Histological and Immunohistochemical Evaluation for the Effect of Pilocarpine and Quercetin on Gamma-irradiated Parotid Salivary Glands

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**Introduction**

During the last decades, there has been a substantial improvement in the treatment of head and neck cancer and radiotherapy is currently a widely used and important part of this treatment (El-Faramawy et al., 2013). In comparison with the surgical procedures, it shows better results because it can be used as a curative, adjuvant, neoadjuvant and palliative type of treatment and is often used in the conservative approaches, with protocols that preserve the organs and tissues (Marta et al., 2014). The major oral surrounding tissues that suffer from side effects when interacting with ionizing radiation from radiotherapy are the salivary glands. These tissues usually receive secondary doses from those delivered to the head and neck tumors, and although they have a low mitotic rate, salivary glands are considered to be extremely radiosensitive (Garg & Malo, 1997). While the radiation damage to salivary glands is well known in the clinic by its sides effects, it is not known exactly what mechanism of destruction the ionizing radiation have on salivary glands. It is known that the serous acini are more radiosensitive than mucous acini (Coppes et al., 2002). Existing protection strategies comprise strict dental and oral hygiene, parotid-sparing radiation technique, and pharmacotherapy such as salivary substitutes and sialogogues (Chambers et al., 2004).

**RA DIOTHERAPY** of patients with head and neck tumors usually causes damage to the salivary glands since these are most frequently included in the field of irradiation. This study aims at investigating the possible protective effect of pilocarpine or quercetin against radiation induced parotid gland damage. Forty-five adult male rats were divided randomly into three main groups as follows: Radiation group where the rats were exposed to a single whole body 6Gy γ-irradiation; pilocarpine- and quercetin-radiation groups where the rats injected intraperitoneally by a single dose of pilocarpine (0.2mg/kg) or quercetin (1.25g/kg), respectively, 30min later exposed to a single whole body 6Gy γ-irradiation. The parotid glands were stained with hematoxylin and eosin and immunohistochemical reaction for vascular endothelial growth factor A (VEGF-A). Irradiated glands revealed massive acinar atrophy, degeneration and cytoplasmic vacuolization while, the ducts showed cytoplasmic vacuolizations with loss of regular cell architecture. Pilocarpine or quercetin treatment before radiation exposure offered some protection effect manifested as reduced acinaratrophy, degeneration and vacuolization. The ducts showed mild cellular vacuolizations. Regarding VEGF-A immunoreactivity, gamma irradiated parotid gland showed intense expression and significantly higher area that has persisted up to sixteen weeks. Pilocarpine or quercetin treatment was associated with lesser degree and significantly lower area percent of VEGF-A expressionas compared to the radiation group. Pilocarpine or quercetin has some improvement for the injurious effects of radiation on salivary glands so, they can be used as radioprotective agents before radiotherapy schedules to decrease radiation side effects on salivary glands.

**Keywords:** Gamma-irradiation, Pilocarpine, Quercetin, Parotid, Vascular endothelial growth factor.

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Pilocarpine is a naturally occurring compound derived from the South American shrub, the pilocarpus jaborandi. This alkaloid plant is a cholinergic parasympathomimetic agent that acts as muscarinic agonist with a mild beta-adrenergic activity in addition to its ability to induce smooth muscle contraction and exocrine glands (Ferguson, 1993). Many studies demonstrated the efficacy of pilocarpine for the treatment of xerostomia caused by Sjogren’s syndrome, radiotherapy or drug treatment. Pilocarpine is well tolerated and not only ameliorate patient quality of life, but it also inhibits the complications likewise (Vivino, 2001; Chitapanarux et al., 2008 and Loostrom et al., 2010).

Quercetin, a unique bioflavonoid, is found in fruits, vegetables, grains, bark roots, stem, flowers, tea and others (Crozier et al., 2009). It is considered a powerful antioxidant flavonoids against reactive oxygen species, produced during the normal oxygen metabolism or are induced by the exogenous damage (De Groot, 1994). In addition, it possesses anti-inflammatory (Comalada et al., 2005), vasodilatory (Nishida & Satoh, 2009) and angiogenic effects (Sumi et al., 2013). Quercetin administration, not only improves the radiation-induced impaired salivary secretion, but it may also be an efficient way to maintain healthy salivary secretion (Takahashi et al., 2015). So, the purpose of this study is to investigate the efficiency of pilocarpine and quercetin as protector agents against radiation induced parotid glands damage.

Experimental

Forty-five adult male Sprague- Dawley rats weighing 150-250g were used in this study. The experiment was performed after accommodation period in the laboratory environment for 7 days. The rats were housed in a room with controlled temperature (25°C±2°C), humidity (50%±5%), about 12h light/dark cycle and were fed on chew and water ad libitum. The experimental protocol followed the rules and regulations of the animal experimental studies approved by Ethical Committee of Faculty of Dental Medicine for Boys, AL-Azhar University, including their facilities diet and method of scarification.

The whole body irradiation of animals was performed at the National Center for Radiation Research and Technology (NCRRT), Cairo, Egypt, using Gamma cell 40. All animals were exposed to a dose of 6Gy (137Cs) at dose rate of 0.43Gy/min.

The animals were divided randomly into three main groups (n= 15). Radiation group (R) where the rats were exposed to a single whole body 6Gy γ-irradiation. Pilocarpine-radiation group (PR) and Quercetin-radiation group (QR) where the rats were injected intraperitoneally with a single dose of pilocarpine (0.2mg/kg) (Sigma Chemical Co., St. Louis, MO), quercetin (1.25g/kg) (Sigma Chemical Co., St. Louis, MO) dissolved in 1ml of normal saline, 30min later they were exposed to a single whole body 6Gy γ-irradiation.

At 1, 4 and 16 weeks after radiation exposure, 5 rats from each group were euthanasilously decapitated. Then, the parotid glands were carefully dissected out and immediately fixed in neutral calciformol solution for 72h. The salivary glands were embedded in paraffin wax and paraffinized tissue sections were cut at 5-7 μm thick and prepared for hematoxylin and eosin stain for histological evaluation and immunohistochemical reaction using the avidin-biotin technique for demonstration of the Vascular Endothelial Growth Factor A (VEGF-A) (Okada et al., 1999) were used to detect the vessels changes. The immunostained sections were digitized using a Zeiss Mirax automated slide scanner with an objective of ×20 magnification. The VEGF-A expression was evaluated in 5 fields/slide, three slides/group using a score corresponding to the area percent of the positive cells. The digital image analysis was performed by importing the Mirax files into the image analyzer software Visiopharm integrater system (VIS).

The data were tabulated, coded then analyzed using the computer program SPSS (Statistical package for social science) version 17.0 for analysis of the immune expression of VEGF. The measured values were expressed as mean values ± standard deviation. The statistical importance of the difference in these values between the different groups was estimated using one-way analysis of variance (ANOVA). A (P) value less than 0.05 was considered significant.

Results

Histological examination

One week post-irradiation

Gamma-irradiated parotid glands expressed many destructive changes including a massive loss of acinar architecture, sever acinar atrophy,
vacuolization and degeneration. From the cellular point of view, the acini showed nuclear pleomorphism and hyperchromatism. The destructive manifestations of the ducts included loss of cellular architecture, cell vacuolization and perinuclear hallow (Fig. 1 A). The parotid glands of pilocarpine-radiation group showed a relatively preserved acinar outline and architecture. Some acini showed a mild acinar atrophy with a widen interacinar space and few degenerated acini were also detected while, the nuclei were normal in size and shape and the duct system demonstrated almost intact architecture (Fig. 1 B). The parotid glands of the quercetin-radiation group revealed almost a normal outline and architecture of the acini. Moderate acinar atrophy and degenerative changes were confined to discrete areas in the gland. The acinar cells showed normal- sized and stained nuclei while, the ducts were normal with slight architectural distortion (Fig. 1 C).

Four weeks post-irradiation
The irradiated parotid glands showed slightly less intense degenerative changes compared to those seen after one week post- irradiation. The acinar vacuolization was markedly reduced, moderate acinar atrophy were seen throughout the gland, while the degeneration of acini was restricted to less and discrete areas. The nuclear pleomorphism and hyperchromatism were persisted to slightly less distributed extent. The ducts showed some cellular vacuolization, while the others demonstrated a loss of normal architecture and degeneration (Fig. 2 A). The pilocarpine-radiation treated parotid glands demonstrated a better integrity of the structural architecture of the gland than after one week post-irradiation. The acinar atrophy was only restricted to few acini, while the acinar degeneration was almost absent, whereas, some acinar cells were still exhibiting nuclear pleomorphism. The ducts showed an intact architecture in the majority of the gland, while scanty ducts showed a structural breakdown. The glands also exhibited a slight periductal inflammatory cell infiltration and intense fibrosis (Fig. 2 B). The glands of the quercetin-radiation group demonstrated more or less normal acini with a slight widening of interacinar space while, few acinar cells appeared with nuclear pleomorphism and the ducts showed almost intact architecture (Fig. 2 C).

Sixteen weeks post-irradiation
Gamma-irradiated glands have relatively restored the normal acinar architecture, but with moderate acinar atrophy and absence of acinar degeneration. The nuclear changes were markedly diminished and manifested as a slight nuclear hyperchromatism. Most of the ducts showed a normal cellular architecture with absent ductal cell vacuolization and degeneration (Fig. 3 A). The parotid glands of pilocarpine-radiation group exhibited tilt toward the normal architecture. The acini demonstrated normal outline and architecture. The nuclei showed normal size, shape and staining. The ducts showed intact architecture and well-arranged ductal cells (Fig. 3 B). In the quercetin-radiation group, the glands showed relatively preserved acinar outline and architecture. The nuclei of the acinar and ductal cells were uniform in size and stain while, the ducts also have preserved architecture in most areas of the gland with neither vacuolization nor degeneration (Fig. 3 C).
Immunohistochemical examination

One week post-irradiation

Regarding the radiation group, there were an intense positive immunoreactivity to VEGF expression in the duct, whereas the serous cells showed a moderate to severe reaction. The mean area percent of VEGF expression was 21.38±3.1. However, there was a moderate immunoreactivity to VEGF in the ductal and most of the acinar cells of the parotid glands of pilocarpine-radiation group. The mean area percent of VEGF expression (18.25±3.3) was significantly lower than those of the radiation group. on the other hand, the glands of the quercetin-radiation group showed similar results to the previous one, where the immunoreactivity to VEGF in both the ductal and acinar cells were moderate. The mean area percent of VEGF expression (17.01±4.1) showed a significant decrease compared to the radiation group. It was worth noting that no significant difference occurred in respect to VEGF expression between both treatments groups (Fig. 4).

Four weeks post-irradiation

There was an intense immunoreactivity to VEGF in the ducts of the gamma-irradiated glands. Some acini expressed a moderate VEGF reaction while the others were negatively reactive. The mean area percent of VEGF was 17.69±3.9. The glands of pilocarpine-radiation group had a mild to moderate VEGF reaction in the ductal cells, while the acinar cells exhibited a negative reaction. The mean area percent of VEGF expression (12.91±2.9) was significantly decreased as compared to the radiation group. For the quercetin-radiation group, the whole gland demonstrated a moderate VEGF reactivity. The mean area percent of VEGF (14.5±2.4) was significantly reduced compared to the radiation group. Similar to the previous date, no significant difference between treatments groups could be detected (Fig. 4).
Sixteen weeks post-irradiation

The whole gland of radiation group showed a mild immunoreactivity to VEGF and the mean area percent of VEGF was 10.54±4.6. The glands of the pilocarpine-radiation and quercetin-radiation groups revealed a mild VEGF reactivity that was restricted to the ductal cells only while the acinar cells showed a negative reaction. The mean area percentages of VEGF were 6.42±4.6 and 6.82±3.4, respectively which were significantly decreased compared to the radiation group without a significant difference between both groups (Fig. 4).

Discussion

Radiation therapy to the head and neck area is one among causes of impaired salivary secretion (Moutsopoulos, 1994). Salivary glands were the most affected organ by radiation in this area however, radiation induced alterations which were variable among the different types of the glands. The present study demonstrated many degenerative changes in the parotid salivary glands when they become exposed to radiation. The cytoplasmic vacuoles detected in the present study in either the acinaror duct cells of parotid glands were comparable to those demonstrated by Sagowski et al. (2004). The observed atrophic changes of serous acini were also documented by Onodera et al. (2006). The acinar atrophic changes had been attributed to radiation-induced apoptosis (Guchelaar et al., 1997), or could be attributed to mast cells activation and release of their secretory products (Henriksson et al., 1994). The detected nuclear pleomorphism and hyperchromatism were also demonstrated in the nuclei of irradiated submandibular glands (Urek et al., 2005).

In the present study, the pilocarpine and quercetin administration before radiation exposure have ameliorated, but did not completely prevent radiation-induced pathological alterations along the experimental periods. The radioprotective effect of pilocarpine and quercetin treatment was manifested by the decreased atrophic changes, acinar cells degeneration, nuclear pleomorphism and hyperchromatism. The relative preservation of the parotid gland architecture due to pilocarpine administration could explain the improved salivary gland function in rats when degranulation was stimulated by pilocarpine administration prior to radiation exposure (Nagler, 2003). Clinically, the administration of pilocarpine during radiation therapy has reduced the development of xerostomia for months even after discontinuation of pilocarpine (Zimmerman et al., 1997).

The improvement of post-radiation xerostomia was more obvious in patients with lower baseline saliva secretion rates (Johnson et al., 1993). In the same way, the quercetin was found to suppress effectively radiation-induced salivary gland impairment as well as to reinforce the normal salivary gland secretion. This could be attributed

Fig. 4. A histogram representing the mean area percent of VEGF expression in parotid glands of different groups throughout the experiment (* means significant difference against radiation group).
to inhibition of oxidative stress and inflammation associated with radiation exposure, enhancement of calcium uptake and up regulation of aquaporin 5 expression (Takahashi et al., 2015).

In the present study, the irradiated parotid glands showed an intense expression of VEGF. Whereas either the pilocarpine or quercetin administration has reduced the intense expression of VEGF induced by gamma-irradiation exposure. A similar result was obtained by Witte et al. (1989) who found that radiation has induced the release of endothelial cell-derived growth factors. They added that these factors may be involved in the pathogenesis of both the early vascular damage and the late fibrosis which represents a prominent feature of late radiation damage in normal tissues.

Conclusions

Gamma irradiation has deleterious effects on the histological structure of the parotid salivary glands. Pilocarpine, a parasympathomimetic agonist, has considerably improved the injurious radiation effects on salivary glands. In addition, quercetin, a natural radioprotective agent, has a relative protective effect against radiation induced damage. Therefore, they could be used before radiotherapy schedules to decrease radiation side effects on salivary glands.

Disclosure of Interest

The authors report no conflicts of interests.

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تقييم هستولوجي و مناعى لتأثير البيلوكاربين و الكويرسيتين على الغددالنكفية المشعة

بأشعة جاما

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أجريت هذه الدراسة بهدف إيضاح تأثير كل من البيلوكاربين والكويرسيتين على علاج الأضرار الوظيفية المستحدثة بالأشعة على الغدة النكافية. وقد اعتمد البحث على فحص هذه الغدة هستولوجي ومناعيا. وقد أجرى هذا البحث على 45 فأراً ذكرًا أبيضًا. تم تقسيمهم إلى 3 مجموعات بكل مجموعة 15 فأراً وهي كالآتي:

- المجموعة الأولى: حيث تم تعرض جسم الفأر كله لأشعة جاما بجرعة 1 مجم/كجم من البيلوكاربين ثم تعرض جسم الفأر كله لأشعة جاما بجرعة 6 مجم/كجم من البيلوكاربين.
- المجموعة الثانية: حيث تم حقن الفئران داخل الغشاء البريتوني بجرعة 0.2 مجم/كجم من البيلوكاربين ثم تعرض جسم الفأر كله لأشعة جاما بجرعة 6 مجم/كجم من الكويرسيتين ثم تعرض جسم الفأر كله لأشعة جاما بجرعة 1.25 مجم/كجم من الكويرسيتين.

استخرجت الغدة كلها بمجرد نهاية الفحص. ثم وُجهت إلى قسم علم الأiouية الدموية في الجامعة للعشر الزمانية، وتم تعبير عن النتائج كالتالي:

- قد تم الكشف عن عامل النمو للأوعية الدموية في الغددPlanet.

وكانت النتائج متناوبة في البعض من المجموعات، حيث ظهرت بعض الخلايا الافرازية ضامرة أو متحللة تماماً مع وجود فجوات داخل الخلايا والأنويات. وخصوصاً في المجموعة التي عُلِجت بالبيلوكاربين والكويرسيتين. وقد تمت صباغة الشرائح بالهيماتوكسلين والأيوسين بغرض تقييمها هستولوجي. الأبيض من الأوعية الدموية مشبعة بالخلايا المناعية. أما بالنسبة لبيئات النسيج الطبيعي، فقد تبين أن الخلايا المناعية يوجد فيها الكثير من الخلايا الثانوية، وكثير منها يتواجد في الخلايا الافرازية. وخصوصاً في المجموعات التي عُلِجت بالبيلوكاربين والكويرسيتين. أما بالنسبة لبيئات النسيج الطبيعي، فقد تبين أن الخلايا المناعية يوجد فيها الكثير من الخلايا الثانوية، وكثير منها يتواجد في خلايا النكفية المشعة. وخصوصاً في المجموعات التي عُلِجت بالبيلوكاربين والكويرسيتين. وقد تبين أن الخلايا المناعية يوجد فيها الكثير من الخلايا الثانوية، وكثير منها يتواجد في خلايا النكفية المشعة. وخصوصاً في المجموعات التي عُلِجت بالبيلوكاربين والكويرسيتين. وقد تبين أن الخلايا المناعية يوجد فيها الكثير من الخلايا الثانوية، وكثير منها يتواجد في خلايا النكفية المشعة. وخصوصاً في المجموعات التي عُلِجت بالبيلوكاربين والكويرسيتين. وقد تبين أن الخلايا المناعية يوجد فيها الكثير من الخلايا الثانوية، وكثير منها يتواجد في خلايا النكفية المشعة. وخصوصاً في المجموعات التي عُلِجت بالبيلوكاربين والكويرسيتين. وقد تبين أن الخلايا المناعية يوجد فيها الكثير من الخلايا الثانوية، وكثير منها يتواجد في خلايا النكفية المشعة. وخصوصاً في المجموعات التي عُلِجت بالبيلوكاربين والكويرسيتين. وقد تبين أن الخلايا المناعية يوجد فيها الكثير من الخلايا الثانوية، وكثير منها يتواجد في خلايا النكفية المشعة. وخصوصاً في المجموعات التي عُلِجت بالبيلوكاربين والكويرسيتين. وقد تبين أن الخلايا المناعية يوجد فيها الكثير من الخلايا الثانوية، وكثير منها يتواجد في خلايا النكفية المشعة. وخصوصاً في المجموعات التي عُلِجت بالبيلوكاربين والكويرسيتين. وقد تبين أن الخلايا المناعية يوجد فيها الكثير من الخلايا الثانوية، وكثير منها يتواجد في خلايا النكفية المشعة. وخصوصاً في المجموعات التي عُلِجت بالبيلوكاربين والكويرسيتين. وقد تبين أن الخلايا المناعية يوجد فيها الكثير من الخلايا الثانوية، وكثير منها يتواجد في خلايا النكفية المشعة. وخصوصاً في المجموعات التي عُلِجت بالبيلوكاربين والكويرسيتين. وقد تبين أن الخلايا المناعية يوجد فيها الكثير من الخلايا الثانوية، وكثير منها يتواجد في خلايا النكفية المشعة. وخصوصاً في المجموعات التي عُلِجت بالبيلوكاربين والكويرسيتين. وقد تبين أن الخلايا المناعية يوجد فيها الكثير من الخلايا الثانوية، وكثير منها يتواجد في خلايا النكفية المشعة. وخصوصاً في المجموعات التي عُلِجت بالبيلوكاربين والكويرسيتين. وقد تبين أن الخلايا المناعية يوجد فيها الكثير من الخلايا الثانوية، وكثير منها يتواجد في خلايا النكفية المشعة. وخصوصاً في المجموعات التي عُلِجت بالبيلوكاربين والكويرسيتين. وقد تبين أن الخلايا المناعية يوجد فيها الكثير من الخلايا الثانوية، وكثير منها يتواجد في خلايا النكفية المشعة. وخصوصاً في المجموعات التي عُلِجت بالبيلوكاربين والكويرسيتين. وقد تبين أن الخلايا المناعية يوجد فيها الكثير من الخلايا الثانوية، وكثير منها يتواجد في خلايا النكفية المشعة. وخصوصاً في المجموعات التي عُلِجت بالبيلوكاربين والكويرسيتين. فقد تبين أن الخلايا المناعية يوجد فيها الكثير من الخلايا الثانوية، وكثير منها يتواجد في خلايا النكفية المشعة. وخصوصاً في المجموعات التي عُلِجت بالبيلوكاربين والكويرسيتين. وقد تبين أن الخلايا المناعية يوجد فيها الكثير من الخلايا الثانوية، وكثير منها يتواجد في خلايا النكفية المشع