Angiogenesis inhibition by green propolis and the angiogenic effect of L-lysine on bladder cancer in rats

Inibição da angiogênese pela própolis verde e efeito angiogênico da L-lisina no câncer de bexiga em ratos

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ABSTRACT

PURPOSE: To determine the effects of water-soluble derivative of green propolis in bladder cancer angiogenesis in rats given N-butyl-(4-hydroxybutyl) nitrosamine (BBN).

METHODS: Nine groups were established, where six of them (Groups 1 to 6), the animals received 0.05% BBN in their drinking water for 14 weeks. From the 32nd to the 40th week, Groups 1, 2, 3 and 4 were treated respectively with water, L-lysine (300 mg/kg/day), celecoxib (30 mg/kg/day) and propolis (300 mg/kg/day). Groups 5 and 6 were given propolis and L-lysine from the 1st to the 40th week (150 mg/kg/day). Microvascular density was determined by histological sections stained for the marker CD-31 and analyzed with specific software.

RESULTS: The microvascular density in bladder carcinomas was lower (p<0.01) in rats receiving propolis than in controls given carcinogen only. On the other hand, the microvascular density of tumors in rats receiving carcinogen and L-lysine for 40 weeks from the beginning of carcinogen treatment was significantly higher (p<0.01) than in the corresponding controls.

CONCLUSION: Water-soluble derivative of propolis inhibits angiogenesis in BBN-induced rat bladder cancer, while L-lysine treatment stimulates angiogenesis if initiated concurrently with BBN.

Key words: Lysine. Propolis. Angiogenesis Inhibitors. Urogenital Neoplasms. Rats.

RESUMO

OBJETIVO: Determinar os efeitos da própolis verde solúvel em água na angiogênese de câncer de bexiga em ratos que receberam n-butil-(4-hidroxiibutil) nitrosamina (BBN).

METODO: Nove grupos foram estabelecidos, onde em seis destes (grupos de 1 a 6) os animais receberam BBN a 0,05% em água...
de beber por 14 semanas. Na 32ª semana das 40 semanas, os grupos 1, 2, 3 e 4 foram tratados respectivamente com água, *L* lisina (300 mg/kg/dia), celecoxibe (30 mg/kg/dia) e própolis (300 mg/kg/dia). Os grupos 5 e 6 receberam própolis e *L* lisina da 1ª a 40ª semana (150 mg/ kg/dia). A densidade microvascular foi determinada por cortes histológicos corados pelo CD-31 e analisados por programa de computador específico.

**RESULTADOS:** A densidade microvascular em carcinomas de bexiga foi menor com p<0,01 nos ratos que receberam própolis do que nos carcinomas do grupo controle que recebeu apenas carcinógeno. Por outro lado, a densidade microvascular de tumores de ratos que receberam carcinógeno e *L*-Lisina por 40 semanas desde o início do carcinógeno foi significamente maior com p<0,01 que a densidade microvascular dos tumores de seu respectivo grupo controle.

**CONCLUSÃO:** A própolis verde solúvel em água inibiu a angiogênese em câncer de bexiga induzido pelo BBN, enquanto a *L*- lisina estimulou a angiogênese quando iniciada juntamente com o BBN.


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**Introduction**

Angiogenesis is a physiological or physiopathological process involving the growth of new blood microvessels from pre-existing capillaries\(^1\), induced by a range of chemical factors, some of which are related to genetic changes, termed angiogenic and antiangiogenic factors.

Propolis is a resinous substance that honey bees collect from botanical sources and mix with their own saliva. It is used as a sealant for unwanted open spaces and protects the hive against microorganisms. Green propolis is found in Southeastern Brazil and is so called because of the presence of green pigment collected from the shrub species *Baccharis dracunculifolia*.

Water-soluble derivative of propolis (WSDP) is produced by extracting soluble propolis in water using 8% *L*-lysine\(^2\). *L*-Lysine is a strictly essential amino acid required for collagen synthesis, tissue growth and repair, and production of antibodies, hormones and enzymes. Patented by Nicolov, the process uses *L*-lysine to extract propolis, making it possible to obtain a range of compounds with pharmacological action in a practical, water-soluble form.

Researchers have reported anti-inflammatory\(^4\), immunomodulatory\(^5,6\), antitumor\(^7\), antioxidant\(^8-10\) and antimetastatic\(^5\) effects of WSDP. The discovery of the anticarcinogenic effects of propolis encouraged efforts to identify, isolate and synthesize constituents of propolis and to study their effects and mechanisms of action, including artepillin\(^11\), caffeic acid\(^12\), caffeic acid phenethyl ester\(^13\), and in the present study, we used an in vivo model to evaluate the effect of intragastric administration of green propolis (in the form of WSDP) on angiogenesis in BBN-induced rat bladder cancer.

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**Methods**

The experiment was previously approved by the Ethics Committee for Animal Research under protocol # 03/06 and was carried out in accordance with the guidelines of the Animal Testing.

The study used 92 five-week-old female Wistar rats accommodated in polyethylene cages (5 or 6 per cage) at 22°C with a 12/12-hour circadian cycle. The animals had access to food (Biobase\(^®\) pelleted ration) and water *ad libitum*.

**Experimental procedures**

The exclusion criteria were presence of carcinoma with extensive necrosis, autolysis or failure to develop carcinoma. The analysis included 49 animals. Microvascular density was determined for 31 (six animals from Group 1 and five animals from each of Groups 2, 3, 4, 5 and 6). See experimental design Figure 1.
N-butyl-N-[4-hydroxybutyl] nitrosamine

BBN was purchased from TCI America (Portland, OR). The chemical formula is \( \text{CH}_3(\text{CH}_2)_3\text{N}[(\text{CH}_2)_4\text{OH}] \) (Chemical Abstract Service #3817-11-6). The product was stored at 4°C. A 0.05% concentration of BBN was dissolved in water and was supplied \textit{ad libitum} for 14 weeks in dark bottlers.

**Biosafety norms**

The laboratory animal facility and the handling of animals and materials followed WHO guidelines for use of nitrosamines in experimental research\textsuperscript{14,15} and MSDS (Material Safety Data Sheet) specifications.

**Water-soluble derivative of propolis (WSDP)**

Green propolis was collected from hives in Passa Quatro, Southern Minas Gerais (Brazil), from September to October. WSDP was prepared following the procedures described by Nicolov et al.\textsuperscript{2} with slight modifications. Propolis was extracted in ethyl alcohol three times, concentrated in a water bath, submitted to liquid-liquid partitioning with petroleum ether and, subsequently, with ethyl ether, and then again concentrated in a water bath. The extract was added to an 8% \( \text{L-lysine} \) solution at 51-60°C in a proportion of 100g propolis to 1500 ml \( \text{L-lysine} \) solution. The \( \text{L-lysine} \)-based extract was lyophilized and cold-stored in small sterilized vials appropriate for administration of 150 or 300 mg/kg body weight (Figure 2).

**Antioxidant activities**

The freeze-dried propolis WSDP and the ethanol extract of propolis in natura in concentrations of 1000, 250 and 62.5 ppm were evaluated for DPPH (1,1-diphenyl-2-picrylhydrazyl) radical scavenging activities and compared with the controls for vitamin C and Trolox (a water-soluble derivative of vitamin E).

**L-Lysine**

The amino acid \( \text{L-lysine monohydrochloride} \) (\( \text{C}_6\text{H}_{14}\text{N}_2\text{O}_2\cdot\text{HCl} \); CAS #657-27-2) was prepared for extraction in a distilled water solution at 8% and administered at doses of 150 and 300 mg/kg. Celecoxib was administered orally as a positive control at a dose of 30 mg/kg body weight.

**Ultrasound scanning**

At 31 and 40 weeks, all animals were anesthetized peritoneally with 80 mg/kg ketamine and 8 mg/kg xylazine and submitted to ultrasound scanning of the bladder in transversal and sagittal sections using a 12-MHz linear transducer (Philips®).

**Immunohistochemistry**

Histological 5-\( \mu \)m sections of bladder tumors were placed on slides coated with organosilane (3-aminopropyltriethoxysilane/SIGMA). Following deparaffination and hydration, antigen retrieval was performed with 1 mM ethylenediamine tetraacetic acid (EDTA) at pH 8 in a pressure cooker for 8 minutes. Endogenous peroxidase activity was blocked with 3% \( \text{H}_2\text{O}_2 \) for 10 minutes. Samples were incubated overnight in a moist chamber at 4°C with primary antibody PECAM-1 (M-20): sc-1506 (Santa Cruz Biotechnology, Inc.) diluted to 1:50. Subsequently, samples were rinsed with phosphate buffer saline (PBS) and incubated with secondary biotinylated swine anti-goat/rabbit/mouse antibody (yellow reagent in the streptavidin-biotin kit; LSAB kit/HRP by Dako) in a moist chamber at room temperature for 30 minutes. Samples were again rinsed in PBS for 1 minute followed by incubation for 30 minutes with red streptavidin-peroxidase reagent. The reaction was revealed using 10% 3,3’-diaminobenzidine tetrahydrochloride (DAB) and hydrogen peroxide, and then counterstained with Carazzi’s hematoxylin.

**Image acquisition and quantification of angiogenesis**

Images of histological samples were acquired in standard fashion using a light microscope (BX4, Olympus Optical Co. Ltd.,...
Japan) fitted with a digital camera (C7070 Wide Zoom, Olympus Imaging America Inc., USA). The procedure included an initial scan of the tumor at 40X magnification in order to identify areas with increased vascular density (hot spots)\(^6\). Three hot spots selected for each tumor were then color-scanned at 200x magnification. Images were saved as Windows\(^8\) bitmaps measuring 512 x 384 pixels (1 pixel = 24 bits, RGB color scheme). Images were processed with a morphometric analysis software developed specifically for this purpose\(^7\). The system was previously adjusted to the color spectrum produced by the staining techniques employed. This step enabled the software to automatically identify and segment (single out) blood microvessels in both full images and user-defined regions. However, interactive segmentation (making changes in segmentation parameters) remained an option whenever the automatic mode was deemed inadequate. Following segmentation, the software provided the microvascular density of areas of interest (Figure 3). The area density was calculated by dividing the area of microvasculature by the total area analyzed.

**FIGURE 3** - Identification and segmentation of vessels and cells immunostained with calculation of microvessel density: total area occupied by microvessels divided by the total area of the field analyzed.

**Descriptive statistics**

The quantitative variables were initially analyzed with the Kolmogorov-Smirnov pretest to verify the normality of distribution. Having observed normality in all cases, means and standard deviations were calculated for the descriptive statistics, and the data were analyzed with parametric tests. Pairwise comparisons were made between groups with one-way analysis of variance combined with Tukey’s multiple comparison test\(^8\). In all cases, the \(\alpha\)-probability of a type-I error was established as 0.05 (5 %). Thus, two-tailed \(p\)-values less than 0.05 were considered statistically significant. Statistics and preparation of figures were carried out with the help of the software GraphPad Prism\(^8\) v. 5.00 for Windows\(^8\) (GraphPad Software, San Diego, CA, USA, 2007).

**Results**

**Characterization of propolis**

The lyophilized propolis was submitted to high-performance liquid chromatography (HPLC) and quantified (Table 1, Figure 4).

**TABLE 1 - Quantification of markers identified in the lyophilate (mg compound/g powder).**

<table>
<thead>
<tr>
<th>Compound</th>
<th>mg/g</th>
<th>%</th>
</tr>
</thead>
<tbody>
<tr>
<td>3-[4-hydroxy-3 (oxo-butenyl) phenyl acrylic] acid</td>
<td>0.385</td>
<td>0.04</td>
</tr>
<tr>
<td>3-prenyl-4-hydroxycinnamic acid</td>
<td>0.453</td>
<td>0.04</td>
</tr>
<tr>
<td>caffeic acid</td>
<td>2.450</td>
<td>0.24</td>
</tr>
<tr>
<td>Caffeoy quinic acid</td>
<td>1.128</td>
<td>0.11</td>
</tr>
<tr>
<td>p-coumaric acid</td>
<td>35.99</td>
<td>3.60</td>
</tr>
<tr>
<td><strong>Total</strong></td>
<td>40.41</td>
<td>4.04</td>
</tr>
</tbody>
</table>

**FIGURE 4** - High Performance (pressure) Liquid Chromatography (HPLC).

**Antioxidant activities**

DPPH radical scavenging activity of freeze-dried propolis (WSDP) and propolis ethanol extract in natura, at concentrations of 1000, 250 and 62.5 ppm show similar values to the antioxidant activity of the controls Trolox (T) and Vitamin C (Figure 5).
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Ultrasonography

At weeks 31 and 40, tumors were identified on ultrasonographic images (all groups), determining the beginning of treatment (groups 1, 2, 3 and 4) and tumor development (Figure 6).

Statistical findings

Findings were expressed as mean ± standard deviation of triplicate measurements of six animals from Group 1 and five animals from each of Groups 2, 3, 4, 5 and 6. Pairwise comparisons were made between groups with one-way analysis of variance combined with Tukey’s multiple comparison test (Figure 7).

Discussion

The chemical composition of propolis is strongly dependent on the flora in the area where the hive is located, bee type, climate, temperature and season19,20.

The dose chosen for our experiment (150 mg/kg body weight) was based on previous studies by other researchers investigating the anticarcinogenic effects of WSDP extracted from propolis collected in Minas Gerais5,7,8.

Comparisons between measures of microvascular density in Groups 1, 2, 3 and 4 show that tumor angiogenesis was significantly inhibited in Group 4 (propolis), in contrast to Group 2 (L-lysine) and Group 1 (controls receiving BBN only).

Although Group 4 displayed a lower density than Group 2, there was no statistically significant difference between Group 2 and Group 1. Since Group 2 is statistically similar to Group 1 and both differ from Group 4, and considering that propolis in Group 4 was extracted with the aid of L-lysine, two observations may be made: propolis inhibited angiogenesis even when L-lysine was used in its extraction, and L-lysine apparently did not affect microvascular density in this tumor progression group, nor did it counteract the effect of propolis (Group 4).
Compared to Group 1 (control), by week 40 the microvascular density was equally reduced in Group 5 and Group 4 in spite of the fact that the animals in these groups were started on WSDP at different times and doses. In other words, propolis inhibited angiogenesis regardless of whether it was administered at baseline (concurrently with BBN administration) or only later, in the tumor progression stage (starting week 32).

The findings of the present study differ from those of the first study published on antiangiogenesis induced by artepillin C extracted from propolis from Minas Gerais. In that study, the ethanol extract contained 1.6 mg/g caffeic acid, 27.4 mg/g p-coumaric acid and 43.9 mg/g artepillin C, but only the last constituent was able to inhibit tube formation in 2D culture of human umbilical vein epithelial cells (HUVEC). The authors also reported that propolis inhibited angiogenesis in vivo in sarcoma 180. In our study, artepillin C was not detected in WSDP, but a small amount (0.385 mg/g WSDP) of the derivative of artepillin C o 3-[4-hydroxyl-3 (oxo-butenyl) phenyl] acrylic acid. What component would then be responsible for the antiangiogenic effects observed in Groups 5 and 4?

In an elegant and detailed study using caffeic acid and a synthetic analog, Jung and coworkers observed inhibition of tube formation in 3D culture of HUVEC (in vitro) and in vivo, through the inhibition of angiogenesis in human kidney tumor implanted in nude mice. They also reported a decrease in vascular endothelial growth factor (VEGF) and a delay in tumor growth by suppression of STAT3 phosphorylation and HIF-1 expression downregulating the production of HIF-1-mediated VEGF.

The present findings suggest that propolis may contain compounds with antiangiogenic activity other than artepillin C. This would partly explain why artepillin C-free lyophilized propolis inhibited angiogenesis in the present study. It has been demonstrated that a number of compounds found in propolis around the world, such as caffeic acid phenethyl ester, caffeic acid and artepillin C, display antiangiogenic properties in vitro in HUVEC culture and in vivo in certain types of carcinoma. These authors demonstrated in a recent study that the ethanolic extract of Brazilian propolis suppresses the phosphorylation of ERK 1/2 and inhibits tube formation in cultured HUVEC. The ethanolic extract of propolis induces the activation of caspase 3 leading to apoptosis. Utilizing the same in vitro model, it was observed that propolis in increasing concentrations can suppress the expression of integrin β 4 and protein P53 and reduce the action potential of the mitochondrial membrane, leading to apoptosis in the endothelial cell.

The WSDP propolis used in this study showed strong antioxidant activity. The antioxidant effect may be responsible for the inhibitory action of angiogenesis?

Researchers investigated the angiogenic and antioxidant activities of various components from propolis: artepillin C, caffeic acid, phenethyl Ester, p- coumaric acid, galagin, apigenin, kaempferol, pinocebrin and quercetin. The effects of these components were tested on in vitro models of angiogenesis, tube formation and growth of HUVECs and antioxidants activities by DPPH. Propolis components such as caffeic acid phenethyl ester, quercetin artepillin C, galagin and kaempferol possessed strong antiangiogenic and antioxidant activities. They found a correlation between antioxidant activity and antiangiogenic activity, and so they concluded that these components may be responsible, at last, in part, to the antiangiogenic activity of propolis.

When comparing Groups 1, 6 and 5, the microvascular density was found to be much higher in tumors of animals started on L-lysine at baseline (Group 6) compared to controls (Group 1) or animals started on WSDP at baseline (Group 5), suggesting that L-lysine is angiogenic if administered concurrently with BBN. This is confirmed by comparing Groups 1, 6 and 2. However, the double dose of L-lysine administered from week 32 on (Group 2) did not significantly affect microvascular density in relation to controls (Group 1).

Three amino acids have been described as promoting bladder cancer, namely DL tryptophan, L-leucine and L-isoleucine. This and the fact that WSDP is extracted with the aid of an amino acid led to the inclusion in this study of a group treated with L-lysine. In fact, to our knowledge, no other study on WSDP has included a control group receiving L-lysine. Moreover, this is the first report of the effects of L-lysine on BBN-induced bladder cancer.

It is intriguing that, although the dose of L-lysine in Group 2 is twice that in Group 6, angiogenesis was not stimulated. Is this because the dose was different, or were the angiogenic effects observed in Group 6 due to the gene expression of a more aggressive and thus more invasive and faster-growing tumor, effects produced by L-lysine during initiation/promotion of carcinogenesis of the bladder epithelium? In paper accept for publication, based on the same experimental model shows that L-lysine was able to stimulate carcinoma multiplicity and invasion in BBN-induced rat bladder cancer.

Another important observation is that while the present findings indicate a significant angiogenic effect of L-lysine in Group 6, this effect did not overcome the inhibitory effect of propolis on angiogenesis in Group 5, although the amount of L-lysine in the propolis group was the same (150 mg/kg).
Celecoxib, a selective Cox-2 inhibitor has been shown to effectively decrease tumor angiogenesis and reduce tumor growth of a variety of experimental primary tumors\(^9\)\(^{,10}\). Celecoxib reduced microvascular density (positive control Group), but the difference was not statistically significant in relation to negative controls (Group 1).

**Conclusions**

Green propolis (WSDP) administered intrastratically at 150 mg/kg body weight (40 weeks) beginning concurrently with BBN, or at 300 mg/kg beginning at week 32 of carcinogenesis, inhibits angiogenesis in BBN-induced rat bladder cancer, and that L-lysine administered intrastratically at 150 mg/kg body weight (40 weeks) concurrently with BBN produces angiogenic effects in this model of bladder tumor.

**References**


Acknowledgments

To Professor Maria Cristina Marcucci (Bandeirante University of Sao Paulo-UNIBAN) for typifying and quantifying our propolis samples and Professor Francisco Hélio Rôla from the Department of Physiology and Pharmacology at UFC Medical School for his helpful suggestions and critical comments.

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Received: March 22, 2012
Review: May 23, 2012
Accepted: June 21, 2012
Conflict of interest: none
Financial source: none

Research performed at Laboratory of Experimental Surgery, Department of Surgery, Faculty of Medicine, Federal University of Ceará (UFC) Brazil. Part of PhD thesis, Postgraduate Program in Surgery. Tutor: Prof Manoel Odorico de Moraes.