Serum harvested from heifers one month post-zeranol implantation stimulates MCF-7 breast cancer cell growth

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Received August 3, 2010; Accepted September 15, 2010

DOI: 10.3892/etm.2010.155

Abstract. Breast cancer is a serious disease in the US. Numerous risk factors have been linked to this disease. The safety of using growth promoters, such as zeranol, remains under debate due to the lack of sufficient in vitro and in vivo evidence. Using CellTiter 96® Aqueous Non-Radioactive Cell Proliferation assay, real-time PCR and Western blot analysis, we evaluated the effects of sera harvested from experimental and control heifers before and after one month of zeranol implantation on the growth of human breast cancer cell line MCF-7 as well as the involved mechanisms. We found that sera harvested from the heifers following one month of zeranol implantation were more mitogenically potent in stimulating the proliferation of MCF-7 cells when compared to sera harvested from the same heifers before zeranol implantation and the control heifers. Further investigation found that dextran-coated charcoal suppressed the stimulating effect of the sera on MCF-7 cell growth. The mechanisms involved in the MCF-7 cell proliferation stimulated by zeranol-containing sera may include up-regulation of cyclin D1 and down-regulation of p53 and p21 expression at the mRNA and protein levels in the cells. The results suggest that the consumption of beef products containing biologically active residues of zeranol or its metabolites is a risk linked to breast cancer development. Further investigation is required in order to clarify this critical issue.

Introduction

Breast cancer remains a serious health issue in the US. It is estimated that more than one-fourth of all cancer patients in 2009 were breast cancer patients, and breast cancer is the second leading cause of cancer-related mortality. According to clinical cancer statistics, approximately 182,500 new cases are diagnosed each year, with 40,500 patients succumbing to breast cancer in 2009 (1). Epidemiological studies suggest that there are many risk factors associated with breast cancer including age, relative body weight, the number and timing of reproductive events and lactation, exogenous and endogenous hormone concentrations, exposure to radiation, alcohol consumption and family history of breast cancer (2,3). More risk factors are still being discovered.

Zeranol (Z), which is marketed as Ralgro® (Merck-Schering-Plough Corp., Kenilworth, NJ, USA), is synthesized from the mycotoxin zearalenone and is a nonsteroidal agent with potent estrogenic activities. It has been widely used as a growth promoter in the US beef feedlot industry to accelerate weight gain, improve feed conversion efficiency and increase the lean meat-to-fat ratio (4,5). Previous investigations revealed that, although Z has adverse effects on the human breast, the use of Z as a growth promoter was safe for humans (6). However, the safety of using Z as a growth promoter has been debated ever since it was approved by the US Food and Drug Administration (FDA). Due to these health concerns, the European Union has banned imports of beef products from animals that have been administered any of the six growth-promoting hormones from the US. Subsequently, the US government challenged the ban with the World Trade Organization in early 1996. Recently, it has received increasing attention due to speculations concerning the possible etiological role of Z in breast cancer development. Zearalenone and Z are able to bind to the active site of human estrogen receptor (ER) α and ER β in a similar manner as 17 β-estradiol (E2) (7). Epidemiological investigation found that the sperm quality in sons of ‘high beef consumers’ was lower than that in males whose mothers ate less beef during their pregnancy (8).

It has been reported that a higher red meat intake in adolescence increases the risk of premenopausal breast cancer (9). Our laboratory previously reported that Z was able to transform human normal breast epithelial cells and increase human breast cell growth in a dose-dependent manner (10). Z has the ability to down-regulate estrogen-regulated human...
breast cancer candidate suppressor gene, protein tyrosine phosphatase γ expression (11). More recently, we found that the growth of pre-adipocytes derived from heifer ears implanted for two months with 72 mg Z was approximately 12-fold faster than that from the control heifers, and the response of the former cells to Z treatment was more sensitive compared to the latter. Following the investigation, the expression of cyclin D1 was up-regulated and p53 down-regulated in the former cells (12). Our preliminary data showed that 2.5% of Z-containing serum (ZS) harvested from heifers 60 days post Z implantation (72 mg) was capable of transforming human normal breast epithelial cell line MCF-10A to neoplastic breast cancer cells after a 21-day culture. Additionally, we showed that leptin, which plays a role in breast cancer development in obesity, induces human breast cancer epithelial cell sensitivity to Z (13). Therefore, it is crucial to clarify whether the consumption of beef products with residues of biologically active Z or its metabolites has any relationship with breast cancer.

The present study investigated the effects of biological samples directly harvested from heifers implanted with Z on the proliferation of the human breast cancer cell line MCF-7 as well as the underlying mechanisms. Our experimental results, for the first time, revealed that sera harvested from the heifers after one month of Z implantation significantly stimulated MCF-7 cell growth compared to sera harvested from the same heifers before Z implantation and the control heifers. The stimulatory effect on MCF-7 cells appears to be through the up-regulation of cyclin D1 and down-regulation of p53 and p21 expression at the mRNA and protein levels in MCF-7 cells. Further investigation in primary cultured human normal and cancerous breast epithelial cells is currently in progress in our laboratory. Our results suggest a potential risk of consuming beef products with biologically active Z or its metabolites in breast cancer initiation, promotion and progression.

Materials and methods

Animal treatment and blood sampling. The experimental design and sample collection method have been previously described (12). Briefly, 100 ml of blood was harvested from the Z-implanted heifers at day 0 (ZS-D0, prior to Z implantation), and day 30 post-Z implantation (ZS-D30) and from the control heifers at day 0 (NZS-D0) and day 30 (NZS-D30). The sera were immediately transferred to our laboratory and stored at 4°C overnight. After the clot from the sides of the tube was carefully loosened using a glass Pasteur, the serum was separated by centrifugation of the blood in a 50-ml centrifuge tube at 4000 rpm for 20 min at 4°C. The separated serum was sterilized through a 50-ml conical filter tube (Nalge Nunc International, New York, NY, USA) and stored at -20°C. Part of the sterilized ZS-D30 and NZS-D30 was treated with dextran-coated charcoal (DCC, dextran T-70, Sigma; charcoal, Sigma, USA) and stored at -20°C.

Cell culture. The MCF-7 cells were purchased from the American Type Culture Collection (ATCC, Manassas, VA, USA) and cultured in phenol red-free Dulbecco’s modified Eagle’s medium and Ham’s F12 medium (1:1) (DMEM/F12) (1.05 mM CaCl$_2$) containing 10% fetal bovine serum (FBS) and an antibiotic-antimycotic (100 U/ml penicillin G sodium, 100 µg/ml streptomycin sulfate and 0.25 µg/ml amphotericin B) (Invitrogen, Carlsbad, CA, USA) in a 75 cm$^2$ culture flask in a humidified incubator (5% CO$_2$, 95% air, 37°C). When the cells reached 85-90% confluence, they were subcultured in 75 cm$^2$ culture flasks at a ratio of 1:3 flasks as described above. The cells were dissociated using 3 ml of 0.25% trypsin-5.3 mM EDTA (Invitrogen) in PBS for 1 min at 37°C. The trypsinization was neutralized by the addition of 10 ml of culture medium with 10% FBS. The dissociated cells were collected and transferred to a 15-ml centrifuge tube and centrifuged at 1200 rpm for 5 min. The supernatant was discarded, and the cell pellets were resuspended in the culture medium with 10% FBS and subcultured in 75 cm$^2$ culture flasks.

Cell proliferation assay. The MCF-7 cells were seeded into 96-well plates at a density of 3,000 viable cells per well in 100 µl DMEM/F12 medium supplemented with 10% FBS and incubated at 37°C overnight. The medium was then replaced with 100 µl DMEM/F12 supplemented with 0.2% bovine serum albumin (BSA, Sigma, USA), and the plate was incubated at 37°C for a further 24 h. The cells were then treated with 0.5, 2.5 and 12.5% ZS-D0 and ZS-D30 or NZS-D0 and NZS-D30 in DMEM/F12 medium supplemented with 0.2% BSA for 24 h. The cells treated with DMEM/F12 medium supplemented with 0.2% BSA were the control groups. Treatment of ZS-D30 and NZS-D30 with or without DCC pre-treatment at 0, 0.02, 0.1, 0.5 and 2.5% in cultured medium was also carried out. Following 24 h of treatment, a cell proliferation (MTS) assay was performed as described in the manufacturer's protocol (Promega, Madison, WI, USA).

Cell treatment for RNA isolation and cDNA synthesis. A total of 10$^4$ viable MCF-7 cells/well were seeded in 6-well plates in 5 ml DMEM/F12 medium with 10% FBS and cultured overnight. The medium was replaced with DMEM/F12 supplemented with 10% DCC-treated FBS, and the cells were cultured overnight again. Then, MCF-7 cells were treated with 0, 0.2, 1 and 5% of ZS-D0, ZS-D30, NZS-D0 and NZS-D30 in culture medium for 24 h. After treatment, total RNA was isolated from each well using 1 ml TRIzol® reagent (Invitrogen) according to the manufacturer's instructions. RNA concentration was measured using an ND-1000 spectrophotometer (Thermo Fisher Scientific Inc., USA). RNA (1 µg) from each treatment group was reverse transcribed with 200 units M-MLV Reverse Transcriptase (Invitrogen) at 37°C for 50 min and then at 70°C for 15 min in the presence of 1 µl 10 mM dNTP (10 mM each dATP, dGTP, dCTP and dTTP at a neutral pH) (Invitrogen), 1 µl 50 µM random hexamer (Amersham Bioscience, Buckinghamshire, UK), 10 µl 5X First Strand buffer, 5 µl 0.1 M DTT and 1 µl RNase inhibitor (Invitrogen) in a total volume of 50 µl in a gradient master-cycle (Eppendorf®, USA).

Quantitative real-time PCR. Real-time PCR was conducted to measure cyclin D1, p53 and p21 expression. The PCR conditions were optimized for each primer pair and performed in Stratagene Mx3005p (Agilent Technologies, TX, USA).
Newly synthesized cDNA (2 µl) was used as a template for the reaction in a total volume of 20 µl reactants, which included 10 µl of 2X real-time PCR master mix (Applied Biosystems, Warrington, UK), 3 µl ultra-pure water and 5 µl of primer mixer. The reactants were first incubated at 95˚C for 10 min, and then 40 cycles of amplification were carried out with each cycle consisting of denaturing at 95˚C for 30 sec, annealing at 55˚C for 1 min and elongation at 72˚C for 1 min. A dissociation curve was created at the completion of PCR to ensure that the reaction produced the correct products as anticipated. The primer sequences for cyclin D1 were 5'-ttG GttacaGtaGcGtaG-3' (sense) and 5'- ttataGtaGcGtatcGtaGG-3' (antisense). The primer sequences for p53 were 5'-GacaatGGcaGcatctac-3' (sense) and 5'-GaaGGtGtaatcaGtctcc-3' (antisense). The primer sequences for p21 were 5'-GG aaGGaaGcaGGaaGac-3' (antisense) and 5'- aGcaGaGatacaaGGaaGG-3' (antisense). The primer sequences for 36B4 were 5'- acatGctcaacatctccc-3' (sense) and 5'-GcGGcacttctcctGctcc-3' (antisense). The results of the relative mRNA expression (cyclin D1, p53 and p21 to 36B4) in the MCF-7 cells were analyzed using the ΔΔCt method (14).

Western blot assay. The MCF-7 cells were separately seeded in a 100-mm dish (1x10^6 cells/dish) in DMEM/F12 medium supplemented with 10% FBS and cultured overnight. The medium was then replaced with DMEM/F12 supplemented with DCC-stripped 10% FBS. After being cultured for 24 h, the cells were treated with 0, 0.2, 1 and 5% of ZS-D30 and NZS-D30 in culture medium for 24 h. Protein extraction, measurement of concentration, separation and Western blot analysis were carried out as previously described (15). For immunoblotting, the following primary antibodies were used: rabbit polyclonal antibodies against cyclin D1, p53 and p21 (Cell Signaling Technology, Inc., Beverly, MA, USA) and goat polyclonal antibody against β-actin (Santa Cruz Biotechnology, Santa Cruz, CA, USA). The secondary antibody for cyclin D1, p53 and p21 detection was ECL™ anti-rabbit IgG linked to horseradish peroxidase (Amersham Biosciences, Buckinghamshire, UK) and a donkey anti-goat IgG HRP for β-actin detection (Santa Cruz Biotechnology). Images were captured using FujiFilm LA-3000 image system (FujiFilm Medical Systems USA, Inc., TX, USA). The densities of specific bands were quantified by ImageQuant software (Molecular Dynamics, Sunnyvale, CA, USA).

Results

ZS exhibits stronger stimulation of MCF-7 cell proliferation than NZS. In order to investigate whether ZS exhibits biological activity on MCF-7 cell growth, MCF-7 cells were treated with sera harvested from control and experimental heifers before and after Z implantation. Following 24 h of treatment, it was discovered that all the sera stimulated MCF-7 cell growth in a dose-dependent manner as compared to the control group (Fig. 1). No statistical difference was noted between NZS-D0 and NZS-D30 regarding their effects on MCF-7 cell proliferation at the same concentrations (Fig. 1A). However, a significant difference was found between ZS-D0 and ZS-D30 at the same concentration (Fig. 1B). ZS-D30 significantly stimulated the proliferation of MCF-7 cells compared to ZS-D0 at all doses. These data suggest that after one month of Z implantation, the serum contains certain biologically active components resulting in a more significant effect on the stimulation of MCF-7 cell proliferation.

DCC treatment suppressed the stimulatory effects of ZS-D30 on MCF-7 cell proliferation. In order to confirm whether ZS-D30 contains biological active components, ZS-D30 and NZS-D30 were treated with DCC, which has been reported to effectively strip hormones and growth factors from serum (16), and the effect on MCF-7 cell growth was evaluated. As shown in Fig. 2A, there was no significant difference in the proliferation of MCF-7 cells after treatment with NZS-D30.
stripped or not using DCC. However, the ZS-D30 not stripped by DCC significantly increased MCF-7 cell growth, compared to the MCF-7 cells treated with ZS-D30 stripped by DCC. These results revealed that ZS-D30 stripped by DCC removed the biologically active components thus suppressing the stimulatory effects of ZS-D30 on MCF-7 cell growth.

ZS-D30 up-regulates cyclin D1 and down-regulates p53 and p21 mRNA expression in MCF-7 cells. Previous results showed that zearanol treatment regulated the mRNA expression of cyclin D1 and p53 in pre-adipocytes derived from heifers two months after implantation of Z (12). Yuri et al found that treatment with 50 nM Z accelerated MCF-7 cell growth through down-regulated p21 expression in the cells (17). To explore the mechanisms involved in MCF-7 cell growth stimulated by ZS-D30, the mRNA expression of cyclin D1, p53 and p21 in MCF-7 cells after treatment with different concentrations of ZS-D30 as well as NZS-D30 was investigated using real-time PCR. Treatment with NZS-D30 for 24 h did not significantly alter the mRNA expression of cyclin D1, p53 or p21 in MCF-7 cells (Fig. 3). However, ZS-D30 treatment significantly up-regulated cyclin D1 and down-regulated p53 and p21 mRNA in a dose-dependent manner.

Discussion

Zeranol has been widely used as a growth promoter in certain countries, including the US and Canada, since it accelerates weight gain, improves feed conversion efficiency and increases the lean meat-to-fat ratio particularly in cattle. A previous investigation found that the use of Z in cattle was safe for human consumption (18). However, these results were generated using and based on outdated techniques. With the development of molecular biology and medicine, it was demonstrated that these previous findings may be inconclusive. This scenario has previously occurred with diethylstilbestrol (DES). DES is an estrogen that was approved by the US FDA in 1947 for the prevention of miscarriage and was widely prescribed by doctors in the USA until 1971 and in European countries until the late 1970s (19). Researchers also reported that DES generally increased the weight gain of cattle by 15% and feed conversion improvement was normally approximately 10% (20). Additionally, DES increases leanness (20). Due to its growth stimulation and improved feed utilization, orally administered DES for cattle was approved by the US FDA in 1954. Later, an investigation found that DES was not only associated with cancer in women to whom DES was prescribed during their pregnancy, but also in their daughters (21-23). These findings led the FDA to remove oral DES for cattle from the market in 1972 and implants the following year. A previous investigation among workers occupationally exposed to Z was conducted due to a variety of reported breast symptoms, including sharp pain, tingling, burning, aching and irritation. In addition, two former workers (one male, one female) had boys aged under 5 years who developed gynaecomastia and presented unusual growth spurts. These two boys were exposed to Z through their parents’ work clothes. The symptoms in the two boys abated after their parents changed work to control the Z exposure (6). The investigation clearly illustrated the relationship between breast symptoms and exposure to Z, particularly in children since they...
may be more sensitive to Z than adults. Human exposure to Z occurs through ingestion of beef products that contain pharmacological active residuals or its metabolites. Current research is raising concerns regarding the relationship between the consumption of beef products and adverse health risks associated with breast cancer development. Our previous data revealed that pre-adipocytes derived from heifers two months post-Z-implantation were sensitive to Z treatment. This prompted us to investigate whether biological samples such as serum and meat extracts harvested from the heifers after Z implantation have an effect on human breast cancer cell growth.

The present study revealed that the sera harvested from the heifers one month post-Z implantation significantly stimulated MCf-7 cell proliferation compared to the sera harvested from the same heifers prior to Z implantation as well as that from the control heifers. The results imply that the residues of biologically active Z or its metabolites contained in the sera exhibit a stimulatory effect on MCf-7 cells, and their activities can be suppressed by DCC treatment. The preliminary data also showed that the muscle extracts derived from the heifers one month post-Z implantation stimulated MCf-7 cell proliferation. We detected Z concentrations in the muscles, and the concentrations in sera will be detected using ELISA method once its antibody has developed. It is difficult to claim that the consumption of beef products with residues of biologically active Z or its metabolites is a risk factor in breast cancer. More evidence is required in order to clarify this crucial health issue. Further investigation of the impacts of biologically active Z and its metabolites on primary cultured human normal breast epithelial cells, stromal cells, pre-adipocytes and stem/progenitor cells is in progress in our laboratory.

Cell cycle regulation plays a very important role in mammary gland development and carcinogenesis. Numerous researchers have found that cyclin D1, p53 and p21 are associated with cell cycle regulation in breast cancer initiation, promotion and progression. Cyclin D1 is one of the most frequently overexpressed proteins and one of the most
commonly amplified genes in breast cancer (24,25). It is able to regulate the growth of estrogen-responsive tissues by activating the estrogen receptor in a ligand-independent manner (26). The tumor suppressor p53 is able to regulate cell proliferation and apoptosis. An imbalance between cell proliferation and apoptosis results in a rapid increase in cell numbers, the most prominent characteristic of tumors. Normal breast epithelial cells induce p53-dependent apoptosis and p53-independent cell cycle arrest of breast cancer cells (27). Tumor angiogenesis is considered a multi-pathway process, while p21 (WAF1/Cip1) is a cyclin-dependent kinase inhibitor involved in cell division and survival. It is activated by p53 and is a downstream effector for p53 function by inducing G1 arrest when normal breast cells are exposed to DNA-damaging agents (28). The expression of the p21 gene has been found to be regulated by estrogen in estrogen receptor-positive human breast cancer cells (29).

We demonstrated that ZS-D30 increased cyclin D1 and p16-nk4a expression in MCF-7 cells at the mRNA and protein levels, when compared to NZS-D30. This may partially explain why ZS-D30 significantly stimulated MCF-7 cell proliferation. Other mechanisms involved in cell cycle regulation may also exist, but further investigation is required to elucidate them. The data show that certain as yet undefined growth factors that are responsible for the stimulatory action of MCF-7 cell proliferation may be secreted into the blood circulation of heifers upon Z implantation. We attribute the stimulatory effect of ZS on MCF-7 cell growth to the implantation of Z.

In conclusion, sera directly harvested from heifers one month post-Z implantation exhibited a potent stimulatory effect on MCF-7 cell growth that was mediated through an increase in cyclin D1 and a decrease in p53 and p21 expression at the mRNA and protein levels. Our results, to a certain degree, suggest the association between the biological samples derived from heifers implanted with Z and the potential adverse health risk of breast cancer. We require further evidence to clarify this critical health issue.

Acknowledgements

We thank Dr Walter R. Threftall and his team members for collecting the biological samples from the control and experimental heifers. We also thank manager Martin Mussard and his team members for taking care of the heifers in the beef cattle barn located in the Ohio State university livestock Facilities. We extend our thanks to all the members for their support and cooperation. We also thank Dr Walter R. Threlfall and his team members for taking care of the beef cattle barn and collecting the biological samples from the control and experimental heifers. We also thank manager Martin Mussard and his team members for taking care of the heifers in the beef cattle barn located in the Ohio State university livestock Facilities. This research was supported by NIH Grant R01 ES015212.

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