Lipid-soluble polyphenols from sweet potato exert antitumor activity and enhance chemosensitivity in breast cancer

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Polyphenols are abundant in vegetables and fruit. They have been shown to have various antitumor, antioxidant, and anti-inflammatory effects. Here, we extracted the lipid-soluble fraction of polyphenols from fermented sweet potato (Ipomoea batatas). These lipid-soluble polyphenols mainly contained caffeic acid derivatives with strong antioxidant ability, which we hypothesized to affect diseases for which oxidative stress is a factor, such as cancer. We therefore investigated the antitumor and chemosensitizing effects of lipid-soluble polyphenols on E0771 murine breast cancer cells. The lipid-soluble polyphenols accumulated in the cells’ cytoplasm due to its high lipophilicity, and reduced reactive oxygen species through its strong antioxidant activity. The lipid-soluble polyphenols also arrested the cell cycle at G0/G1 by suppressing Akt activity, and enhanced the cytotoxicity of anticancer agents. In this model, lipid-soluble polyphenols inhibited tumor growth and enhanced the efficacy of chemotherapy drugs. These results suggest the potential of lipid-soluble polyphenols as a functional food to support cancer therapy.

Key Words: polyphenol, breast cancer, functional food, chemotherapy, antioxidant

Polyphenols reportedly have antitumor, antioxidant, anti-inflammatory, and cardiovascular-protective effects,1-4 which are thought to result from modulation of molecular targets and signaling related to cell survival, proliferation, differentiation, migration, and angiogenesis.5 Therefore, polyphenols have the potential to become natural anticancer agents.

Various polyphenols with antioxidant activity are present in tubers of Ipomoea batatas, known as the sweet potato.6,7 Fermented potato pulp also contains alkyl ferulates, a kind of polyphenol that is lipid-soluble and has antioxidant activity.8 Therefore, we fermented sweet potatoes and extracted the lipid-soluble polyphenol fraction. Lipid-soluble polyphenols (PPL) mainly contain caffeic acid derivatives such as caffeic acid ethyl ester and caffeic acid cetyl-ester, as strong antioxidant components, which are reported to inhibit growth of some cancer cells, and to inhibit lipid oxidation.9 PPL is a novel polyphenol that is expected to have high cellular affinity and absorbability.

Among woman around the world, breast cancer (BC) is the most commonly diagnosed cancer the leading cause of cancer death.10 Surgical resection is the first treatment choice for BC, with chemotherapy and radiotherapy typically used to eliminate residual tumor cells, suppress tumor growth, and prevent recurrence. Although these adjuvant therapies improve overall survival and quality of life for patients with BC,11 they often have adverse effects, such as myelosuppression from carboplatin,12 neuropathy from vincristine,13 and dermatitis from radiotherapy.14 To reduce these side effects, chemosensitizers and radiosensitizers have been studied.15,16 The discovery of bioactive natural products with antitumor and chemo-sensitizing effects might therefore provide complementary strategies for treating BC.

In this study we evaluated the antitumor effects of PPL on BC and the effects of combining PPL with chemotherapy drugs. We found PPL to function as an antioxidant with antitumor effects despite its low toxicity, both in vitro and in vivo, and to enhance the efficacy of chemotherapy agents.

Materials and Methods

Reagents. The PPL was manufactured by Genuine R&D Co., Ltd. (Tokyo, Japan). The amount of caffeic acid derivatives in PPL was analyzed by TLC densitometry method as described previously (Table 1).9 We purchased carboplatin (CDDCA; Cat. No. C2538) from Sigma-Aldrich, (St. Louis, MO); vincristine sulfate (VCR; Cat. No. V5254) from LKT Laboratories, Inc. (St. Paul, MN); N-Acetyl-l-cysteine (NAC; Cat. No. 015-05132) and 2,2-diphenyl-1-picrylhydrazyl radical (DPPH; Cat. No. 044150) from Wako Pure Chemical Industries (Osaka, Japan); and acenitromine [MeCN; Cat. No. 00433-95; spectral grade; used (as received) as a solvent for DPPH] from Nacalai Tesque, Inc. (Kyoto, Japan).

Cell culture. We maintained mouse BC E0771 cells (Cat. #94A001, CH3 BioSystems, NY) in RPMI-1640 medium (Cat. No. 189-02025, Wako Pure Chemical Industries) supplemented with 10% fetal bovine serum (FBS) and 10 mM HEPES buffer solution (Cat. No. 345-06681, Wako Pure Chemical Industries) at 37°C, 5% CO2.

We maintained human BC MDA-MB-231 cells (Cat. No. ATCC® HTB-26™, ATCC, Manassas, VA) in DMEM (Cat. No.

| Table 1. The amount of caffeic acid derivatives in PPL |
| --- |
| Caffeic acid derivatives (%) |
| Caffeic acid ethyl ester | 2.51 |
| Caffeic acid cetyl ester | 0.8 |

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043-30085, Wako Pure Chemical Industries) supplemented with 10% FBS at 37°C, 5% CO2.

**Confocal laser microscope.** E0771 cells were seeded in 35-mm, poly-lysine-coated, glass-bottomed dishes (Cat. D111310, Matsunami Glass Ind., Ltd., Osaka, Japan) and cultured overnight. After treatment with 50 μM PPL for 48 h, fluorescence images of live cells were taken with a Leica TCS SP5 confocal laser microscope (Leica, Heidelberg, Germany). PPL was visualized at Excitation/Emission = 331 nm/427 nm.

**Intracellular reactive oxygen species (ROS) detection.** Intracellular oxidative stress was evaluated by staining with 2′,7′-dichlorofluorescin diacetate (DCFDA; Cat. No. D6883, Sigma-Aldrich), as previously reported.

**DCFDA for 30 min at 37°C.** After probing with horseradish peroxidase (HRP)-conjugated antibodies, membranes were separated by SDS-PAGE and transferred onto a PVDF membrane (Immobilon-P, Millipore Corp., Bedford, MA). Proteins were probed with specific antibodies diluted in 5% NAC for 24 h. After treatment, cells were stained with 20 μM DCFDA for 30 min at 37°C. Cells were analyzed for fluorescence intensity using an EC800 Analyzer (Sony Biotechnology, Tokyo, Japan).

**DPPH assay.** Rates of DPPH-scavenging reactions by PPL or NAC were determined by monitoring absorbance changes at 519 nm from DPPH, using a stopped-flow technique on a Unisoku RSP-1000-02NM spectrophotometer at 25°C.(18) The pseudo-first-order rate constants (kobs) were determined by a least-squares curve fit. The first-order plots of ln(A–A∞) vs time (A + A∞) were denoted as the absorbance at the reaction time and the final absorbance, respectively, and were linear until three or more half-lives were seen with correlation coefficients >0.999. In each case, rate constants derived from at least three independent measurements agreed within an experimental error of ±5%.

**Clonogenic survival assay.** A clonogenic survival assay to determine the cytotoxicity of PPL was performed as previously described.(19) Cells were trypsinized, diluted, counted, and seeded into 60-mm dishes at densities of 200–5,000 cells/dish before being allowed to adhere in a 37°C incubator. After 16 h, cells were incubated with PPL, antioxidant agents, or antioxidant agents + PPL or NAC for 48 h. Cells were then allowed to grow in a humidified 5% CO2 atmosphere at 37°C for 5 days before being fixed with methanol and stained with 0.03% crystal violet solution (Wako Pure Chemical Industries). Colonies that contained more than 50 cells were scored as surviving cells. The survival curves and IC50 were calculated using Prism8 Software (GraphPad Software Inc., La Jolla, CA).

**Cell-cycle analysis.** To analyze cell-cycle perturbation, DNA was stained with propidium iodide and the cells were analyzed by flow cytometry.(20) Cells were incubated with PPL for 48 h. We resuspended 0.5 × 106 cells in PBS and fixed them in ice-cold 70% ethanol for at least 6 h. Fixed cells were washed in PBS twice, and incubated with propidium iodide staining solution (50 μg/ml propidium iodide, 5 μg/ml RNase I, 137 mM NaCl, 2.7 mM KCl, 8.1 mM Na2HPO4, and 1.47 mM KH2PO4) at 37°C for 30 min. The DNA content of at least 10,000 cells/sample was analyzed using an EC800 Analyzer (Sony Biotechnology) and FlowJo software (BD Biosciences, CA).

**Western blotting.** After probing with horseradish peroxidase (HRP)-conjugated secondary antibodies, bound antibodies were detected with Immobilon Western HRP substrate. Densitometry was performed using ImageJ software (NIH, Bethesda, MD). For Western blotting, we used anti-Akt (Cat. No. #4069S, Cell Signaling Technology, Beverly, MA), anti-Akt-pSer473 (Cat. No. #4060S), anti-β-Actin-Peroxidase (Cat. No. A3854, Sigma-Aldrich), and goat anti-rabbit IgG(H+L) (Cat. No. #A16110, Thermo Fisher Scientific, Tokyo, Japan) as HRP-conjugated secondary antibodies. Immobilon Western HRP substrate was purchased from Millipore (Billerica, MA).

**General animal methods.** All animal experiments were performed according to the established guidelines of the “Law for the Care and Welfare of Animals in Japan,” and approved by the Animal Experiment Committee of Azabu University (No. 190507-3). Mice were housed in plastic cages in an air-conditioned room at 24°C in a 12 h light-dark cycle (light on at 7:00 am) with food and water available ad libitum in SPF conditions. Tumor-bearing mice were ethically sacrificed when tumor volume reached 1,500 mm3 (or ~1.5 cm in any dimension); tumor burden became >10% of the body weight; tumors ulcerated, became necrotic or infected; a tumor impeded the animal’s ability to move its limbs and assume normal body postures; or an animal became moribund, cachectic, or unable to obtain food or water. At the end of experiments, animals were anesthetized with 2% isoflurane and sacrificed by cervical dislocation. Death was determined by confirming that dislocation took place if the animal’s head hung loosely from its body.

**Tumor transplantation and experimental therapy.** Female C57BL/6/N mice, aged 6–8 weeks, were maintained and bred in the animal facility of Azabu University. For allograft transplantation, two million E0771 cells were inoculated subcutaneously into the mammary gland of each mouse. Tumor volume was calculated as V (mm3) = π/6 × length × width2 (length: largest tumor diameter; width: perpendicular tumor diameter). When tumor volume reached 100–200 mm3 (8 days after transplantation), PPL suspended in methyl cellulose (Cat. No. 131-17811, Wako Pure Chemical Industries) was orally administered to the mice (6 mg/kg/day) for 7 consecutive days; and CBDCA was intraperitoneally administered at a single dose (120 mg/kg/day) on Day 8 after transplantation.

**Immunohistochemistry.** Histological analysis was performed as previously described.(22) Tumor tissues were excised and fixed with 4% buffered formaldehyde, embedded in paraffin, and sliced into 5-μm sections. After being deparaffinized, slides were immersed in citrate buffer and continued to antigen activation at 120°C for 2.5 min using a pressure cooker. Next, slides were immersed in 3% hydrogen peroxide solution for 20 min to remove endogenous peroxidas. After blocking non-specific binding sites, slides were probed with anti-cleaved caspase-3 (Cat. No. #9664, Cell Signaling Technology) overnight. Slides were then incubated with HRP-conjugated secondary antibody (Cat. No. 414321, Nichirei Bioscience, Tokyo, Japan). After coloring with a DAB substrate kit with hydrogen peroxide (Cat. No. SK-4100, Vector, Burlingame, CA), slides were stained with hematoxylin and sealed after dehydration. Images were taken with a BZ-X700 microscope (Keyence, Osaka, Japan).

**Statistical analysis.** All values are expressed as means ± SD of 3–5 independent experiments. Differences between groups were evaluated by Student’s t tests (two-sided) or Tukey-Kramer tests. P<0.05 was considered significant.

**Results**

**PPL reduces intracellular oxidative stress and has strong antioxidant activity.** To evaluate PPL cytotoxicity, we used clonogenic survival assays. E0771 cells were treated with PPL for 48 h. The IC50 of PPL against E0771 cells was approximately 116 μM (Fig. 1A). As PPL can emit 427 nm fluorescence from 331 nm excitation light, we visualized intracellular PPL uptake by a confocal laser microscope. As shown in Fig. 1B, PPL was localized in the cytoplasm of E0771 cells. Cellular ROS levels
were measured after 24 h of PPL or NAC treatment. PPL eliminated more intracellular ROS, and at lower concentrations, than did NAC (Fig. 1C).

Antioxidant activity of PPL was estimated using the 2,2-diphenyl-1-picrylhydrazyl radical (DPPH), which is frequently used as a reactivity model for ROS. Upon mixing PPL with DPPH in MeCN, the absorption band at 519 nm due to DPPH immediately decreased, with a clear isosbestic point at 423 nm. This indicates that PPL efficiently scavenged DPPH. The decay of absorbance at 519 nm obeyed pseudo-first-order kinetics, when the concentration of PPL was higher than a 10-fold excess of the DPPH concentration. The pseudo-first-order rate constant ($k_{obs}$) linearly increased with the PPL concentration ([PPL]). The slope of the linear plot indicated that the second-order rate constant ($k$) for the reaction between PPL and DPPH was 8.2 M$^{-1}$ s$^{-1}$ in MeCN at 25°C (Fig. 1D). The $k_{obs}$ values for the reaction of NAC + DPPH in MeCN at 25°C (determined in the same manner) was much smaller than those for the PPL reaction with DPPH, although no linear correlation between the $k_{obs}$ values and [NAC] was observed; therefore, the $k$ value could not be determined. These results suggest that PPL is a strong antioxidant with low cytotoxicity, and exerts intracellular antioxidant activity by being taken into cells through lipophilicity.

PPL induces cell-cycle arrest at the G$_0$/G$_1$ phase by suppressing Akt. To evaluate the effect of PPL on the cell cycle, cellular DNA contents were analyzed by PI staining and flow cytometry. After PPL treatment for 24 h or 48 h, cells at the G$_0$/G$_1$ phase increased, cells at the S phase decreased slightly and cells at the G$_2$/M phase decreased significantly (Fig. 2B). To elucidate the mechanism of cell-cycle progression, we examined Akt protein expression, which is related to cell-cycle regulation. In accordance with cell-cycle arrest, Western blotting showed phosphorylated Akt (Ser473) was significantly downregulated after PPL treatment (Fig. 2C and D: left side). As well as E0771, PPL decreased phosphorylated Akt (Ser473) in human BC MDA-MB-231 (Fig. 2C and D: right side).
PPL enhances the efficacy of chemotherapy. To examine the chemo-sensitizing effect of PPL, E0771 cells were treated with anti-cancer agents and PPL. Whereas the IC_{50} of CBDCA alone was 8.89 μM, it was 5.66 μM when combined with PPL. In contrast, NAC significantly reduced the cytotoxicity of CBDCA (Fig. 3A, Table 2). PPL also decreased the IC_{50} of VCR (VCR alone: 7.64 nM, VCR + PPL: 7.08 nM); and NAC significantly reduced the cytotoxicity of VCR (Fig. 3B, Table 3). These results suggest that PPL, unlike NAC, enhances the cytotoxicity of anticancer agents.

PPL suppresses tumor growth and enhances chemotherapy effects. To evaluate the effect of PPL on BC growth in vivo, mice allografted with E0771 cells were treated with PPL. As shown in Fig. 4A, PPL significantly inhibited tumor growth (Day 22, control: 1,470.2 ± 79.7 mm^3; PPL: 973.1 ± 85.9 mm^3). The group treated with PPL had a longer median survival time compared with the control group (Fig. 4B). Furthermore, PPL had no effect on the weight of E0771 tumor-bearing mice (Fig. 4C). These results suggest that PPL has an antitumor effect with low toxicity.
Colony formation assay shows PPL enhances cytotoxicity of anticancer agents. (A) Colony formation assay in cells simultaneously treated for 48 h with 20 μM PPL + CBDCA, or 5 mM NAC + CBDCA, at the indicated CBDCA concentrations. (B) CFA in cells simultaneously treated for 48 h with 20 μM PPL + VCR or 5 mM NAC + VCR at the indicated VCR concentrations. Error bars: SD, *p<0.05, **p<0.01.

Table 2. IC<sub>50</sub> of CBDCA, alone and combined with PPL or NAC

|          | IC<sub>50</sub>(μM) |
|----------|---------------------|
| CBDCA    | 8.89                |
| PPL + CBDCA | 5.66               |
| NAC + CBDCA | 20.28              |

CBDCA, carboplatin; IC<sub>50</sub>, half-maximal inhibitory concentration; NAC, N-acetyl-L-cysteine.

Table 3. IC<sub>50</sub> of VCR, alone and combined with PPL or NAC

|          | IC<sub>50</sub>(nM) |
|----------|---------------------|
| VCR      | 7.64                |
| PPL + VCR | 7.08               |
| NAC + VCR | 8.65               |

VCR, vincristine.

PPL suppresses transplanted E0771 tumors. Tumors were formed by injecting E0771 cells into mammary fat pads of C57Bl/6N mice. (A) E0771 tumor-bearing mice were treated with PPL for 7 consecutive days, and their tumor volumes were measured. (B) Kaplan–Meier survival curves for E0771 tumor-bearing mice treated with PPL. Control group treated with methyl cellulose by oral administration. Survival end point was set when tumor volume = 1,500 mm<sup>3</sup>. Survival curves of individual groups were compared by log-rank tests. (C) Body weights for E0771 tumor-bearing mice treated with PPL. Error bars: SE, *p<0.05, **p<0.01.
As PPL increased cytotoxicity of anticancer agents in vitro, we evaluated the effect of PPL and CBDCA on transplanted E0771 tumors. As shown in Fig. 5A, CBDCA decreased tumor growth, but the combination of PPL + CBDCA inhibited tumor growth to a greater extent (Day 22, CBDCA: 657.5 ± 194.4 mm³, PPL + CBDCA: 312.2 ± 62.5 mm³). The combination group also had longer median survival time (Fig. 5B, Table 4). In addition, immunohistochemical analysis showed that PPL increased tumor cell apoptosis to approximately 157% compared with controls—similar to CBDCA (approximately 149%)—and the combination of PPL + CBDCA induced the highest rate of tumor cell apoptosis to approximately 238% (Fig. 5C and D). These results suggest that PPL exerts antitumor effects and sensitizes cancer cells toward chemotherapy in this model.

**Table 4.** P values for survival rates of each treatment group

| vs                  | p value |
|---------------------|---------|
| Control vs PPL      | 0.0008  |
| Control vs CBDCA    | 0.0267  |
| Control vs PPL + CBDCA | 0.0004 |
| PPL vs CBDCA        | 0.8969  |
| PPL vs PPL + CBDCA  | 0.0067  |
| CBDCA vs PPL + CBDCA| 0.0193  |

CBDCA, carboplatin.
Discussion

Polyphenols are micronutrients found in various vegetables and fruit. Some polyphenols can reportedly affect degenerative diseases such as cancer and cardiovascular diseases. In this study, PPL showed cytotoxicity and cell death by removing ROS induced by CBDCA or VCR. It is well documented that inhibition of PI3K/Akt pathway enhances chemosensitivity. Therefore, the increase in cytotoxicity of CBDCA and VCR from PPL is thought to depend more on Akt dephosphorylation than on the removal of intracellular ROS. In the E0771 tumor model, tumor volume was significantly decreased under PPL treatment, but body weight was not affected (Fig. 4). In addition, as shown in Fig. 5, PPL enhanced the antitumor effect of CBDCA by inducing apoptosis. Therefore, PPL could be a safe anticancer and chemo-sensitizing agent.

Our data provide evidences for the antitumor effects of PPL in BC, and for its sensitizing effect for cancer drugs, which suggests that PPL could be a useful complement to cancer treatment. Some caffeic acid derivatives reported to exhibit antioxidant and antitumor activity and to enhance cytotoxicity of antitumor agents. However, the effect of caffeic acid derivatives contained in PPL is not clear. Therefore, further studies are needed to determine the candidate compound of Akt inhibition, and their possible clinical applicability.

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Conflict of Interest

PPL was manufactured by Genuine R&D Co., Ltd. (Masakatsu Miyanabe, Shimohu Hiraki, Xiaolin Luo). Other authors declare no conflict of interest.

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