



Allelopathic properties of *Calliandra haematocephala* Hassk. extracts and fractions as an alternative for weed management in quinoa and rice crops

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Abstract

In this study, aqueous extracts of *Calliandra haematocephala* Hassk. leaves and inflorescences were tested on seeds of quinoa (*Chenopodium album* L.) and rice (*Oryza sativa* L.), and on some of the most noxious-associated weeds, *Chenopodium album* L. and *Holcus lanatus* L. in quinoa, and *Echinochloa crus-galli* (L.) P. Beauv., *Echinochloa colona* L., *Eclipta prostrata* L. and *Rottboellia cochinchinensis* (Lou.) W.D. Clayton in rice. The objectives were to identify extract concentrations at which 50 and 90% of germination ($GR_{[50,90]}$) and radicle elongation ($RR_{[50,90]}$) were inhibited, to fractionate inflorescence extracts for facilitating identifying the chemical group causing allelopathic effects, and to evaluate the fraction showing the stronger weed suppression effects and the least crop damage. Increasing extract concentration rates (0, 6.25, 12.5, 25, 50 and 100% crude extract) were applied to seeds of target crops and weeds. Flower extracts at rates < 0.30 produced $GR_{[50]}$ and $RR_{[50]}$ on *H. lanatus*, and $GR_{[90]}$ and $RR_{[90]}$ in *C. album*, while quinoa seeds were not affected. Rice and its target weeds were minimally affected by flower extracts, whereas radicle elongation of all species was significantly reduced. A concentration rate > 0.52 caused the $RR_{[50]}$ on rice and all its target weeds. Fractions were quantitatively and qualitatively analysed to detect phytochemical groups, using specific chemical reagents and thin-layer chromatography (TLC). The fraction F3 from aqueous flower extract showed a high content of flavonoids, assumed as the potential allelochemical substance. Total flavonoid content in F3 was quantified as 2.7 mg of quercetin per g F3, *i.e.*, 12.8 mg of quercetin per g of inflorescence material. Additionally, field equivalent extract rates obtained from the harvested fresh inflorescence biomass could be determined. These rates ranged between 90 and 143 mL l⁻¹ of F3 aqueous fraction, while for ethanol F3 were 131 mL l⁻¹. Our results are encouraging for finding sustainable and ecologically friendly alternatives for weed management in crops of high nutritional value, contributing also to counteract the growing problem of herbicide resistance.

Keywords Dose–response · Bioassay · Germination reduction · Radicle elongation reduction · Flavonoids · Quercetin · Field equivalent rate

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Introduction

The advent of synthetic pesticides in the 1930s and 1940s shaped agriculture as it is known today (Hall et al. 2000; Rattner 2009). Crop yields have increased remarkably with the use of synthetic fertilizers and pesticides. Almost one of every ten people in the world become ill due to the consumption of contaminated food or water, due to microbial factors and chemical contamination, the latter involving pesticide residues (World Health Organization 2017). Aspects referred to water and soil pollution and their persistence in the environment due to the misuse of pesticides are also well documented (Schwarzenbach et al. 2010; Morrissey et al.

2015; Yadav et al. 2015). Vonberg et al. (2014), based on monitoring data of a shallow aquifer exposed to the effects of intensive agriculture in western Germany, reported the presence of atrazine (in concentrations above $0.1 \mu\text{g l}^{-1}$) after 20 years of being banned. Therefore, the need of alternative crop management strategies, such as the use of natural products, is evident. Natural pesticides may show advantages, assuming that their persistence in the environment is considerably shorter than those synthetically produced.

Weeds cause huge yield losses around the globe. Weed control strategies vary largely among cropping systems, environmental conditions and farm economic factors (Rueda-Ayala et al. 2010), but mainly includes the application of synthetic herbicides (Thill et al. 1991). Continuous use of the few available active ingredients of herbicides has enhanced the selection of resistant weed populations. These populations have increased immensely in the last decades, thus demanding that weed management strategies evolve accordingly (Weis et al. 2012; Colbach et al. 2016).

Alternative substances for weed control can also be of natural origin. The primary source for many organic herbicides is chemicals released by plants, which have direct or indirect detrimental effects on germination, growth or development of other plants (Webber et al. 2012; Koocheki et al. 2013). Different natural products have already been tested and marketed. However, their efficacy is generally low, compared with synthetic products (Cordeau et al. 2016). The phytotoxic effect of aqueous extracts of different plant materials used as bioherbicides have been widely explored, from in-vitro bioassays to greenhouse and field experiments (de Albuquerque et al. 2011; Bhadoria 2011; Rueda-Ayala et al. 2015). Allelopathic effects may be attributed to phenolic compounds found in one or various plant parts, and these effects can be strengthened by mixing compounds of different plant species (Reigosa and Pazos-Malvido 2007). Puig et al. (2018) identified allelopathic activity and the chemical composition of different species of the genus *Eucalyptus* (Myrtaceae). Phytotoxic effects of *E. globulus* aqueous extracts affected the germination and radicle elongation of some crops and weeds with specific modes of action on different target species. Similarly, (Souto et al. 1994) identified allelopathic effects on four forestry species. The soils associated with those studied species appeared to be responsible for the toxic effects, indicating release of allelopathic compounds by the plant roots.

Many growth inhibitors found in plants have been effective for pathogen and weed control (Xuan et al. 2005). The high number of allelopathic compounds existing in various plant species need to be assessed, to determine their potential as alternatives to synthetic herbicides, *i.e.*, as bioherbicides. From leaves of *Calliandra* species, some compounds with insecticidal properties have been identified, *e.g.*, nonproteic imino acids (Romeo 1984). Raja et al. (2017)

synthesized nanoparticles from leaf extracts of *Calliandra haematocephala* Hassk. with antibacterial activity. To the best of our knowledge, nothing is known about allelopathic properties of this plant against weeds in cropping systems. Prior to this study, a high suppressive effect was observed on germination of numerous weed species when soils were covered by vegetative and inflorescence structures (mulch) of *C. haematocephala* shrubs. Furthermore, if some seeds did germinate, they became withered a couple of days later, even when water availability was sufficient. These facts motivated our interest on testing plant extracts of this shrub species.

Testing, validating and adapting the use of bioherbicides to the local conditions and important crops of every agricultural region should be carried out. In this study, aqueous and ethanol-based extracts from leaves and inflorescence of *C. haematocephala* were prepared with the aim of determining the presence of allelopathic compounds. The specific objectives were, (i) to identify the extract concentrations at which 50 and 90% of germination and radicle elongation were inhibited in seeds of quinoa (*Chenopodium quinoa* Willd.) and rice (*Oryza sativa* L.), including their associated most problematic weeds in the Ecuadorian context, (ii) to fractionate inflorescence extracts for facilitating identifying the chemical group causing those effects, and (iii) to evaluate the fraction showing the stronger weed suppression effects and the least crop damage.

Materials and methods

The study was carried out in the laboratories of the Ecuadorian Agency for Quality Assurance in Agriculture, AGRO-CALIDAD ($0^{\circ} 12' 52'' \text{ S}$; $78^{\circ} 24' 40'' \text{ W}$), near Quito, Ecuador. Experiments were implemented in two phases: (1) detection of allelopathic effects in *C. haematocephala* extracts, and (2) fractionation of those extracts for chemical screening and identification of the possible involved substances. Leaves (645 g) and inflorescences (1110 g) were collected from *C. haematocephala* shrubs (10 plants) on neighboring gardens, during 2015, 2016 and 2017 (Table 1). Prior to extraction, all plant material was washed with a 1% solution of Sodium hypochlorite, rinsed with deionized water, dried in a stove at 40°C for 24 h, then stored at -20° until use.

Detection of allelopathic properties

Aqueous extracts were obtained separately from leaves and from inflorescence of *C. haematocephala*. Different amounts of solvent per plant part, according to the requirements of each experiment, were used to obtain the extracts. Leaves of *C. haematocephala* are drier than inflorescences, thus they required more solvent to produce a uniform extract structure.

Table 1 Details for the preparation of *C. haematocephala* crude aqueous extracts: plant material used, fresh weight, volume of solvent applied per 50 g fresh weight, amount of extract obtained, concentration and experimental use on target plant species, *C. album* (CHEAL), *H. lanatus* (HOLLA), *E. colona* (ECHCO), *E. crus-galli* (ECHCG), *E. prostrata* (ECLAL) and *R. cochinchinensis* (ROOEX)

Experiment		Plant material*			Extract retrieved		Target species	
Nr.	Type	Structure	Weight (g)	Solvent (mL)	Amount (mL)	Concentration (g mL ⁻¹)	Crop	Weeds
1	Dose-response Bioassay	Leaf	328	20	107	2.5	Quinoa	CHEAL HOLLA
2	Dose-response Bioassay		313	20	102	2.50	Quinoa	CHEAL HOLLA
3	Dose-response Bioassay	Flower	285	15	141	3.33	Quinoa	CHEAL HOLLA
4	Dose-response Bioassay		276	15	136	3.33	Quinoa	CHEAL HOLLA
5	Dose-response Bioassay		434	15	238	3.33	Rice	ECHCG ECHCO ECLAL ROOEX
6	Fractionation [†] and Bioassay	Flower	53	18	67	2.78	Quinoa	CHEAL HOLLA
7	Fractionation [†] and Bioassay		53	90 [‡]	85	0.56	Quinoa	CHEAL HOLLA

*Collected in 2015 (exps. 1 and 3), in 2016 (exps. 2, 4, 6 and 7) and 2017 (exp. 5)

[†]Aqueous and ethanol-based crude extracts were partitioned into five fractions (F1–F5), and tested together with the crude extract (F0) and the corresponding blank, deionized water or 10%-ethanol solution

[‡]A 70%-ethanol solution was used instead of deionized water (see “Fractionation of flower extracts”)

In the case of extractions for fractioning, it was required to have a more diluted crude extract, to facilitate the phytochemical screening during the qualitatively analyses. A similar method as Rueda-Ayala et al. (2015) was used, but applying less deionized water to generate a highly concentrated stock solution (crude extract). Details for retrieving both, inflorescence (hereafter ‘flower extract’) and leaf extract and for carrying out bioassays is shown in Table 1 and Figure A1. Plant material, in 50 g at a time, were macerated and ground with liquid nitrogen, then dispersed in deionized water and Ultra-Turrax mixed for 1 min. This mixture was filtered on a vacuum pump through a Büchner funnel with qualitative filter paper. The remaining material was further pressed to retrieve the liquid and then filtered. The collected mixture was centrifuged at 3500 rpm for 15 min and the supernatant removed to purify the extract. Depending on the amount of deionized water added, extract concentrations varied (Table 1).

Allelopathic properties were studied via dose-dependent bioassays. Two economically important crops in the Ecuadorian context, including their most noxious weed species, were used as targets: quinoa (*Chenopodium quinoa* Willd., cultivar Tulcanaza), *Chenopodium album* L. (CHEAL) and *Holcus lanatus* L. (HOLLA), representing

the Andean region, and rice (*O. sativa* L.), *Echinochloa colona* L. (ECHCO), *Echinochloa crus-galli* (L.) P. Beauv. (ECHCG), *Eclipta prostrata* L. (ECLAL) and *Rottboellia cochinchinensis* (Lou.) W.D. Clayton (ROOEX), representing the coastal region. EPPO-code is given according to the EPPO Global Database (<https://gd.eppo.int>). Concentration rates, 1 (‘100%-extract’), 0.5, 0.25, 0.125 and 0.0625 plus a control (0), were derived by adding a dilution series with deionized water to the 100% crude extracts. Biochemical effects of each concentration on seed germination and seedling radicle elongation of the aforementioned target crops and weeds were tested. Certified crop seeds were acquired from the National Institute of Agronomic Research (INIAP) Santa Catalina, Quito, while weed seeds were collected from cultivated fields, without weed control. Two dose–response bioassays were implemented in quinoa during 2015, using the six extract concentrations, and two during 2017 in rice, using various concentrations of flower extract. In addition, the flower extract was fractionated (see “Fractionation of flower extracts”) and used on two bioassays during 2016 in quinoa and target species.

For the bioassays, 20 seeds per target species were placed into petri dishes with filter paper; 1 dish per tested species. All petri dishes were sterilized with ethanol 70% and UV

radiation for 30 min. Portions of 3 mL extract were applied onto the seeds and each petri dish was covered, sealed with Parafilm® and kept inside a growth chamber at 12 h/12 h day/night length and 24 °C/18 °C. Treatments were applied according to a factorial arrangement of the leaf and flower extracts, the six concentrations and the target species (quinoa and corresponding weeds, on experiments 1–4; rice and corresponding weeds on experiment 5) in a completely randomized design with three replications. Germination of weed seeds was initially tested on a sample of 100 seeds per species. These seeds were then introduced in a recipient with water, to eliminate non-viable seeds, which were assumed to be the ones floating. The initial germination percentage ranged between 30 and 60%. The same procedure was used to select the seeds for the bioassays, and only 10 germinated seeds were randomly selected for assessment of radicle length. Seed germination and radicle elongation of target species were assessed 4 weeks later, when all seedlings germinated in the control experimental units (concentration 0). Seedling germination of *H. lanatus* was delayed one more week in all extract concentrations > 0%.

Fractionation of flower extracts

For the second experimental phase, two types of crude extracts were prepared: aqueous and ethanol-based (Table 1). Extracts based on 70% ethanol and aqueous extracts were derived from inflorescence only. Cleaned and dried inflorescence were cut in small pieces, and 50 g were placed in a beaker glass with 150 mL 70%-ethanol solution, covered with aluminium foil and stored in darkness for 48 h. The mixture was filtered on a vacuum pump through a Büchner funnel with qualitative filter paper, then centrifuged at 3500 rpm for 15 min and the supernatant was removed to purify the extract. The extract solution was concentrated to 85 mL volume using a rotary evaporator and stored for fractionation and bioassays. Aqueous extracts were prepared as previously described (see “[Detection of allelopathic properties](#)” and Table 1). The extracts were fractionated to produce compounds of similar polarities or molecular sizes, as a starting step towards the identification and isolation of the active compounds causing allelopathic effects. At least 27 mL of each crude extract were used without further dilution in the bioassays as F0 (3 mL per petri dish, 3 target species and 3 repetition samples). Five fractions were generated, using the remaining extract amount, divided in three parts of equal volume, as explained below. The different fractions were obtained via various methods described in Seidel (2005), Otsuka (2005), Evans and Evans (2019) and Ribeiro et al. (2018).

Fractionation with chloroform (F1): A 5% hydrochloric acid solution was applied drop by drop to $\frac{1}{3}$ of the crude extract

until the pH level dropped to 2–3. Then, a 20% sodium hydroxide solution was added drop by drop, to rise the pH to 10–12. This solution was washed with 100 mL chloroform for three times to investigate the presence of alkaloids. The organic phase was concentrated by evaporation to dryness in a rotary evaporator and later reconstituted with sterile water to 50 mL. A 10% portion of this fraction was preserved for qualitative analysis, together with the resulting aqueous phase.

Fractionation with petroleum ether (F2): Petroleum ether (100 mL) was added three times through a separation funnel to another $\frac{1}{3}$ of the crude extract, to investigate presence of triterpenes and sterols. The organic phase was evaporated to dryness and reconstituted with sterile deionized water to 50 mL; a 10% portion and the aqueous phase were preserved for qualitative analysis and further fractionation.

Fractionation with methanol/water (F3 and F4): A 50:50 methanol/water mixture was added to the remaining aqueous phase from F2, for investigation of flavonoids, anthraquinones, tannins, saponins and reducing sugars. This solution was left for maceration for 24 h, then filtered on a vacuum pump and the solid part was solved again in chloroform. The residue was evaporated to dryness and reconstituted with sterile deionized water to 50 mL (F3), and the 10% part was preserved for qualitative analysis. The liquid residue from the filtration was also evaporated and reconstituted with sterile water to 50 mL (F4), preserving the 10% part for qualitative analysis.

Fractionation with lead acetate and chloroform (F5): The last $\frac{1}{3}$ of crude extract was precipitated by adding a 20% lead acetate drop by drop, and then vacuum filtered after which the solid residue was disposed off. The liquid phase was extracted with chloroform, dried with anhydrous sodium sulfate and filtered by gravity. These solvents were used to examine the existence of sesquiterpene lactones and coumarines (Tsao and Deng 2004). The resulting chloroform phase was evaporated to dryness, reconstituted with sterile deionized water to 50 mL and the 10% part preserved for qualitative analysis.

Bioassays with fractions

All obtained fractions (F1–F5), crude extracts (F0) and blanks (sterile deionized water and 10% ethanol) were tested on target species to determine the effects on germination and radicle elongation, via bioassays (Table 1, experiments 6 and 7). The implemented factorial experiment contained two extraction methods, ethanol and aqueous-based (factor method), six fractions (F0–F5) plus one control blank per

extraction method (factor fractions) and three target plant species (factor target species). The experiment was arranged in a completely randomized design with three repetitions. Management of bioassays and variable assessments were carried out similarly as for the dose–response bioassays (see “[Detection of allelopathic properties](#)”).

Qualitative and quantitative analysis

All fractions were analysed by thin-layer chromatography (TLC) for identification of the most important phytochemical groups with allelopathic properties. The mobile phases and developer substances used for TLC identification are described in Table 2. Further qualitative phytochemical group screening was applied, using specific tests and reagents (see “[Qualitative and quantitative analysis of fractions](#)”). A standard calibration curve with quercetin was built to quantify flavonoids on the aqueous fraction F3 (Ivanova-Petropulos et al. 2010; Hossain et al. 2013). Quercetin was measured in *C. haematocephala* flower extract, because flavonoids have been found in the form of quercetin in mimosaceae and fabaceae species (Sulaiman and Balachandran 2012; da Silva et al. 2015). F3 was chosen, because it showed high inhibiting effects on germination and radicle elongation of target species. Standard ethanol solutions (5 mL) at different concentrations were prepared, 2.0, 1.0, 0.5, 0.25, 0.125 and 0.0625 mg mL⁻¹. A sample of F3 was dissolved in ethanol, at a ratio 1/10 in weight (0.5/5.0 g); 1.25 mL ethanol and 0.075 mL sodium nitrite (5%) were added to both, the fraction sample and the standards, and left to rest for 6 min. Then, 0.15 mL aluminium chloride (10%) were added to the mixture, left to rest for 5 min, added 0.5 mL sodium hydroxide (1 M). The mixture was filled with ethanol to a volume of 2.5 mL. Thirty min later, the fraction sample and standards were

evaluated with the spectrophotometer at 510 nm. The fraction density was determined by pycnometry.

The fraction density of major activity was determined by Eq. (1), in which $M1$, $M2$, and $M3$ represent the weights of empty pycnometer (11.2 g), pycnometer with deionized water (12.19 g), and pycnometer with the fraction sample (12.22 g), respectively, and using a known density of deionized water at 24° ($\rho_{H_2O} = 0.9973$ g mL⁻¹). The quantity of flavonoids within the fraction of major activity (F3) was calculated with Eq. (2). The constants $a = 1.24$ and $b = -0.06$ came from the calibration curve (Figure A2); $A = 1.583$ was the determined sample absorbance and C is the concentration of flavonoids. This resulting C value was divided by the used sample weight (0.498 g) to calculate mg of quercetin per g of fraction (C_f). Total concentration of flavonoids was quantified by Eq. (3), as mg quercetin per g inflorescence material.

$$\rho_f = \frac{M3 - M1}{M2 - M1} \times \rho_{H_2O} \quad (1)$$

$$C = \frac{A + b}{a} \quad (2)$$

$$C_{f \text{ tot}} = C_f \cdot \rho_f \cdot \frac{50 \text{ mL}}{10.65 \text{ g inflorescence}} \quad (3)$$

Statistical analysis

Treatment effects on germination percentage (log transformed) and radicle length were determined using ANOVA for all bioassays (not shown). Effects of increasing extract concentration dose were described by nonlinear regression,

Table 2 Mobile phases and developer substances used for qualitative analysis by TLC of Phytochemical groups in the crude flower extracts of *C. haematocephala*

Fraction	Group	Mobile phase	Developer	References
F1	Alkaloids Methanol (19:1)	Chloroform Reagent	Dragendorff's	Sherma (2000)
F2, F4	Sterols and Terpenes	Petroleum-ethyl Acetate (80:20)	Acetic anhydride + Sulfuric acid + Ethanol	Sherma (2000)
F3, F4	Flavonoids Acid–water (4:1:5)	1-Butanol-acetic	UV/fluorescence	Ivanova-Petropulos et al. (2010)
F3, F4	Anthraquinones	Benzene-ethyl Acetate–acetic Acid (75:24:1)	Potassium Hydroxide	Sherma (2000)
F5	Sesquiterpene Acetone (60:40)	Dichloromethane- Hydroxide	Potassium	Ivanova-Petropulos et al. (2010)

using a log-logistic model with three parameters, Eq. (4), where $f[x(b, d, e)]$ is the expected target–plant response; parameter b denotes the relative slope around e , which is the dose reducing 50% of germination or radicle elongation (GR₅₀ or RR₅₀, respectively); d is the upper asymptote limit and x is the extract concentration (Streibig 1980; Ritz and Streibig 2005). Regression models were tested for lack-of-fit ($\alpha = 0.05$) and via graphical agreement between fitted values and residuals (Belz and Piepho 2012). Model selection was according to the lowest value of AIC (Akaike information criterion). Parameter estimates from non-linear models were accompanied by 95% confidence intervals (95% CI), which allow making a direct evaluation of how precise estimates were and easily determine statistical differences when intervals do not overlap, contrary to what is possible by providing only standard errors (SE). All analyses were done with the statistical software R version 3.5.1 – ‘Feather Spray’, The R Foundation for Statistical Computing (R Core Team 2018). Dose–response curves of asymmetrical sigmoid form were fitted with the R package `drm` (Ritz et al. 2015). Only the case of leaf extract effect on germination of target species was defined by the dose-dependent log-logistic function with four parameters, Eq. (5), which derives from Eq. (4) with a lower asymptote limit c . Furthermore, the doses causing 50 or 90% inhibiting effect on germination (GR_[50,90]) and radicle elongation (RR_[50,90]) were calculated.

$$f[x(b, d, e)] = \frac{d}{1 + \exp\{b[\ln(x) - \ln(e)]\}} \quad (4)$$

$$f[x(b, c, d, e)] = c + \frac{d - c}{1 + \exp\{b[\ln(x) - \ln(e)]\}} \quad (5)$$

The bioassays to study the allelopathic potential from fractions (see “[Fractionation of flower extracts](#)”) were analysed by ANOVA to identify the main effects of the factorial experiment, extraction method \times fractions \times target plant species. Successive F tests were applied to reduce model complexity. Tukey ($\alpha = 0.05$) Honest Significance Difference (HSD) to identify treatment differences, including the general average (AVG), the coefficient of variation (CV) and the HSD value. Further post hoc analysis was applied to the factorial interactions (Ottenbacher 1991), to study whether each fraction interacted differently to each target species according to the extraction method.

Results

Dose-dependent allelopathic effects

ANOVA and successive F test showed strong inhibitory effects of increased concentration ($P < 0.0001$) of both,

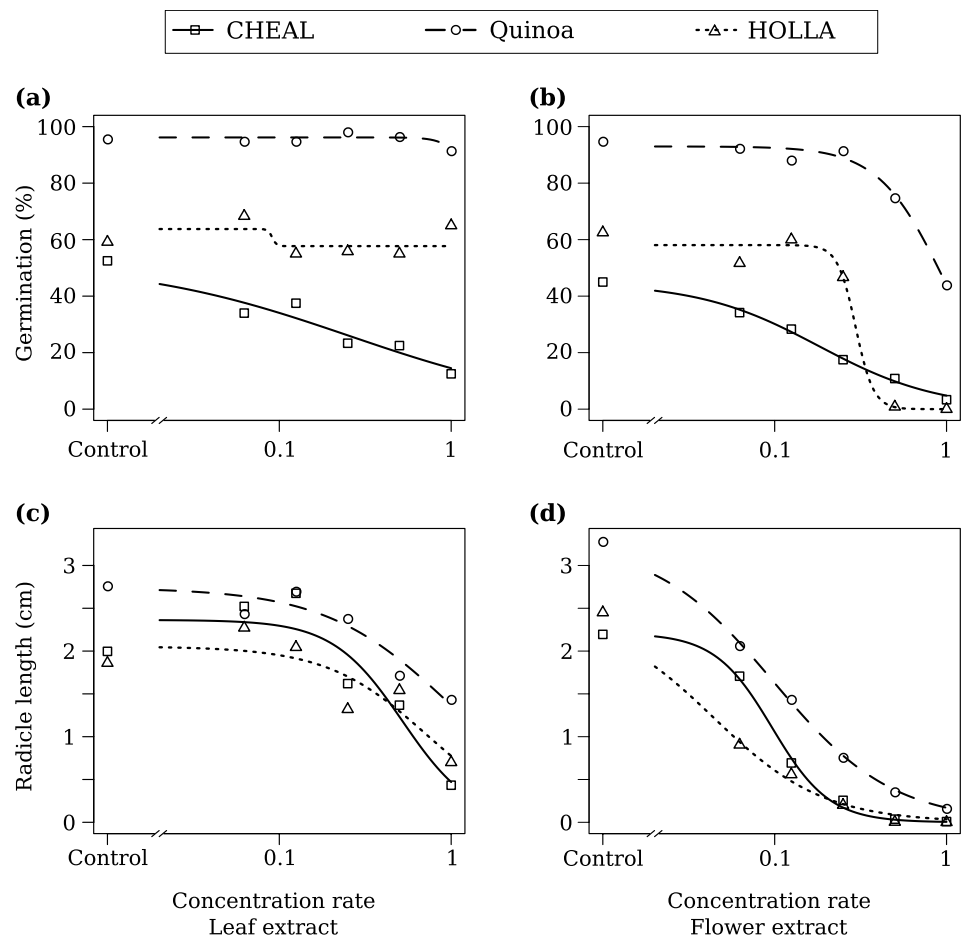
leaf and flower *C. haematocephala* extracts on the target crops quinoa and rice, as well as, on their corresponding target weeds. More intense effects were generally generated by flower compared with leaf extracts. These effects prevailed for bioassays on quinoa during 2015 and 2016, without any significant difference between years ($P > 0.1$), thus results are presented averaged over years. Lack-of-fit tests and residuals of predicted against fitted values (not shown) determined a good fit of data to the nonlinear models (Figs. 1 and 2).

Quinoa, *H. lanatus* and *C. album*

Seed germination was minimally affected by *C. haematocephala* leaf extract, while radicle elongation was reduced somewhat more (Fig. 1a, c). Apparently, the grass weed HOLLA was visibly more affected than the broad-leaved weed CHEAL or the quinoa crop. However, calculated values for 50 and 90% germination (GR_[50,90]) and radicle elongation (RR_[50,90]) reduction parameters and their corresponding confidence intervals for all target species varied extremely (Table 3). Concentration rates of leaf extracts ranged from 0.09 and 6.44 for reflecting the 50 or 90% inhibiting effect on the target species, correspondingly, and the 95% CI vary from negative (-31.57) to extremely high positive (44.45) GR_[50,90] and RR_[50,90] values (Table 3). Additionally, a slight hormetic effect was observed for the leaf extract on radicle elongation of HOLLA and CHEAL (Fig. 1c). Nevertheless, the log logistic model (4) with three parameters was preferred over a Brain-Cousens model with three parameters plus a hormesis parameter (Belz and Piepho 2012; Ritz et al. 2015), because it allowed calculation of GR_[50,90] and RR_[50,90] values and AIC values were equal for both models.

Flower extract caused a much stronger inhibitory effect than that of leaf extract (Fig. 1b, d and calculated GR_[50,90] and RR_[50,90] values in Table 3). A selective inhibitory effect towards weed species could be observed, which was more prominent on germination (Fig. 1b) than on radicle elongation (Fig. 1d). Extract concentration rates of 0.18 and 0.30 caused 50% germination inhibition (GR_[50]) on HOLLA and CHEAL seeds, while a rate of nearly 0.95 was required to achieve GR_[50] on quinoa seeds (Table 3; Figures A3, A5). No significant inhibition of radicle elongation resulted from flower extract; concentration rates between 0.04 and 0.10 were required to achieve RR_[50] on all target species (Table 3). A specific inhibitory effect of flower extract towards the target weeds was also seen for 90% radicle elongation reduction (Figures A4; A6). A concentration rate of 0.22 for both, HOLLA and CHEAL caused RR_[90] (Table 3), while for the same effect on quinoa, a rate of 0.57 was required. However, overlapping 95% CIs suggest that quinoa was affected as much as the target weed species.

Fig. 1 Dose–response of increased *C. haematocephala* extract concentration on target crop quinoa and weeds *C. album* (CHEAL) and *H. lanatus* (HOLLA). **a** Leaf extract and **b** flower extract on germination; **c** leaf extract and **d** flower extract on radicle elongation. Corresponding estimates for 50 and 90% germination reduction ($GR_{[50,90]}$) and radicle elongation reduction ($RR_{[50,90]}$) in Table 3



Rice, *E. crus-galli*, *E. colona*, *E. prostrata* and *R. cochinchinensis*

Flower extract of *C. haematocephala* had a weak germination inhibiting effect on seeds of rice and its target weeds, compared with quinoa and target weeds. Overlapping confidence intervals suggested no differences among species (Fig. 2a; Table 3). However, a tendency of selective effects towards weeds prevailed, as rice seed germination was maintained at around 40%, while only < 20% of the weeds germinated at the 100%-extract concentration. Conversely, radicle elongation was affected by increasing flower extract concentration, being significant for target weeds ECHCG, ECHCO and ECLAL ($RR_{[50]} < 0.21$). Seeds of rice and ROOEX were less affected ($RR_{[50]} > 0.44$; Fig. 2b; Table 3). Flower extract concentration rates > 0.52 (CI: 0.14–1.83) was required to reduce 90% radicle length (Table 3), without significant differences for rice and its target weeds. In contrast,

overlapping 95% CIs suggested that radicle elongation inhibition of quinoa was affected by flower extract, as much as its target weed species.

Allelopathy of obtained chemical fractions on target species

Germination and radicle elongation of the target species were affected differently by the tested fractions, crude extract and tested blanks. The highest germination percentage was obtained with the deionized water blank, averaging nearly 70% for all target species. The general ANOVA showed significant germination inhibiting effects for all factors and their interactions ($p < 0.001$), except for the factor extraction method ($p = 0.08$). The crude extract (F0) and the fraction containing flavonoids (F3) were ranked as the most inhibiting substances against of all target species (average germination 53.1 and 45.8%, respectively). This effect was significantly stronger than

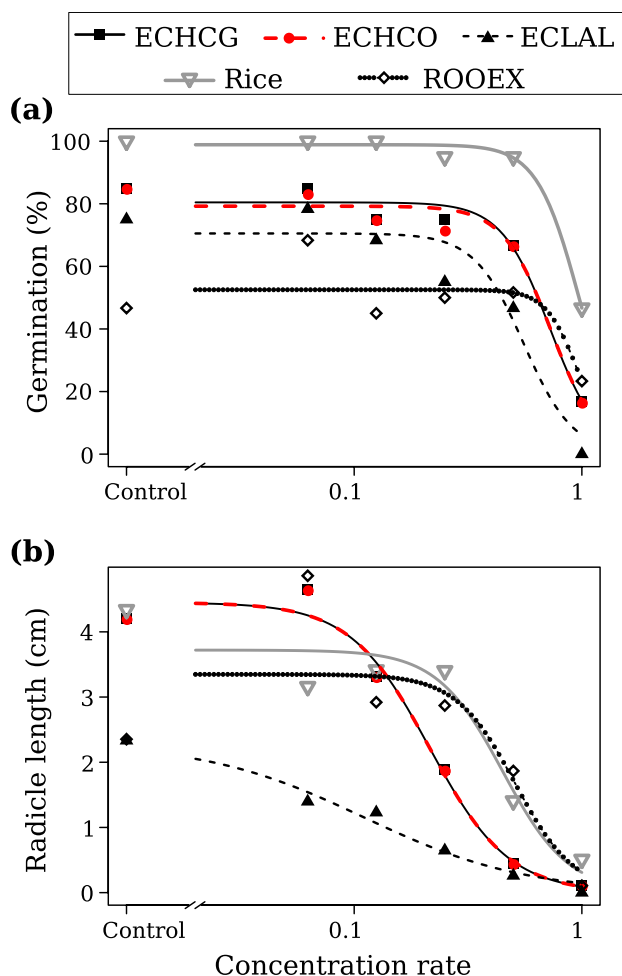


Fig. 2 Dose–response of increased *C. haematocephala* flower extract concentration on target crop rice and weeds *E. crus-galli* (ECHCG), *E. colona* (ECHCO), *E. prostrata* (ECLAL) and *R. cochinchinensis* (ROOEX). **a** Germination inhibiting effect; **b** Root growth reduction effect. Corresponding estimates for 50 and 90% germination reduction ($GR_{50,90}$) and radicle elongation reduction ($RR_{50,90}$) in Table 3

the other fractions, according to Tukey HSD comparison (average germination = 59.8%, CV = 15.9% and HSD = 9.6). In addition, some selective effects were detected for the target species (HSD= 5.0). Germination of quinoa (76.4%) was less affected than its target weeds, and within them, CHEAL was significantly less affected ($\approx 60.1\%$) than HOLLA ($\approx 43\%$).

The interaction between extraction method and fractions on germination was explained by F0 and F3 having a more intense allelopathic effect (Fig. 3; Figures A7, A8). However, the ethanol-based extracts apparently produced a weaker inhibition than the aqueous fractions ground with liquid nitrogen (Fig. 3a). From the aqueous fractions, F0 and F3 significantly reduced germination by 68.3 and 52.8%, respectively (Tukey HSD= 15.5). Within the ethanol-based fractions, germination was more affected by the crude extract

F0 (40%) and fraction F3 (41.2%), although insignificantly. Moreover, 10%-ethanol blank seemed to affect germination more than the fractions themselves, with a 44% reduction.

Fractions interacted differently with the target species, significantly reducing the germination of weeds, without affecting quinoa (Fig. 3b; Tukey HSD = 20.2, CV = 15.9%). The crude extract (F0) tended to reduce germination of quinoa (66.7%) more than F1–F5 ($> 70\%$), whereas F3 enhanced its germination (80.8%). F0 and F3 reduced markedly germination of both target weeds, where HOLLA was more affected than CHEAL (Fig. 3b).

The general ANOVA indicated strong effects of extraction method, fractions, target species and interactions between methods against fractions and methods against target species ($p < 0.001$). Root growth of all target species was reduced by the crude extract F0 and fraction F3 (Fig. 4), from both extraction methods. Ethanol-based extracts showed the strongest inhibitory effect (Fig. 4a). The average radicle length possible under the 10% ethanol blank and ethanol-based F1, F2, F4 and F5 was nearly 0.8 cm, while under the aqueous blank and corresponding fractions was above 1.2 cm (AVG = 0.99 cm, CV = 29.96; LSD = 0.10). Root growth of quinoa was low for the deionized water blank, but radicle length of the target weeds was diminished at a higher level, particularly with fractions F0 and F3 (Fig. 4b). Root length was the lowest for F0 (0.22 cm) and F3 (0.36 cm), compared with the rest of fractions and blanks (1.12–1.40 cm; Tukey HSD = 0.30). Similar to germination, radicle length of target species was selectively reduced by fractions (HSD= 0.62). quinoa and CHEAL were less affected by F3 (0.36 and 0.48 cm, respectively) and F0 (0.28 cm both species); HOLLA appeared to be more sensitive to F3 and F0, with 0.23 and 0.09 cm, respectively.

Qualitative and quantitative analysis of fractions

Using the chemical tests with reagents and TLC, it was possible to detect and quantify allelochemicals in the aqueous and ethanol-based extracts (Table 4; Figure A9). Among the identified phytochemical groups in the aqueous extracts were sesquiterpenes, flavonoids, hydrosoluble tannins, reducing sugars and coumarines (Figure A10). The same phytochemical groups were identified in the ethanol-based extracts, however, with less accuracy and quality. Aqueous extracted F3 produced the most pronounced allelopathic effects. Preparation of F3 allowed determining the flavonoid content in the flower extract through quantification of quercetin, using Eq. (3). Since the fraction density was calculated in 1.03 g mL^{-1} , and that 10.65 g inflorescence were required to produce 50 mL of F3, the total quercetin content was determined as 2.67 mg per gram F3, corresponding to a concentration of 12.82 mg quercetin per gram inflorescence.

Table 3 Estimates for 50 and 90% germination reduction ($GR_{50,90}$) and radicle elongation reduction ($RR_{50,90}$) for the leaf and flower *C. haematocephala* extracts tested on target crops quinoa, rice and target weeds CHEAL, HOLLA, and ECHCG, ECHCO, ECLAL and ROOEX

Plant part	Target species	Estimates (95% CI lower/upper)			
		GR_{50}	GR_{90}	RR_{50}	RR_{90}
Leaf	Quinoa	2.37 [†] (-16.15/20.88)	3.34 [†] (-25.91/32.58)	1.00 (0.36/1.63)	6.38 (-6.62/19.39)
	CHEAL	0.26 [†] (-0.48/0.99)	6.44 [†] (-31.57/44.45)	0.52 (0.32/0.72)	1.46 (0.47/2.44)
	HOLLA	0.09 [†] (-2.19/2.37)	0.10 [†] (-2.11/2.30)	0.72 (0.29/1.14)	3.10 (-1.20/7.40)
Flower	Quinoa	0.95 (0.79/1.12)	2.48 (1.14/3.82)	0.10 (0.07/0.13)	0.57 (0.29/0.85)
	CHEAL	0.18 (0.07/0.29)	1.06 (0.02/2.11)	0.10 (0.07/0.12)	0.22 (0.12/0.33)
	HOLLA	0.30 (0.18/0.42)	0.38 (0.00/0.77)	0.04 (0.02/0.07)	0.22 (0.07/0.38)
Flower	Rice	0.98 (0.85/1.10)	1.56 (0.62/2.51)	0.44 (0.31/0.59)	0.94 (0.14/1.73)
	ECHCG	0.72 (0.57/0.88)	1.24 (0.79/1.70)	0.21 (0.14/0.28)	0.52 (0.25/0.80)
	ECHCO	0.73 (0.58/0.89)	1.25 (0.78/1.71)	0.21 (0.14/0.28)	0.52 (0.24/0.80)
	ECLAL	0.56 (0.37/0.74)	0.96 (-0.05/1.98)	0.11 (-0.02/0.24)	0.66 (-0.50/1.83)
	ROOEX	0.96 (0.73/1.20)	1.38 (-0.56/3.32)	0.51 (0.32/0.69)	0.98 (0.28/1.69)

All parameters were calculated from Eq. (4), except where stated otherwise; 95% confidence intervals in parentheses

[†]From Eq. (5)

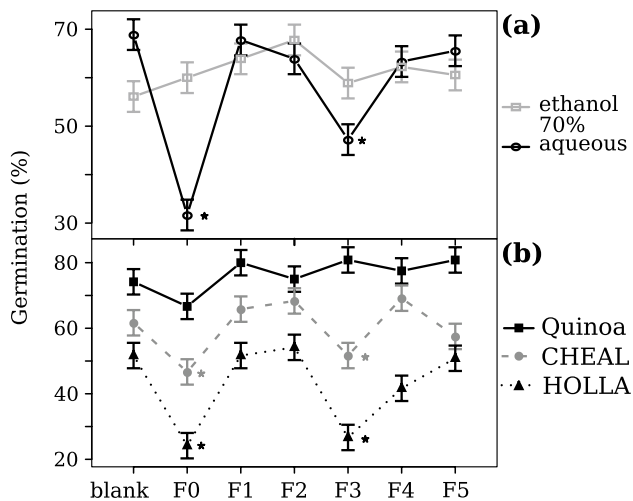


Fig. 3 Effect of the interactions between *C. haematocephala* extract fractions (F0–F5) and the extraction methods, ethanol 70% and aqueous-based, including their corresponding blanks on the germination of target species quinoa, *C. album* (CHEAL) and *H. lanatus* (HOLLA). **a** Influence of extraction method; **b** Effect by target species. The horizontal lines at each observed point is the standard error; significant differences are shown with the symbol *

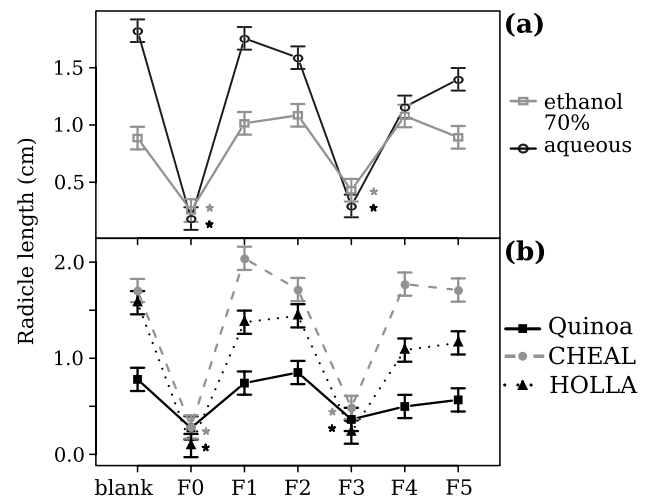


Fig. 4 Effect of the interactions between *C. haematocephala* extract fractions (F0–F5) and the extraction methods, ethanol 70% and aqueous-based, including their corresponding blanks on radicle elongation of target species quinoa, *C. album* (CHEAL) and *H. lanatus* (HOLLA). **a** Influence of extraction method; **b** Effect by target species. The horizontal lines at each observed point is the standard error; significant differences are shown with the symbol *

Discussion

Allelopathic effects on target species

Chemical compounds extracted from many plant species have shown allelopathic properties, which give a great hope in the development of non-synthetic alternatives for weed control. Most studies have found allelochemicals mainly in aqueous extracts (Chon et al. 2003; Tsao et al. 2002; Xuan

et al. 2005; Hill et al. 2007; Reigosa and Pazos-Malvido 2007; Shrestha 2009; Lin et al. 2010; Zhang et al. 2010; Rueda-Ayala et al. 2015; Amari et al. 2014; Puig et al. 2018), describing almost exclusively inhibitory effects towards germination and root elongation. The aqueous extracts of *C. haematocephala* used in this study evidenced a high content of chemical compounds that inhibited the development of target species more than ethanol based extracts. Moreover, a selective effect on weeds without hampering the crop seedlings were observed (Fig. 1). Mixture of extracts from

Table 4 Results of the phytochemical screening of fractions from *C. haematocephala* flower extracts, obtained after two methods ethanol (10%) and aqueous (plus grinding with liquid nitrogen)

Fraction	Phytochemical group	Extraction method		Chemical test
		M1: ethanol 70%	M2: aqueous	
F1	Alkaloids	–	–	
F2	Sterols	–	–	
	Sesquiterpenes	+	+	TLC [†]
	Terpenes	–	+	Zack's
F3 and (F4) [‡]	Flavonoids	+ (–)	+ (–)	Shinoda's
	Anthraquinones	– (–)	– (–)	
	Tannins	+ (–)	+ (–)	Ferric chloride
	Saponins	– (–)	– (–)	
	Reducing sugars	+ (–)	+ (–)	Fehling's reagent
F5	Lactones	+	+	Baljet's
	Coumarines	–	+	Ferric hydroxamate
	Cardiac glycosides	–	–	

(Sherma 2000; Seidel 2005; Otsuka 2005; Ivanova-Petropulos et al. 2010; Evans and Evans 2019; Ribeiro et al. 2018). See also Figure A9

[†]Sesquiterpenes were identified after TLC in both extract methods

[‡]All results for F4 were negative

different plant structures or plant species have shown higher phytotoxicity than individual extract compounds (Chon et al. 2003). Results in the present study showed much stronger germination and development inhibiting effects by flower than leaf *C. haematocephala* extracts (Fig. 1).

Some alternative solvents, such as methanol and ethyl acetate produce plant extracts more concentrated with allelochemicals than aqueous extracts (Hill et al. 2007), for instance compounds from methanolic extracts of *Everniastrum sorocheilum*, *Usnea roccellina*, and *Cladonia confusa* have shown growth inhibitory effects on *Trifolium pratense* (Nieves et al. 2011). Conversely, ethanol-based extracts in this study showed a much lower allelopathic effect on germination (Fig. 3) and radicle elongation (Fig. 4) than aqueous extracts. Aqueous extracts have shown a joint action of several contained compounds at once (Puig et al. 2018).

Apparently, allelochemicals in *C. haematocephala* are highly concentrated in the above-ground plant structures, although in this study, root extracts were not investigated. Leaf extracts reduced seed germination and radicle elongation of quinoa and its target weeds (Fig. 1a) less efficiently than flower extracts. Flower extracts showed strong inhibiting effects on germination, and even stronger inhibiting effects on radicle elongation (Fig. 1b, d). Hill et al. (2007) found a selective inhibiting effect of allelochemicals on weed species, due to their smaller seed size than that of crops. In this study, quinoa seedlings (*C. quinoa*) were little affected, while germination and radicle elongation of *C. album* (CHEAL) was strongly inhibited, even though there was no great difference in seeds size between these two species. Thus, other factors than seed size affected

their sensitivity. This selectivity of *C. haematocephala* extracts could offer an advantage in the search of new formulas of bioherbicides for pre-emergence and early emergence use, provided that the involved compounds could be accurately identified. A similar effect occurred with the flower extracts applied to rice and its target weeds, where rice germination was much less affected than that of the target weeds (Fig. 2a). However, the selectivity of flower allelochemicals was somewhat lower for radicle elongation reduction.

Fractions and effects on target species

Although the same phytochemical groups, except coumarines, were identified in both, aqueous and ethanol-based extracts, the latter yielded lower quantity and quality of allelochemicals. There is evidence that thermolabile and volatile organic compounds, commonly found in plant extracts, are affected by the ethanol maceration (Bart 2011). This effect occurs because the sample remains exposed for longer time to the extracting agent ethanol compared with water. Conversely, grinding the plant sample with liquid nitrogen produces a immediate rupture of the cell wall and cell structures, liberating immediately the metabolites and essential oils, thus reducing losses. Furthermore, the aqueous fractionation to get F3 was extracted from $\frac{1}{3}$ of the crude flower extract (F0), using first petroleum ether and adding a 50:50 methanol/water mixture. These processes allowed separating the residual liquid and to ease the determination of flavonoids, hydro-soluble tannins and reducing sugars in a more efficient

way. The total flavonoid concentration (regarded as mg quercetin per g of inflorescences) was 12.8 mg, is comparable with other plant extracts with allelopathic properties. Inhibiting effects of leaves and flowers in *Zygophyllum album* and *Thymelaea hirsuta* have been quantified as 4.1 mg quercetin per g of the tested plant structure (Amari et al. 2014; Mnafigui et al. 2012, 2016).

Practical implications of *C. haematocephala* flower extracts

For flower *C. haematocephala* extracts to become a realistic alternative for weed control, the equivalent extract concentration that can be obtained from the harvested fresh inflorescence biomass under field conditions should be determined, more specifically, the field equivalent rate of the F3 fraction. In this study, the fraction F3 was obtained from aqueous and ethanol flower extracts (Table 1, Exps. 6 and 7, respectively). To produce 50 mL of F3, 13.33 mL of aqueous crude extract were required with a concentration of 2.78 g mL⁻¹, while 18.33 mL were required with an ethanol crude extract concentration of 0.56 g mL⁻¹. Similarly, 9.26 mL of F0 were calculated as needed for obtaining F3 in Exps. 3, 4 and 5 (Table 5), at 3.33 g mL⁻¹ F0 concentration.

Using a similar methodology as proposed by Hill et al. (2007), the field equivalent rates of F3 were calculated (Table 5) with $FER = (EP)/(ADW)$, where *FER* is the field equivalent rate; *E* is the amount of F3 fraction (mL) obtainable with the total weight of harvested inflorescence per experiment; *P* is the petri dish volume used (64 cm³) on the bioassays; *A* is the area harvested of *C. haematocephala* inflorescence; *D* is the hypothetical depth at which inflorescence material might be incorporated in the soil (15 cm), assuming a simultaneous decomposition; and *W* corresponds to the F3 volume applied to each petri dish. Field equivalent rates of aqueous F3 fractions ranged between 90 and 143 mL l⁻¹, while for ethanol, F3

were 131 mL l⁻¹. With these estimations, and knowing that 10.65 g inflorescence yielded 50 mL F3 (aqueous), the field rates in this study were within the range of concentrations and amount of harvested fresh inflorescence biomass (i.e., 19 to 31 g l⁻¹). However, the extract concentrations achievable from decomposition in the soil will be lower than under laboratory conditions (Hill et al. 2007), results in the present study show a great potential of *C. haematocephala* inflorescence extract for germination and development of weed species, with a high likelihood to be species specific (Figs. 1b, d and 2a; Table 3).

In conclusion, the market supply of bioherbicides is rapidly increasing with to social trends. However, formulation of bioherbicides is still a huge challenge. Less than 10% of available biopesticides are bioherbicides, while weeds are still the major cause of yield loss in current agriculture (Charudattan 2001). Promising inhibitory effects on seed emergence have been observed from essential oil extracts of eucalyptus (Zhang et al. 2010; Puig et al. 2018) or Black walnut (Shrestha 2009), the latter being already commercially formulated as a bioherbicide that limits completely the growth of *Conyza canadensis* at a concentration of 33%. The flower extracts identified and tested in this study already fulfill some requirements to be acceptable and efficient biopesticides, such as weed specificity, minimal effect on crops, easy and cheap elaboration. However, other important aspects such as environmental effects, including the production chain, or non-toxicity towards humans and non-target living organisms still require further investigation and validation. Nevertheless, the results of this study are encouraging for finding sustainable and ecologically friendly alternatives for weed management in crops of high nutritional value, as quinoa and rice, contributing also to counteract the growing problem of herbicide resistance.

Table 5 Field equivalent rates (FER) estimated on the approximate F3 fraction obtainable from F0 *C. haematocephala* flower extracts tested in the bioassays

Exp.	Area harvested (m ²)	<i>Calliandra</i> height* (m)	Fractions retrieved		Field equivalent rate (mL l ⁻¹)
			F0 to 50 mL F3 (mL)	F3 approx. † (mL)	
3	1.20	2.10	9.26	761.34	90.23
4	0.80	2.10	9.26	734.34	130.55
5	1.80	2.80	9.26	1285.10	101.54
6	0.25	1.70	13.33	251.31	142.97
7	0.25	1.70	18.33	231.86	131.90

This amount was calculated on a basis of F0 required to produce 50 mL of F3. FER were calculated using a similar methodology as Hill et al. (2007). Values in experiments 6 and 7 are true estimates

* Taller shrubs often had more flowers, thus yielding more flower fresh weight per area harvested

† Approximate F3 amount calculated from the total crude extract (F0) retrieved at the corresponding experiments in Table 1

Author contribution statement Conceptualization, design of experiments, methodology for detection of allelopathic properties, statistical analyses, writing—original draft preparation and editing: VPR-A. Methodology for fractionation, qualitative and quantitative analyses of fractions: LR-G, PV-J, DB and LG. Performed bioassays: BH and IT. Draft co-writing and review: LR-G, PV-J, BH, IT, MH and DA. Supervision: VPR-A, LR-G, PV-J and BH.

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Compliance with ethical standards

Conflict of interest The authors declare that they have no conflict of interest.

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