ROLE OF NITRO-OXIDATIVE STRESS IN THE PATHOGENESIS OF EXPERIMENTAL RAT PERIODONTITIS

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Abstract

Background and aims. Periodontitis is a common chronic adult condition that implicates oxidative damage to gingival tissue, periodontal ligament and alveolar bone. This study aimed at assessing the association between the nitro-oxidative stress and the periodontal tissues destructions in experimental rat periodontitis.

Methods. Periodontitis was induced in 15 male albino rats by repetitive lesions to the gingiva adjacent to the inferior incisors, performed daily, for 16 days. On D1, D3, D6, D8, and D16 the onset and evolution of periodontitis were monitored by clinical and histopathological examinations; blood was collected and serum nitro-oxidative stress was evaluated through total nitrites and nitrates, total oxidative status, total antioxidant capacity, and oxidative stress index.

Results. The results demonstrated that there was a graded and continuous increase in serum levels of total nitrites and nitrates, total oxidative status and oxidative stress index, which was consistent with the severity of periodontal destructions during periodontitis progression. However, total antioxidant capacity was not significantly influenced by the disease progression.

Conclusions. In experimental rat periodontitis, the systemic nitro-oxidative stress was associated with the severity of periodontal destructions assessed clinically and histopathologically. Therefore, systemic nitro-oxidative stress parameters might be used as diagnostic tools in periodontitis.

Keywords: nitro-oxidative stress, experimental periodontitis, pathogenesis, rat
periodontal pathogenic microorganisms is the recruitment
and activation of polymorphonuclear neutrophils (PMN)
and macrophages [3,4]. Activated PMN produce a large
amount of reactive oxygen species (ROS), which lead
to local oxidative stress and mediate periodontal tissues
destructions [5,6].

Recently, abundant evidence has shown that
periodontal inflammation is highly associated with
several inflammation-related systemic diseases, such as
cardiovascular disease, chronic respiratory diseases, and
diabetes mellitus; moreover, the common denominator of
all these diseases is the implication of oxidative stress in
their pathogenesis [7-11].

However, little is known about the exact mechanism
of periodontitis development, including the correspondence
between the severity of periodontal destructions and the
serum levels of oxidative stress parameters.

It has been hypothesized that oxidative stress arising
from periodontal lesions may be an important cause of
systemic inflammation. Studies investigating the oxidative
stress biomarkers reported high levels in the peripheral
blood of periodontitis patients compared with healthy
subjects [12,13]. But the dynamic of oxidative stress
biomarkers changes during the progression of periodontitis
has not been yet assessed.

Firstly, we aimed at assessing the nitro-oxidative
damage during the progression of experimental rat
periodontitis. Secondly, we investigated the association
between the severity of periodontal tissues destructions
and serum levels of total nitrites and nitrates (NOx), total
oxidative status (TOS), total antioxidant capacity (TAC),
and oxidative stress index (OSI).

**Materials and methods**

**Chemicals**

Analytical grade chemicals were used exclusively.
Trolox (6-hydroxy-2,5,7,8-tetramethylchroan-2-carboxylic
acid), N-(1-Naphthyl) ethylenediamine dihydrochloride
(NEDD), xylenol orange [o-cresosulfonphthalein-3,3-
bis(sodium methyliminodiacetate)], ortho dianisidine,
vanadium (III) chloride (VCl₃), hydrogen peroxide (H₂O₂),
methanol, diethyl ether sulphanilamide (SULF) and ferrous
ammonium sulphate were purchased from Sigma-Aldrich
(Germany) and Merck (Germany).

**Experimental design**

The experiment was performed on fifteen adult
male Wistar-Bratislava albino rats, weighing between
200 and 250 g that were bred in the Animal Facility of
Iuliu Hatieganu University of Medicine and Pharmacy.
The rats were kept in a room with controlled temperature
(21±1°C) and humidity (50-55%) and a 12h light-12h dark
cycle. Animals were fed with standard pellet (Cantacuzino
Institute, Bucharest, Romania) basal diet and water *ad
libitum*.

Periodontitis was induced in rats by repetitive
acute lesions to the gingival epithelium around the inferior
incisors. The rats were anesthetized by intramuscular
injection of 50 mg/kg body weight (b.w.) ketamine and 20
mg/kg b.w. xylazine. The lesions were performed with a
Gracey curette placed into the gingival sulcus and moved
circularly to detach the junctional epithelium from the
tooth surface. These interventions were repeated daily,
for sixteen days and the progression of the periodontal
inflammation was monitored daily (D1 to D16).

On D1, D3, D6, D8, D16, clinical and histopathological
examinations were performed, and blood samples were
harvested for serum nitro-oxidative stress tests.

The experimental protocol was approved by the
Institutional Animal Ethical Committee of the Iuliu
Hatieganu University of Medicine and Pharmacy, Cluj-
Napoca (approval no. 107/06.03.2015). At completion of
the study, rats were euthanized by cervical dislocation.

**Clinical examination**

Two features were evaluated: the periodontal
inflammation and gingival bleeding (GB). Periodontal
inflammation was assessed by the aspect of the gingiva:
color, consistency and volume. The scores were the
followings: 0 – no signs of inflammation; 1 – mild; 2 –
moderate; 3 – severe; 4 – periodontitis. GB was assessed
upon probing the gingival sulcus with a periodontal probe,
and the scores were: (−) – absent or (+) – present; when GB
occurred upon gentle removal of the debris on the gingival
surface or spontaneously, it was scored with (++) and (+++)
respectively.

**Histopathological examination**

On D1, D3, D6, D8 and D16, five rats were sacrificed
and the frontal parts of the mandibles were harvested.
The specimens were immersed in 10% buffered formalin
solution for fixation. The gingiva adjacent to the inferior
incisors (including the free gingiva, the interdental papilla
and the attached gingiva) was prelevated and submitted to
further routine histological processing. The tissues were
embedded in paraffin blocks and serial 5μ sections in the
bucco-lingual direction were obtained. The histological
sections were stained using Goldner’s trichrome technique
and examined under an Olympus BX41 light microscope
for descriptive evaluation. The following aspects were
assessed: the necrosis and ulceration of the gingival
epithelium, destruction of the extracellular matrix in the
chorion, the types of inflammatory cells and the
microcirculatory changes.

**Oxidative stress evaluation**

Blood was collected by retro-orbital puncture,
without anticoagulant; coagulated blood was centrifuged
and the serum was submitted to nitro-oxidative stress test.
TOS, TAC and NOx were measured in the serum; OSI was
also calculated.

Serum samples were filtered through 10-kDd filters
(Sartorius AG, Goettingen, Germany) and contaminant
proteins were extracted with a 3:1 (v:v) solution of
Nitric oxide (NO) synthesis was indirectly determined using the Griess reaction. Nitrate was reduced to nitrite by combining 100 μL of filtered and extracted serum supernatant with 100 μL of 8 mg/mL VCl$_3$; then, the Griess reagents were added, 50 μL of SULF (2%) and 50 μL of NEDD (0.1%). The sample was incubated 30 minutes at 37°C and the absorbance was read at 540 nm. Serum NOx concentration (expressed as nitrite μmol/L) was determined using a sodium nitrite-based curve [14].

TOS was measured using a colorimetric assay that measured the oxidation of ferrous ion to ferric ion in the presence of various reactive oxygen species in an acidic medium [15]. Then, ferric ions were detected by reaction with xylene orange. Assay measurements were standardized with hydrogen peroxide (H$_2$O$_2$) used as the oxidative species, and the results are expressed in μmol H$_2$O$_2$ Equiv./L.

TAC was measured using a colorimetric assay that monitored the rate of hydroxyl radical production by the Fenton reaction the changes in the absorbance of colored dianisidyl radicals [16]. Upon addition of a serum sample, the antioxidant present in the serum suppressed the oxidative reactions initiated by hydroxyl radicals. Inhibition of dianisidyl oxidation prevented the subsequent colour change, thus measuring the serum TAC. This assay was calibrated using trolox and results were expressed as mmol trolox Equiv./L.

OSI was calculated as the ratio of the TOS to the TAC: OSI (Arbitrary Unit) = TOS (μmol H$_2$O$_2$ Equiv./L) / TAC (nmol trolox Equiv./L) [17].

The spectroscopic measurements were performed using a Jasco V-530 UV-Vis spectrophotometer (Jasco International Co., Ltd., Tokyo, Japan).

Statistical methods

All results were expressed as mean ± standard deviation (SD). Normal distribution was assessed using Shapiro-Wilk test. Statistical comparisons between the moments were made using one-way ANOVA, followed by Tukey’s post hoc test. $p$-values < 0.05 were regarded as statistically significant. Pearson’s and Spearman’s correlation tests were performed in order to evaluate statistical correlation. Data was analyzed using R 3.0.2 - software environment for statistical computing and graphics.

Results

Clinical and histopathological examination

On D1, clinically, gingiva had a normal appearance; periodontal inflammation score = 0, GB: (-). The gingiva was pale pink, with a smooth surface; the gingival sulcus was virtual and gingiva was firmly attached to the cervical zone of the teeth and to alveolar bone. Histologically, the gingival epithelium and lamina propria showed no signs of inflammation (Figure 1).

On D3, clinically mild gingivitis was observed; periodontal inflammation score = 1, GB: (+). The gingiva was pale pink, with a smooth surface; the gingival sulcus was virtual and gingiva was firmly attached to the cervical zone of the teeth and to alveolar bone. Histologically, the gingival epithelium and lamina propria showed no signs of inflammation (Figure 1).

On D3, clinically mild gingivitis was observed; periodontal inflammation score = 1, GB: (+). The gingiva showed discrete oedema and erythema. On the gingival surface, the superficial linear incisions induced experimentally were visible. Histopathologically, the initial lesion was characterized by the following features: epithelium underwent necrosis in the zone where it was detached from the surface of the tooth; in the lamina propria, the inflammatory reaction was characterized by moderate infiltration with PMN, vasodilatation and slight, mainly perivascular oedema (Figure 2).

Figure 1. Clinical and histological aspect of the gingiva on D1. A: healthy gingiva. B: the gingival stratified squamous parakeratinized epithelium and the lamina propria consisting of connective tissue. (Goldner’s trichromic stain).
On D6, clinically moderate gingivitis could be seen; periodontal inflammation score = 2, GB: (+). Gingival oedema and erythema increased, the gingival sulcus was a real and narrow space, as the gingiva was detached from the cervical zone of the teeth. The ulcerations were covered by debris of which detachment during examination caused bleeding. Histopathological examination identified the early lesion: the zone of necrotic epithelium was wider, and the underlying connective tissue was exposed. In the lamina propria, adjacent to the necrotic epithelium, the inflammatory infiltrate with PMN and macrophages was denser; the oedema was diffuse, but more intense in some areas, where the extracellular matrix lysis occurred (Figure 3).

On D8, clinically severe gingivitis was diagnosed; periodontal inflammation score = 3, GB: (++). Gingival erythema and oedema were more intense, the ulceration was deeper, covered by a larger amount of debris. Moreover, the wide gingival detachment and retraction caused the root exposure on a length equal to 1/3 of the clinical crown of the teeth. Histopathologically, the established lesion consisted of necrosis extended both into the epithelium and into the superficial layer of lamina propria. In the profound layers, the areas where the extracellular matrix lysis occurred were occupied by intense oedema and diffuse hemorrhage. The cells in the inflammatory infiltrate were mainly PMN, macrophages and lymphocytes (Figure 4).

On D16, clinically periodontitis was observed; periodontal inflammation score = 4, GB: (++++). Gingiva was cyanotic, increased in volume, and the increased gingival fragility resulted in spontaneous bleeding. The gingival ulcerations were profound and covered by abundant debris. Moreover, the gingiva was severely detached from the teeth and the root exposure was approximately equal to the clinical crown length. Histopathologically, the advanced lesion was characterized by extensive necrosis of the epithelium. In the lamina propria; the extracellular matrix underwent intense lysis and was replaced by the inflammatory infiltrate with PMN, macrophages and lymphocytes, oedema and hemorrhage; in some areas, necrosis also occurred in the connective tissue. The surface of the lesion was covered by abundant fibrinoid deposits, enclosing numerous PMN, macrophages, lymphocytes and degenerated erythrocytes (Figure 5).

The evaluation of the oxidative stress
We assessed the serum levels of NOx, TOS, TAC and OSI during periodontitis progression (Table I).

Analyzing the progression of the serum parameters in D1, D3, D6, D8, and D16, excepting TAC which did not change significantly (p>0.05), NOx, TOS and OSI constantly increased: abruptly until D3, and smoothly thereafter (Figure 6).

NOx serum levels were significantly lower on D1 compared with D3, D6, D8, and D16 (p<0.001). Additionally, NOx serum levels were significantly lower on D3, D6, and D8 compared with D16 (p<0.001).

TOS serum levels were significantly lower on D1 compared with D3 (p < 0.05), D6 (p < 0.05), D8 (p < 0.001), and D16 (p < 0.05). Moreover, TOS serum levels were significantly higher on D8 compared with D3 and D6 (p < 0.05).

OSI serum levels were significantly lower on D1 compared with D3 (p < 0.05), D6 (p < 0.05), D8 (p < 0.001), and D16 (p < 0.05). Additionally, TAC serum levels were significantly higher on D8 compared with D3 and D6 (p < 0.05).

During the 16 experimental days OSI was highly correlated with TOS (r = 0.99).

On D6, NOx was correlated with TOS (r = 0.65), and OSI (r = 0.64).
Figure 3. Clinical and histological aspect of the gingiva on D6. A: moderate gingivitis: gingiva is inflamed and detached from the teeth; the ulcerations were covered by debris. B: early lesion: more extensive epithelial necrosis, diffuse oedema and denser inflammatory infiltrate with PMN and macrophages. (Goldner’s trichromic stain).

Figure 4. Clinical and histological aspect of the gingiva on D8. A: severe gingivitis: gingival inflammation and oedema were more intense; retraction and dental root exposure could be noticed; larger ulcerations were covered by debris. B: established lesion: areas of necrosis in the epithelium and lamina propria, diffuse oedema and hemorrhage, dense inflammatory infiltrate with PMN, macrophages and lymphocytes. (Goldner’s trichromic stain).

Figure 5. Clinical and histological aspect of the gingiva on D16. A: periodontitis: gingival cyanosis, ulcerations covered by debris and spontaneous bleeding; extensive gingival retraction and destruction of the underlying alveolar bone and periodontal ligament. B: advanced lesion: the necrotic epithelium was replaced by fibrinoid deposits; in the lamina propria, dense inflammatory infiltrate, oedema and diffuse hemorrhage, and areas of necrosis could be observed. (Goldner’s trichromic stain).
Discussion
The results of the study proved that there was a good correlation between the clinical, histological and nitro-oxidative stress parameters during the progression of experimental rat periodontitis.

Periodontitis involves complex pathogenic mechanisms that are part of the host immune-inflammatory response to the presence of microbial plaque [18,19]. These mechanisms are characterized by the production of cytokines, enzymes and free radicals in the periodontal tissues, which results in progressive gingival inflammation, periodontal ligament destruction, alveolar bone resorption, and eventually lead to teeth loss [20]. Moreover, the chronic periodontal inflammation is associated with increased systemic nitro-oxidative stress, responsible for numerous disorders such as rheumatoid arthritis, chronic kidney dysfunction, ischemic heart disease, premature births and others [21,22].

Nitro-oxidative stress is induced by an excessive production of ROS [23], associated with increased synthesis of NO by the inducible nitric oxide synthase (iNOS) [24]. The increased production of reactive species might be associated with a decrease in antioxidant capacity. This is why the nitro-oxidative stress should be evaluated by measuring ROS, NO, and TAC [25].

In the present study, in order to assess the severity and progression of experimental rat periodontitis, we employed global nitro-oxidative stress tests. NO synthesis was measured indirectly by the quantification of serum nitrites and nitrates [26], ROS were measured by the

Table 1. Comparison between time points during periodontitis progression.

| Moment | NOx (µmol/L) | TOS (µmol Equiv H$_2$O$_2$/L) | TAC (mmol Equiv TROLOX/L) | OSI |
|--------|--------------|-------------------------------|---------------------------|-----|
| D1     | 34.111±1.826 | 31.1±1.703                    | 1.09±0.001                 | 0.285±0.015 |
| D3     | 96.029±2.493 | 39.235±2.508                  | 1.0905±0.001              | 0.359±0.023 |
| D6     | 101.95±3.34  | 40.93±1.914                   | 1.0902±0.001              | 0.375±0.017 |
| D8     | 101.773±3.234| 46.354±1.286                  | 1.0898±0.001              | 0.425±0.01 |
| D16    | 114.409±6.948| 44.216±3.962                  | 1.0897±0.001              | 0.405±0.036 |

Values are expressed as Mean ± SD of five determinations. NOx = total nitrites and nitrates; TOS = total oxidative status; TAC = total antioxidant capacity; OSI = oxidative stress index, in 5 moments: D1- day 1, D3 – day 3, D6 – day 6, D8 – day 8, D16 – day 16.

Figure 6. Nitro-oxidative stress parameters changes during 16 days of periodontitis progression. NOx = total nitrites and nitrates; TOS = total oxidative status; TAC = total antioxidant capacity; OSI = oxidative stress index, in 5 moments:D1- day 1, D3 – day 3, D6 – day 6, D8 – day 8, D16 – day 16.
assessment of TOS, and antioxidant capacity was assessed by TAC [14-16, 25]. Since the final result depends on the proportion between ROS and the anti-oxidant mechanisms, determination of OSI was necessary [16].

Our study was the first to assess the nitro-oxidative stress parameters as indicators of the progression and severity of periodontal destructions in experimental rat periodontitis. Previous studies evaluated the injury only at the end of the experiment [27,28].

The clinical and histopathological examinations recorded, stage by stage, the structural changes throughout the initiation and progression of gingivitis and periodontitis [29]. These findings were correlated with the serum levels of NOx, TOS, TAC and OSI, which are considered to be markers of the nitro-oxidative stress [30,31] and may be used to stepwise evaluate the progression of the disease [32].

At baseline, the absence of periodontal inflammation was associated with reduced serum levels of NOx, TOS and OSI.

Clinically, on D3, the gingiva showed discrete signs of inflammation, which were consistent with the histopathological changes. Gingival erythema and oedema were the result of the vasodilatation and increased vascular permeability, whereas bleeding was the consequence of capillary fragility. The incisions induced experimentally disrupted the epithelial barrier and initiated the penetration of bacteria into the lamina propria and the consequent recruitment and activation of PMN and macrophages. Serum levels of NOx, TOS and OSI significantly increased compared with D1.

On D6, the gingival erythema and oedema were more intense, due to increased vascular dilatation and permeability. The epithelial ulceration was wider, allowing a higher bacterial invasion into the lamina propria. The inflammatory process extended, as a larger number of inflammatory cells were attracted into the area adjacent to the necrotic epithelium; moreover, the first signs of extracellular matrix lysis occurred. Epithelium detachment at the cervical line of the teeth initiated the periodontal ligament damage. Serum levels of NOx, TOS and OSI were not significantly different from D3, but significantly increased compared with D1.

Furthermore, on D8, destruction of both epithelium and lamina propria intensified, and the necrotic connective tissue was replaced by a dense inflammatory infiltrate and diffuse haemorrhage. The gingival retraction and detachment from the tooth surface suggested the ongoing periodontal ligament damage and alveolar crest resorption. Serum levels of TOS and OSI were significantly increased compared with D6 and D1; moreover, NOx was significantly higher compared with D1.

Finally, on D16, the rats displayed clinical signs of periodontitis: severe gingival inflammation accompanied by cyanosis and spontaneous bleeding. The histopathological features characteristic for advanced lesion associated with periodontitis consisted in: necrosis in the epithelium and lamina propria, lysis of the extracellular matrix, diffuse oedema, haemorrhage and infiltration with lymphocytes, macrophages and neutrophils. Additionally, extensive gingiva detachment and retraction indicated that the supporting tissues also underwent degenerative changes. Since the periodontal ligament and the alveolar bone were affected, the periodontal inflammation was no longer confined to the gingiva, therefore gingivitis developed into periodontitis. NOx significantly increased compared with D8 and D1, whereas TOS and OSI significantly increased compared with D1.

The clinical and histopathological features described in our experimental rat periodontitis were consistent, in evolution, with the specific aspects reported in human chronic periodontitis [33, 34]. On the other hand, our findings regarding the association between periodontitis and the high levels of NOx, TOS and OSI were similar to other experimental and clinical studies [27,28,35,36].

Recent evidence showed a link between the activation of phagocytes (PMN and macrophages) and the pathways leading to periodontal damage [37-39]. Periodontal inflammation is initiated by the chemotactic signals released from the plaque bacteria, which result in the recruitment of PMN into the sulcular epithelium and the gingival crevice [4,6]. As inflammation progresses, a vast number of phagocytes (represented mainly by PMN, but also including macrophages) will be attracted into the periodontal tissues [40]. Our observations indicated that during the experimental rat periodontitis, PMN infiltrate increased constantly.

The key to neutrophil-mediated tissue destruction resides in the homeostatic imbalance caused by three mechanisms associated with PMN response to the oral biofilm: the impaired neutrophils, the hyperactive neutrophils and chronic recruitment and activation of normal neutrophils [41]. The weight of evidence supports two main ideas regarding the neutrophils dysfunction: genetically impaired or defective phenotypes are associated with a predisposition to aggressive periodontitis whereas hyperactive, primed neutrophils may be implicated in chronic periodontitis [41]. Recently, it was suggested that the control of periodontal pathogens can be accomplished through the chronic activation and extended longevity of normal neutrophils over many years, and thus, the increased prevalence of periodontitis with age can be explained [42]. Considering all these, it is important that in our experiment there was a mostly PMN infiltrate, explaining the progressive increase of the nitro-oxidative stress.

Neutrophils are thought to promote oedema due to their interaction with the vascular endothelium during diapedesis and through the production of various mediators, such as: arachidonic acid derivatives and chemokines [43]. Additionally, neutrophils release ROS and serine proteases capable of damaging the vascular endothelium [44,45].
In attempt to eliminate pathogens, host defence cells release various factors, which can broadly be included in two categories: enzymes and oxidative burst products [46]. Even though these factors are effective in destroying bacteria, increased levels of enzymes cause substantial damage to the periodontal tissues, whereas high levels of free radicals generate oxidative stress [6,47,48].

In vitro studies demonstrated that free radicals released by neutrophils and macrophages are capable to damage resident cells and to degrade the main constituents of extracellular matrix in the periodontal tissues. Intracellularly, ROS induce lipid peroxidation through activation of cyclooxygenases and lipooxygenases [35], and DNA damage through strain breaks and hydroxylations [31]. In glycosaminoglycans, free radicals induce chains depolymerisation, whereas in proteoglycans, they modify the amino acid functional groups leading to core protein fragmentation [49].

Moreover, ROS cause oxidation of enzymes [46], stimulate release of proinflammatory cytokines from macrophages by activating Nuclear Factor xB (NF-xB) and are also involved in alveolar bone resorption [50,51,46].

In our study, the clinical and histopathological dynamic of periodontitis was correlated with the variations in the serum levels of NOx, TOS and OSI: higher nitro-oxidative stress was associated with more severe gingival injury. Our results were consistent with other studies [12, 13,52,53].

After the first interventions on the gingiva, there was an important increase in serum levels of NOx, TOS and OSI, suggesting the initiation of local and systemic nitro-oxidative stress. Additionally, in the following days, these parameters were significantly higher compared with the baseline, proving the role of nitro-oxidative stress in the pathogenesis of experimental rat periodontitis.

The comparison between the days showed that NOx, TOS and OSI progressively increased; occasionally, there were significant differences between consequent observations, made after a period of two days (D1 - D3 and D6 - D8), but in most cases, the significant increases could be observed when there were longer periods of time between examinations (D3 - D8, D6 - D16 and D8 - D16).

At completion of the study, the most severe periodontal destructions were consistent with the highest serum levels of nitro-oxidative stress parameters. However, TAC was not influenced by the progression of the periodontal inflammation. These findings were different from the observations of a number of clinical studies that reported lower serum TAC in periodontitis patients compared with the healthy controls [46,54,55]. A recent meta-analysis by Liu et al. discussed serum levels of oxidative biomarkers and indicated that TAC, MDA (malondialdehyde), and NO were strongly associated with periodontitis [36], whereas SOD levels were not different from healthy subjects. Data reported by most studies showed that TAC levels were lower, whereas NO and MDA levels were significantly higher in peripheral blood of periodontitis patients [12,13,54,55,56].

Our results showed a correlation between the tissue damage and NOx, TOS and OSI, suggesting that these nitro-oxidative stress markers could be used as diagnostic tools for periodontitis. However, further studies are needed in order to validate the relationship between periodontitis and systemic oxidative stress.

Conclusions

In experimental rat periodontitis, the severity of periodontal damage assessed by clinical and histopathological examinations was correlated with the serum levels of nitro-oxidative stress parameters. Serum levels of NOx, TOS and OSI steadily increased throughout the progression of experimental rat periodontitis, but TAC was not significantly influenced by the disease progression.

Nitro-oxidative stress parameters might be used as diagnostic tools in periodontitis.

Acknowledgement

This paper was published under the frame of European Social Found Human Resources Development Operational Programme 2007-2013, project no. POSDRU/159/1.5/S/138776.

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