Chitin increases the angiogenesis in chorioallantoic membrane model in the presence of testosterone and progesterone

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ABSTRACT

Aims: Angiogenesis is a process of generating new blood vessels from preexisting vessels. In adults, it is activated in pathologic conditions. Chitin is an organic molecule which is used in scaffold technology in tissue engineering. Growth hormones such as testosterone and progesterone are used in scaffold structure for induction of angiogenesis. No literature was found about the angiogenic roles of chitin/testosterone/progesterone. Here, chitin was analyzed in the presence of testosterone/progesterone to find out its possible role on angiogenesis.

Methods: Chitin was obtained from shrimp shells in our laboratory. The angiogenic effects of chitin/testosterone/progesterone were analyzed on chick embryo chorioallantoic membrane (CAM) model. Six different study groups were prepared (control group-Group 1, testosterone applied group-Group 2, progesterone applied group-Group 3, chitin/testosterone applied group-Group 4, chitin/progesterone applied group-Group 5, chitin/testosterone/progesterone applied group-Group 6). Hormones were used in different concentrations. The angiogenic role of selected molecules was clarified according to the total differentiation score of angiogenesis (TDSA) results in all groups. In obtaining of TDSA results, Knighton’s protocol was applied.

Results: TDSA was 6±0.1 in testosterone applied group, 5±0.2 in progesterone applied group, 7±0.1 in chitin/testosterone applied group, 5±0.1 in chitin/progesterone applied group, 7±0.1 in chitin/testosterone/progesterone applied group. In all groups, TDSA results were statistically significant. These results represented the angiogenic role of chitin in the presence of testosterone and progesterone (p<0.05).

Conclusions: Our results support the angiogenic roles of chitin in the presence of testosterone and progesterone. Chitin, testosterone and progesterone can be used in scaffold technology together.

Introduction

Angiogenesis (capillary vascularization) is an interesting process which new blood vessels form from pre-existing vessels. It physiologically occurs only in embryonal period. In adult period, it occurs in pathologic conditions like cancer and wound healing. Because of its positive role in wound healing process, angiogenic factors are used in tissue engineering studies. Different hormones such as testosterone and progesterone are used as angiogenic factors in scaffold technology in artificial skin formation studies (1, 2).

Chicken embryo chorioallantoic membrane (CAM) is a complex structure composed of branched blood vessels in chicken eggs. Fertilized eggs CAM part is similar to placenta in mammals. It is a good “in vivo” culture media for the angiogenesis studies (3, 4). For this purpose, chicken fertilized embryos have been used in neovascularization studies since the early 1970’s (5). Up to now, angiogenic roles of different organic/inorganic molecules had been analyzed in CAM studies. The angiogenic roles of growth factors, hormones, anticancer agents, antibiotics and antibodies were clarified by using CAM in literature (6, 7, 8). Today we know that hormones such as testosterone and progesterone stimulate the angiogenesis. Therefore, testosterone and progesterone are used in tissue scaffold structure (9, 10).

Chitin is a cationic compound which is found in the shells of sea creatures such as crabs, lobsters and shrimps. It is used in scaffold technology in tissue engineering studies (11, 12). Chitosan is a natural fiber which is extracted from chitin. It is made by treating the chitin shells of shrimp and other crustaceans with an alkaline substance, like sodium hydroxide. Chitin and chitosan are used in scaffold technology because of their biodegradable, biocompatible and nontoxic specialties (11). In literature, no manuscript was found about the angiogenic role of chitin in the presence of testosterone and progesterone.

Here, chitin was obtained from shrimp shells in our laboratory. Testosterone and progesterone solutions were prepared in different concentrations. Chitin in the presence of testosterone and progesterone was analyzed by using CAM methodology. Our manuscript is a unique example which explains the angiogenic role of chitin/testosterone/progesterone together in literature.
Methods

This study was performed with the ethical decision of University of Health Science, Gülhane Health Science Institute, Animal Experiments Local Ethics Committee (February 27, 2018-Etk-2018/06).

Chitin Extraction from Shrimp Shells: Chitin was obtained from shrimp cells in our laboratory by using Kocer’s methodology (13). First, shrimp cells were cleaned with distilled water (Figure 1. A). For the deproteinization of shrimp cells, NaOH solution (3.5% w/v) was used. It was applied on shrimp shells for 2 hours at 65°C (10 ml solution for 1 gr shrimp shell). For the demineralization of shrimp cells, HCl solution (1 M) was used. It was applied on shrimp shells for 30 minutes at room temperature (15 ml solution for 1 gr shrimp shells). For the neutralization of shells, protein and mineral wastes were removed from the shrimp shells by washing them with distilled water. This neutralization step is important for “in vivo” cell culture studies (14). After neutralization step, chitin molecules were dried in the oven at 60° C for one night (Figure 1. B).

Preparation of Testosterone and Progesterone Solutions: Testosterone (Sustanon iv. 250 mg/ml) was obtained from Schering-Plough Company. Progesterone (Progestan iv. 25 mg/ml) was obtained from KocakFarmaCompany. Physiologic serum (SF) was used for dilutions in testosterone and progesterone solutions. Testosterone solutions were prepared in 20 nmol/L, 50 nmol/L, 100 nmol/L concentrations (16). Progesterone solution was prepared in 5 µmol/L, 10 µmol/L, 20 µmol/L concentrations (15).

Preparation of Fertilized Chicken Eggs: Atak-S type fertilized chicken eggs were obtained from the Chicken Production Farm of Başkent University. For our study, one hundred and twenty fertilized chicken eggs were selected carefully. Fertilized chicken eggs were kept one day in a dark room. At the end of 24 hour, they were transferred to a CO2 incubator (Sanyo MCO-19AIC CO2 170L Incubator). Fertilized chicken eggs were kept hold in an incubator for six days (at 37°C heat, 85-90% humidity). At the end of the 6th day, the egg shells were cleaned with an antiseptic solution. The shells of eggs were opened carefully with tweezers. As known, vascularization in chick embryo CAM begins at the 5th and the 6th days (17). Therefore, the CAMs of fertilized chicken eggs were visualized and photographed on the 6th day for observing the increase in angiogenesis.

Application of Chitin/testosterone/progesterone on CAM assay: All the fertilized chicken eggs were separated into six different groups for the application of chitin/testosterone/progesterone (Group 1- Control group, Group 2- testosterone applied group, Group 3- progesterone applied group, Group 4- chitin/testosterone applied group, Group 5- chitin/progesterone applied group, Group 6- chitin/testosterone/progesterone applied group). On the 6th day, CAMs were observed and photographed for the neovascularization. Then, SF solution was added as 250 µl in control group (Group 1). Chitin was applied in its natural form (Figure 1. B). Testosterone and progesterone solutions were applied as 250µl for each condition in chitin/hormone applied groups (Groups 2, 3, 4, 5, 6). Then, eggs were covered with parafilm. Parafilm application prevented the eggs from moisture loss and kept the sterilization of the CAMs. At the end, all of the eggs were placed back into the CO2 incubator.

Twenty four hours later (on the 7th day), the parafilms were opened. All the CAMs were analyzed for neovascularization. CAMs were visualized and photographed for observing the changing in angiogenesis.

Methods used in Angiogenesis Measurement: For finding the increase in angiogenesis, Knighton et al.’s scoring methodology was used (17). On the 6th and the 7th days, the increases in vascular structure on CAM were visualized and photographed. Three different observers noted the increases in vascular structure and numbered due to Knighton et al.’s scoring methodology. As known, the observations on the 6th and 7th days were enough to score the vessels (17, 18, 19). The response rate in vascular structure was graded as zero (0), one (1), two (2). Zero (0) means no change in vessel formation. One (1) means a minimal increasing in density and length in vessel formation on CAM. Two (2) reflects a maximal increase in density and length in vessel formation on CAM (17). Mean values were obtained from the different observers results. They were reflected the total differentiation score of angiogenesis (TDSA). TDSA results were represented the increase in angiogenesis (17, 18, 19) (Table 1).

Statistics analyses: TDSAs were obtained by having mean

Table 1. The total differentiation score of angiogenesis (TDSA) results which were seen in control, chitin, testosterone, progesterone applied groups.

<table>
<thead>
<tr>
<th>Study Groups</th>
<th>n</th>
<th>TDSA</th>
<th>p Values</th>
</tr>
</thead>
<tbody>
<tr>
<td>Group 1 Control</td>
<td>8</td>
<td>0</td>
<td></td>
</tr>
<tr>
<td>Group 2 Testosterone</td>
<td>11</td>
<td>6±0.1</td>
<td>p &lt; 0.05</td>
</tr>
<tr>
<td>Group 3 Progesterone</td>
<td>9</td>
<td>5±0.2</td>
<td>p &lt; 0.05</td>
</tr>
<tr>
<td>Group 4 Chitin + Testosterone</td>
<td>8</td>
<td>7±0.1</td>
<td>p &lt; 0.05</td>
</tr>
<tr>
<td>Group 5 Chitin + Progesterone</td>
<td>8</td>
<td>5±0.1</td>
<td>p &lt; 0.05</td>
</tr>
<tr>
<td>Group 6 Chitin + Testosterone + Progesterone</td>
<td>9</td>
<td>7±0.1</td>
<td>p &lt; 0.05</td>
</tr>
</tbody>
</table>

n: Number Of Evaluated Eggs / TDSA: Total Differentiation Score of Angiogenesis
value results of scores. Student t tests results (p values) represent two group comparisons of TDSA among control and chitin/testosterone/progesterone applied groups (Table 1).

Results

Neutral form of chitin is suitable for “in vivo” studies and/or tissue engineering studies (14). So, neutral form of chitin was obtained from shrimp shells in appropriate condition and used in our experiments (Figure 1B).

TDSA Results for Testosterone and Progesterone applied groups on CAM model: In control group (Group 1), TDSA value was found as zero (0) (Table 1, Figure 2). No obvious increase was observed in CAM assay in 6th and 7th day results in main and capillary vessel formations (Figure 2, Figure 3A). In testosterone applied group (Group 2), TDSA was found 6±0.1 (Table 1, Figure 2, Figure 3B). TDSA was observed as 5±0.2 in progesterone applied group (Group 3) (Table 1, Figure 2). So, testosterone and progesterone increased the main and capillary vessel formations in density and length in eye observations. The increases in TDSA values in Groups 2, 3 emphasize the increase in main and capillary vessel formations as seen in Figure 3B. The TDSA results were statistically significant in groups 2 and 3 (p≥0.05) (Table 1). Similar TDSA values were observed in the uses of hormones in different concentrations.

Results for Chitin/Testosterone and Chitin/Progesterone applied groups on CAM model: In group 4 (chitin/testosterone applied group), TDSA value was found as 7±0.1 (Table 1 and Figure 2). This TDSA result represents the angiogenic stimulator role of chitin/testosterone together. Chitin/testosterone increased the main and capillary vessel formations in density and length (Figure 3C). The TDSA result in group 4 was statistically significant (p≥0.05) (Table 1). Similar TDSA values were observed in the uses of testosterone in different concentrations.

In group 5 (chitin/progesterone applied group), TDSA value was found as 5±0.1 (Table 1 and Figure 2). High TDSA result represents the angiogenic roles of chitin/progesterone together. Chitin/progesterone also increased the main and capillary vessel formations in density and length on CAM assay. The TDSA result in group 5 was also statistically significant (p≥0.05) (Table 1). Similar TDSA values were observed in the uses of testosterone in different concentrations.

Results for Chitin/Testosterone/Progesterone applied group on CAM model: In group 6, chitin was used with different testosterone/progesterone solutions. TDSA was found as 7±0.1 in this group, which represents the stimulator role of chitin/testosterone/progesterone together. Chitin/testosterone/progesterone treatment increased the main and capillary vessel formations in density and length on CAM assay (Table 1, Figure 2, Figure 3D). The TDSA result obtained in group 6 was statistically significant (p≥0.05) (Table 1). Similar TDSA values were observed in the uses of testosterone in different concentrations.

We observed that chitin/testosterone/progesterone stimulate angiogenesis on CAM model. In different concentrations of hormones, similar angiogenic effects were found. May be the smallest hormone concentrations used in our experiments (20 nmol/L for testosterone, 5 μmol/L for progesterone) were enough for having angiogenic effect on CAM assay.

Discussion

Angiogenesis is a process of new vessel formation. Nowadays in medicine it is important to encourage angiogenesis especially for ischemic diseases and wound healing (20, 21). Wound healing is a complex and dynamic process of replacing devitalized and missing cellular structures and tissue layers. Angiogenesis plays a crucial role in wound healing by forming new blood vessels from preexisting vessels by invading the wound clot and organizing to a micro vascular network throughout the granulation tissue. This dynamic process is highly regulated by signals from both serum and the surrounding extracellular
matrix environment. Vascular endothelial growth factor, angiopoietin, fibroblast growth factor and transforming growth factor beta are amongst the potent angiogenic cytokines in wound angiogenesis (22). Therefore, potent angiogenic factors generally use in tissue regeneration studies in tissue engineering and regenerative medicine (especially in manufacturing of artificial skin) (23). Today, growth factors are used in scaffold structure for stimulating the angiogenesis in wound healing studies. In our experiment, testosterone and progesterone were analyzed for angiogenesis with chitin which one generally uses in scaffold structures in tissue engineering (24).

CAM model of chick embryos is a useful in-vivo working model for angiogenesis because of its' high sensitivity (25). In literature, we know that the hormones and growth factors stimulate angiogenesis in certain tissues (26, 27). Despite of these findings, no study has been observed in literature about the angiogenic role of chitin/testosterone/progesterone together. In our manuscript, the possible positive angiogenic effects of testosterone and progesterone were analyzed on CAM in study groups 2 and 3 (only testosterone or only progesterone). We observed the stimulatory effects of testosterone/progesterone hormones which support literature findings (Table 1, Figure 2, Figure 3B) (26, 27).

Chitin is abundant in invertebrates and fungi as an important structural molecule. It is a primary component of cell walls in fungi, the exoskeletons of arthropods, such as crabs, lobsters and shrimps and insects (27, 28). Chitin has proved useful for several medicinal, industrial and biotechnological purposes. In biotechnology, it generally uses in scaffold structure (29). In our experiments, we obtained chitin from shrimp shells in our laboratory (Figure 1A). At the end, we neutralized them with distilled water. In this way they were formed as useful for “in vivo” studies (Figure 1B).

The angiogenic role of chitin in the presence of testosterone/progesterone was analyzed in study groups 4 and 5 (Chitin and testosterone or progesterone). Excessive vessel formations were observed on CAM in this panel (Table 1, Figure 3C). In group 6 (Chitin and testosterone and progesterone)angiogenesis was increased by chitin in the presence of testosterone/progesterone (Table 1, Figure 3D). Our manuscript is the first example in literature which represents angiogenic role of chitin/testosterone/progesterone together.

In our study, testosterone and progesterone hormones were used in different concentrations. Similar results were obtained in highest and lowest hormone concentrations. No extra angiogenic role was observed in high hormone dosages in hormone and chitin/hormone groups (Groups 2, 3, 4, 5, 6). We suppose that the lowest hormone dosages used in our experiments were enough in angiogenic role on CAM.

Chick embryo CAM model results support angiogenic roles of chitin, testosterone and progesterone. Chitin can be used easily in scaffold technology with testosterone and progesterone.

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Conflict of Interest

No conflict of interest exists related to our submission.

References


