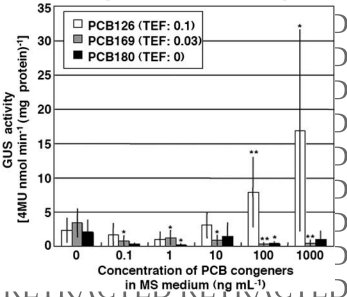


Fig. 1. GUS activity in leaves of the transgenic to bacop plant XD4V-26 cultured for 2 weeks on MS medium containing PCB126, PCB169, or PCB180. Values are means \pm SDs (PCB126, n = 9 - 12 PCB169, n = 6 - 8; PCB180, n = 3 - 9). Significant difference (Student's t-test): **, p < 0.01; *, p < 0.05. RETRACTED RETRACTED

value) for 2 weeks, and then leaves were subjected to GUS assay. As shown in Figure 1, a dose-dependent increased GUS activity was clearly observed when the tobaccoplants were treated with PCB126, with significant differences at 100 and 1000 ng mD (P < 0.01 and P < 0.05, respectively, relative to no treatment with PCB126, Student's t-test). The GUS activity at 1000 ng mL (was 7.4 times higher than that in the plants treated with DMSO alone. In contrast, no dose-dependent increased GUS activity was observed when treated with PCB169 and PCB180. Background GUS activity was detected in tobacco plants treated with DMSO and was significantly decreased by the treatment with increasing doses, of PCB169 and PCB180 (P < 0.05). Thus, the present study indicated that the transgenic tobacco plant XD4V-26 showed induced GUS activity in response to the against PCB126 and reduced GUS activity in response to the antagonists PCB169 and PCB180.

GUS activity and uptake-of PCDDs, PCDFs, and TED Co-PCBs in the transgenic tobacco plant XD4V-26 cultured on the actually contaminated soils FTRA CTED

The transgenic tohacco plant XD4V-26 was cultured in pots containing the soils actually contaminated with PCDDs, PCDFs, and Co-PCBs. The GUS activity in their leaves was significantly induced after 5 weeks of culture on the soils containing 360 pg-TEQ g⁻¹ of these congeners (Fig. 2a). On the other hand, the GUS activity in the tobacco plants cultured on the uncontaminated soils containing 1H pg-TEQ g⁻¹ was not induced so high. The transgenic

tobacco plant XD4V-26 was also cultured in the soils containing 510 and 5100 pg-TBQ g for 33 days. The leaves were subjected to assay of GUS activity, and the aerial parts were analyzed in HRQC/HRMS for PCDD, PCDF, and Co-PCB congeners. The GUS activity in the tobacco plants dultured on the soils containing \$100 pg-TEQ g was twice higher than that in the plants cultured on the soils of 310 pg-TEQ g (Fig. 2b). The concentrations of PCDDs, PCDFs, and Co-PCBs in the plants cultured on the soils containing 5100 pg-TEQ g were higher than those of the plants cultured on the soils of 510 pg-TEQ g (Fig. 2c). Particularly, the concentrations of A23,4,5,6,7,8 octaCDD and 2,3',4,4',5-pentaCB (PCB118) in the plants were the highest, followed by 2,3,3',4,4'-pentaCB (PCB105) and others (Fig. 3).

RETRACTED RETRACTED Correlation between GUS activity and TEQ values of RCDD, RCDF, and Go-POB Caken up into the TED transgenic tobacco plant XD4V-26

There was a significant positive correlation [0.681 (Person's correlation coefficient), P < 0.01, Fig. 4] between the GUS activity and TEQ values of PCDDs, PCDFs, and Co-PCBs in the tobacco plant. In contrast, the transgenic tobacco plants grown on the uncontaminated soils that contained low level of PCDDs, PCDFs, and Co-PCBs including toxic and non-toxic congeners, accumulated as indicated by 0.5 to 1.0 pg-TEQ g⁻¹, although the GUS activity was very low. These results suggested that the induced GUS activity may be due to TEF values of the congeners, although the uptake of congeners of PCDDs, PCDF sand Co-PCBs in the plants seemed to be under passive diffusion mecharism. The effects of certain congeners in which a TIFF value was not assigned, were examined on GUS activity in the transgenic plants. The relative amounts are defined as the amounts of PCDD and PCDF congeners without TEF value in the plants divided by the amounts of PCDD and PCDF congeners with TEF values of 0. These relaive amounts were negatively correlated with GUS activity 1-0.486 (Person's correlation coefficient), P < 0.05, data not shown]. These results suggested that relatively low GUS activity in the transgenic plants grown in the uncontaminated soils seemed to be due to antagonistic effects of certain congeners in which TEF values were not assigned.

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The transgenic tobacco plant XD4V-26 carrying the recombinant mouse AhR-mediated GUS reporter gene expression system exhibited a dose-dependent GUS activity towards PCB426 (TEF = 0.1), but not towards PCB169 (TEF = 0.03) and PCB180 (TEF was not assigned,). It suggested that the plant took up PCB126, which activated the recombinant mouse AhR XD4V, and then induced GUS activity. Uptake of PCB169 and PCB180 also occurred

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Transgenic tobacco plant for dioxin assay
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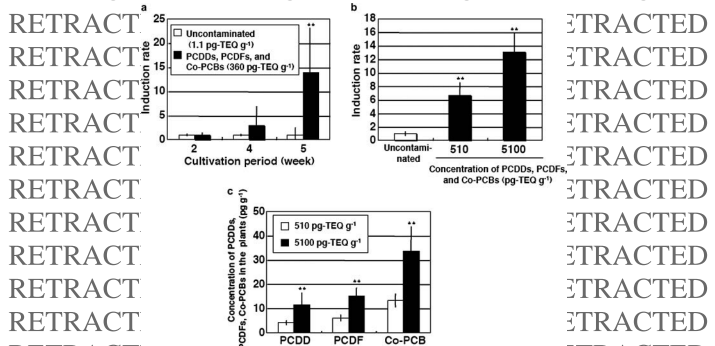


Fig. 2. Time-dependent (a) and dose-dependent (b) GUS activity, and concentrations of PCDD, PCDF, and Co-PCB congeners (c) in the transgenic tobacco plant XID4V26 cultured on the actually contaminated soils. The transgenic plant XID4V26 was cultured on the soils containing 510 or 5100 pg-TEQ g⁻¹ for 33 days. The induction rate is expressed as the GUS activity in the transgenic plants cultured on the contaminated oils divided by the activity in the plants cultured on the uncontaminated soils. Values are means \pm SDs (a: uncontaminated, n = 11; dioxins 300 pg-TEQ g⁻¹, n = 16-17; b: uncontaminated, n = 4; 310 pg-TEQ g⁻¹, n = 8; 5100 pg-TEQ g⁻¹, n = 8

but those compounds did not contribute to induce GUS activity because of very low TEF value and almost no toxicity towards mammals, respectively. Instead, background GUS activity, probably due to endogenous AlaR digands such as indoleacetic acid, was suppressed by PCB169 and PCB180 Goncurrent treatment with PCB126 and PCB180 tended to suppress GUS activity induced by PCB126 (data not shown). It was reported that certain congeners with low or not assigned TEF values have shown antagonistic activity toward AlaR in rat primary hepatocytes: PCB153 (TEF was not assigned), decreased the induction of CYPIA1 by the treatment with 2,3,7,8-TCDD, but PCB77 (TEF = 0.0001) and PCB156 (TEF = 0.0003) did not. Utwas also reported that di-, tri-, and tetra-ortho-substituted PCBs suppressed the activation of AlaR by PCBs with no or one ortho chlorine substitution. In It was reported that certain persistent organic politicants (POPs), including pp-dichloro-diphenyl-trichloroethane (DDT) and dieldrin, were antagonistic toward AlaRa (ILL2) In transgenic Arabidopsis plants carrying genes encoding a recombinant guinea pig AlaR (and the GUS reporter, p,p(-DDT), pip-dichloro-diphenyl-dichloroethane (p,p-DDE), and p,p-dichloro-diphenyl-dichloroethane (p,p-DDE), and p,p-dichloro-diphenyl-dichloroethane (p,p-DDE) decreased the GUS activity induced by MC[13] Antagonism between the agonist MC and dieldrin or p,p-DDT was also ob-

RETRACTED RETRACTED Served in an in vitro assay using an Ah immunoassay. The present assay responded to both agonists and antagonists among PCB congeners. Therefore, the GUS activity in the assay may reflect the relative toxicity level of a mix-ture of PCB congeners. It was suggested that the relative toxicity level of a mixture of congeners of PCDD, PCDF, and Co-PCB should not be simply determined by TEQ as the sum of concentrations of congeners multiplied by their TEF values, since TEQ does not count antagonistic effects of certain congeners. ED RETPAY Was enly tured on the soils containing dioxins and dioxin-like com-pounds for 5 weeks, the GUS activities in the leaves were time- and dose-dependently increased. The transgenic plant was thus able to monitor the levels of both 360 pg-TEQ g⁻¹ and 5100 pg-TEQ g⁻¹. The environmental standard (less than 1000pg-THQ g⁻¹) for dipxins and dioxin-like compounds in soils in Japan was possibly detected in the transgenic plant XD4V-26 within 5 weeks of planting. However the GUS activity of plants grown in 5100 pg-TEQ g⁻¹ was not 10 lines Aigher than that of plants grown in 510 pg TEQ g⁻¹. This suggests that the tobacco plants may not be able to accumulate very high concentrations of these congeners and may reach the plateau between 510 and 5100 pg-TEQ g ?. Moreover the different levels of uptake of the

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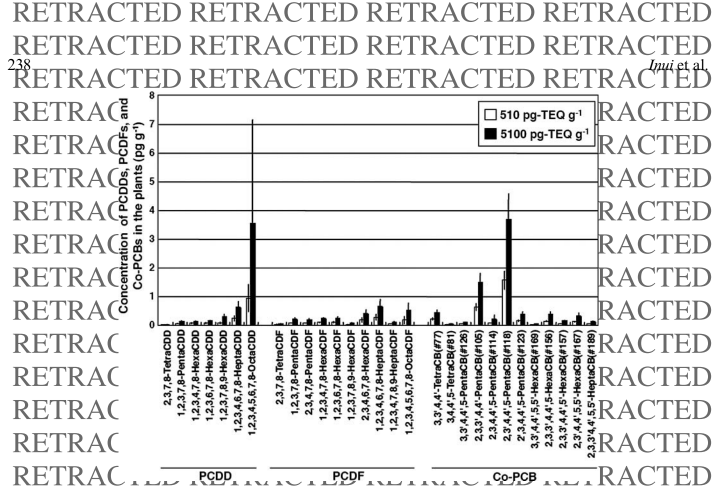


Fig. 3. Concentrations of PCDD, PODE and Co-PCB congeners in the transgenic Advacco plant XD49-26 cultured on the containing instead soils containing 510 and 5100 pg-TEQ g — for 33 days. Values are means ± SDs (510 pg-TEQ g —, n = 8; 5100 pg-TEQ

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congeners in the plants may be due to the mass and growth stages of the plants.

The amounts of uptake of dioxins and dioxin-like compounds in the tobacco plants by passive diffusion were not

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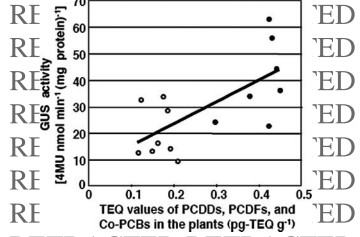


Fig. 4. Correlation between GUS activity and TEQ value in the transgenic tobacco plant XD4V-26 cultured on the actually contaminated soils containing 510 (open circle) and 3100 (closed circle) pg-TEQ g^{-1} for 33 days. Pearson's correlation coefficient: r = 0.681

so large as compared with that of zucchin pants. [0.13] It was reported that root exudates such as low-molecular-weight organic acids [1.16] and blosurfactants [7.18] may be involved in the uptake of hydrophobic compounds such as POPs. By the use of these compounds, the transgenic tobacco plants seem to increase the uptake of dioxins and dioxin-like compounds in soils. The this study, transgenic tobacco plant XD4V-26 carrying genes encoding a recombinant mouse AbR and the GUS reporter successfully detected residues of PCDD, PCDP, and Go-PGB congeners in the contaminated soils within 5 weeks. These results suggested that the assay using the plant seems to be useful for on-site assays of these residues in contrast, cultivation of the transgenic plants on samples of the contaminated soils in the closed laboratory may provide some advantages such as public acceptance of the use of transgenic plants, and possible control of environmental conditions, including temperature, drought, and day length, all of which may influence on the stable GUS assay.

Acknowledgments.

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