

Research Article

Transdermal Nicotine Application Attenuates Cardiac Dysfunction after Severe Thermal Injury

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Background. Severe burn trauma leads to an immediate and strong inflammatory response inciting cardiac dysfunction that is associated with high morbidity and mortality. The aim of this study was to determine whether transdermal application of nicotine could influence the burn-induced cardiac dysfunction via its known immunomodulatory effects. **Material and Methods.** A standardized rat burn model was used in 35 male Sprague Dawley rats. The experimental animals were divided into a control group, a burn trauma group, a burn trauma group with additional nicotine treatment, and a sham group with five experimental animals per group. The latter two groups received nicotine administration. Using microtip catheterization, functional parameters of the heart were assessed 12 or 24 hours after infliction of burn trauma. **Results.** Burn trauma led to significantly decreased blood pressure (BP) values whereas nicotine administration normalized BP. As expected, burn trauma also induced a significant deterioration of myocardial contractility and relaxation parameters. After application of nicotine these adverse effects were attenuated. **Conclusion.** The present study showed that transdermal nicotine administration has normalizing effects on burn-induced myocardial dysfunction parameters. Further research is warranted to gain insight in molecular mechanisms and pathways and to evaluate potential treatment options in humans.

1. Introduction

Severe burn injury activates a multitude of immunologic defense mechanisms, one of these being the massive production of proinflammatory mediators [1, 2]. Looking at the organ level, one pivotal consequence of burn trauma is cardiac dysfunction [3, 4]. For its pathogenesis the focus of scientific interest has been on proinflammatory cytokines such as TNF- α , IL-1 β , and IL-6, which have been shown to either alone or in concert exert cardiodepressive effects [1, 3, 5]. Elevated levels of IL-1 β , IL-6, and TNF- α corresponded to increased levels of creatinine kinase [5]. Various proinflammatory cytokines demonstrate peak concentrations 12 to 24 hours after trauma [1].

There is a link between the central nervous system and immunologic mechanisms [6]. The vagus nerve axis of parasympathetic activity represents one of the mechanisms that can induce anti-inflammatory effects. Its activation may contribute to preventing a hyperactivation of immune subsystems and reaction overshoot [7, 8]. Macrophages have been identified to be the key target cells for the parasympathetic anti-inflammatory effect of vagus nerve activation. A reason for this is the vagus nerve innervation of most internal organs [6, 9]. Also in other organs such as the heart, the release of TNF- α and IL-1 β was inhibited directly via stimulation of the vagus nerve [10]. Tracey and colleagues identified this signal transduction pathway and named it

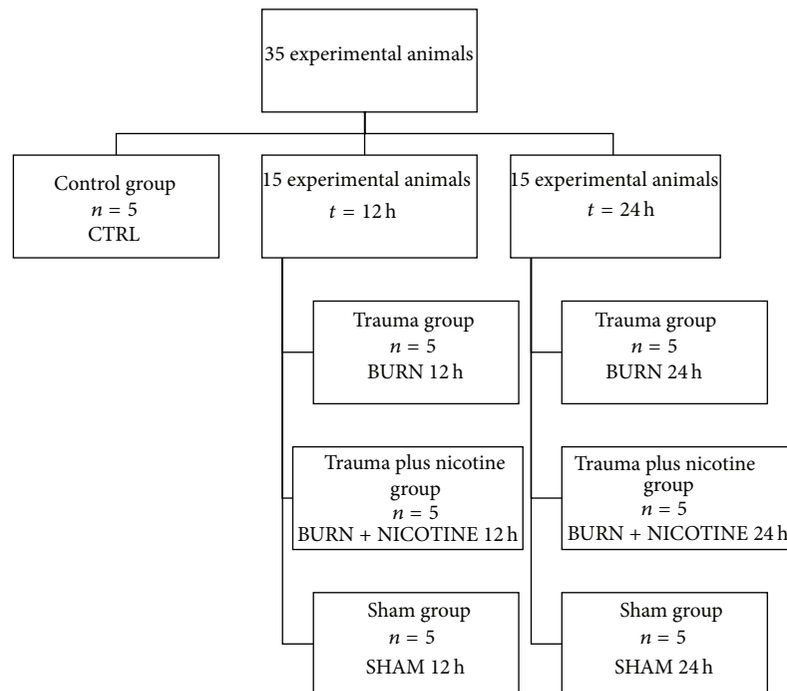


FIGURE 1: Experimental groups. The classification of 35 experimental animals into seven experimental groups with five experimental animals each is illustrated. Catheter results 12 h and 24 h after experimental burn trauma are compared to control group.

“cholinergic anti-inflammatory pathway” because of its primary neurotransmitter acetylcholine [6, 11]. Mechanistic studies have shown that the molecular basis for this pathway is the nicotinic acetylcholine receptor (nAChR), which is also present in the sympathetic part of the autonomic nervous system [6, 12]. Nicotine, a receptor agonist, can stimulate the parasympathetic anti-inflammatory mechanisms via the nAChR [12, 13]. Nicotine is well known as one of hundreds of components of tobacco smoke [14, 15]. However, it has also been used pharmacologically. Transdermal application and absorption of nicotine have been used as a noninvasive application method in a plethora of studies, verifying its anti-inflammatory potential [16–18]. Additionally nicotine application in experimental animal models of rheumatoid arthritis and autoimmune myocarditis, other pathologies based on an overwhelming inflammation, provided evidence for a potential clinical relevance of the immunomodulatory effect of nicotine [19, 20].

The present experimental study was designed to evaluate whether the transdermal application of nicotine is feasible and to evaluate potential beneficial effects on myocardial function after severe thermal injury.

2. Material and Methods

The university committee for the use and protection of animals and the Lower Saxony State Office for Consumer Protection and Food Safety approved the present study (study protocol # 05/1052). We created seven study groups with five animals each. This included one control group (CTRL). Microtip catheterization was done after 12 h or 24 h according

to the respective treatment. That resulted in the additional six groups BURN 12 h and BURN 24 h, BURN + NICOTINE 12 h, and BURN + NICOTINE 24 h and SHAM 12 h and SHAM 24 h (Figure 1). The study design, the burn trauma, and nicotine application were previously described [21].

2.1. Myocardial Function. Microtip catheterization of the left ventricle was carried out in experimental animals of the CTRL group without any previous treatment and in the BURN groups, BURN + NICOTINE groups, and SHAM groups 12 or 24 hours after experimental burn injury or sham trauma, respectively. The right carotid artery was surgically exposed by midline sternotomy. Using microsurgical techniques, a vascular incision large enough to induce the microtip catheter was created (Fa. Millar Instruments, Houston, TX, USA). The correct catheter position was confirmed by the appearance of the characteristic pressure curves and fluoroscopy imaging (Figure 2). Cardiac actions were then recorded for 5 minutes using commercially available analysis software (Chart 5, AD Instruments GmbH, Spechbach, Germany).

2.2. Statistical Analysis. We used Prism 5 software for statistical analysis (GraphPad Inc., La Jolla, CA) performing Analysis of variance (ANOVA) followed by Tukey’s post hoc test. Statistical significance was set at $p < 0.05$. The results are illustrated as means \pm standard deviation (SD).

3. Results

3.1. Blood Pressure. The results of the microtip catheter for systolic and end-diastolic blood pressure measured 12 and 24

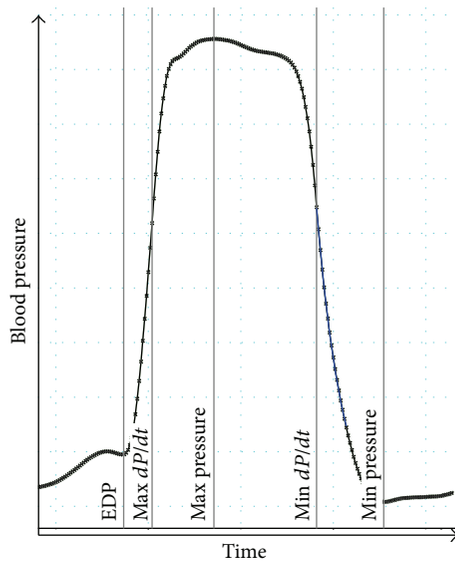


FIGURE 2: Representative blood pressure curve to illustrate measured parameters. Relevant values are marked. From left to right: end-diastolic pressure (EDP) and maximum blood pressure rise per time (Max dP/dt), systolic blood pressure (Max pressure) and maximum blood pressure drop per time (Min dP/dt). Additionally the minimum blood pressure is illustrated (Min pressure).

hours after burn trauma showed significantly decreased values for the BURN groups compared to the respective CTRL groups. For both physiological parameters analyzed, nicotine administration increased BP levels significantly compared to the untreated BURN groups except for end-diastolic blood pressure after 24 h. However, the baseline BP value of the control group was not completely reestablished by nicotine application.

After time $t = 12$ h post burn injury, the mean values of the BURN group for the systolic blood pressure were significantly decreased compared with values measured in the CTRL group (Figure 3(a), 91.1 ± 3.8 mmHg versus 113.4 ± 10.3 mmHg, black versus white bar, $p < 0.001$). Nicotine partially normalized the effect of the burn trauma and lead to a systolic blood pressure of 106.3 ± 5.5 mmHg (Figure 3(a), striped bar, $p < 0.001$). The same trend was found at $t = 24$ h after experimental burn procedure. The average systolic BP level of BURN 24 h group of 93.6 ± 5.1 mmHg was restored by nicotine application to 101.2 ± 8.6 mmHg (Figure 3(b), black versus striped bar, $p < 0.001$).

The results of the end-diastolic blood pressure are shown in Figures 3(c) and 3(d). Rats subjected to burn trauma developed significantly lower end-diastolic blood pressure than their CTRL counterparts ($p < 0.001$). The nicotine administration resulted in a significantly higher end-diastolic pressure 12 h after burn trauma whereas the difference after 24 h was statistically not relevant (Figures 3(c) and 3(d), striped versus black bars, $p < 0.001$ after 12 h and not significant after 24 h).

3.2. Systole. For the assessment of cardiac function during systole the duration of systole and contractility were measured in terms of maximum blood pressure rise per second.

The duration of the systole was on average 0.083 s in CTRL group (Figures 3(a) and 3(b), white bar). For duration of systole of BURN groups we found mean values of 0.096 s after 12 h and 0.098 s after 24 h, whereas the values of BURN + NICOTINE groups were significantly decreased (Figures 4(a) and 4(b), black versus striped bars, $p < 0.001$).

A significant increase of maximum blood pressure rise per second was observed when results of BURN groups were compared to BURN + NICOTINE groups (Figures 4(c) and 4(d), black versus striped bars, $p < 0.001$). Interestingly, SHAM and BURN + NICOTINE groups displayed slightly higher values for cardiac contractility than CTRL animals (Figures 4(c) and 4(d), grey and striped bars versus white bar). The BURN 12 h group demonstrated a value for maximum blood pressure rise per seconds of 4586 ± 859.7 mmHg/s and the BURN 24 h remained at a low level with 4653 ± 451.4 mmHg/s. In contrast, the BURN + NICOTINE groups revealed 5878 ± 576.3 mmHg/s and 6316 ± 649.9 mmHg/s, respectively, (Figures 4(c) and 4(d), striped bars versus black bars, $p < 0.05$).

3.3. Diastole. Based on the assessment of cardiac function during systole we determined the duration of diastole and the maximum blood pressure drop per second (relaxation) for the assessment of cardiac function during the diastolic phase of the cardiac cycle. With regard to the relaxation it is important that a higher negative value indicates a faster relaxation.

The duration of diastole after burn injury was on average 0.96 after 24 h and 0.99 s after 12 h (Figures 5(a) and 5(b), black bars). When burn-injured animals underwent nicotine treatment, duration of diastole was slightly increased compared to burn alone animals (Figures 5(a) and 5(b), striped versus black bars, not significant).

Animals subjected to burn trauma showed a markedly slower relaxation compared to their SHAM and CTRL counterparts (Figures 5(c) and 5(d), black versus grey and white bars, $p < 0.001$). Nicotine treatment induced a relaxation similar to sham animals (Figures 5(c) and 5(d), black versus striped bars, $p < 0.001$).

3.4. Heart Rate. Concerning the heart rate no significant differences were found 12 and 24 h after burn injury. The heart rate values ranged from 309.3 ± 3.2 beats/min (Figure 6(b), black bar) up to 315.4 ± 23.4 beats/min (Figure 6(b), striped bar).

4. Discussion

Intense burn trauma imposes life-threatening consequences to the victim. The multifactorial pathogenesis of burn-induced cardiac dysfunction has been extensively evaluated and described. This study was conducted to evaluate whether the known immunomodulatory effect of nicotine can improve cardiac function after burn trauma.

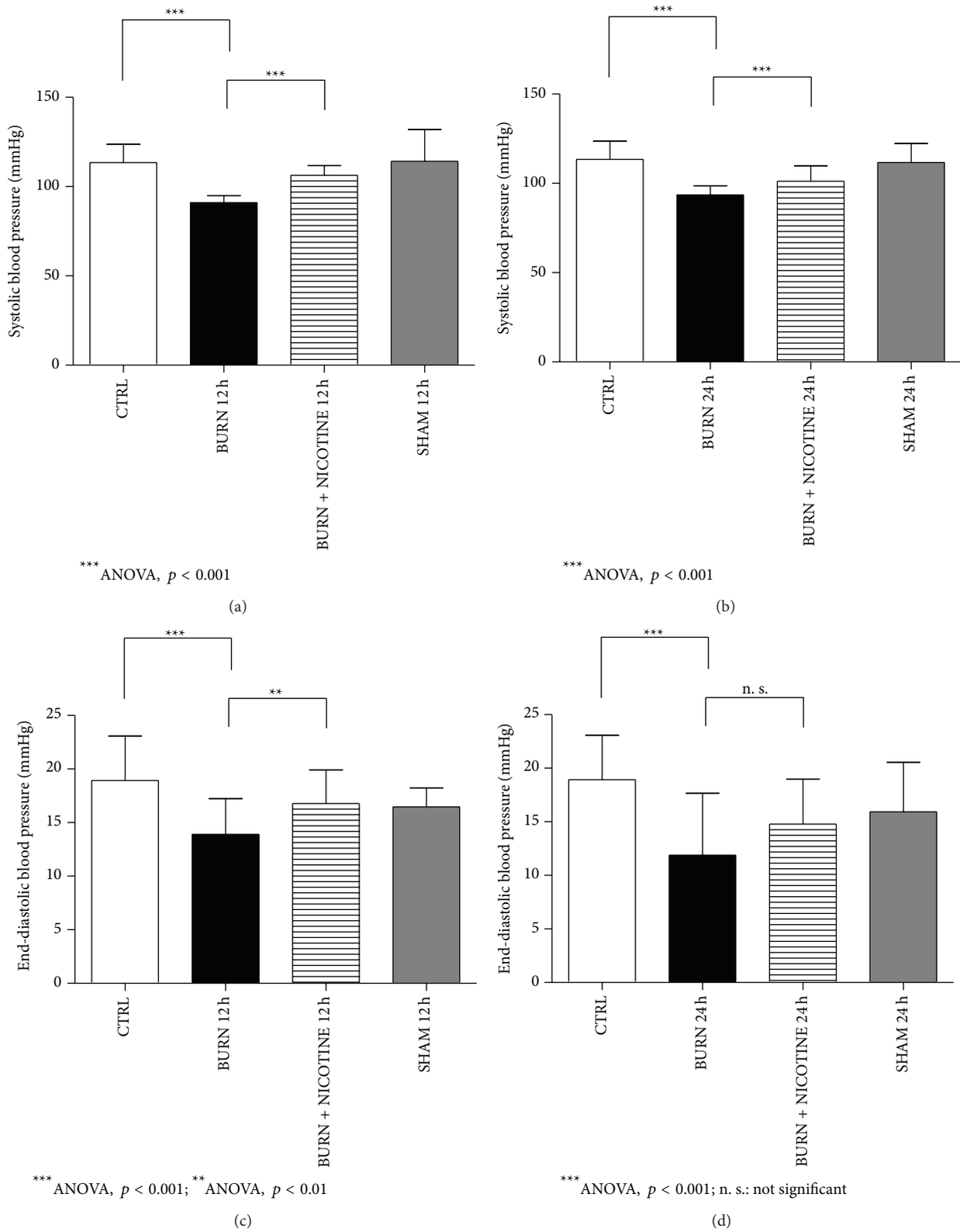


FIGURE 3: Blood pressure parameters. (a) and (b): systolic blood pressure. (c) and (d): end-diastolic blood pressure. Concerning the experimental design, note decreased BP levels in the BURN groups in contrast to the CTRL group ($p \leq 0.05$). Transdermal nicotine application at least partially restored this effect. BP values are expressed as means \pm SD. Each bar represents $n = 5$ experiments.

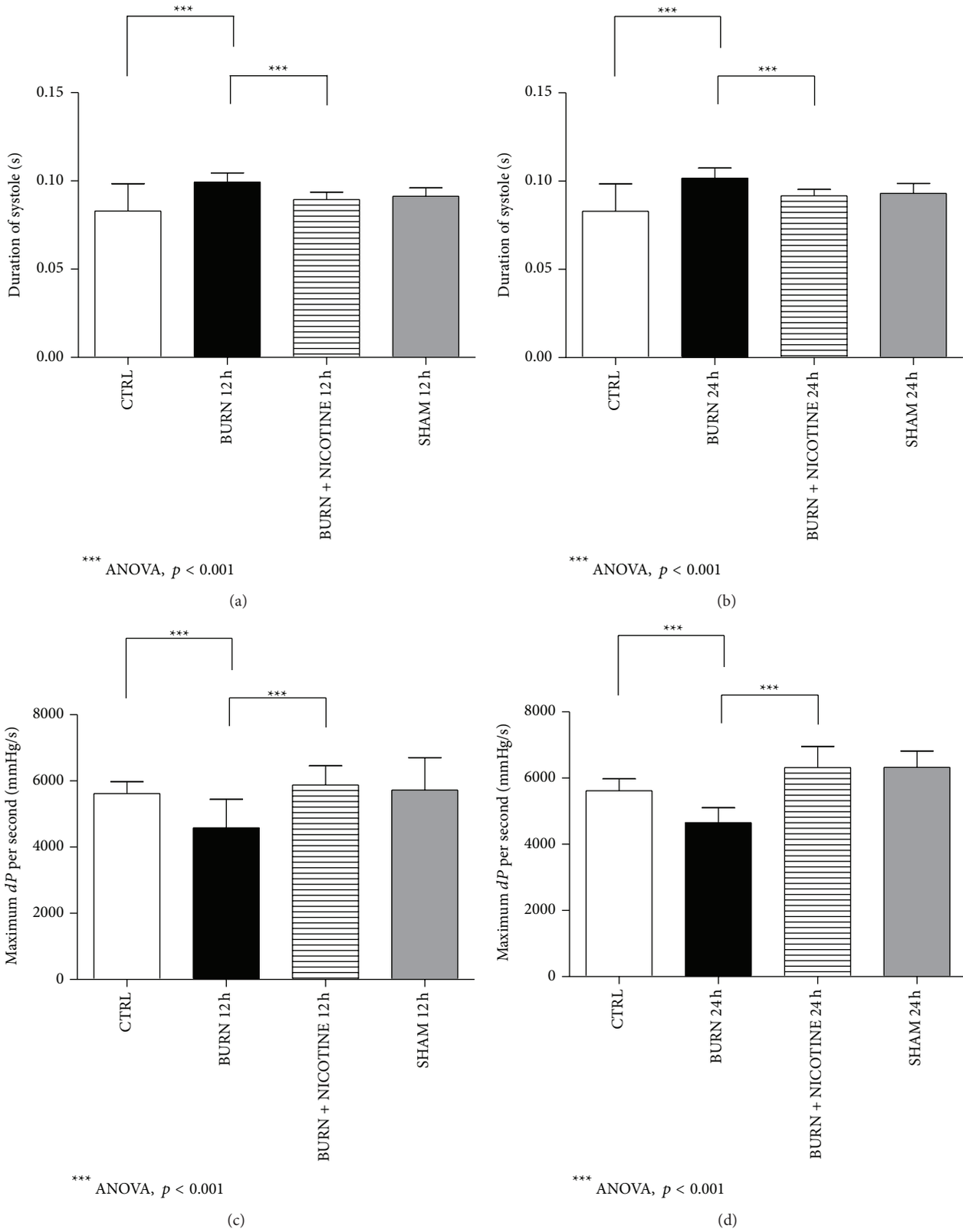


FIGURE 4: Systole. (a) and (b): duration of systole. The systole duration increased in experimental burn group (black versus white bar, $p < 0.05$). The BURN + NICOTINE groups revealed significantly lower duration of systole values than their burn alone counterparts (striped versus black bar, $p < 0.05$). (c) and (d): maximum increase of blood pressure per second (contractility). Reduced contractility was found for the BURN group (black versus white bar, $p < 0.05$). Nicotine administration showed restored contractility (striped versus black bar, $p < 0.05$). All values are expressed as mean + SD. Each bar represents $n = 5$ experiments.

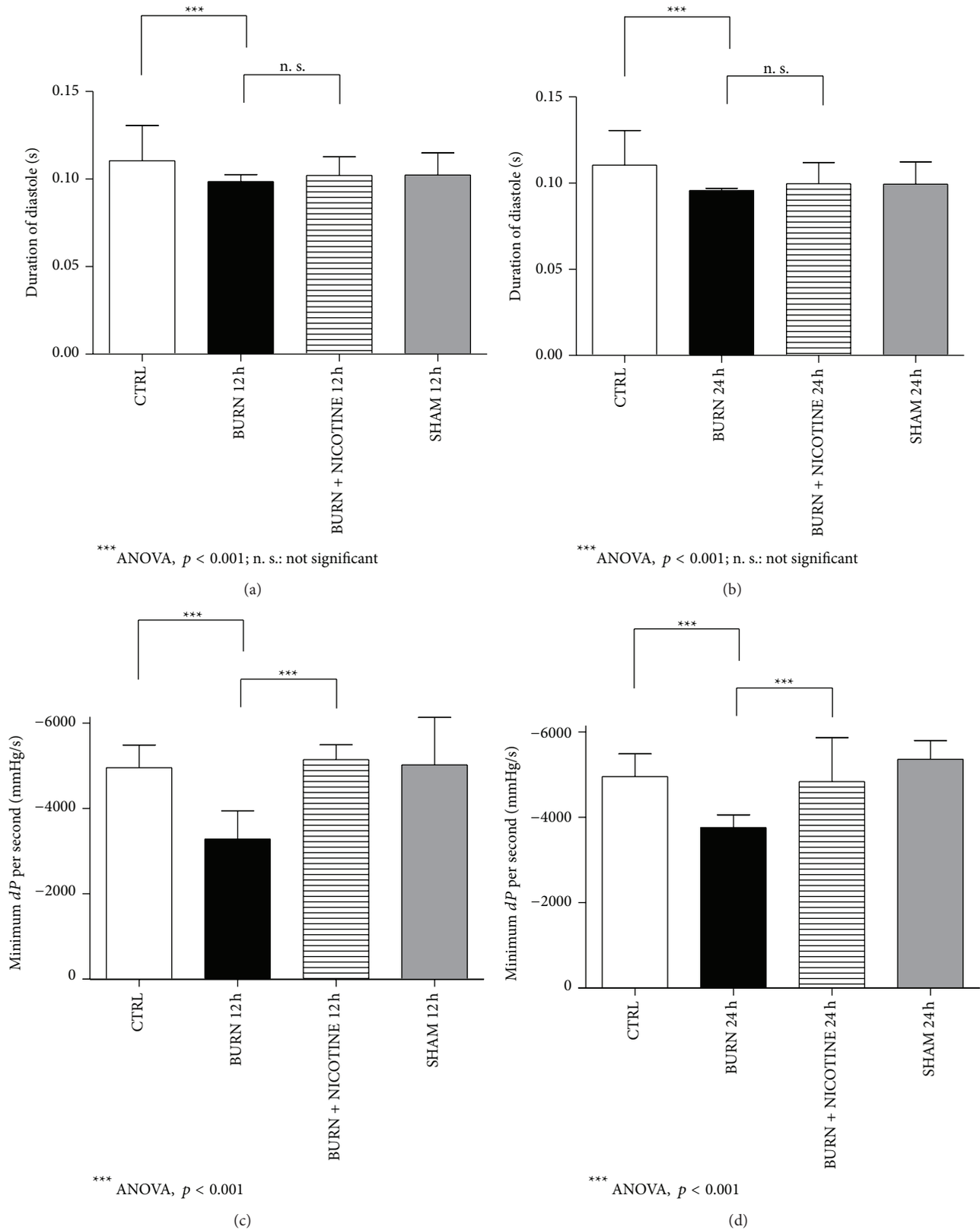


FIGURE 5: Diastole. (a) and (b): duration of diastole. The BURN group revealed significantly shorter diastole duration than the CTRL group (black versus white bar, $p < 0.05$). After posttraumatic nicotine application this effect is slightly reduced although differences were not statistically relevant (striped versus black bar, *not significant*). (c) and (d): minimum blood pressure difference per second (relaxation). Note that the scale is negative. The difference between the BURN and BURN + NICOTINE animals is marked (black versus striped bars, $p < 0.05$). Interestingly, both SHAM and BURN + NICOTINE groups display higher values than the control group. All results are expressed as mean + SD. Each bar represents $n = 5$ experiments.

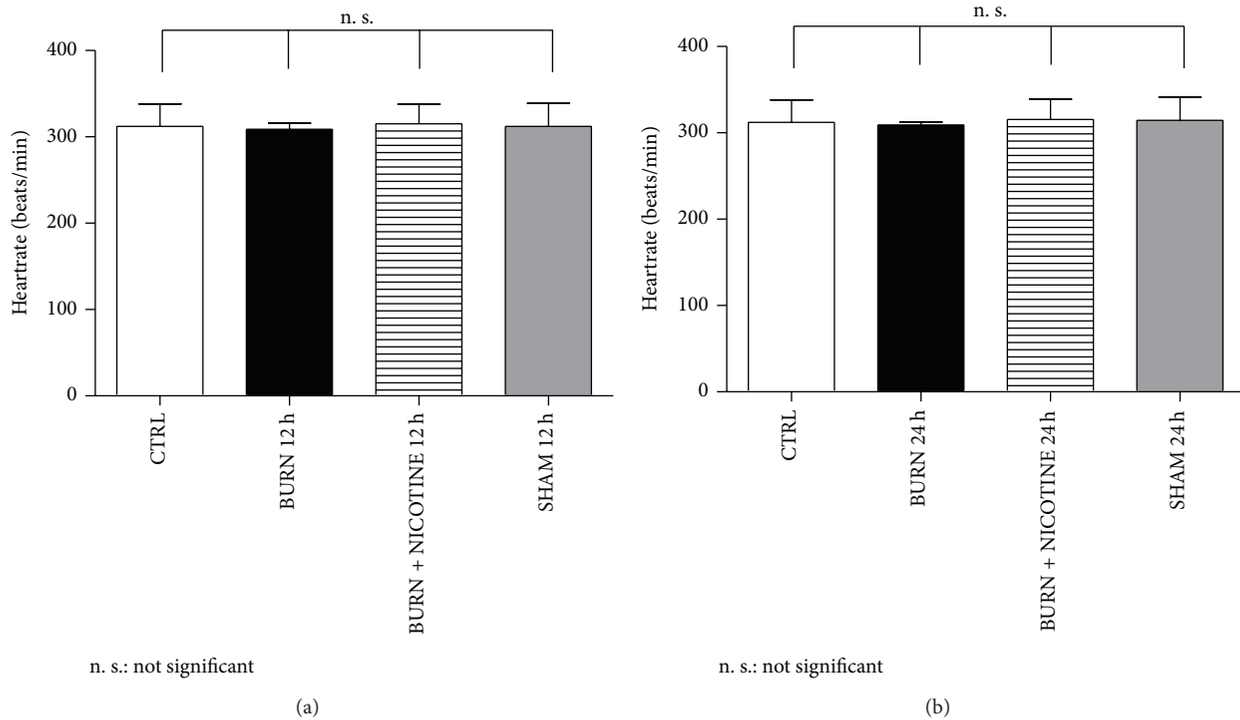


FIGURE 6: Heart rate. (a) and (b): heart rate. In the analysis of heart rate, no significant differences between the groups of experimental animals were found. The values are expressed as mean + SD. Each bar represents $n = 5$ experiments.

Previous experiments of our research group were looking at direct electrical stimulation of the vagus nerve and we could prove attenuation of proinflammatory cytokine production after electrostimulation of the parasympathetic axis [10, 22]. Additionally several studies used transdermal nicotine application to stimulate the parasympathetic axis because nicotine administration has been shown to induce anti-inflammatory effects [16, 17, 21, 23]. Clinically, this has been demonstrated in patients suffering from colitis ulcerosa [17, 23]. But in contrast, nicotine has adverse effect on Crohn's disease [23]. The reason for this ambiguous reaction in inflammatory bowel diseases is still not understood in entirety [23]. Still nicotine itself is seen doubtfully as an additive aspect in the treatment after burn trauma especially due to described adverse effects on wound healing [24, 25]. This points up the importance of further experiments to distinguish the positive anti-inflammatory effects of nicotine from adverse local effects on wound healing.

The present study showed that the administration of nicotine partially restores or normalizes decreased systolic blood pressure following burn injury. The results published by the groups of Sambol and Adams also revealed decreased systolic blood pressure at 24 hours after burn trauma [4, 26]. However decreasing the burn-induced increased activity of cardiodepressive cytokines like $\text{TNF-}\alpha$ and IL-1 thereby leading to normalization of blood pressure after a burn trauma [4, 26]. Thus, there is a negative correlation between circulating proinflammatory cytokine concentrations of $\text{TNF-}\alpha$ and IL-1 and the systolic blood pressure [5].

The left ventricular systolic blood pressure has a direct relationship to the arterial blood pressure that is based on the intravascular blood volume, stroke volume, and the total peripheral resistance. Burn trauma affects all of these three parameters. Capillary leakage results in severe fluid loss and decreased intravascular blood volume; the decreased contractility of the ventricle generates reduced stroke volume and circulating proinflammatory cytokines such as IL-1 β result in total peripheral resistance loss [3, 27]. Nicotine effectively reduces the concentration of vasodilatory cytokines such as IL-1 β and thereby leads to an increased total peripheral resistance [12]. In addition, the concentrations of $\text{TNF-}\alpha$, IL-1 β , and IL-6 are reduced which diminishes and, at best, abolishes their negative impact on cardiac contractility [3]. Stimulation of sympathetic neurons via nicotine application might support the restoration of systolic blood pressure. Additionally the volume resuscitation after burn trauma could contribute to the normalization of systolic blood pressure levels. However, sole fluid therapy is insufficient to abolish burn-induced cardiac dysfunction [28], as also seen in our BURN group.

For end-diastolic blood pressure levels burn injury also resulted in decreased levels. This parameter is essentially determined by the intravascular volume and contractile force of the left ventricle. So it may be deduced that the increase of the end-diastolic blood pressure of BURN + NICOTINE groups compared to BURN groups is a result of an impact on left ventricular function. This would be contradictory to the measurements of the other parameters

of myocardial function in the present study, indicating a positive effect of nicotine administration on left ventricular function. A second explanation for the increase in end-diastolic blood pressure according to nicotine application is based on the posttraumatic intravascular volume. The large burn-induced volume loss followed to the capillary leak results in a decreased intravascular volume and consecutively in a decreased end-diastolic blood pressure. In this context, Ipaktchi et al. were able to show that immunosuppressive therapy after burn trauma reduces the capillary leakage significantly [29]. This group used the specific inhibition of mitogen-activated protein kinases that has been shown to be a signal transduction element of the proinflammatory signaling cascade. This pathway is also inhibited by nicotine, which was shown by Oke and Tracey [30]. The improved end-diastolic blood pressure may result secondarily via the Frank-Starling mechanism.

The BURN groups showed significantly higher values for duration of systole compared to the control group. This was accompanied by significantly decreased values for the maximum blood pressure rise per second and a shorter duration of diastole. Previous studies also revealed a burn-induced contractile dysfunction [4, 26]. Additionally burn trauma affected diastolic cardiac function. Again, nicotine could attenuate the burn-induced negative effect: The BURN + NICOTINE and SHAM groups revealed increased levels compared to the BURN group and even to control group. Comparative data for this purpose does merely exist to our knowledge. Only the work of Adams and colleagues described the same effect of burn trauma [4]. Their and our results reveal that thermal trauma leads to systolic and diastolic cardiac dysfunction. As a result of our experiments, the application of nicotine has demonstrated, with few exceptions without significant differences, a positive effect on systolic and diastolic function after burn trauma.

Heart rate showed no significant differences. This is consistent with the results of several authors [5, 10, 26, 31]. Adams and colleagues found a steady heart rate in a burn model 24 h after trauma [4]. Similarly, Sambol, Maass, and our own results of burn models showed no changes in heart rate [5, 10, 26]. It is noteworthy that nicotine administration does not induce bradycardia. Bradycardia would be expected from stimulation of the vagal nerve. Interestingly however, in the literature there are data that nicotine has a more tachycardic effect [15, 17]. However, an additional effect of nicotine on sympathetic neurons cannot be excluded and could be an explanation for this aspect.

5. Conclusions

In the present study, we found that in a rat model of severe burn injury the trauma causes decreased blood pressure and a decrease in contraction and relaxation velocity of the heart. Adding on to the current knowledge base, further experiments will be done to explore the potent anti-inflammatory mechanisms of activation of the parasympathetic neural system. Our encouraging results that led to restoration of

nearly normal myocardial function parameters have already prompted us to follow up on this path.

Whether nicotine may also affect the lethality of severe burn trauma was not the subject of these investigations. This would be of great importance for further studies.

Conflict of Interests

The authors declare no conflict of interests or financial incentives in the production of this paper.

References

- [1] G. G. Gauglitz, J. Song, D. N. Herndon et al., "Characterization of the inflammatory response during acute and post-acute phases after severe burn," *Shock*, vol. 30, no. 5, pp. 503–507, 2008.
- [2] J. R. Saffle, J. J. Sullivan, G. M. Tuohig, and C. M. Larson, "Multiple organ failure in patients with thermal injury," *Critical Care Medicine*, vol. 21, no. 11, pp. 1673–1683, 1993.
- [3] J. W. Horton, "Left ventricular contractile dysfunction as a complication of thermal injury," *Shock*, vol. 22, no. 6, pp. 495–507, 2004.
- [4] H. R. Adams, C. R. Baxter, and S. D. Izenberg, "Decreased contractility and compliance of the left ventricle as complications of thermal trauma," *American Heart Journal*, vol. 108, no. 6, pp. 1477–1487, 1984.
- [5] D. L. Maass, J. White, and J. W. Horton, "IL-1 β and IL-6 act synergistically with TNF- α to alter cardiac contractile function after burn trauma," *Shock*, vol. 18, no. 4, pp. 360–366, 2002.
- [6] K. J. Tracey, "Physiology and immunology of the cholinergic antiinflammatory pathway," *The Journal of Clinical Investigation*, vol. 117, no. 2, pp. 289–296, 2007.
- [7] T. Kawada, T. Yamazaki, T. Akiyama et al., "Vagal stimulation suppresses ischemia-induced myocardial interstitial norepinephrine release," *Life Sciences*, vol. 78, no. 8, pp. 882–887, 2006.
- [8] L. M. Hoesel, A. D. Niederbichler, J. Schaefer et al., "C5a-blockade improves burn-induced cardiac dysfunction," *The Journal of Immunology*, vol. 178, no. 12, pp. 7902–7910, 2007.
- [9] C. J. Czura and K. J. Tracey, "Autonomic neural regulation of immunity," *Journal of Internal Medicine*, vol. 257, no. 2, pp. 156–166, 2005.
- [10] A. D. Niederbichler, S. Papst, L. Claassen et al., "Burn-induced organ dysfunction: vagus nerve stimulation attenuates organ and serum cytokine levels," *Burns*, vol. 35, no. 6, pp. 783–789, 2009.
- [11] D. Altavilla, S. Guarini, A. Bitto et al., "Activation of the cholinergic anti-inflammatory pathway reduces NF-kappaB activation, blunts TNF-alpha production, and protects against splanchnic artery occlusion shock," *Shock*, vol. 25, no. 5, pp. 500–506, 2006.
- [12] W. J. de Jonge and L. Ulloa, "The alpha7 nicotinic acetylcholine receptor as a pharmacological target for inflammation," *British Journal of Pharmacology*, vol. 151, no. 7, pp. 915–929, 2007.
- [13] R. P. A. Gaykema, I. Dijkstra, and F. J. H. Tilders, "Subdiaphragmatic vagotomy suppresses endotoxin-induced activation of hypothalamic corticotropin-releasing hormone neurons and ACTH secretion," *Endocrinology*, vol. 136, no. 10, pp. 4717–4720, 1995.

- [14] N. L. Benowitz, J. Hukkanen, and P. Jacob III, "Nicotine chemistry, metabolism, kinetics and biomarkers," in *Nicotine Psychopharmacology*, vol. 129 of *Handbook of Experimental Pharmacology*, pp. 29–60, Springer, 2009.
- [15] N. L. Benowitz, "Drug therapy. Pharmacologic aspects of cigarette smoking and nicotine addiction," *The New England Journal of Medicine*, vol. 319, pp. 1318–1330, 1988.
- [16] R. Kalra, S. P. Singh, J. C. Pena-Philippides, R. J. Langley, S. Razani-Boroujerdi, and M. L. Sopori, "Immunosuppressive and anti-inflammatory effects of nicotine administered by patch in an animal model," *Clinical and Diagnostic Laboratory Immunology*, vol. 11, no. 3, pp. 563–568, 2004.
- [17] W. J. Sandbom, "Nicotine therapy for ulcerative colitis: a review of rationale, mechanisms, pharmacology, and clinical results," *American Journal of Gastroenterology*, vol. 94, no. 5, pp. 1161–1171, 1999.
- [18] C. E. Richardson, J. M. Morgan, B. Jasani et al., "Effect of smoking and transdermal nicotine on colonic nicotinic acetylcholine receptors in ulcerative colitis," *QJM*, vol. 96, no. 1, pp. 57–66, 2003.
- [19] M. A. van Maanen, M. C. Lebre, T. van der Poll et al., "Stimulation of nicotinic acetylcholine receptors attenuates collagen-induced arthritis in mice," *Arthritis and Rheumatism*, vol. 60, no. 1, pp. 114–122, 2009.
- [20] C. Leib, S. Göser, D. Lüthje et al., "Role of the cholinergic anti-inflammatory pathway in murine autoimmune myocarditis," *Circulation Research*, vol. 109, no. 2, pp. 130–140, 2011.
- [21] L. Claassen, S. Papst, K. Reimers et al., "Inflammatory response to burn trauma: nicotine attenuates proinflammatory cytokine levels," *Eplasty*, vol. 14, article e46, 2014.
- [22] A. D. Niederbichler, S. Papst, L. Claassen et al., "Burn-induced organ dysfunction: vagus nerve stimulation improves cardiac function," *Eplasty*, vol. 10, p. e45, 2010.
- [23] M. Guslandi, "Nicotine treatment for ulcerative colitis," *British Journal of Clinical Pharmacology*, vol. 48, no. 4, pp. 481–484, 1999.
- [24] S. Xanthoulea, A. Deliaert, A. Romano, S. S. Rensen, W. A. Buurman, and R. R. van der Hulst, "Nicotine effect on inflammatory and growth factor responses in murine cutaneous wound healing," *International Immunopharmacology*, vol. 17, no. 4, pp. 1155–1164, 2013.
- [25] C. T. Selçuk, B. Özalp, M. Durgun et al., "The effect of hyperbaric oxygen treatment on the healing of burn wounds in nicotized and nonnicotized rats," *Journal of Burn Care and Research*, vol. 34, no. 4, pp. e237–e243, 2013.
- [26] J. T. Sambol, J. White, J. W. Horton, and E. A. Deitch, "Burn-induced impairment of cardiac contractile function is due to gut-derived factors transported in mesenteric lymph," *Shock*, vol. 18, no. 3, pp. 272–276, 2002.
- [27] C. R. Baxter and J. F. Waeckerle, "Emergency treatment of burn injury," *Annals of Emergency Medicine*, vol. 17, no. 12, pp. 1305–1315, 1988.
- [28] J. W. Horton, J. White, and C. R. Baxter, "The role of oxygen-derived free radicals in burn-induced myocardial contractile depression," *The Journal of Burn Care & Rehabilitation*, vol. 9, no. 6, pp. 589–598, 1988.
- [29] K. Ipaktchi, A. Mattar, A. D. Niederbichler et al., "Attenuating burn wound inflammatory signaling reduces systemic inflammation and acute lung injury," *Journal of Immunology*, vol. 177, no. 11, pp. 8065–8071, 2006.
- [30] S. L. Oke and K. J. Tracey, "From CNI-1493 to the immunological homunculus: physiology of the inflammatory reflex," *Journal of Leukocyte Biology*, vol. 83, no. 3, pp. 512–517, 2008.
- [31] A. D. Niederbichler, L. M. Hoesel, M. V. Westfall et al., "An essential role for complement C5a in the pathogenesis of septic cardiac dysfunction," *The Journal of Experimental Medicine*, vol. 203, no. 1, pp. 53–61, 2006.



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