Polyglycolic acid/chitosan glue and apoptosis of endometriotic cells

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Objective: To induce apoptosis of endometriotic cells of patients with endometriosis.

Design: To demonstrate that polyglycolic acid/chitosan glue directly inhibits cell proliferation by inducing apoptosis.

Setting: University hospital infertility center.

Patient(s): Twelve women who visited the center for infertility therapy.

Intervention(s): Polyglycolic acid/chitosan glue was applied into primary endometriotic cells; the manipulated cells were collected 1–4 days after polyglycolic acid/chitosan glue treatment.

Main Outcome Measure(s): Primary endometriotic cell cultures from eutopic endometriotic tissue were established. The effect of the novel biological glue, polyglycolic acid/chitosan glue A, on endometrial cells in vitro was examined. The different stages of apoptosis were analyzed using flow cytometry with fluorescein isothiocyanate conjugate (FITC)-annexin V and propidium iodide staining.

Result(s): The growth inhibitory effects of polyglycolic acid/chitosan glue A on endometrial cells were found to be dose-response and time dependent. Less than 15% viability was detected in cultures containing 2,000 μg of polyglycolic acid/chitosan glue A after 4 days of treatment. Induced apoptosis and caspase activity were revealed. The caspase-3 activity increased 2.2-fold with 4 days of culture with 2,000 μg of polyglycolic acid/chitosan glue A.

Conclusion(s): This is the first study to demonstrate that polyglycolic acid/chitosan glue directly inhibits cell proliferation by inducing apoptosis, thus suggesting that this new biological glue may be useful for endometriosis therapy.

Key Words: Endometriosis, polyglycolic acid, chitosan, apoptosis

Endometriosis is described as the presence of endometrial tissue in locations outside the endometrial (uterine) cavity. Endometriosis is commonly found in the cul-de-sac (behind the uterus), the rectovaginal septum (the tissue between the rectum and the vagina), on the surface of the rectum, the fallopian tubes, ovaries, the uterosacral ligaments, the bladder, and the pelvic sidewall (1, 2). Generally, endometriosis in the rectovaginal septum is more likely to penetrate deeply into the underlying structures. In most cases, endometriotic cells can be found in the fluid behind the uterus of patients during menstruation. None of the mentioned theories seems to explain all cases. Unfortunately, we do not know its causes. The estimated worldwide population that suffers from endometriosis varies wildly; at least 10% of the female population suffers from the disease, which makes endometriosis one of the most common diseases affecting women’s health (1–3).

Although there is no effective treatment for endometriosis, a variety of treatment options have been established. Several treatment goals include relief or reduction of pain, suppression of endometrial growth, safeguarding or restoration of fertility, and avoiding recurrence of the disease. Danazol (4), birth control pills, Zoladex, and Norplant have not proven effective as either primary or adjunctive therapy (combined with surgery) for endometriosis-related infertility. Medical treatments for endometriosis might cause inflammation; surgical correction has fewer side effects and is mainly used to reduce endometriosis-related pain.

Endometriosis and cancer are similar in several aspects such as cell invasion, unrestrained growth, development of new blood vessels, and a decrease in the number of cells undergoing apoptosis. Endometriotic cells were identified with lower susceptibility to apoptosis and more surviving genes (5). Surviving may antagonize caspase-3-mediated apoptosis and subsequently promote the development of endometriosis (6).

In this study, growth inhibition effects of a new type of biological glue composed of polyglycolic acid (PGA) on cultured endometrial cells were observed. This biological glue was designed and processed with two significant characteristics: biodegradability and cell growth inhibition (7, 8).
Two types of biodegradable biomaterials, chitosan and PGA polymer, are used as ingredients of the glue. By using PGA, we can easily manipulate the biodegradation rate, and chitosan inhibits the growth of tumor-like cells. Chitosan is a binary polyheterosaccharide of N-acetylglucosamine and glucosamine with a $\beta 1\rightarrow 4$ linkage. This polyheterosaccharide structurally resembles glycosaminoglycans consisting of long-chain, unbranched, repeating disaccharide units (9–11). Chitosan was previously demonstrated to have anticancer activity, immune-enhancing effects, wound healing, and antimicrobial activity (9–11). Chitosan-induced cell apoptosis was proposed by caspase-3 activation, which plays a critical role in the initiation and execution of apoptosis (12). The advantages of this glue are that it is nontoxic and biodegradable, and it induces apoptosis of tumor-like cells. These characteristics make it a promising biomaterial for the treatment of endometriosis. To the best of our knowledge, there is no information in the literature on how PGA/chitosan (P/C) glue affects endometrial cell growth, or on its application for the treatment of endometriosis.

**MATERIALS AND METHODS**

**Culture Media**

Cell culture medium consisted of M199 and F12 (1:1) supplemented with P (5 $\mu$g/mL, Sigma, St. Louis, MO), insulin-transferrin-sodium selenite medium supplement (ITS), 100 U penicillin, 0.1 mg of streptomycin, and amphotericin B (0.25 $\mu$g/mL). Collagenase (Sigma, St. Louis, MO) was dissolved as a 2-mg/mL solution in Dulbecco’s modified Eagle’s medium-high glucose (DMEM-H) medium (Life Technologies, Rockville, MD).

**Endometriotic Tissue and Preparation of Primary Endometriotic Cells**

Endometriotic tissue samples were obtained from patients undergoing laparoscopy for unexplained infertility with known endometriosis. Twelve tissue samples were collected at the Department of Obstetrics and Gynecology, Taipei Veterans General Hospital. Before collection of the endometriotic tissues, consent was obtained from each of the donors for analysis of possible molecular diagnosis and utilization. Tissue samples were obtained from the proliferative phase (days 5–14) from women who had received no hormonal therapy within the prior 30 days. Tissue samples (approximately 1–2 g) preserved in DMEM-H transfer medium were transported to the laboratory. The tissue was rinsed in Hanks’ balanced salt solution (HBSS) to remove blood and cell debris. After gentle centrifugation at 600 × g for 5 minutes, the tissue was collected and placed into a plastic tissue culture dish (Corning-Costar, Cambridge, MA). Then, the tissue was minced into 1-mm$^2$ fragments and digested with collagenase for 2.5 hours at 37°C. The digested tissue was vigorously pipetted and placed on a stacked sterile wire sieve assembly with a no. 100 wire cloth sieve, followed by a no. 400 wire cloth sieve (37 $\mu$m). Two types of endometrial cells, glandular epithelium and stroma, were collected by sequential sieves (13). Primary epithelial cells were cultured in P/C glue-coated culture plates or noncoated tissue culture polystyrene culture plates containing cell culture medium.

**Glue and Culture Plate Preparation**

In this study, we prepared two types of P/C glue. Type 1 was named P/C glue A, which was prepared by mixing a PGA solution (24% wt) and a chitosan solution (2% wt). The other was P/C glue B for which PGA/chitosan films were resolubilized in phosphate-buffered saline (PBS) solution for 1 week; the pH was adjusted using HBSS. Serial doses of P/C spray A were prepared by dilution with culture media. In the P/C glue A group, we sprayed glue A on the surface of a cell monolayer. In P/C glue B/A treatment, we sprayed P/C glue B for 30 minutes and then sequentially sprayed glue P/C A on the cell monolayer. Four types of P/C glue A (containing 2,000, 1,000, 600, and 100 $\mu$g/mL P/C glue A) were prepared and sprayed to the surface of the cell monolayer in this study.

**Growth Inhibition Assay**

Four types of P/C glue A (2,000, 1,000, 600, and 100 $\mu$g/mL) were prepared and subjected to cell culture. Aliquots of 2.5 × $10^5$ cells/mL were seeded in complete culture medium for 24 hours, and then cultured with four types of P/C glue A or P/C glue B/A. After 4 days of incubation, treated cells were washed with medium and then subjected to a growth inhibition assay using the cell proliferation assay kit of the WST-1 method (Chemicon International, Temecula, CA). This colorimetric quantification based on the cleavage of the tetrazolium salt, WST-1, by mitochondrial dehydrogenase measured the absorbance by a microplate reader-SpectraMax (Molecular Devices, Sunnyvale, CA) at a 450-nm wavelength. The reference wavelength was 600 nm.

**Morphological Analysis of P/C Glue-Treated Cells**

An aliquot of 2.5 × $10^5$ cells was seeded in complete culture medium for 24 hours, and then cultured with four types of P/C glue A. After 4 days of incubation, treated cells were washed with medium and then incubated with the fluorescent probes at 37°C, after which they were visualized using an Olympus Fluoview 2-laser confocal microscope (IX-70, Olympus Optical, Tokyo, Japan). Thus, a combination of nonyl acridine orange (NAO) (14) and propium iodine (PI) (Molecular Probes, Eugene, OR) (15, 16) was used for fluorescence imaging analysis of cell viability.

**DNA Fragmentation Analysis**

Cells were cultured under monitored conditions for 24 hours, after which the medium was replaced with fresh medium containing various amount of P/C glue A or glue B and cultured for 72 hours. Treated cells were scraped off and
prepared for DNA extraction. DNA fragmentation was measured using a Suicide-Track DNA ladder isolation kit (Calbiochem, Darmstadt, Germany). The DNA ladder fragments were separated by electrophoresis on a 1.5% agarose gel in 1× TAE buffer at 100 V for 1 hour and stained with ethidium bromide.

**Flow Cytometry**

A combination of fluorescein isothiocyanate conjugate (FITC)-annexin V and PI was also used for simultaneous flow cytometric analysis of the stages of cell apoptosis (15, 16). Cells treated with or without P/C glue A were harvested and washed with PBS. Aliquots of 1 × 10^6 cells were gently stained with FITC-annexin V and PI in the dark for 15 minutes at room temperature. After staining, cells were washed with PBS and applied to flow cytometric analysis. All analyses were performed by FACScan (Becton Dickson, San Jose, CA) equipped with a single 488-nm argon laser. A minimum of 30,000 cells per sample was analyzed. Debris was gated out based on light-scatter measurements. Data were acquired in the list mode, and the relative proportions of cells within different areas of the fluorescence profile were quantified using the LYSYS II software program (Becton Dickson, Franklin Lakes, NJ).

**Assessment of Caspase-3 Activity**

Caspase-3 activities of cell lysates were determined using the “EnzChek” Caspase-3 Activity Assay kit with the Z-DEVD-AMC substrate. The fluorescence was detected using a microplate reader-Spectra MAX (Molecular Devices) with an excitation wavelength of 355 nm and an emission wavelength of 460 nm. All fluorescence was normalized with the soluble protein amount of cell lysate. Protein concentrations were determined with a Bio-Rad Protein Assay reagent (Bio-Rad Laboratories, Hercules, CA). The relative ratio of caspase-3 activity was defined as the activity of treated cells divided by that of the untreated cells.

**RESULTS**

**Growth Inhibitory Effect of P/C Glue A**

The reduction in cell viability occurred in a dose-dependent manner after 96 hours of incubation with P/C glue A. Cell viability was 95% in cultures containing 100 μg/mL P/C glue A, and less than 15% in cultures containing 2,000 μg/mL of P/C glue A (Fig. 1A). In addition, the cell viability was 88% in cultures containing P/C glue B/A (treated with P/C glue B for 30 minutes and then treated with P/C glue A) (Fig. 1A). The growth inhibitory effect was reduced in cells incubated with P/C glue B/A. The P/C glue B was found to protect cultured cells from growth inhibition by P/C glue A.

**Induction of Cell Death by P/C Glue A**

Using two fluoresceins, NAO and PI, provided further verification of cell death. The NAO was used to stain the mitochondria of viable cells, whereas DNA staining by PI was used to identify dead cells. The staining pattern of viable cells was NAO-negative and PI-negative, whereas it was NAO-positive and PI-positive in apoptotic or necrotic cells (Fig. 2). The differential staining pattern of treated cell was obtained after 4 days of culture with tissue culture polysty-
rene, and 1,000, 1,500, or 2,000 μg/mL P/C glue A. The reduced NAO and increased PI staining patterns were revealed in the P/C glue A-treated cells in a dose–response manner. In Figure 2D, most of the cells were stained red, indicating that a large proportion of the endometriotic cells were in the late apoptotic stage after treatment with 2,000 μg/mL P/C glue A. Furthermore, DNA fragmentation was detected in endometrial cells exposed to 1,000 or 1,500 μg/mL P/C glue A for 4 days (data not shown).

Analysis of Apoptotic Cells by Flow Cytometry
The staining pattern of viable cells was annexin V-negative and PI-negative, whereas it was annexin V-positive and PI-negative in early apoptotic cells, and both were positive in apoptotic or necrotic cells. The percentages of early apoptotic fractions were 80%, 95%, and 82% with 4, 7, or 8 days of treatment, respectively. Late apoptosis or necrosis was only detectable in the group at 8 days of culture. Induction of apoptosis was reduced when cells were pretreated with P/C glue B (Fig. 3). Percentages of the early apoptotic fractions were 1% and 3% of cells treated with P/C glue B/A for 1 and 4 days, respectively. Cells treated with P/C glue B/A remained viable.

Caspase-3 Activities
We examined the affects of caspase-3 on treated endometriotic cells. After 4 days of treatment, caspase-3 activity had increased 1.3-fold in 1,000 μg P/C glue A-treated cells (Fig. 4, column 3) compared with cells in the control group (column 1), and 2.2-fold in cells treated with 2,000 μg P/C glue A (column 5). There was no significant difference in caspase-3 activity between cells treated with P/C glues B/A (column 7) and the control group. In columns 2, 4, 6, and 8, we added a caspase-3 inhibitor into cells before treatment as in columns 1, 3, 5, and 7. Activation of caspase-3 was diminished by supplementation with the caspase-3 inhibitor.

According to our data, P/C glue A possibly induced apoptosis through caspase-3 activation in endometriotic cells.

DISCUSSION
This study reports the novel biological activities of P/C glue. Recently there have been several studies showing that chitosan is widely used in medical and pharmaceutical fields because of its low toxicity and biological effects (17–20). The watersoluble chitosan oligomer (WSCO) appears to have anticancer
activity, immune-enhancing effects, wound healing, and antimicrobial activity (9–11). Furthermore, WSCO is able to inhibit proliferation of human leukemia (HL)-60 cells and induce differentiation of those cells. Previous studies showed that chitosan, as a cationic polymer, interacts with cell membranes, causing lysis of red blood cells and growth inhibition of melanoma cells (21–25). When chitosan is dissolved in solution, the amino groups in the N-deacetylated subunits confer a highly positive charge density. These amino groups have intrinsic pKa values of 6.5, and thus, they behave as a polyglycan at both acidic and neutral pH values (26). Chitosan binds mammalian cells by interacting with surface glycoproteins (27). Based on the characteristics of chitosan, we hypothesized that a possible interaction of the positive charge of chitosan with cell surface proteins leads to ionic changes in cell membranes. This interaction is important in maintaining cell integrity, related functions, and cell growth and differentiation (28).

In our study, we addressed the biological effects of blended glue. Its unique characteristics, such as nontoxicity,
Caspase-3 activities were determined in polyglycolic acid/chitosan (P/C) glue A-treated cells. In column 1, 1,000 µg/mL of P/C glue; column 2, 1,000 µg/mL of P/C glue A with a caspase-3 inhibitor; column 3, 1,500 µg/mL of P/C glue A; column 4, 1,500 µg/mL of P/C glue A with a caspase-3 inhibitor; column 5, 2,000 µg/mL of P/C glue A; column 6, 2,000 µg/mL of P/C glue A with a caspase-3 inhibitor; column 7, 2,000 µg/mL of P/C glue B/A, and column 8, 2,000 µg/mL P/C glue B/A with a caspase-3 inhibitor. Values shown are the mean (± SD) of three independent assays performed in triplicate.

In this study, we demonstrated that P/C glue offers growth inhibitory activity in endometriotic cells. This growth inhibition was due to induction of cell apoptosis by P/C glue A. The P/C glue A-treated cells showed nuclear fragmentation and chromatin condensation, which indicate cell apoptosis. DNA showed fragmentation after cells had been treated for 4 days with 2,000 µg of P/C glue A (data not shown). Nearly 80% of cells treated with 2,000 µg of P/C glue A for 4 days underwent apoptosis (Fig. 3). Recently many chemotherapeutic agents have been shown to activate apoptotic mechanisms, which lead to drug-induced cell death (31). Activation of caspases plays a crucial role in the biological events associated with apoptosis (32, 33). In particular, caspase-3 is reported to be the executioner, which can cleave the DEVD-substrate into a fluorescent product. As shown in Figure 4, caspase-3 activity in P/C glue-treated cells was significantly elevated. The caspase-3 activity in the 2,000-µg P/C glue-treated cells was increased approximately 2.2-fold compared to untreated cells. We suggest that the growth inhibitory activity of P/C glue A is by induction of apoptosis through caspase-3 activation. Caspase-3 activity was not changed in P/C glue B-treated cells. In addition, the growth inhibitory effect of P/C glue A was reduced in cells treated with a caspase-3 inhibitor, suggesting that caspase-3 activation is required for P/C glue A-induced apoptosis. The P/C glue B may adhere to the cell membrane and evoke a biological change in the cell membrane. This result suggests that P/C glue B might block the interactions of P/C glue A with cell membranes that induce apoptosis.

The P/C glue directly inhibits cell proliferation by inducing caspase-3-mediated apoptosis. Takimoto et al. (12) had previously demonstrated that chitosan induces apoptosis through caspase-3 activation, which leads to activation of caspase-8 detached from the “death domain,” the cytoplasmic sequence of death receptors. Given its molecular size and cationic property, chitosan might not directly enter into the cell, but rather interact with the cell membrane. When exposed to anticancer agents, the majority of endometriosis cells appear to express death receptors such as Fas-Fas ligand (FasL), tumor necrosis factor receptor-1 (TNF-R1), and TNF-related apoptosis-inducing ligand (TRAIL) receptors, which are activated by or participate in the death program (12, 22). Chitosan may induce cell death through an interaction with death receptors and sequential activation of the caspase cascade. Moreover, clinically relevant concentrations of diverse anticancer agents such as cisplatin, doxorubicin, mitomycin, and camptothecin, have been shown to induce Fas/FasL expression in a number of cancer cell lines and endometriotic cells (33–35). It is possible that many endometriotic cells not only become resistant to Fas-mediated apoptosis, but also acquire the ability to use this pathway to their advantage by launching a “Fas counter-attack” against the host’s immune system (36). Because P/C modulates a death receptor expression and promote endometriotic cells to loss resistant to Fas–caspase-8–caspase-3-mediated apoptosis.

In our previous study (30), a mixture of PGA and chitosan was prepared using solvents of low toxicity to create novel, biocompatible, degradable, and modifiable hybrid materials. Fibroblast cells cultivated in these materials proliferated. In this study, we also monitored the pH changes during the P/C glue degradation and examined effects caused from the degradable products. We also checked the telomerase activity and Ki-67 level in the P/C glue A- or P/C glue A/B-treated endometriotic cells. None of the cytotoxicity and tumorigenicity was found in the P/C glue-manipulated cells (data not shown).

In the future, studies should be conducted to clarify the second messenger to the apoptosis pathway activated by P/C.
Fertility and Sterility

REFERENCES