Anti-infective properties of medicinal plants from the Baja California peninsula, Mexico for the treatment of *Fusarium oxysporum* f. sp. *basilici* in organic sweet basil (*Ocimum basilicum*)

Propiedades anti-infectivas de plantas medicinales de la península de Baja California, México para el tratamiento de *Fusarium oxysporum* f. sp. *basilici* en albahaca orgánica (*Ocimum basilicum*)

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**ABSTRACT**

Certified-organic farming systems in Baja California Peninsula and Northwest Mexico are nationally and globally recognized, especially due to the production of vegetables and aromatic herbs under protected agriculture systems. Based on the background of some species of the flora of Baja California Sur (BCS) to inhibit a diversity of microorganisms, the effect of 22 medicinal plants of the region was explored to know the *in vitro* activity against the fungus *Fusarium oxysporum* f. sp. *basilici* isolated from basil (*Ocimum basilicum* L.). The plants processed as crude ethanolic and aqueous extracts were analyzed in duplicate (three replicates) evaluating the inhibition of mycelial growth and spore germination. In mycelial inhibition test, all plants extracts (1000 mg L\(^{-1}\)) showed an effectiveness of 11 to 40% to inhibit *F. oxysporum*. The most effective plant extracts according to 50% effective inhibition dose (ED\(_{50}\)), were *Larrea tridentata*, *Hymenoclea monogyra* and *Lippia palmeri* with an ED\(_{50}\) of 220, 303 and 3000 mg L\(^{-1}\), respectively. Tukey’s PostHoc tests indicated that *H. monogyra* and *L. tridentata* are ten times (ED\(_{50}\)<300 mg L\(^{-1}\)) more effective than *L. palmeri* (ED\(_{50}\) 3000 mg L\(^{-1}\)). In addition, the dose-response trend analyzes according to the logarithmic-logistic model (*drc* packages), showed the maximum slope values between 100 and 1000 mg L\(^{-1}\). In the spore germination inhibition tests, most ethanolic extracts (5000 mg L\(^{-1}\)) showed an effectiveness between 21 and 80%. The results of this study demonstrated that the inhibitory potential of these plants used in BCS traditional medicine are a viable alternative for the control of *F. oxysporum* f. sp. *basilici* in organic basil production systems.

**Keywords**

organic agriculture • crown rot • fusariosis • Larrea • Lippia • crude extracts • wild oregano • BCS
Medicinal plants of BCS against *Fusarium oxysporum* in basil

**RESUMEN**

Los sistemas de agricultura orgánica-certificada en la Península de Baja California y el noroeste de México son reconocidos a nivel nacional y mundial, por la producción de verduras y hierbas aromáticas bajo los sistemas de agricultura protegida. Con base en los antecedentes de algunas especies de la flora de Baja California Sur (BCS) para inhibir una diversidad de microrganismos, se exploró el efecto de 22 plantas medicinales de la región para conocer la actividad *in vitro* contra *Fusarium oxysporum* f. sp. *basilici* aislado de la albahaca (*Ocimum basilicum* L.). Los extractos etánolicos y acuosos crudos se analizaron por duplicado (tres réplicas). En las pruebas de inhibición del crecimiento micelial, todos los extractos (1000 mg L⁻¹) mostraron una efectividad entre el 11 al 40% para inhibir *F. oxysporum*. Los extractos más efectivos de acuerdo con la dosis efectiva de inhibición al 50% (ED₅₀) fueron *Larrea tridentata*, *Hymenoclea monogyra* y *Lippia palmeri* con una ED₅₀ de 220, 303 y 3000 mg L⁻¹, respectivamente. Las pruebas PostHoc de Tukey indicaron que *H. monogyra* y *L. tridentata* son diez veces (ED₅₀ <300 mg L⁻¹) más efectivas que *L. palmeri* (ED₅₀ 3000 mg L⁻¹). Además, los análisis de tendencia dosis-respuesta de acuerdo con el modelo logístico mostraron los máximos valores de pendiente entre 100 y 1000 mg L⁻¹. En la inhibición de germinación de esporas, la mayoría de los extractos (5000 mg L⁻¹) mostraron una efectividad entre 21 y 80%. Los resultados de este estudio demostraron que el potencial inhibitorio de estas plantas utilizadas en la medicina tradicional de BCS son una alternativa viable para el control de la fusariosis causada por *F. oxysporum* f. sp. *basilici* en los sistemas de producción orgánica de albahaca.

**Palabras clave**

agricultura orgánica • pudrición de la corona • fusariosis • Larrea • Lippia • extractos crudos • orégano • BCS

**INTRODUCTION**

Sweet basil (*Ocimum basilicum* L.) is a popular herb used both fresh and dry as flavoring and antioxidant in food and pharmaceutical industries (22). Fresh basil has been the most demanded herb in the United States of America (USA) markets for the gourmet cooking industry, and ranks first among aromatic herbs used by Californian restaurants (4). Most of the basil production takes place in the state of Baja California Sur (BCS) in Mexico, where approximately 700 t are produced by around 200 specialized farms under the certified-organic agriculture scheme (41).

Basil crop is susceptible to a number of fungal diseases caused mainly by two large groups; one related with leaf spots produced by the genera *Cercospora*, *Curvularia* and *Alternaria*, and the second one is related to soil-borne rot diseases caused by the genera *Phytophthora*, *Pythium*, *Rhizoctonia* and *Fusarium* (33). Thus, fusarium disease can produce wilt and crown rot symptoms, when caused by *F. oxysporum* f. sp. *radicis-lycopersici* (FORL) (28), or premature defoliation, vascular wilt and crown rot when associated to *F. oxysporum* f. sp. *basilici* (FOB) (20). It is known that the spread of *Fusarium* in the American continent is so wide. In Argentina, high infection values were observed in soybean seedlings when infected with *F. graminearum* isolated from maize crop residues (5). In Mexico, some variants of *F. oxysporum* have been found associated with corky-root in coffee (19) and several species of *Fusarium* associated with avocado diseases (31). FOB colonizes the xylem of the host plant, resulting in its blockage and decomposition, with the consequent appearance of the most common and recognizable symptoms, (36). Fusarium disease was widely detected and caused a significant problem in cultivated basil in the USA for two decades. This is currently reported to be one of the most persistent diseases of this crop worldwide (45). Essential oils and tincture preparation are commonly used in a traditional medicine to treat minor infections (10). In organic agriculture, the extracts are commonly utilized as repellants for a wide range of pests (21) and, to a lesser extent, as inhibitors of several pathogenic *Fusarium* species (1, 9).

In BCS, the most important production areas, such as San José del Cabo, Todos Santos and La Paz, different *Fusarium* species have become the predominant problem in shade mesh or open cultivation (38). This occurs, mainly because the only solvents allowed in organic agri-
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culture are water and ethanol, which necessarily focus on the search for active molecules that may have activity with these solvents. In this sense, a variety of secondary metabolites and diverse compounds of aqueous and ethanolic fractions with antifungal activity have been described (18). However, despite the promising field of research on medicinal extracts with antimicrobial properties for agriculture (11, 24, 34), most of the ethnobotanical and biological heritage of BCS medicinal plants has focused on their potential use on bacteria and yeasts of clinical interest (12, 26).

Aware of the antifungal potential of rained plant resources in the southern part of the Baja California peninsula, the objective of this research was to explore the potential of 22 species of medicinal plants with known antimicrobial activity, as a source of antifungal compounds for the treatment of *Fusarium* wilt diseases in the production of organic basil in BCS, Mexico.

**MATERIALS AND METHODS**

**Collection of plant material**

Twenty-two plants used in traditional medicine with antimicrobial properties were selected and collected from different areas of the southern desert region of the Baja California in Mexico. They were grouped according to their traditional behavior and ethnobotanical uses. Table 1 summarizes the species name, botanical identification and ethno-pharmaceutical data. The Voucher specimens were deposited in the Herbarium of the Centro de Investigaciones Biológicas del Noroeste (CIBNOR) in La Paz, BCS, México. All plant material was collected during the 2012 rainy season (August to October). The vegetative material was only from leaves that were transported and pressed with a botanical press and dried at room temperature.

**Fungus isolation and molecular identification**

Isolation was performed from plants collected in El Pescadero, BCS with symptoms associated with fusariosis disease according to Summerell et al. (2003). The presumptive *Fusarium* species were characterized based on colony growth on *Fusarium*-selective medium. Potato dextrose agar (PDA) medium was used to determine colony morphology, growth rates and pigmentation. For the production of chlamydospores, macro- and microconidia, the medium CLA (carnation leaf agar) was used (43, 47). For molecular identification, total DNA from the fungus recovery from agar medium was obtained as template in polymerase chain reaction (PCR). One microliter (50 ng μL) of the supernatant was amplified with the primer pair Bik 1 (5’-TTCAAGAGCTAAAGGTCC-3’) and Bik 4 (5’-TTTGACCAAGATAGATGCC-3’). PCR conditions were according to the standard procedures for specific detection of FOB (7).

**Extracts preparation**

Aqueous and ethanolic extracts were prepared. The aqueous extract was obtained by grinding 10 g of air-dried powder in 100 mL of sterile distilled water, which was boiled on slow heat for two hours. The macerate was sieved with a five-layer muslin cloth and centrifuged at 4000 g for 30 min. The collected supernatant was filtered with Whatman filter paper No. 2 (GE Healthcare, UK) and autoclaved at 121 °C and 15 lb pressure. The resulting solution of 100 g L\(^{-1}\) was considered as mother stock. For ethanol extracts, the leaves were dried at room temperature for two weeks and then ground into powder with a blender: Then, 50 g of the powdered plant material were soaked in 200 mL (95% ethanol) at room temperature for five days. Each of the solvent extracts was shaken daily with a shaker for regular infusion. The filtrates obtained were evaporated using a rotary vacuum vapor distillation (Buchi Rotavapor R-114, Labortechnik, AG) at 60 °C, re-suspended in 20 mL of ethanol and fixed at a concentration with 100 g L\(^{-1}\) as maximum in test concentration and stored in sterile glass bottles at 5 °C until use (12).

**Preliminary inhibition screening**

To investigate the inhibition potential of the selected medicinal plants, two different methods were used. The first was the sensi-disc diffusion method (SDD) selected to perform the general screening of the 22 selected plants at 1000 mg L\(^{-1}\). This type of test provided a quick and inexpensive way to obtain reliable quantitative results. The three most active extracts in SDD
were selected for screening in the agar-dilution method (ADL), which allowed the sample to dissolve the agar. Therefore, its concentration was constant throughout the plate and was used to evaluate the effective doses (ED$_{50}$) of mycelial growth inhibition in response to 50% of the expected effect (7). Inoculum concentrations were prepared and adjusted to $10^6$ CFU mL$^{-1}$.

**Table 1.** Ethnobotanical data of plants collected from the Baja California peninsula in Mexico, according to the ethno-pharmacological use.

**Tabla 1.** Datos etnobotánicos de las plantas recolectadas de la península de Baja California de México, según el uso farmacológico.

| Family/plant species | Common name/native name | Voucher$^1$ | Ethno pharmacological uses                          |
|----------------------|-------------------------|-------------|---------------------------------------------------|
| Anacardiaceae        | Cytocarpa edulis        | AS-218      | Disinfection of gums                               |
| Cyrtocarpa edulis    | Plum tree/ciruelo del monte |           |                                                    |
| Aristolochiaceae     | Aristolochia monticola | AP-269      | Intestinal parasites, amoeba and skin infections   |
| Brandegee            | Dutchman’s pipe/hierba del indio |         |                                                    |
| Asteraceae           | Ambrosia ambrosoides    | AB-111      | Antiseptic and antibiotic                          |
| (Cav) Payne          | Canyon ragweed/chicura  |             |                                                    |
| Ambrosia ludoviciana | Triangle leaf bursage or donkeybush/estafate | AP-21   | Amoebae and parasites, insecticide                  |
| subsp mexicana       |                         |             |                                                    |
| Anaphalis margaritacea | Western pearly everlasting/ gordolobo | AP-17 | Stomach parasites, antiseptic                       |
| (L.) Benth.          |                         |             |                                                    |
| Artemisia absinthium | Master herb or ajenjo macho/ prodigiosa | AS-134 | Antiseptic, parasites and anthelmintic             |
| L.                   |                         |             |                                                    |
| Baccharis glutinosa  | Douglas’ false willow/ guatamote | AS-271 | Antiseptic for infections in sores and wounds       |
| Pers.                |                         |             |                                                    |
| Perityle californica | Rock daisy/manzanilla del monte | AB-250 | Preventive of venereal diseases                    |
| Benth.               |                         |             |                                                    |
| Haploppappus sonorensis | Haploppappus/hierba del pasmo | AS-46 | Preventive for minor infections, antiseptic        |
| (A. Gray) S.F. Blake | California trixis/hierba de Santa Lucia | AP-185 | Stomach parasites                                  |
| Trisix californica   |                         |             |                                                    |
| var. peninsularis   |                         |             |                                                    |
| & A. Gray.           |                         |             |                                                    |
| Hymenoclea monogyra  | Singlewhorl or burrobrush/ romerillo | AB-47 | Tetanus disease                                    |
| Torr.               |                         |             |                                                    |
| Bignonaceae          | Calabash or cirian/ Teomate | AP-131 | Parasites                                         |
| Crescencio alata     |                         |             |                                                    |
| Kunth                |                         |             |                                                    |
| Boraginaceae         | Borago officinalis      | AB-345      | Measles and smallpox                               |
| L.                   | Borege or cooltankard t.aiwot/borrajá |         |                                                    |
| Heliotropium curassavicun (L.) var. ooultum (Heller) | Wild heliotrope/hierba del sapo, berro, María Luisa | AS-23 | Antiseptic and to wash bounds                       |
| Euphorbiaceae        | Acalypha comonduana     | AP-150      | Parasites and antiseptic                           |
| Mill.                | Cooper leaf/hierba sanalotodo |         |                                                    |
| Ricinus communis     | Castor beans/higuerrilla | AS-179 | Antiseptic, antibiotic, acaricial, insecticidal   |
| L.                   |                         |             |                                                    |
| Euphorbia nutsan (Lag.) | Nodding spurge/ hierba golondrina | AP-38 | Parasites                                         |
| Jatropha cinerea     | Arizona netlespurge/ lomboy | AS-27 | Antiseptic                                         |
| (Ortega) Muell.-Arg. |                          |             |                                                    |
| Nyctaginaceae        | Mirabilis jalapa       | AS-52       | Infected wounds                                    |
| L.                   | Common four o’clock/ maravilla |         |                                                    |
| Pedaliaceae          | Proboscidea althaeofolia (Roxb) Benth | AS-161 | For the treatment of infected wounds               |
| Verbenaceae          | Lippia palmeri S. Watson | AF-44 | Antimicrobial                                      |
| Larrea tridentata    | (Seesé & Moc. Ex D.C.)  | AS-50.1     | Parasites, antiseptic                              |
| Wild or Mexican oregano/ oregano del monte |         |             |                                                    |

$^1$ Key letters in voucher were according to the reference use; antibiotic (AB), antifungal (AF), antiseptic (AS), parasite prevention (AP), other traditional uses (TU).

$^1$ Las letras de las claves de la ficha botánica son de acuerdo a la referencia de uso; antibiótico (AB), antifúngico (AF), antiséptico (AS), prevención de parásitos (AP), otros usos tradicionales (TU).
**Sensi-disc diffusion (SDD) method**

After preparing the PDA medium, *F. oxysporum* strains were incubated at 25 °C to allow mycelial development from seven to 10 days. Consequently, six-millimeter cylindrical sections (explants) were cut from the fungal-medium, and used as inoculum for testing. The explant containing the test microorganism was inoculated in the center of the previously prepared sterile agar bed (15 mL) and contained in the Petri dishes. Radially, four diffusion paper discs (0.5 cm in diameter) were placed on each agar plate previously impregnated with 1000 mg L⁻¹ of each extract. Subsequently, the diameters of the inhibition zones were measured, and the inhibition percentage of radial growth was calculated. Growth inhibition was established in ranges with values from 0.1 to 10% (−), 11 to 20% (±), 21 to 40% (+), 41 to 80% (++), and values ≥ 81% (+++).

**Agar dilution method (ADL)**

The best treatments for SDD with an activity between 21 and 40% (++) in all the tests evaluated were tested by the ADL method to obtain ED₅₀ and ED₉₀. The doses were determined logarithmically based on 2, 4, 8, 20, 40, 80, 200, 400, 800, 2000, 4000, 8000 and 20000 mg L⁻¹. The plates were inoculated at the center with 5 mm of fungal culture disc and then incubated at 28 °C for seven days. The effectivity of the extracts was calculated as mycelial growth inhibition using the formula inhibition percentage (%) = [(control growth-sample growth) / (control growth) x 100]. As controls, 0.1% benomyl (1000 mg L⁻¹) was used according to the *in vitro* doses of systemic (250 to 1000 mg L⁻¹) and non-systemic fungicides (1000 to 2000 mg L⁻¹)(29). Treatments using only ethanol and water (1 mg mL⁻¹ each) were included in the controls. The treatments had four repetitions and were carried out in duplicate.

**Spore germination inhibition**

Based on the literature, spore germination inhibition studies were carried out by the ADL method using only the ethanol plant extracts (29). The stock solutions (S) were prepared as ethanolic extracts at a concentration of 5 mg mL⁻¹. Fungal spores (~500) from one-week-growing cultures, together with one drop of each plant extract at each concentration, were placed on a glass slide and incubated (25 ±2 °C) for 24 h. The spores were stained with cotton blue and mounted in lactophenol; 100 μL were placed in a Neubauer chamber (Geleromics, Grenoble, FR) and observed under a light microscope. Inhibition percentage was calculated with the formula: percentage of spore germination inhibition (SGI) = [(A-B) / A) x 100, where A = Number of germinated spores in the control-number of germinated spores in the treatment, B = number of germinated spores in control (42).

**Statistical analysis**

The dose-response analysis was performed with a generalized log-logistic model using R Statistical Computing of drc packages (40). To fit the dose-response model, ED₅₀ was determined according to the three-parameter log-logistic function (LL.3) where the lowest limit was zero. Data were expressed as the mean (± SE). The lack-of-fit was determined by one-way analysis of variance (ANOVA). Tukey HSD test was used for multiple comparisons between treatments (extracts) with a previous evaluation of normality (Shapiro–Wilks test) and homogeneity of variance (Levene’s test). Statistical significance in spore germination was analyzed by Student’s t-test. All experiments were conducted in duplicate.

**RESULTS AND DISCUSSION**

**Fungus identification and disease determination**

The fungus, consistently isolated from the vascular tissue, was preliminarily confirmed as *F. oxysporum* based on the morphological characteristics of the mycelia and conidia grown in CLA medium. The characteristics that describe the species were observed, from the pale pink pigmentation of the mycelium to the observation of microconidia, macroconidia and globose chlamydospores (47). The pathogenic fungus was identified as *Fusarium oxysporum* f. sp. *basilici* based on the specific amplification of the expected 943 bp with Bik 1 and Bik 4 primers. No amplification was obtained from DNA extracted from healthy seedlings. In addition, the symptoms of vascular wilt, crown and root rot observed in the field were similar to those reported in other basil production systems in the world (17, 36, 37, 46).
**In vitro screening of mycelial growth activity**

Most of the aqueous-ethanol extracts tested in initial screening by SDD method showed an inhibition effectiveness similar to those reported with other filamentous fungi worldwide (7, 8, 11, 15, 18, 28, 30) with inhibition ranges between 1000 mg L\(^{-1}\) for mycelial growth (29, 42). Table 2 shows that all selected plants (24) showed at least moderate activity (21 to 40%) in mycelial growth. Among the most promising crude extracts with potential to be used in organic agriculture against *F. oxysporum* were *L. tridentata*, *H. monogyra* and *L. palmeri*, with mostly inhibitions from 41 (++) to ≥ 81% (+++) with water and ethanol solvents (1000 mg L\(^{-1}\)).

The next most effective plants with values from 21 to 40% were *Aristolactea monticola*, *Cyrtocarpa edulis*, *Haploppappus sonorensis* and *Ambrosia ambrosoides*. The ethanol extracts of these selected plants significantly showed higher activity compared to aqueous extracts. Thus, ethanol and aqueous extracts with the highest activity were selected for further analysis of 50%-effective doses (ED\(_{50}\)) and spore inhibition tests.

Table 2. *In vitro* screening of mycelial radial growth and spore germination inhibition of the most effective extracts by the SDD method against *Fusarium oxysporum* f. sp. *basilici* after six days.

| Plant species                      | Range of inhibition (%) \(^1\) | Radial growth mycelium \(^2\) | Spores \(^3\) | Ethanol | Water | Ethanol |
|------------------------------------|---------------------------------|-------------------------------|--------------|---------|-------|---------|
| *Larrea tridentata*                | ++                              | ++                            | +++          |         |       |         |
| *Hymenoleca monogyra*              | ++                              | +                             | +++          |         |       |         |
| *Lippia palmeri*                   | ++                              | ++                            | ++           |         |       |         |
| *Trixis californica var. peninsularis* | +/-                             | +/-                           | +++          |         |       |         |
| *Aristolactea monticola*           | +/-                             | +/-                           | +/+-         |         |       |         |
| *Cyrtocarpa edulis*                | +/+-                            | +/-                           | ++/+-        |         |       |         |
| *Haploppappus sonorensis*          | +                               | +/-                           | +/+-         |         |       |         |
| *Ambrosia ambrosoides*             | +/-                             | +/-                           | ++/-         |         |       |         |
| *Baccharis glutinosa*              | +/-                             | -                             | +/+-         |         |       |         |
| *Crescencia alata*                 | +/-                             | -                             | +/-         |         |       |         |
| *Artemisia absinthium*             | +                               | -                             | +/+-         |         |       |         |
| *Heliotropium curassavicun*         | +/-                             | +/-                           | +/-          |         |       |         |
| *Ricinus communis*                 | +/-                             | +/+-                          | -            |         |       |         |
| *Euphorbia nutans*                 | +/-                             | +/+-                          | ++           |         |       |         |
| *Jatropha cinerea*                 | +/+-                            | -                             | +/+-         |         |       |         |
| *Mirabilis jalapa*                 | +/+-                            | -                             | +/+-         |         |       |         |
| *Proboscidea althaefolia*          | -                               | +                             | ++           |         |       |         |
| *Acalypha comonduana*              | -                               | +                             | +            |         |       |         |
| *Anaphalis margaritacea*           | +/-                             | +/+-                          | -            |         |       |         |
| *Borago officinalis* L*            | +/-                             | +/+-                          | -            |         |       |         |
| *Ambrosia ludoviciana* (Cav.) Payne| +/-                             | +/+-                          | -            |         |       |         |
| *Perityle californica* Benth       | +                               | +                             | -            |         |       |         |
| *Benomyl* (1000 mg L\(^{-1}\))    | +++                             | +++                           | +++          |         |       |         |
| Control 1 (ethanol 70%)            | -                               | -                             | -            |         |       |         |
| Control 2 (water)                  | -                               | -                             | -            |         |       |         |

1 The inhibition symbology (+, +) in the table was according to the observed ranges: (-) ≤ 10% growth inhibition, (±) 11 to 20%, (+) 21 to 40%, (+++) 41 to 80%, and (++++) ≥ 81% (n = 6). 2 All extracts for mycelial inhibition were extracted from the leaves at standard concentrations of 1000 mg L\(^{-1}\). Spore germination inhibition test was carried out at concentrations of 5 mg mL\(^{-1}\). 3 Spore germination inhibition test was carried at concentrations of 5 mg mL\(^{-1}\).

**Spore germination inhibition analysis**

At least 17 plant extracts showed some inhibitory effect from 11 to 20% at low doses of 5 mg mL\(^{-1}\). The best treatments for the inhibition of the fungus were *L. tridentata*, *H. monogyra* and *Trixis peninsularis*. They were the most effective with inhibition above 81% (+++). Similar response at 5 mg mL\(^{-1}\) has been observed in other ethanol extracts against the *in vitro* *F. oxysporum* test (6, 29). The extracts of *L. palmeri*, *Heliotropium curassavicun*, *Euphorbia nutans* and *Proboscidea althaefolia* showed an inhibition of sporulation...
in a range from 41 to 80%, although the effect on mycelial inhibition was moderate (11 to 20%). The tendency of progressive increase in sporulation inhibition with the increase of concentrations was also observed (table 2, page 239).

**Effectiveness of dose response (ED_{50})**

Model parameters of ED_{50} indicated 50% of effective biocontrol (dose-response). ED_{50} response of the most effective plant extracts by the ADL method were *L. tridentata*, *H. monogyra* and *L. palmeri* with values ranging from 220 to 3000 mg L\(^{-1}\) (table 3). The three tested extracts showed significant differences among the effective doses (one-way ANOVA, F = 136.6, P < 0.001). The PostHoc Tukey HSD test indicated that ED_{50} of the *L. tridentata* (ED_{50} = 220 mg L\(^{-1}\) and *H. monogyra* (ED_{50} = 303 mg L\(^{-1}\)) were more efficient to inhibit the fungus (figure 1A, page 241). The predictive trend of the three extract curve analysis, focusing on dose-response at the logarithmical concentration studied (10-10000 mg L\(^{-1}\)), were more pronounced in *L. tridentata* than *H. monogyra* and *L. palmeri* (figure 1B, page 241). However, no significant differences were observed between ED_{50} of *L. tridentata* and *H. monogyra* (P > 0.05). This result indicated that *L. tridentata* was 13.6 times more effective than *L. palmeri* and 1.3 times more than *H. monogyra*.

Table 3. Percentage of inhibition of the radial mycelial growth of *Fusarium oxysporum* f. sp. *basilici* of the extracts with the highest antimicrobial activity in the SDD tests, showing the effective dose (ED_{50}), logarithmical concentration and the fiducial limits

| Logarithmical concentration (mg L\(^{-1}\)) | Mycelial radial inhibition (%) | Lippia palmeri | Larrea tridentata | Hymenoclea monogyra |
|------------------------------------------|-------------------------------|----------------|------------------|---------------------|
| 40                                       | 5.25±0.4                      | 33.91±1.74     |                  |                     |
| 60                                       | 10.36±.38                     | 30.78±0.68     |                  |                     |
| 80                                       | 9.69±0.86                     | 42.81±1.13     | 36.02±0.93       |                     |
| 200                                      | 14±0.23                       | 52.66±1        | 46.77±0.81       |                     |
| 400                                      | 26.65±0.81                    | 65.16±2.11     | 54.3±0.94        |                     |
| 600                                      | 32.97±0.59                    | 73.28±0.66     | 59.01±1.06       |                     |
| 800                                      | 40.38±0.36                    | 78.25±0.9      | 63.98±0.51       |                     |
| 2000                                     | 53.97±3.01                    | 87.81±0.37     | 73.12±1.09       |                     |
| 4000                                     | 68.64±3.52                    | 100            | 79.03±1.69       |                     |
| 6000                                     | 79.27±0.45                    | 100            | 84.5±0.41        |                     |
| 8000                                     | 85.87±0.61                    | 100            | 84.11±1.07       |                     |
| ED_{50}                                  | 3000.28±979.75                | 220.87±31.01   | 303.46±33.73     |                     |
| P                                        | 0.008                         | 0.0022         | 0.0026           |                     |

The ED_{50} of the BCS plant extracts were lower than those of other of similar reported species (*Origanum* spp.), extracted as essential oils, where ED_{50} was from 450 to 8000 mg L\(^{-1}\) (3). The literature has reported that the effectiveness *in vitro* of the aqueous-ethanol extracts of medicinal plants has a wide range from 100 to 5000 mg L\(^{-1}\) to inhibit fungi of different genus; *Aspergillus*, *Colletotrichum*, *Phytophthora*, *Penicillium*, *Botrytis*, *Alternaria* and *Fusarium* (15, 35). On the other hand, the formulation of botanical fungicides with good potential to inhibit a wide range of filamentous fungi ranges from 1000 to 2000 mg L\(^{-1}\) (44).

In spore germination assays, the extracts of *L. tridentata*, *H. monogyra* and *T. peninsularis* were the most effective to inhibit spore germination at 5000 mg L\(^{-1}\) by more than 81%, followed by the extracts of *L. palmeri*, *H. curassavicun* and *P. althaefolia*, which presented values of 41 to 80% (table 2, page 239). The results of this study were in agreement with Obongoya *et al.* (2010) who showed the phytotoxic effect selecting crude extracts of some Mexican plants against soil-borne *F. oxysporum* f. sp. *phaseoli* ranging from 2.5 to 10 mg mL\(^{-1}\).
This research showed that three species used in traditional medicine in BCS could potentially control fusarium disease. However, for this particular species (*F. oxysporum* f. sp. *basilici*), very few studies have documented the antifungal activity, and the most documented are especially related to antimicrobial activity with some bacteria of clinical interest (12, 26). By far, *L. tridentata* locally known as "gobernadora" (governor), creosote bush or chaparral, is one of the most documented species with inhibitory effects. For approximately half a century, *L. tridentata* has been the most studied plant in Mexico and the world and reported for the treatment of more than 50 diseases (2). The effect of this species as anti-parasitic, antiseptic and control of foot fungi and bacteria and treatment of kidney and gynecological infections are well documented (16). One of the most active ingredients, nordhydroguaiaretic acid (NDGA) and numerous lignans have been characterized. This species was found to be the most promising in this study based on values of 220 mg L\(^{-1}\) and values ≥ 81% to inhibit sporulation at 1000 mg L\(^{-1}\). Oregano is one of the most interesting cases from its nomenclature, which can be referred to as Mexican or wild oregano (*L. palmeri*, Verbenaceae family) or in the world known as common oregano (*Origanum vulgare*; Lamiaceae family). The microbial properties of both species are well documented, and it appears to be an interesting promise because of its availability as an agricultural domesticated species (27, 32). In addition to their wide use in the food industry, active compounds, such as eugenol, carvacrol, carvone, p-cymene and thymol inhibit mycelial growth and sporulation of different types of *Fusarium* species in a variety of crops (13). The effectiveness of...
L. palmeri as an aqueous-ethanol extract has been observed in other medicinal and culinary plants in the state of BCS, such as sage (Salvia officinalis) and where this type of extraction maximizes antifungal components (10). On the other hand, its multiple use as an export crop (41) could be interesting for the industry as essential oil or dry for the food industry or for the formulation of natural antifungal solutions for agricultural use (8). Hymenoclea monogyra seems to be the least documented; however, different species of the genus have shown antimicrobial properties due to the presence of flavonoids and sesquiterpene lactones. H. monogyra has been reported against enteropathogenic bacteria in BCS and in other regions of northwest Mexico (25). A diversity of active compounds can be extracted from medicinal plants using ethanol-water methods, which makes it an interesting model for the semi-pilot scale extractions, providing confidence for the optimization processes in the formulation of botanical fungicides.

Mexico is a country with an agricultural vocation where certified agriculture activities in the production of vegetables and aromatic herbs occur mainly in the states of Sinaloa, Sonora and BCS (23). The most important regions of the peninsula are mainly in Ensenada, which is the largest municipality with an organic agricultural vocation. Other producing areas in the southern part of the peninsula, such as Santo Domingo, Valle de Vizcaino, La Paz and San Jose del Cabo are known for the production of annual vegetables and for the certified organic production of aromatic herbs (14). Basil is the most commercialized gourmet herb species that generates a good economic performance (39). Although losses due to fusarium in this crop have not been estimated in BCS, the pressure of the disease and the lack of botanical fungicides for its control are a constant risk for production systems. This study has reported F. oxysporum f. sp. basilici associated with crown and root rot and wilt diseases in basil grown in BCS for the first time. Crude aqueous-ethanol extracts of L. tridentata, L. palmeri and H. monogyra, can efficiently inhibit mycelial growth and spore germination of F. oxysporum f. sp. basilici with potential use in phytosanitary management programs for organic basil production.

CONCLUSIONS

The in vitro effectiveness of Fusarium-growth inhibition (ED$_{50}$ ranges from 220 to 3000 mg L$^{-1}$) and inhibition of sporulation (80% at doses of 5000 mg L$^{-1}$) of native medicinal plants (L. tridentata, H. monogyra and L. palmeri) from the Baja California peninsula, opens the possibility of using them as an ethanol-aqueous extract for long-term control of the fusarium disease caused by F. oxysporum f. sp. basilici in organic basil in BCS, Mexico and in other producing areas of the world.

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