

STORAGE OF THE OIL OF *Moringa oleifera* AFFECTS YOUR TOXICITY ABOUT *Tetranychus* *urticae*?

Influence of storage time/duration on the toxicity of *Moringa oleifera* (Family) oil to *Tetranychus urticae* (Family)

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ABSTRACT

The objective of this study was to evaluate the storage time of *Moringa oleifera* oil on the acaricidal activity on *Tetranychus urticae*. Was used amber bottle for storage of oil which remained in a room at 25°C. The storage times considered in the experiment were 0, 30, 60, 90 and 120 days after extraction. At each time a suspension at the concentration of 3% (v/v) was applied on the mite. A completely randomized design with 5 treatments (storage times) was used, containing 8 replicates, composed of 12 females per replicate. The application was carried out by spraying. Mortality data were submitted to the sphericity test and then to analysis of variance, followed by non-linear regression. The analysis of variance revealed that the time factor of storage significantly affected the mortality of the mite, according to an exponential model. Mortality was increasing, reaching a mean of $74.16 \pm 8.37\%$ at 120 days. The oil of *M. oleifera* is promising for the control of *T. urticae*, improving the acaricidal activity over time.

Keywords: *Keywords: mite; alternative control; storage time; Moringa.*

1. INTRODUCTION

Phytophagous mites, such as *Tetranychus urticae* Koch (Acari: Tetranychidae), are pests of crops of economic importance, such as cotton, soybean, tomato, papaya, strawberry and others [1 - 5]. When not controlled, these organisms can cause damage to crops? Or crops and farmers? farmers. For most crops, the chemical method has been the main tool to combat this mite. The products used generally have molecules of wide spectrum, eliminating even natural enemies, besides possessing high residual power [5, 6].

Faced with this, the search for healthier acaricides?products- has intensified research that provides less aggression to agroecosystems. Among them, studies related to predatory mites [7, 8], entomopathogenic fungi [2, 9] and plants with insecticides property have has gained prominence [4]. Extracts, fixed and essential oils from vegetables have been studied in several insect pests and vectors of diseases, providing promising results [4, 10 - 12].

Moringa oleifera Lam, Moringaceae, is a plant studied for several purposes, from biodiesel production [13], to insecticidal activity on disease vectors [14 - 17] and even on *T. urticae* [18]. However, the form of action and effects of the substances present in the oil are still little

34 explored on agricultural pests. In addition, information on the storage time and temperature
35 of the oil, types of storage containers of these extracts and oils [13], in order to preserve the
36 insecticidal / acaricidal characteristics, are deficient.

37 The objective of this study was to evaluate the action and duration of the acaricidal effect of
38 the oil of *M. oleifera*, stored in amber glass, for the control of *T. urticae*, under laboratory
39 conditions.

40

41 2. MATERIAL AND METHODS

42

43 2.1 ~~Rearing~~ ~~Creation~~ and maintenance of *Tetranychus urticae*

44 The mite creation was established in the entomology sector of the Federal Institute of
45 Espírito Santo, Itapina Campus, Colatina-ES (IFES-Campus Itapina), in *Canavalia*
46 *ensiformis* plants cultivated in pots without any phytosanitary treatment. The vessels were
47 packaged in wooden cages (50 x 50 x 100 cm), coated with anti-aphid screen and with a
48 front opening closed by removable glass. ~~Rearing was carried out~~ ~~Creation was maintained~~
49 in air-conditioned rooms regulated at $25 \pm 1^\circ\text{C}$, relative humidity $70\% \pm 10$ and 12h
50 photophase.

51 2.2 Extraction and storage of moringa oil

52 Moringa seeds were collected at the IFES-Campus Itapina and subjected to the extraction of
53 the oil by cold pressing. After this procedure, the oil was filtered through a fine mesh screen
54 and stored in an amber glass container in an air-conditioned room with temperature of $25 \pm$
55 1°C , relative humidity $70\% \pm 10$ and photophase of 12h.

56 2.3 Bioassays

57 For the experiment, aqueous suspensions of 3% (v/v) moringa oil were used, which
58 corresponded to the highest concentration soluble in water. As solvent, distilled water plus
59 Tween[®] 80 adhesive spreader (0.05% v/v) was used. Thereafter, the mixture was left under
60 stirring (magnetic stirrer) for 30 minutes at room temperature. The storage times for the oil
61 considered in the experiment were 0, 30, 60, 90, 120 days after extraction.

62 Leaf discs of *C. ensiformis* ($\varnothing = 4$ cm) were introduced into Petri dishes (10 x 1 cm),
63 containing cotton moistened around this to maintain leaf turgescence and avoid mite
64 leakage. For each plate a leaf disc was considered, which constituted a repetition. Twelve
65 mite females were transferred per replicate.

66 To perform the applications, an airbrush (Model SW-130K) was used, connected to a
67 compressor calibrated at a constant pressure of 25 psi. The application suspension volume
68 was 3 ml per replicate. Afterwards the plates were conditioned in an air-conditioned room
69 (temperature of $25 \pm 1^\circ\text{C}$, RH of $70 \pm 10\%$ and photophase of 12h).

70 The acaricidal effect was evaluated 24, 48 and 72 hours after the application, registering the
71 mortality of the individuals. ~~For the As~~-control treatment, only the solvent was used, and
72 these mortality values were used to correct the treatments using the formula proposed by
73 Abbott [19].

74 2.4 Statistical analysis

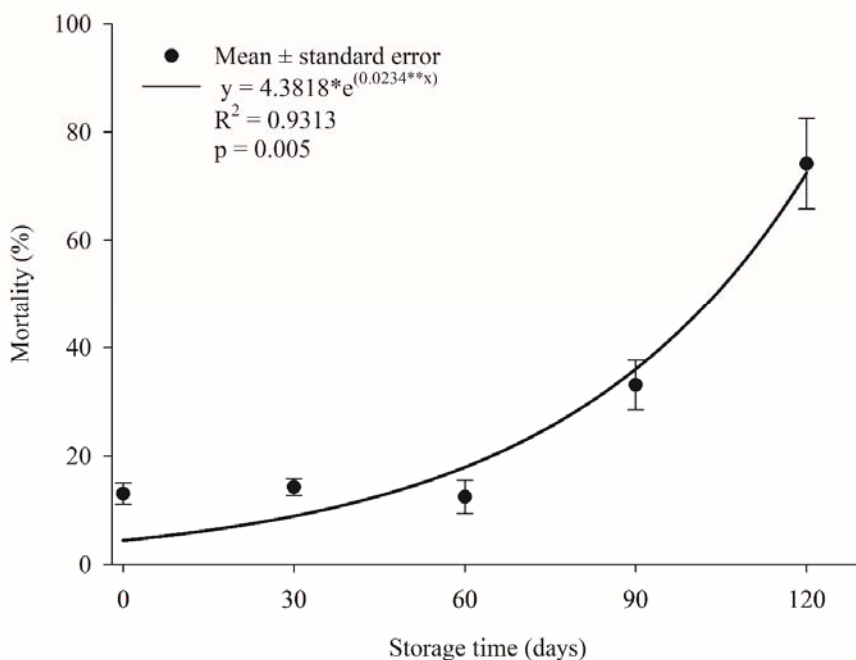
75 A completely randomized design with 5 treatments (storage time) was used, with 8
76 replications, using the accumulated mortality data in 72h. Since the treatments are time-
77 repeated measures, the data were submitted to the Mauchly [20] test to verify the sphericity,
78 as recommended by Huynh & Feldt [21] for time-repeated measurements [22]. After
79 checking the sphericity, we performed the analysis of variance followed by non-linear
80 regression analysis in R software version 3.4 [23].

81

82 3. RESULTS AND DISCUSSION

83

84 The covariance matrix showed sphericity ($W = 0.61$, $p = .38$). Based on this principle, the
85 analysis of variance was performed. It was verified that the time factor was adjusted to the
86 exponential model ($F = 55.22$; $p < .01$) (Figure 1). There was an increase in the mortality of
87 the mite with the increase of the storage time, reaching $74.16 \pm 8.37\%$ of mortality with 120
88 days of storage.



89

90 Figure 1. Mortality of *Tetranychus urticae* treated with *Moringa oleifera* oil with different
91 storage times. ** and * significant regression coefficient at the 1 and 5% level, respectively.

92 The oil from moringa seeds is a potential for the management of *T. urticae*. The increase in
93 mortality with the advancement of storage time was an interesting ~~and even unexpected~~
94 result. Such a ~~result may be surprise is~~ due to the fact that many products lose their
95 potential/effectiveness due, perhaps, to some process of deterioration and / or oxidation. On
96 the other hand, *M. oleifera* oil can maintain some properties, such as density and viscosity,
97 and increase the percentage of free fatty acid for up to 24 months [13].

98 Toxicity activity of *M. oleifera* in arthropods was reported by other researchers in
99 coleopterans [24 - 27], dipterans [14 - 17], lepidopterans [28 - 30] and mites [18].

100 On *Aedes aegypti* (L.) (Diptera: Culicidae) the aqueous extract of the seed presented
101 larvicidal action and ovicidal, being able to cause 100% mortality after 24 h of exposure [16].
102 For the malaria vector, *Anopheles stephensi* Liston (Diptera: Culicidae), the use of methane
103 extract from moringa seeds, provided larval and pupal mortality [15]. In *Sitophilus oryzae* (L.)
104 (Coleoptera: Curculionidae), *Oryzaephilus mercator* (Faur) (Coleoptera: Cucujidae) and
105 *Ryzopertha dominica* (Fabr.) (Coleoptera: Bostrichidae) methanoic extract caused mortality
106 of more than 90% of insects [26]. Aqueous extracts of moringa seeds presented high toxicity
107 to *T. urticae*, resulting in a lethal concentration for 50% of the population around 12.39%
108 [18]. These results demonstrate that, independent of the solvent used, *M. oleifera* oil is toxic
109 to different groups of insects and to the mite.

110 Moringa oil is a fixed type, as are soybean oils (*Glycine max* - Fabaceae), canola (*Brassica*
111 *napus* - Brassicaceae) and castor bean (*Ricinus communis* - Euphorbiaceae). Thus, the
112 choice of the solvent is crucial for the solubilization and homogenization of the suspension
113 so that it does not alter its properties [31]. Therefore, the use of the Tween® adhesive
114 spreader provides these characteristics without affecting the potentiality of the oil, as well as
115 reported in papers used with *R. communis* oil [10, 11].

116 The moringa studies show that plants of this genus are rich in lectin [32], α - and γ -
117 tocopherols, glycosylates, nitriles, glycosides, quercetin, canferol, rhamnosides,
118 isothiocyanates and steroids [33]. In addition, the oil has as main component oleic acid (\approx
119 78.0%) [34].

120 Lectin is a type of protein, which prevents the process of digestion and absorption of
121 nutrients in the insects, causing death by malnutrition [30, 32]. Seeds of *M. oleifera* contain
122 cMoL (coagulant *M. oleifera* Lectin) and WSMoL (Water-Soluble *M. oleifera* lectin) lectins
123 that promoted mortality in *A. aegypti* larvae [30, 35]. At the concentration of 1% (m/m), lectin
124 cMoL caused mortality of pupae of *Anagasta kuehniella* (Zeller) (Lepidoptera: Pyralidae) and
125 delayed the total development cycle of surviving insects [28].

126 In the case of fatty acids, larvicidal and anti-nutritive activity of oleic acid have been reported
127 on *A. aegyptii*, *Helicoverpa zea* Boddie (Lepidoptera: Noctuidae), *Lymantria dispar* L.
128 (Lepidoptera: Lymantriidae), *Culex quinquefasciatus* Say (Diptera: Culicidae) [36, 37]. Faced
129 with this, it can be seen that *M. oleifera* presents lethal and anti-nutritive activity, as well as
130 sublethal effect.

131

132 4. CONCLUSION

133

134 It is concluded that the oil of *M. oleifera* shows acaricidal activity to *T. urticae* and that the
135 Storage time can influence positively made possible its toxicity to the mite. Oil stored at 120
136 days is most suitable for use in mite control.

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139 **COMPETING INTERESTS**

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141 Authors have declared that no competing interests exist.

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