Sciatic nerve-conditioned medium with the addition of *Croton blanchetianus* Baill essential oil promotes morphological plasticity in spinal cord cultured cells

Meio condicionado de nervo ciático com a adição de óleo essencial de *Croton blanchetianus* Baill promove plasticidade morfológica em cultura de células de medula espinal

El medio acondicionado del el nervio ciático con la adición de aceite esencial de *Croton blanchetianus* Baill promueve la plasticidad morfológica en las células cultivadas de la médula espinal

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Abstract
Spinal cord injury is a devastating condition that leads to significant disabilities. The treatment for this problem is a challenge in neuroscience, and it is necessary to combine different strategies to obtain functional recovery outcomes. There are many herbal natural products, such as *Croton blanchetianus* Baill (CB) essential oil, a Brazilian semi-arid bush with neuroprotective substances tested in regeneration processes and synaptic plasticity. Thus, this study analyzed the cellular plasticity of spinal cord neurons and glial cells in the presence of sciatic nerve-conditioned medium (SNCM) before the addition of CB essential oil. Cell morphology was assessed over 96 hours, and immunocytochemistry analyses were conducted for GFAP, GAP-43 and NeuN. Photomicrographs were made by scanning electron microscopy (SEM). Morphological analysis showed evident trophic development in the groups that received CB essential oil (P>0.000), immunoreactivity for GFAP, GAP-43 and NeuN and the plasticity of these cells were confirmed by SEM. This pioneer study about the plasticity of spinal cord neurons and glial cells opens new possibilities and techniques with essential oils for cell therapy in the presence of SNCM, which promoted neuroprotective action.

Keywords: Essential oil; Sciatic nerve; Trophism; Neuroprotection.

1. Introduction
Spinal cord injury (SCI) is defined as a central nervous system injury that results in a large functional deficit. It is a devastating condition that leads to significant disabilities because patients with SCI develop many complications and multiple organ dysfunctions (Chariker et al., 2019). In veterinary practice, SCI often culminates in euthanasia of the affected animal, and therefore physical and emotional
stress on owners (Bergman, 2000; Dumont et al., 2001). In addition to primary mechanical trauma to the spinal cord (SC) axons, there is rupture of blood vessels and cell membrane, resulting in a cascade process leading to a progressive secondary injury which can lead to edema, ischemia, inflammation, and cytokine production. Consequently, oxidative stress and glial scarring cause irreversible tissue necrosis and neuronal death (Huang et al., 2007; Lin et al., 2011; Ormond et al., 2014).

For this reason, when investigating therapeutic strategies, it is imperative to explore interventions that minimize these effects and promote morphological and functional recovery (Marcos et al., 2016; Souza et al., 2010). The major negative effect on cells by the secondary lesion of SCI is the release of free radicals that attack the cell membrane and this mechanism, in turn, modify cell components such as unsaturated fatty acids by lipid peroxidation process (Cemil et al., 2010; Huang et al., 2007).

SC treatment after the injury is a challenge for neuroscience, considering that recent studies show that the combination of different strategies is necessary to obtain functional recuperation results (Houle et al., 2013). Therefore, the search for new alternatives and research in regeneration is important (Mothe & Tator, 2012; Ren & Young, 2013; Varma et al., 2013).

Different therapeutic modalities have been suggested to avoid changes leading to neuronal death, in the neonatal or adult phase. Neuroprotection is defined as any strategy and mechanism that protects the central nervous system (CNS) from neuronal injury. Thus, there are currently numerous natural plant-derived products that have been reported as neuroprotective in traditional medicine (Elufioye et al., 2010).

Many studies indicate the efficiency of essential oils in reducing pain, anxiety and stress symptoms in animal models and humans alike, with different CNS disorders (Lee et al., 2011; Tankam & Ito, 2013). They are substances derived from secondary metabolism and present considerable complexity with hundreds of organic compounds, such as monoterpenes, sesquiterpenes and phenylpropanoids, which confer their organoleptic characteristics (Barbosa et al., 2016; Bizzo et al., 2009; Souza et al., 2010). The family Euphorbiaceae has in the genus Croton several species for which neuroprotective activity on the nervous system was evidenced. Also, compounds isolated from Croton species exert a wide range of biological activities such as anti-inflammatory, antifungal activity, neurite growth-promoting activities (Xu et al., 2002) as well as astrocyte and microglial activation and differentiation (Costa et al., 2012). *Croton blanchetianus* Baill. (CB), is a plant from the Brazilian semi-arid and has botanical synonyms like *Croton sonderianus* Müll. Arg. and *Croton alagoensis* Müll. Arg. (Angélico et al., 2011; de Lima & Pirani, 2008; I. M. M. Oliveira et al., 2015).

Taking into account the neuroscience necessity to understand and provide therapeutic strategies that enable central axonal regeneration and considering the potential neuroprotective effect of CB essential oil, this study aims to evaluate their effects on the CNS cell plasticity.

2. Methodology

This is quantitative and experimental study carried out under controlled conditions (Sampieri et al., 2013).

2.1 Plant material

Air-dried plants of cultivated *Croton blanchetianus* were obtained from Universidade do Estado do Rio Grande do Norte (UERN). Essential oil was obtained by hydrodistillation with clevenger apparatus. Then, 750 ml of distilled water (1:10, w/v) was added. Extraction was performed for three hours. Next, the essential oil was collected and placed into a vial with anhydrous sodium sulfate to remove any water. Then, vials were stored at 4°C for later analysis. The characterization of a sample of *C. blanchetianus* essential oil using gas chromatography–mass spectrometry revealed a high concentration of azulene, caryophyllene and β-elemene. For the purpose of study, 0.1 µL of the essential oil was diluted in 100 µL of
phosphate buffered saline (PBS) 0.2 M. This solution has the concentration 0.01 µL C. blanchetianus essential oil.

2.2 Animals

Male Wistar rats of an approximate age of 45 days and neonates aged 2 days were used under the approval of the Ethics Committee on Animal Experimentation from UERN, Protocol number 006/17, in accordance with the ethical principles adopted by the Brazilian Society of Laboratory Animal Science and according to law number 11,794, the Arouca law, of the Ministry of Science, Technology and Innovation. After the breast-feeding period, rats were kept in a UERN’s vivarium in separate cages with adequate housing conditions with free access to food and water until they were of appropriate weight and age.

2.3 DMEM Conditioning for Sciatic Nerve Explants

Under laminar flow, 60-mm (P60) culture plates with lids were prepared with 5 mL of Leibovitz-15 medium (L-15: Gibco, USA). Animals were anesthetized (Ketamine and Xylazine of Agener Union 10%) and subjected to extraction of the sciatic nerve after the dorsal region was shaved and for local asepsis was used 2% chlorhexidine. Surgical incisions were made in the posterior region, and the sciatic nerve was removed and placed in P60 with L-15 medium under aseptic surgical conditions with the aid of micro devices (scissors, forceps and retractors). All the excess tissues (muscle, fat, and blood vessels) attached to the nerves were removed under magnification by a SZ61 stereomicroscope (Olympus, Japan). Next, the epineurium of the nerves was removed under magnification via microsurgical techniques. The dissected nerves were segmented into 2 mm length explants. Under laminar flow, the nerve fragments were placed in 60-mm plates with 1.5 mL of low Knockout Dulbecco’s modified Eagle’s medium (DMEM) (Gibco, USA) plus 10% foetal bovine serum and 0.1% gentamicin, with a medium called D-10. Excess medium was removed from the explants so they would not float or submerge. The D-10 medium from these cultures was changed twice per week, and the explants were transferred to a new plate with fresh medium once per week. The medium was changed once and then disposed. This procedure allowed an adequate nutrient supply to the explants and an analysis of their reactivity (Figure 1).

**Figure 1.** Initial cellular migration by the sciatic nerve explants.

The Figure 1 indicates that the cells migrated from their explants, making possible the active compounds production and its exocytosis in culture medium, acting as an extracellular matrix. In this way it was possible collect the conditioned medium for SC neurons treatment. At this stage, SNCM (100 µl/plate) was collected for inoculation into the SC cell culture.
2.4 Isolation and SC Cell Culture

The animals were killed with an anaesthetic overdose of isoflurane inhalation (Isoforine®) (Cristália, Brasil), and then dissected under aseptic conditions for the SC removal. After the removal, the explants were held in a conical tube with L-15. Under laminar flow, 60-mm culture plates were prepared with D-10 cell culture medium. Unattached cells and residual non adherent red blood cells were removed after 24 hours by PBS wash. After 1 week in treatment the cells were resuspended from the plates with trypsin-EDTA and subjected to the 3 cycles of centrifugation with washes. Then, the SC cells were plated in 24-well plates (1×10^6 cells/well) and were observed over 96 hours. With this procedure, it was possible to assess the adhesion and proliferation of SC cells in the following groups: Group 1: SC cells + D-10 medium; group 2: SC cells + SNMC; group 3: SC cells + SNMC + CB essential oil (0,01 μL) and group 4: SC cells + CB. The cell count was done using phase microscopy at 10x magnification with a CKX41 microscope (Olympus, Japan). The 4 groups photomicrographs were performed after 96 hours.

The cell morphology was observed using phase contrast microscopy at 10x magnification. After cell observation at 96 hours of cultivation, we proceeded with immunocytochemistry for neuronal definition and the cellular morphometric analysis (area and perimeter) was performed.

2.5 Immunofluorescence Staining

By day three (96 hours), the cells had adhered to the plates, the medium was removed, and they were washed in two steps of five minutes each in PBS 0.1 M, pH 7.4; the cells were then fixed in paraformaldehyde (PFA) 4% for thirty minutes and washed again in three baths of PBS (five minutes each). Then, the cells were treated with 0.5% Triton X-100 (Sigma) for 10 minutes and washed in PBS again. Subsequently, blocking of nonspecific sites was performed for 30 minutes in a PBS 0.1 M solution containing 0.2% Triton and 1% cattle serum albumin (CSA).

Plates were incubated for 2 hours at room temperature with anti-mouse Glial fibrillary acidic protein (GFAP) (Sigma, 1:400), anti-mouse GAP-43 (Sigma, 1:2600) and NeuN (Millipore, 1:500). Upon completion of this step, cells were washed in PBS (0.1 M, pH 7.4) for five minutes and incubated for 1 hour with an anti-rabbit or anti-mouse secondary antibody produced in donkey (Jackson, USA) conjugated with AlexaFluor 594 or FITC fluorophore and kept under refrigeration with absence of light. After secondary incubation, cells were washed with PBS for five minutes and immediately examined with a fluorescence microscope (Eclipse E200, Nikon) and then with a second fluorescence microscope (Eclipse Ni, Nikon). Photomicrographs were taken with Moticam 3.0 and 5.0 (Motic) digital cameras at 4x, 10x and 20x magnification in 9 fields in a predetermined sequence on each plate. The presence of fluorescent staining was recorded in SC cells, taking care to examine the subcellular, cytoplasmic, and nuclear compartments.

2.6 Electronic Scanning Microscopy

After a period of 96 hours, the samples were fixed to stabilize the form, which consists in the process of preserving the original state of such samples in culture, avoiding to the maximum the introduction of artefacts.

The SC cells from each group were fixed in a 2.5% glutaraldehyde solution buffered with 0.1 M sodium phosphate (pH 7.4) and washed in 0.1 M sodium phosphate buffer (pH 7.4). Post-fixation was then performed on 1% osmium tetroxide solution buffered with 0.1 M sodium phosphate (pH 7.4) for two hours. After post-fixation, the samples underwent three washes in the buffer solution and two washes with only distilled water. Subsequently, the samples were dehydrated in ethanol at different concentrations (50%, 70%, 90%, and 100%). The material was mounted on a Stub sample holder and sputtered gold on the plate for observation in a TESCAN® vega 3 LMU scanning electron microscope. The best representative images were processed.
2.7 Statistical Analyses

Two independents calibrated investigators (kappa=0.94) counted cells per field in absolute numbers, using cell cultures of at least 3 different experiments with 10x magnification. The Motic Images Plus 2.0 (Motic) software was used for morphological observation (area in $\mu m^2$ and perimeter in $\mu m$). Image J software was used for cell counting, and Adobe Photoshop CS6.0 (Adobe) software was used to fix the minimum brightness and contrast of the photomicrographs. The database search was built on the SPSS platform software (Statistical Package for Social Sciences) version 22.0, with a subsequent consistency check of typing. After the final database structure was completed, a descriptive analysis of all data was initially performed. The data were statistically compared by analysis of variance (ANOVA) with Tukey and Bonferroni tests and considered significant when $P < 0.05$.

3. Results
3.1 Cellular morphometry analysis

The cellular morphometry analysis results can be seen in Figures 2 and 3.

Figure 2. Neuronal cells morphometry analysis.

(A) Average area of neuronal cells in the SNCM, CB essential oil and SNCM plus CB essential oil groups. Means ± S.E.M. Difference in a ($p = 0.022$), b ($p = 0.000$), c ($p = 0.019$), according to ANOVA and post-test Tukey and Bonferroni. (B) Perimeter of neuronal cells in the SNCM, CB essential oil and SNCM plus CB essential oil groups. Means ± S.E.M. Difference in a ($p = 0.001$), b ($p = 0.003$), according to ANOVA and post-test Tukey and Bonferroni. (C) Axonal length of neuronal cells in the SNCM and SNCM plus CB essential oil groups. Means ± S.E.M. Difference in a ($p = 0.001$), b ($p = 0.000$), c ($p = 0.000$), d ($p = 0.001$), according to ANOVA and post-test Tukey and Bonferroni. Source: Authors.
When comparing the mean areas of neuronal and glial cells observed at 96 hours, the Figure 2 graphic A shows that group 3 was superior to group 1 \((p \leq 0.01)\), group 4 was superior to groups 1 and 2 \((p \leq 0.01)\) for neuronal cells. It reveals evident trophic development in the groups that received CB essential oil, when compared to the control group (D10) and in the group that received SNCM in the presence of CB essential oil, when compared to the group with SNCM (D10 + MC). For glial cells, Figure 3 graphic A shows group 3 was superior to group 4 \((p = 0.001)\) and group 4 was superior to groups 1, 2 and 3 \((p \leq 0.001)\). It indicates evident trophic development in the group that received SNCM in the presence of CB essential oil, when compared to the control group (D10), group that received SNCM and group that received only CB essential oil.

When comparing the mean perimeter of neuronal and glial cells observed at 96 hours, Figure 2 graphic B shows that groups 3 and 4 were superior to group 1 \((p \leq 0.001)\) for neuronal cells. There is evident trophic development in the group that received CB essential oil, when compared to the control group (D10). For glial cells, in the Figure 3 graphic B group 3 was superior to group 4 \((p = 0.001)\) and group 4 was superior to groups 2 and 3 \((p \leq 0.001)\). Trophic development is evident in the group that received SNCM in the presence of CB essential oil, when compared to the control group (D10), group that received SNCM and group that received only CB essential oil.

For the mean axonal length of neuronal cells observed at 96 hours, Figure 2 graphic C displays that groups 2, 3 and 4 were superior to group 1 \((p \leq 0.001)\), group 4 was superior to group 2 \((p = 0.001)\). This indicates evident trophic development in the group that received CB essential oil, when compared to the control group (D10) and in the group that received SNCM in the presence of CB essential oil, when compared to the group with SNCM (D10 + MC).

### 3.2 Immunofluorescence Staining

The Immunofluorescence staining photomicrographs are grouped in the Figure 4.
The groups treated with SNCM in the presence of CB essential oil showed more evident immunoreactivity for GFAP (Fig. 4A, 4B, 4C and 4D), GAP-43 (Fig 4E, 4F, 4G and 4H) and NeuN (Fig. 4I, 4J, 4K, 4L) compared to the other groups. GFAP marking demonstrates cells treated with control group (4A), SNCM (4B), CB essential oil (4C) and cells treated with SNCM + CB essential oil (4D) and demonstrates more reactive astrocytes in groups treated with SNCM and CB essential oil. Those for GAP-43 (4E, 4F, 4G and 4H) demonstrates a higher trophic effect in groups treated with SNCM and SNCM + CB essential oil while for NeuN (4I, 4J, 4K and 4L) demonstrates a higher trophic effect in groups treated with SNCM and SNCM + CB essential oil.
3.3 Electronic Scanning Microscopy

The plastic and trophic effect on cultured cells was verified in the presence of CB essential oil. Over 96 hours, glial and neuronal cells developed plasticity when compared to the D10 medium as in the Figure 5.

**Figure 5.** Photomicrographs by scanning electron microscopy.

The Figure 5 demonstrates the spinal cord cells plasticity, especially cells belonging to the groups treated with CB essential oil (8A, 8B, 8C and 8D).

4. Discussion

This work investigated the plastic behavior of SC glial and neuronal cells after 96 hours of treatment with CB
essential oil. A conditioned medium with sciatic nerve fragments was also developed, as well as two control groups (D-10 medium and another with the conditioned medium solely, respectively).

Therefore, an experimental medium was created from nerve fragments that simulated an *in vivo* nerve injury condition where the explants became reactive and naturally secreted factors that enabled a favorable environment for post traumatic nerve regeneration.

Groups treated with CNSM in the presence of CB essential oil showed better results regarding the neuronal and glial area and perimeter and axonal length of SC cells. Furthermore, these results demonstrate that good growth was probably due to the SNCM nutritional contributions, which created a favorable cellular ambient for cell regulation and that promotes growth (Hall, 2001), because Schwann cells retain their biochemical properties during regeneration in the CNS environment (Chen et al., 2005; Guzen et al., 2009, 2012; Schwab, 2002).

CB essential oil probably acted as a neuroprotective and neurotrophic agent, increasing the plastic effect of SC cells. This fact is due to the presence of sesquiterpenes in the CB essential oil, specifically, caryophyllene and β-elemene, found after gas chromatography coupled to mass spectrometry. Caryophyllene has been reported as neuroprotective in models of cerebral ischemia (Choi et al., 2013), convulsion (Liu et al., 2015), glial cell hypoxia (Guo et al., 2014) and Alzheimer's disease (Cheng et al., 2014). A study by Oliveira (2016) found that the caryophyllene present in the *Spiranthera odoratissima* essential oil was able to increase BDNF hippocampal levels, the main regulator of synaptic plasticity in learning and memory (Cunha et al., 2010). In another pioneering study by Ferreira (2015), evaluating the caryophyllene effect in three neurodegeneration models in PC-12 cells, it was found that this secondary metabolite, besides inducing cell differentiation, also induced the formation of longer neurites. Wang and Heinbockel (2018), studying β-elemene, an unoxygenated sesquiterpene, concluded that this compound probably promoted neurite growth *in vivo* and *in vitro*, suggesting that β-elemene may be used as a new treatment strategy for patients with SCI.

The immunofluorescence staining showed that the groups containing CB essential oil had cellular markings for GAP-43 and NeuN. Increased GAP-43 expression after injury indicates that the axon is in a regenerative process. Some reports in the literature relate the action of sesquiterpenes in extracts and essential oils with increased GAP-43 expression. Wang and Heinbockel (2018) found that β-elemene significantly upregulated the level of GAP-43. According to Mullen et al. (1992) and Casella et al. (2004) the high expression and concentration of NeuN indicates a differentiation state or neuronal activity. Therefore, it is evident that there may be some other active compound responsible for the neurotrophic action of CB essential oil. Since essential oils are a phytocomplex, several components present in its constitution may be responsible for its final action (D. R. de Oliveira, 2016).

5. Conclusion

This paper opens interesting perspectives for understanding the role of essential oils as stimulants in the nervous cellular environment and their interaction with a nutritive environment within this environment, inducing morphological changes and enabling the plasticity of ME cells in a neuronal and glial lineage. It is concluded that CB essential oil probably has neuroprotective and neurotrophic activity in SC cells. Further research should be done to clarify phytoconstituent’s mechanism of action in the essential oil since it is unknown and deepen the use of plant-derived natural products in neuroscience studies. This work contributes with knowledge in the search for better therapy techniques applied in diseases and trauma in the nervous system.
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