Penta-O-galloyl-β-D-glucose induces G₁ arrest and DNA replicative S-phase arrest independently of P21 cyclin-dependent kinase inhibitor 1A, P27 cyclin-dependent kinase inhibitor 1B and P53 in human breast cancer cells and is orally active against triple-negative xenograft growth

Yubo Chai¹, Hyo-Jeong Lee²†, Ahmad Ali Shaik¹,³, Katai Nkhata¹, Chengguo Xing³, Jinhui Zhang¹, Soo-Jin Jeong², Sung-Hoon Kim¹,²*, Junxuan Lü*†

Abstract

Introduction: Natural herbal compounds with novel actions different from existing breast cancer (BCa) treatment modalities are attractive for improving therapeutic efficacy and safety. We have recently shown that penta-1,2,3,4,6-O-galloyl-β-D-glucose (PGG) induced S-phase arrest in prostate cancer (PCa) cells through inhibiting DNA replicative synthesis and G₁ arrest, in addition to inducing cell death at higher levels of exposure. We and others have shown that PGG through intraperitoneal (i.p.) injection exerts a strong in vivo growth suppression of human PCa xenograft models in athymic nude mice. This study aims to test the hypothesis that the novel targeting actions of PGG are applicable to BCa cells, especially those lacking proven drugable targets.

Methods: Mono-layer cell culture models of p53-wild type estrogen receptor (ER)-dependent MCF-7 BCa cells and p53-mutant ER-/progesterone receptor (PR)- and Her2-regular (triple-negative) MDA-MB-231 BCa were exposed to PGG for a comprehensive investigation of cellular consequences and molecular targets/mediators. To test the in vivo efficacy, female athymic mice inoculated with MDA-MB-231 xenograft were treated with 20 mg PGG/kg body weight by daily gavage starting 4 days after cancer cell inoculation.

Results: Exposure to PGG induced S-phase arrest in both cell lines as indicated by the lack of 5-bromo2'-deoxy-uridine (BrdU) incorporation into S-phase cells as well as G₁ arrest. Higher levels of PGG induced more caspase-mediated apoptosis in MCF-7, in strong association with induction of P53 Ser¹⁵ phosphorylation, than in MDA-MB-231 cells. The cell cycle arrests were achieved without an induction of cyclin dependent kinase (CDK) inhibitory proteins P21cip¹ and P27kip¹. PGG treatment led to decreased cyclin D1 in both cell lines and over-expressing cyclin D1 attenuated G₁ arrest and hastened S arrest. In serum-starvation synchronized MCF-7 cells, down-regulation of cyclin D1 was associated with de-phosphorylation of retinoblastoma (Rb) protein by PGG shortly before G₁-S transition. In vivo, oral administration of PGG led to a greater than 60% inhibition of MDA-MB231 xenograft growth without adverse effect on host body weight.

Conclusions: Our in vitro and in vivo data support PGG as a potential drug candidate for breast cancer with novel targeting actions, especially for a triple negative BCa xenograft model.
**Introduction**

Breast cancer (BCa) is the major cause of cancer-related deaths for women in the US [1] and other Western countries. Approximately 60% to 70% of BCa cases express estrogen receptors (ERs) or progesterone receptors (PRs) or both, and another approximately 20% of cases have amplified HER-2 proto-oncogene and express high levels of the HER-2 protein [2]. Approximately 15% to 20% of BCa cases are in the category of triple-negative phenotype because of their lack of ER and PR and do not have amplification of HER-2 [2,3]. These patients have a very poor prognosis because, unlike the situation for other types of BCa, there is no clinically validated, molecularly targeted therapy. When surgical and radiation options are no longer applicable to these triple-negative patients, treatment with available cytotoxic and genotoxic chemotherapy drugs produces limited efficacy and significant side effects. There remains a strong and urgent need for safer anti-cancer compounds for the treatment/management of the triple-negative BCa and its metastasis. Novel agents with multiple targeting ability distinct from the known drugable targets could be useful for circumventing the limitations of current treatment options.

Penta-1,2,3,4,6-O-galloyl-β-D-glucose (PGG) is a naturally occurring gallotannin polyphenolic compound in Oriental herbs such as *Galla Rhois*, the gallnut of *Rhus chinensis* Mill, and the root of peony *Paeonia suffruticosa* Andrews [4]. A couple of earlier papers have examined the *in vitro* effects of PGG while using an ER+ estrogen-dependent and p53-wild-type MCF-7 BCa cell culture model [5,6]. Chen and colleagues [5] reported that PGG induced G1 arrest in association with upregulated abundance of cyclin-dependent kinase inhibitor (CDKI) proteins 1A (p21Cip1) and 1B (p27Kip1). Later, the same group showed that PGG decreased ERα and the HER family of membrane tyrosine kinase (EGFR, HER-2, and HER-3) and PI3K/AKT signaling in MCF-7 cells [6]. A close inspection of the experimental designs of these studies revealed a lack of critical time-matched controls, and therefore the conclusions and the validity of the mechanistic work reported are questionable.

In cell culture studies, we recently showed that PGG induces caspase-mediated apoptosis in the human LNCaP prostate cancer (PCa) cells that express wild-type p53 [7]. The caspase-mediated apoptosis induction by PGG was mediated in large part by activation of p53 as established through siRNA (small interference RNA) knockdown and dominant negative mutant approaches [7]. More recently, we showed the induction of cell death with autophagic features (e.g., autophagosome formation and addition of a phosphatidylinositolamine moiety to the microtubule-associated protein 1 light chain 3 [LC3] to a faster moving LC3-II form on electrophoresis) by PGG of p53-null, PTEN-null, (high AKT) PC-3 PCa cells, which did not undergo caspase-mediated apoptosis after exposure to PGG [8]. We have also investigated the cell cycle effects of PGG in these and other PCa cells [9]. We showed for the first time that, irrespective of the p53 and androgen dependence status of the PCa cell lines, PGG exerted a rapid (within 2 hours) and potent (IC50 [half inhibitory concentration] of approximately 6 μM) inhibition of 5-bromo-2′-deoxyuridine (BrdU) incorporation into S-phase cells. In isolated nuclei, PGG inhibited DNA replicative synthesis with an efficacy superior to that of a known DNA polymerase-alpha inhibitor, aphidicolin. We have found, in addition to the S-arrest action, a close association of downregulation of cyclin D1 with G1 arrest induced by PGG. Taken together, our data with PCa cells indicate that PGG induced S arrest, probably through DNA replicative blockage, and induced G1 arrest via cyclin D1 downregulation to contribute to its anti-cancer activity. These results sharply contrasted with the questionable BCa cell culture studies mentioned above [5,6]. Therefore, whether the S and G1 cell cycle arrests and caspase-mediated or autophagic cell death actions of PGG are applicable to BCa cells needs to be experimentally tested.

In addition to PCa cells, PGG was shown to induce G1 cell cycle arrest and apoptosis of leukemia [10,11] to inhibit invasion-related molecules such as matrix metalloprotease-9 in melanoma cells [12] and EGFR signaling [13] and VEGFR2 signaling and angiogenesis *in vitro* and *in vivo* [14], supporting multiple targeting actions. A number of *in vivo* studies by us and others in Lewis lung cancer allograft [14] and PCa xenograft [7,13] models with a dose of 20 or 25 mg/kg every other day have shown anti-cancer efficacy without adverse effect on body weight. These *in vivo* and *in vitro* studies suggest probable anti-cancer activity of PGG against BCa, especially triple-negative BCa. In this report, we evaluated the cell cycle and cell death actions of PGG against MDA-MB231 triple-negative BCa cells and MCF-7 BCa cells, and we established, for the first time, an impressive oral efficacy of PGG against xenograft growth established from human MDA-MB231 cells.

**Materials and methods**

**Chemicals and reagents**

PGG was prepared in-house by methanolysis of tannic acid in accordance with a published method [4,15]. The purity was approximately 99%. For treatment of cells in mono-layer culture, PGG was dissolved in dimethyl sulfoxide (DMSO) as a stock solution. The final DMSO added to cell culture medium was below 0.1%. Anti-
dies, including anti-CDK4, anti-P21\textsuperscript{Cip1}, anti-P27\textsuperscript{kip1}, and anti-ER\alpha, were purchased from Santa Cruz Biotechnology, Inc. (Santa Cruz, CA, USA). Additional antibodies specific for cleaved poly-ADP-ribose polymerase (cPARP) (p89), cyclin D1, p53-Ser\textsuperscript{15P}, pRb-Ser\textsuperscript{795}, pRb-Ser\textsuperscript{807/811}, and pAKT(Ser\textsuperscript{473}) were purchased from Cell Signaling Technology, Inc. (Danvers, MA, USA). Antibody for LC-3 was purchased from MBL International (Woburn, MA, USA).

**Cell culture and treatments**

MCF-7 and MDA-MB231 cell lines were purchased from the American Type Culture Collection (Manassas, VA, USA). No cell line was derived directly from human tumor tissue for the purposes of this study. MCF-7 cells were grown in RPMI-1640 medium supplemented with 10% fetal bovine serum (FBS) without antibiotics in an incubator at 37°C with 5% CO\textsubscript{2}. MDA-MB231 cells were grown in L-15 medium supplemented with 10% FBS without antibiotics in an incubator at 37°C with atmospheric CO\textsubscript{2}. At 24 hours after plating, the medium was changed before starting the treatment with PGG or the other agents. To standardize all PGG/drug exposure conditions, cells were bathed in culture medium at a volume-to-surface area ratio of 0.2 mL/cm\textsuperscript{2} (for example, 15 mL for a T75 flask and 5 mL for a T25 flask).

**Cell growth assay by crystal violet staining**

For the evaluation of the overall inhibitory effect of PGG on cell number, the cells were treated with PGG daily (fresh medium) for 3 days. After treatment, the culture medium was removed and the cells were fixed in 1% glutaraldehyde solution in phosphate-buffered saline for 15 minutes. The fixed cells were stained with 0.02% aqueous solution of crystal violet for 30 minutes. After the washing with phosphate-buffered saline, the stained cells were solubilized with 70% ethanol. The absorbance at 570 nm with the reference filter at 405 nm was evaluated using a microplate reader (Beckman Coulter, Inc., Brea, CA, USA).

**BrdU incorporation and cell cycle measurement**

The protocol was based on our previous publications [9,16]. After the desired experimental treatments, 10 μL of BrdU (9 mg/mL) solution was added to 5 mL of medium for 30 minutes before harvesting the cells. The cells were collected by trypsinization, centrifuged at 1,600g for 6 minutes, fixed with 70% ethanol overnight, and analyzed for cell cycle distribution by propidium iodide/BrdU bivariate flow cytometry.

**Synchronic MCF-7 cell G0/G1 progression model**

MCF-7 cells were seeded in RPMI-1640 medium supplemented with 10% FBS without antibiotics in an incubator at 37°C with 5% CO\textsubscript{2}. Twenty-four hours later, the cells were washed with serum-free phenol-red-free RPMI-1640 medium and then incubated in serum-free phenol-red-free medium for another 24 hours. One flask of cells was reserved as 0-hours baseline control. For the other flasks, serum-free medium was replaced with complete medium (10% FBS) to release cells from G\textsubscript{0} arrest. At selected time points, cells were harvested for flow cytometry to analyze cell cycle distribution. For PGG treatment, cells were released into complete medium and treated simultaneously with serum stimulation or were exposed to PGG after different time periods of G\textsubscript{1} progression until 24 hours, when the cells were collected for cell cycle analyses.

**Immunoblot analyses**

The cell lysate was prepared in ice-cold lysis buffer as described previously [17]. Immunoblot analyses were essentially as described [17], except that the signals were detected by enhanced chemifluorescence with a Storm 840 scanner (Molecular Dynamics, now part of GE Healthcare, Little Chalfont, Buckinghamshire, UK).

**MDA-MB231 xenograft model**

The animal use protocol was approved by the Kyung Hee University Institutional Animal Care and Use Committee and carried out at the Cancer Preventive Material Development Research Center, College of Oriental Medicine, Kyung Hee University (Seoul, South Korea). One million MDA-MB231 cells were mixed with 50% Matrigel (Becton, Dickinson and Company, Franklin Lakes, NJ, USA) and injected (in 100 μL) subcutaneously into the right flank of each 6-week-old female BALB/c athymic nude mouse (NARA Biotech, Deajon, South Korea). Starting 4 days after inoculation, 10 mice per group were given a daily gavage treatment of 2% Tween-80 (vehicle) or 20 mg of PGG per kg body weight. The dosage was based on our PCa xenograft work [7] and lung cancer allograft work [14]. Tumors were measured twice per week with a caliper, and tumor volume was calculated using the following formula: 1/2 \(w_1 \times w_2 \times w_2\), where \(w_1\) represents the larger tumor diameter and \(w_2\) represents the smaller tumor diameter.

**Statistical analyses**

Numerical data were expressed as mean ± standard error or standard deviation (where noted). Statistical analyses were carried out with GraphPad Prism (GraphPad Software, Inc., San Diego, CA, USA) software, and a \(P\) value of less than 0.05 was considered statistically significant. The data were analyzed by analysis of variance followed by Dunnett’s multiple comparison post tests or other appropriate tests.
Results
PGG inhibited MCF-7 and MDA-MB231 breast cancer cell growth and induced caspase-mediated and caspase-independent cell death

To evaluate the growth inhibitory effect on estrogen-dependent BCa cell line MCF-7 and the triple-negative BCa cell line MDA-MB231, we exposed these cells to daily changes of fresh complete medium with increasing concentrations of PGG. After 3 days of daily exposure, PGG decreased the number of MCF-7 and MDA-MB231 cells in a dose-dependent manner, and the IC₅₀ of PGG for both cell lines was lower than 12.5 µM (Figure 1a). The p53-wild-type MCF-7 cells were more sensitive than the p53-mutant triple-negative MDA-MB231 cells to the growth suppression action of PGG at each tested concentration of PGG.

The difference in sensitivity of the two BCa cell lines was associated in part with the propensity for MCF-7 cells to undergo caspase-mediated apoptosis preceded by P53-Ser¹⁵ phosphorylation (P53-Ser¹⁵P) (24 hours) and cleavage of PARP (48 hours) (Figure 1b). In the MDA-MB231 cells, whose mutant P53 phosphorylation was not responsive to PGG treatment, minimal cleavage of PARP (72 hours) was preceded by autophagic features as indicated morphologically by cytosolic vacuolation (48 hours); biochemically by an early (6 hours) increase of phosphorylation of AMPK (AMP-activated protein kinase), a well-known autophagy signaling kinase in response to nutrient deprivation [18]; and by increased phosphatidylethanolamine modification of the microtubule-associated protein 1 light chain 3 (LC3-I) to the faster moving LC3-II form (24 to 48 hours) (Figure 1b). These results therefore mirror our data (on PGG induction of apoptosis and other types of cell death) obtained with LNCaP [7] and PC-3 PCa [8] cells, respectively.

PGG induced S and G₁ arrests in MCF-7 and MDA-MB231 cells

Prompted by our findings of G₁ and S arrests in different PCa cell lines (LNCaP, DU-145, and PC-3) [8,9], which contrasted with the reported G₁ arrest in MCF-7 cells [5], we measured the cell cycle distribution patterns of MCF-7 and MDA-MB231 cells after exposure in complete medium to different concentrations of PGG for 6, 24, and 48 hours.

In both MCF-7 and MDA-MB231 cells, PGG exposure for 6 hours led to a concentration-dependent increase of G₁-phase cells and was accompanied by a decrease of G₂-phase cells (Figure 2). Probably owing to their faster growth, the MB231 cells (Figure 2b) appeared to more readily achieve G₁ arrest than the MCF-7 cells (Figure 2a) in the presence of the lowest PGG concentration tested. The percentage of S-phase cells remained relatively steady in both cell lines. Inspection of the BrdU incorporation index (measured, as we previously described, for 30 minutes of pulse labeling before cell harvest [9]) showed a near-complete blockage of DNA synthesis in the S-phase cells in MB231 cells at all three PGG exposure concentrations, whereas in the MCF-7 cells, a clear concentration dependency on PGG was observed.

For both cell lines, as time progressed to 24 and 48 hours, the lowest concentration of PGG (12.5 µM) was not able to hold the cells arrested in G₁, manifesting as the accumulation of S-phase cells that remained incapable of incorporating BrdU. The higher concentrations of PGG (50 µM) kept cells arrested in G₁ phase and S phase throughout the 6- to 48-hour period (Figure 2). The data therefore support both S arrest and G₁ arrest by PGG in BCa cells, as in PCa cells [9].

PGG did not alter P21Cip¹ and P27Kip¹ expression in breast cancer cells

An earlier report by Chen and colleagues [5] has claimed G₁ arrest and P21Cip¹ and P27Kip¹ induction by PGG in MCF-7 cells, without including critical time-matched controls. We therefore examined these proteins as possible molecular mediators for the G₁ and S arrests. Since we have reported the rapid P53-Ser¹⁵P by PGG treatment in LNCaP PCa cells [7] and have observed P53-Ser¹⁵P in PGG-exposed MCF-7 cells (Figure 1b) and since the P53-P21Cip¹ axis is best known for mediating G₁ arrest by genotoxic stress [19], we focused on the relationship among these proteins in PGG-exposed MCF-7 cells.

We observed that PGG treatment activated P53-Ser¹⁵P at 6 hours with a clear concentration dependency but did not increase the protein abundance of either P21Cip¹ or P27Kip¹ (Figure 3a). Later, we found the same pattern of disengaged P53/P21Cip¹ response (that is, P53-Ser¹⁵P but not upregulated P21Cip¹) in the synchronic MCF-7 model (Figure 4). In MDA-MB231 cells, we did not observe any induction of these two CDK1i proteins by PGG (Figure 3b). These results suggest that PGG induced G₁ arrest in the absence of detectable alterations of P21Cip¹ and P27Kip¹ protein abundance and was independent of P53 function in these BCa cells.

PGG decreased cyclin D¹ abundance in breast cancer cells

In contrast to a lack of expression change of P21Cip¹ or P27Kip¹, PGG treatment significantly decreased the abundance of cyclin D¹ in MCF-7 and MDA-MB231 cells (Figure 3a, c). PGG treatment decreased cyclin D¹ expression as early as 6 hours, and by 12 hours, its expression decreased dramatically. From 24 to 48 hours, there was almost no detectable cyclin D¹ expression in
MCF-7 and MDA-MB231 cells treated with PGG at a high dose (Figure 3a, c).

To test the contribution of cyclin D1 downregulation to the G1 arrest, we made stable transfectants of MCF-7 and MDA-MB231 cells with forced overexpression of cyclin D1 (Figure 5a) (the expression plasmid was kindly provided by Joshua D Liao, The Hormel Institute, Austin, MN, USA). Compared with vector transfectant cells, the cyclin D1-overexpressing MCF-7 cells significantly attenuated PGG-induced G1 arrest (Figure 5b). Similarly,
MDA-MB231 cells overexpressing cyclin D1 partially overcame PGG-induced G\textsubscript{1} arrest (Figure 5c). Instead, PGG exposure of the cyclin D1-overexpressing cells hastened S arrest, without affecting G\textsubscript{2}-phase decline. The data suggest that G\textsubscript{1} arrest and S arrest operate by independent mechanisms in BCa cells, as in PCa cells [9], and that cyclin D1 downregulation by PGG was an important contributor (but perhaps not the sole mediator) to G\textsubscript{1} arrest.

**Defining G\textsubscript{1}-targeting action of PGG in a synchronic MCF-7 model**

To further probe the G\textsubscript{1}-targeting mechanisms of action of PGG without the complication from S arrest, we synchronized MCF-7 cells to G\textsubscript{0} by serum starvation for 24 hours and released the cells into complete medium (this time point was referred to as 0 hours). Cell cycle distribution patterns suggested that the G\textsubscript{1}-phase cells started to transit into S phase between 20 and 22 hours of FBS re-stimulation (Figure 6a).

In this synchronic MCF-7 cell model, inclusion of PGG at the time of serum stimulation (PGG at 0 hours) caused a complete block of G\textsubscript{1}-to-S transition, measured by flow cytometry at 24 hours (Figure 6b). To determine whether the presence of PGG during the early stage G\textsubscript{0}/S progression was necessary for G\textsubscript{1} arrest and to pinpoint the responsible molecular events, we delayed the starting time for PGG exposure in reference to serum

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**Figure 2** The effect of PGG on cell cycle distribution of MCF-7 (a) and MDA-MB231 (b) cells detected by propidium iodide/BrdU-bivariate flow cytometric analyses. Cells were exposed to increasing concentrations of PGG for 6, 24, and 48 hours. BrdU was added for the last 30 minutes to label S-phase cells active in DNA replication. Values are mean ± standard error of the mean (n = 4). Results are from two independent experiments with duplicate values at each concentration. The medium was not changed for PGG exposure of longer than 24 hours. Statistical significance: (a) BrdU incorporation at all three time points, one-way analysis of variance (ANOVA) \( P < 0.0001 \), with Dunnett’s multiple comparison post test \( P \) value of less than 0.01 for 0 versus 12.5, 25, and 50 \( \mu \)M PGG. For G\textsubscript{1}, 6 hours \( P < 0.05 \) for 0 versus 25 and 50 \( \mu \)M PGG; 24 hours \( P < 0.01 \) for 0 versus 12.5, 25, and 50 \( \mu \)M PGG; 48 hours \( P < 0.01 \) for 0 versus 12.5, 25, and 50 \( \mu \)M PGG. For S, 24 hours/48 hours \( P < 0.01 \) for 0 versus 12.5 and 25 \( \mu \)M PGG. (b) BrdU incorporation at all three time points, one-way ANOVA \( P < 0.0001 \), with Dunnett’s multiple comparison post test \( P \) value <0.01 for 0 versus 12.5, 25, and 50 \( \mu \)M PGG. For G\textsubscript{1}, 6 hours \( P < 0.05 \) for 0 versus 12.5, 25, and 50 \( \mu \)M PGG; 24 hours \( P < 0.01 \) for 0 versus 25 and 50 \( \mu \)M PGG; 48 hours \( P < 0.01 \) for 0 versus 50 \( \mu \)M PGG. For S, 24 hours/48 hours \( P < 0.01 \) for 0 versus 12.5 \( \mu \)M PGG. BrdU, 5-bromo-2’-deoxy-uridine; PGG, penta-O-galloyl-D-glucose.
stimulation. As shown in Figure 6b, delaying the starting exposure time to 14 hours (that is, PGG@14 hours) did not lessen the G1-arrest action of PGG. Starting PGG treatment at 16 to 18 hours was less able to prevent G1-to-S transition. These data indicated that the crucial time window for PGG targeting during G1/S progression was 16 to 18 hours after serum stimulation.

PGG decreased cyclin D1 in synchrony with retinoblastoma de-phosphorylation in synchronous MCF-7 cells

In the synchronous MCF-7 model, serum stimulation led to increased cyclin D1 expression (4 hours was the earliest point sampled), which persisted through 20 hours (G1/S transit) (Figure 4). Serum stimulation increased survival signaling, as indicated by AKT phosphorylation, in a temporal pattern similar to that of cyclin D1 and suppressed background level apoptosis as indicated by the decreased cPARP. Serum stimulation decreased ERα, which declined progressively over time. A well-known downstream effector molecule of cyclin-CDK complexes for G1 progression is the retinoblastoma (Rb) protein [20]. Cyclin-CDK complexes phosphorylate Rb to decrease its binding to the E2F transcriptional factor, releasing E2F to activate expression of its target genes for G1/S transition. Indeed, we detected increased Rb phosphorylation at 12 hours at the Ser795 site and 16 hours at Ser807/811 sites prior to the onset of G1/S transition (20 hours).

Exposure of synchronous MCF-7 cells to PGG at the time of serum stimulation did not decrease cyclin D1 until 16 hours, coinciding with decreased Rb phosphorylation at Ser795 and Ser807/811 sites (Figure 4). Although P53-Ser15P was detected by 8 hours of PGG treatment, there was a clear absence of P21Cip1 induction by PGG throughout 20 hours. Increased cPARP was detected by 16 hours, and this was preceded by increased AKT (Ser473) phosphorylation by several hours. PGG treatment did not affect ERα until the 20-hour time point. Given that P21Cip1 abundance was not upregulated
throughout the G₁ phase by PGG, the data suggest that the G₁ arrest was regulated predominantly by the cyclin D-CDK-Rb axis, preventing the release of E2F to promote the passage of the restriction point.

Orally administered PGG suppresses MDA-MB231 breast cancer xenograft growth

The cell culture data presented above suggest probable in vivo anti-cancer efficacy of PGG against BCa growth. Because oral administration is the most practical and non-invasive way to deliver an anti-cancer agent, we evaluated the efficacy of PGG delivered by oral gavage against MDA-MB231 cells injected subcutaneously into the right flank of each female athymic nude mouse at the dosage of 20 mg/kg body weight, starting 4 days after cancer cell inoculation. This dosage of PGG did not exert any adverse effect on body weight of the host nude mice (Figure 7a). PGG treatment led to a significant inhibition of tumor growth rate over time (Figure 7b) and decreased the final tumor size by over 60% at necropsy.

Discussion

As pointed out in the Introduction, there is an urgent clinical need for safe and effective treatment and preventive agents for triple-negative BCa. Our results presented above provide in vitro and in vivo data that support the potential for PGG to be such a promising drug candidate with multiple targeting actions, distinct from known drugable BCa targets such as the ER (for example, estrogen antagonist drug tamoxifen) and HER-2 (for example, inactivating monoclonal antibody herceptin). In cell culture, PGG treatment caused P53-Ser15 phosphorylation (Figures 1, 3, and 4) and caspase-mediated apoptosis (Figures 1 and 4) in MCF-7 BCa cells. In p53-mutant MDA-MB231 triple-negative BCa cells, PGG caused not only apoptosis but also autophagic responses (Figure 1b). We showed that independently of P53 status or ERα status of the BCa cells, PGG induced S arrest and G₁ arrest (Figure 2) without inducing P21Cip1 and P27Kip1 expression (Figures 3 and 4). Our data support cyclin D1 downregulation by PGG as an important mediating event for the G₁-arrest action (Figures 3 to 5). The clear disengagement of P53-Ser15 phosphorylation from the best-known P53 transcriptional target P21Cip1 in PGG-exposed MCF-7 cells remains an interesting question for further investigation.

Our findings are important in two respects. First, they are consistent with recently published results for PCa cells [7-9], suggestive of a treatment applicability of PGG for cancers of other organ sites. The documented ability in this study to generate high-purity PGG in multi-gram quantities from tannic acid will enable us and others to explore the in vivo anti-cancer efficacy of PGG in relevant animal models of cancers of other organ sites. Second, the findings point out the possibility that some published data are highly questionable concerning the action mechanisms of PGG in BCa cells. In contrast to the data published by others [5], our data did not detect a change of P21Cip1 and P27Kip1 expression to be associated with the G₁-arrest action of PGG (Figure 3). We also did not observe a dramatic impact of PGG on ERα abundance or a suppression of AKT phosphorylation (Figure 4), as were claimed [6]. Instead, PGG treatment increased AKT phosphorylation in...
MCF-7 cells (Figure 4), as we have reported for a similar increase of AKT phosphorylation in PC-3 cells by PGG [8]. Although many reasons could be cited for the discrepancies between our data and the previous reports [5,6], their lack of time-matched controls could be the leading cause of confusion and misleading conclusions.

Our in vivo data demonstrated, for the first time, a growth inhibitory efficacy of PGG against triple-negative BCa and supported the oral bioavailability of PGG. The potency of PGG (20 mg per kg body weight) is remarkable, especially considering that PGG was given by the oral route. Furthermore, just the fact that PGG is orally available and therefore can be self-administered by

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**Figure 6** Effect of PGG on G_{0/1}-S progression in synchronized MCF-7 cells. (a) The temporal kinetics of serum-stimulated progression of starvation-synchronized MCF-7 cells. Each time point was the average of duplicate flasks. *P < 0.05; **P < 0.01; ***P < 0.001 versus 0 time. (b) Impact of delaying PGG treatment with reference to serum stimulation on G_{1} arrest. Results are from two independent experiments with duplicate values at each time point. *P < 0.05; **P < 0.01 versus serum-free (SF) or PGG@0h-14 h. CM, complete medium; PGG, penta-O-galloyl-D-glucose.
patients will have a major impact on reducing the health-care delivery cost compared with injection-only drugs (such as paclitaxel) that have to be given by health-care professionals. The data on efficacy and safety of PGG provide an impetus for further studies about the therapeutic and chemopreventive targets and mechanisms of action.

Conclusions
Our cell culture data showed that PGG could induce both G1 and S arrests in BCa cells, regardless of their ER or P53 functional status. Cyclin D1 downregulation by PGG was a mechanism for G1 arrest in BCa cell lines, and the data ruled out P21Cip1 and P27Kip1 for mediating G1 arrest. We demonstrated for the first time that PGG given by oral administration was quite safe to the host nude mice and potent for suppressing a triple-negative BCa xenograft model. The therapeutic and chemopreventive utility of PGG for BCa merits further study.

Abbreviations
BCA: breast cancer; BRDU: 5-bromo-2'-deoxy-uridine; CDK: cyclin-dependent kinase; CDKI: cyclin-dependent kinase inhibitor; CPARP: cleaved poly-ADP-ribose polymerase; DMSO: dimethyl sulfoxide; ER: estrogen receptor; FBS: fetal bovine serum; IC50: half inhibitory concentration; LC3: microtubule-associated protein 1 light chain 3; PCA: prostate cancer; PGG: penta-O-galloyl-β-D-glucose; PR: progesterone receptor; P21Cip1: cyclin-dependent kinase inhibitor 1A; P27Kip1: cyclin-dependent kinase inhibitor 1B; RB: retinoblastoma.

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Author details
1The Hormel Institute, University of Minnesota, 801 16th Avenue NE, Austin, MN 55912, USA. 2Cancer Preventive Material Development Research Center and Institute, College of Oriental Medicine, Kyung Hee University, 1 Hoegi-dong, Dongdaemun-gu, Seoul 131-701, Republic of Korea. 3Department of Medicinal Chemistry, College of Pharmacy, University of Minnesota, 308 Harvard Street SE, Minneapolis, MN 55455, USA.

Authors’ contributions
JL conceived of and coordinated the studies, designed the experiments, and drafted and edited the manuscript. SHK helped to conceive of and coordinate the studies and to design the experiments. VC helped to perform cell culture experiments and statistical analyses, and to draft the manuscript. HJL helped to design the experiments, to carry out the xenograft study, and to draft the manuscript. JZ helped to design the experiments and to perform cell culture experiments. KN helped to perform cell culture experiments and statistical analyses. SU helped to carry out the xenograft study. AAS and CX scaled-up PGG preparation from tannic acid and performed chemical characterization. All authors read and approved the final manuscript.

Competing interests
The authors declare that they have no competing interests.

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