

Propolis gel versus benzydamine in preventing oral mucositis for patients irradiated in head and neck: a preliminary study

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Abstract

Oral Mucositis refers to oral mucosa erythematous and ulcerative lesions and is caused by radiation dispensed in the treatment of malignant tumors of the head and neck. Propolis shows several biological activities such as antimicrobial and anti-inflammatory properties. These biological activities should prevent a mucositis. The aim is to verify the effectiveness of Brazilian green propolis in a mucoadhesive gel, alcohol free, in preventing oral mucositis in patients that underwent radiotherapy. This research is characterized as a preliminary study with a mean duration of 12 weeks of patients' follow-up in two groups. The selection of participant's groups was randomized. The 26 selected patients were distributed between two groups, 13 in the benzydamine group and 13 in the propolis group. Patients were assessed on an average of 4.5 times totaling 116 diagnoses of mucositis. The percentage of patients who had mucositis greater or equal to 2 in this study was 30.6% for the benzydamine group and 29.6% for propolis gel. For these patients, the propolis gel shows a better performance in maintaining lower rates/grades and recovery of patients from the 17th session of radiotherapy. This work shows a great possibility of further study in Phase III due to its good results and acceptance with patients.

Introduction

Patients with malignant tumors on the upper aero digestive way need a multidisciplinary treatment [1]. These tumors treatment can be done through surgery, radiation therapy (RT) and chemotherapy. All these treatments increase the rate of comorbidities that can be reversible or irreversible [2]. Oral Mucositis refers to oral mucosa erythematous and ulcerative lesions observed in patients being treated with radiotherapy (RT) for head and neck cancer [3]. Mucositis is related as severe in 30.0-40.0% of the cases [4,5]. Multinational Association for Support and care of Cancer and the International Society of Oral Oncology (MASCC/ISOO) advocated the use benzydamine as a treatment for mucositis, a topical anti-inflammatory drug, with fluconazole. This protocol requires the combination of two drugs being that benzydamine has alcohol in formulation and may cause irritation and burning, these was the most frequently reported toxicity [6-9]. The term propolis comes from greek pro-"in front of, on the entrance of" and polis-"community or city" [10-12]. The chemical composition of propolis is complex, and some factors such as plant ecology of the region where the propolis was collected and also the genetic variability of the queen bee, can influence on the chemical composition of the resin material [13-15]. Nowadays, it is known that propolis shows several biological activities such as antimicrobial, anti-inflammatory, anesthetic and cytostatic properties [16]. This product activity should prevent mucositis pathophysiology characteristics [6] and properties of propolis [17]. The aim of this preliminary study was the verify the effectiveness of Brazilian green propolis in a mucoadhesive gel, alcohol free, in preventing oral mucositis in patients that underwent RT in head and neck region compared with a benzydamine (0.15%) alcoholic solution.

Patients and methods

This research is characterized as a preliminary study with a mean duration of 12 weeks of patients' follow-up. A gel containing propolis manipulated within the standards required by ANVISA (Brazil, 2000), ISO 9001 and GMP International. Propolis was collected and manipulated according to the technical standards of the Brazilian Pharmacopoeia (2015). For inclusion in this study the patients must have a confirmed diagnosis of malignant lesion in head and neck RT, submitted to a dose higher than 50 Gys. The selection of participants to groups was randomized and conditioned to availability of the patients attend the study during RT [18]. All patients signed informed consent prior to participation and the Ethics Committee on Human Research (COEP/UFMG) approved the study under number CAAE 04328812.1.0000.5149. A total of 70 patients were initially enrolled in the study. Twenty-six patients were included in these studies and divided in two groups: group 1 used benzydamine with fluconazole according MASCC/ISOO and group 2 used mucoadhesive gel with propolis 5%. Flow diagram about the clinical trial design study is shown in Figure 1. All selected patients, were administered on the first day of RT propolis gel-every 8 hours (6 h, 14 h, 22 h). It was used on the first day of RT and it was indicated its administration for another two weeks after the last session of RT. For patients in the control group benzydamine, protocol

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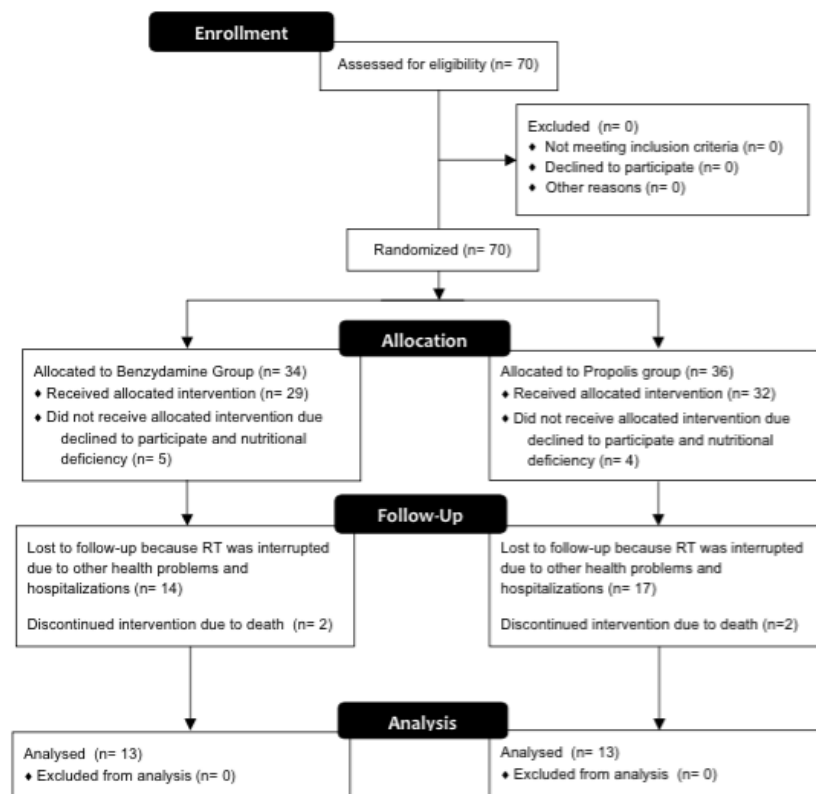


Figure 1. Clinical trials flow diagram.

according to the MASCC/ISOO, it was administered following the same dosage used for the gel containing propolis, however, fluconazole was prescribed only for patients with clinical features of infection with *Candida spp.* All study patients were evaluated every 7 days during the whole period of RT. The World Health Organization (WHO) scale performed the evaluation of the degree of mucositis [7]. The same examiner, using the same initial criteria, made these assessments. These tests were considered: intra-oral examination (soft tissue changes), data collection for the presence or absence of candidiasis.

Statistical analysis

The statistical analysis began by checking the homogeneity of demographic and clinical quantitative variables between the benzydamine and the propolis gel groups with the Mann-Whitney statistical test [19]. Fisher's Exact Test was used for homogeneity of qualitative variables between the groups [20]. This research is a study of completely unbalanced repeated measures. To evaluate all factors that exert influence on the grade of mucositis, two levels of mucositis, "mucositis grade <2" and "mucositis grade ≥2", were set to analyze. However, this study should, in a careful manner, model the structure of the medium, i.e., as the average percentage of the higher degree of mucositis equal to 2, behave along the RT sessions. For this, the Splines method [22] was used with a knot in the in the tenth seventh session of RT, to set graphical views using nonparametric regression via LOWESS method [21]. These segments are called Splines and the location where the lines meet is called knot [22]. Thus, we call the adjusted models as Logistic Marginal Regression Splines, in which we evaluate the significance of the variables of interest and its interaction with the number of sessions on two occasions, before and after the session RT 17. Marginal Logistic Regression Splines was first used to check for

significant differences in the average percentage of the higher degree of mucositis equal to 2, along the RT sessions between the Benzydamine and the propolis groups. To select the set of significant variables, the stepwise selection method variables were used [23]. The stepwise method is defined as a mixture of Backward and Forward methods. The Forward method in this study was carried out via analysis of deviation from univariate regressions, using a p-value of 0.25 for entering variables in the multivariate model. On all variables selected by the forward method, we applied the backward method with a significance level of 5%. The software used in the analysis was the R version 3.0.1.

Results

Epidemiological data and the homogeneity of demographic and clinical quantitative variables between groups are shown in Table 1. Treatment for other systemic changes was reported by 57.7% (n=15) of patients, the most frequent cardiovascular disorders with 73.3% (n=11) of cases. The chemotherapy was performed in 6 (46.2%) benzydamine group and 3 (23.1%) propolis group. The chemotherapy did not interfere in the result as a whole, because the p-variable showed no significant variation (0.179) to say that the groups were different from each other. The number of sessions of RT in primary lesion was 33.39 (±4.04), ranging from 20 to 39 sessions. The total dose of RT in primary lesion was, on average, 6466.1 cGy range 5000-7200 cGy. The extent of mucositis was performed along the RT sessions on average 4.5 times ranging from 2-7 per patient ratings. In total 116 diagnoses mucositis, benzydamine in the group 62 and group 54 in the gel were propolis. The total number of diagnosis (n=116) with mucositis grade 2 or higher was 30.6% in the benzydamine group and 29.6% in the propolis group according to Table 2. To visualize the behavior along the sessions of RT, the mucositis percentage of grade 2 or higher

Table 1. Homogeneity according to clinical and demographic variables between the study groups.

Variables		Benzdamine (n=13)		Propolis (n=13)		p-Value
Gender	Female	3	23.10%	1	7.70%	0.593
	Male	10	76.90%	12	92.30%	
Age-Average (SD)		54.1	-15	53.7	-12.9	0.938
Medical Treatment	No	4	30.80%	7	53.80%	0.428
	Yes	9	69.20%	6	46.20%	
Smoking	No/Ex-Smoker	11	84.60%	10	76.90%	1
	Yes	2	15.40%	3	23.10%	
Alcoholic Beverages	No	12	92.30%	9	69.20%	0.322
	Yes	1	7.70%	4	30.80%	
Histological Type	CCE	9	69.20%	11	84.60%	0.645
	Other	4	30.80%	2	15.40%	
Location	Larynx	2	15.40%	5	38.50%	0.372
	Tongue	4	30.80%	1	7.70%	
	Oropharynx	3	23.10%	2	15.40%	
	Other sites	4	30.80%	5	38.50%	
Chemotherapy	No	3	23.10%	7	53.80%	0.179
	Yes	6	46.20%	3	23.10%	
Number of Primary Lesion sessions - Average (SD)		33.6 (5.9)		33.2 (3.2)		0.289
Total dose of RT Primary Lesion - Mean (SD)		6534.5 (742.1)		6403.3 (477.1)		0.476

Table 2. Degree of mucositis in all evaluations by study group.

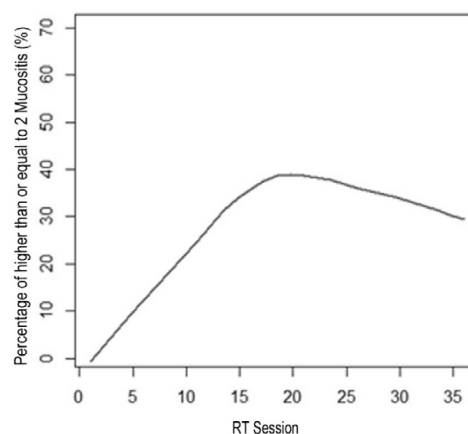
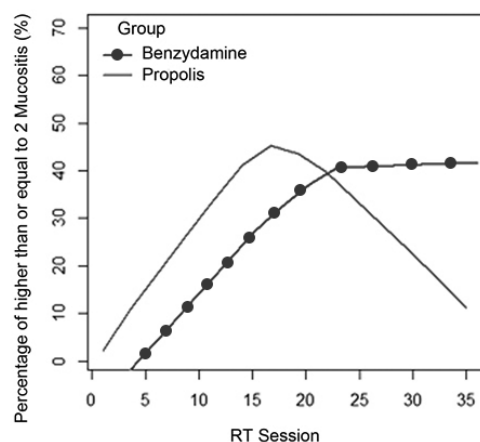
Degree of mucositis	Benzdamine		Propolis	
	N	%	N	%
0	30	48.4%	31	57.4%
1	13	21.0%	7	13.0%
<2	43	69.4%	38	70.4%
2	12	19.4%	8	14.8%
3	6	9.7%	8	14.8%
4	1	1.6%	0	0.0%
≥2	19	30.6%	16	29.6%

between benzydamine hydrochloride and gel containing propolis was used the scatter diagram using nonparametric regression via LOWESS method that can be observed in Figure 2. Thus, it is observed that in both groups there was an increase of the percentage of mucositis with a grade equal to 2 or higher, to the 17th session, and in this interval the propolis group showed a higher percentage (>40%). In Figure 3 we can observe that after the 17th session of RT the propolis group showed a decrease in the percentage of mucositis with a grade equal to 2 or higher, while the benzydamine hydrochloride group tended to stabilize at 40% of mucositis with a grade equal to 2 or higher.

The follow up after RT it was performed for five weeks and in this period patients no had mucositis.

Discussion

Several authors reported that mucositis is multifactorial and changes in the physiology of the mucosa by the action of radiation have different effects acting simultaneously leading to the reported side effects. In order to avoid these changes, the scientific community has been seeking ways to minimize adverse effects. The basic therapy of mucositis should include drugs that should have antifungal, anti-inflammatory, antioxidants and healing effects [3,4,24]. The results on mucositis in this study are compared to several studies marked by MASCC/ISOO [7,9]. The number of patients without or with mucositis and mucositis grade I presented in this study ($\pm 70\%$) agree with other results described in the literature [3,25,26]. The green propolis has in its


Figure 2. Smoothing method through LOWESS to the percentage of mucositis along the RT sessions for both groups.

Figure 3. Smoothing method through the LOWESS method to the percentage of mucositis along the RT sessions by group.

chemical constitution various flavonoids such as galangin and artemillin C, characterizing this compound as anti-inflammatory, antimicrobial and antioxidant [27,28]. Such features described in the literature are important for the prevention of mucositis because when the knowledge of the pathophysiology of mucositis and the knowledge of the products used in choice of treatment of mucositis [6] were associated, it became clear that the propolis gel without alcohol prevented inflammation, reduced free radicals and showed antifungal effects. All these features gathered in a single product make it more advantageous than other products such as benzydamine, which is an alcoholic solution and requires association with fluconazole. Propolis gel also has advantages over products made from salt, considering that the percentage of patients with cardiovascular disorders can't be neglected, because some products use saturated salt in their formulation [29]. Patients with diabetes should also be treated with caution because some authors recommend the use of honey [30,31]. Since some products use saturated salt in their formulation [29] and the percentage of patients with cardiovascular disorders can't be neglected, propolis gel has also advantages over those products. Patients with diabetes should also be treated with caution because some authors recommend the use of honey [30,31].

Conclusion

In this study, we observed a better behavior of the gel containing propolis regarding benzydamine after the 17th session of RT. This has been justified by anti-inflammatory and antifungal characteristics of propolis. Since the mucositis is caused by inflammation and aggravated by the presence of fungus, it is predictable that patients with the continued use of this propolis containing product can have a better oral health. Mucositis is exacerbated by the extent and amount of applied radiation. Observing patients in more advanced stages of the malignant tumors of the head and neck treatment, we note that the presence of mucositis in lower levels is the expected effect for the patients treated with propolis gel. Since the propolis gel have good acceptance by patients, this study points to the necessity of further research in Phase III to obtain more significant results.

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