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Abstract

In this work a novel nano-formulation is proposed to control leaching and volatilization of a broadly used herbicide, dicamba. Dicamba is subject to significant leaching in soils, due to its marked solubility, and to significant volatilization and vapor drift, with consequent risks for operators and neighbour crops. Natural, biocompatible, low-cost materials were employed to control its dispersion in the environment: among four tested candidate carriers, a nanosized natural clay (namely, K10 montmorillonite) was selected to adsorb the pesticide, and carboxymethyl cellulose (CMC), a food-grade biodegradable polymer, was employed as a coating agent. The synthesis approach is based on direct adsorption at ambient temperature and pressure, with a subsequent particle coating to increase suspensions stability and control pesticide release. The nano-formulation showed a controlled release when diluted to field-relevant concentrations: in tap water, the uncoated K10 released approximately 45% of the total loaded dicamba, and the percentage reduced to less than 30% with coating. CMC also contributed to significantly reduce dicamba losses due to volatilization from treated soils (e.g., in medium sand, 9.3% of dicamba was lost in 24 h from the commercial product, 15.1% from the uncoated nanoformulation, and only 4.5% from the coated one). Moreover, the coated nanoformulation showed a dramatic decrease in mobility in porous media (when injected in a 11.6 cm sand-packed column, 99.3% of the commercial formulation was eluted, compared to 88.4% of the uncoated nanoformulation and only 24.5% of the coated one). Greenhouse tests indicated that the clay-based nanoformulation does not hinder the dicamba efficacy towards target weeds, even though differences were observed depending on the treated species. Despite the small (lab and greenhouse) scale of the tests, these preliminary results suggest a good efficacy of the proposed nanoformulation in controlling the environmental spreading of dicamba, without hindering efficacy toward target species.

Keywords

Nanopesticides; dicamba; controlled pesticide leaching; nanocarrier; natural clays

Highlights

natural clay and food-grade biopolymer are used in a novel nanopesticide formulation

natural clays showed effective in reduce environmental drawbacks of agrochemicals

the nanoformulation effectively encapsulated dicamba and controlled its release

the nanoformulation significantly reduced dicamba mobility in porous media

the herbicidal efficacy of dicamba was not hindered by the nanoformulation

1. INTRODUCTION

The use of agrochemicals, including pesticides and fertilizers, is unavoidable for the optimization of crop production, but has numerous drawbacks on both human health and the environment (Damalas and Eleftherohorinos, 2011). Several products have toxic effects on aquatic life and beneficial insects, and a growing number of plant protection products have been found or suspected to be toxic or carcinogenic to humans (DeLorenzo et al., 2001; Mostafalou and Abdollahi, 2017). Herbicides are particularly relevant in this sense, being the agrochemicals most frequently found in superficial waters and groundwater.

The persistency, hydrophilicity, volatility and more in general the affinity of the individual substance to the different environmental matrices and components ultimately control their preferred migration routes and accumulation compartments, and consequently their long-term fate and associated potential risks for human health and ecological receptors. Poorly biodegradable substances tend to progressively accumulate in deep soil, sediments and groundwater, where they remain unaltered for years or even decades (Arias-Estévez et al., 2008; Wauchope, 1978). Highly soluble compounds show a prominent infiltration potential with rain and irrigation (and consequently transport and accumulation in deep soil and groundwater is the preferred migration route), while volatile compounds are prone to significant dispersion in the atmosphere. It has been reported that 10% to 75% of the applied pesticides do not reach the target pest species (Aktar et al., 2009; Pimentel, 2009) and the unused product can therefore migrate towards non-target crops, insects and ultimately spread in the environment (Boutin et al., 2014; Sanchez-Bayo and Goka, 2014; van den Berg et al., 1999).

The major routes available to reduce the environmental impact of pesticides include the development of more efficient crop management strategies (Bongiovanni and Lowenberg-Deboer, 2004; Reichenberger et al., 2007), the identification of new, less toxic and less persistent substances with pesticidal effects (Dayan et al., 2009; Lamberth et al., 2013), and the development of new formulations of existing active ingredients (AIs) (Iavicoli et al., 2017; Sopeña et al., 2009). Nanotechnology can play an important role in finding new pesticide formulations with reduced toxicity and environmental impacts (Moulick et al., 2020; Servin et al.,

2015; Worrall et al., 2018), even though research is still needed to verify several aspects of the actual efficacy and convenience of nano-formulated pesticides, particularly at the field scale (Gomes et al., 2019; Kah et al., 2018a; Singh et al., 2021; Usman et al., 2020). Nanopesticides consist of nanoparticles (NPs), usually referred to as nanocarriers, containing an active ingredient dispersed in a colloidal suspension. Nanopesticides are formulated with different objectives (Adisa et al., 2019; Kah et al., 2019; Kah et al., 2018b): to promote the use of AIs that are less harmful toward non-target organisms, but whose premature degradation may need to be hindered; to optimize pest targeting; to reduce the overall amount of employed chemical substances by improving the delivery of poorly soluble AIs, or conversely by controlling and retarding the release of soluble AIs. Two main types of nano-formulations are currently used: (1) NPs that act as pesticides themselves and (2) NPs that act as a carrier for AIs with pesticidal effects, adsorbed or immobilized on them using different techniques (Kah and Hofmann, 2014; Kah et al., 2018a).

In this work a nano-to-micro-formulation, aimed at reducing environmental spreading of a highly soluble and moderately volatile herbicide, namely dicamba (3,6 dichloro-2 methoxy benzoic acid), was developed. Dicamba is an auxin-like herbicide, registered in 1967, applied in post-emergence to control annual and perennial broadleaf weeds in non-agricultural settings, lawn and turf and in different crops, such as wheat, barley, corn, oats, millet, sorghum, and asparagus (Behrens et al., 2007). Dicamba has been established to be toxic to aquatic organisms and its metabolite was defined as harmful (Authority, 2011). Because of its high efficacy, relative inexpensiveness, relative environmental safety and low risk for weeds to develop resistance, it is used worldwide. Dicamba, as well as other auxinic herbicides, is characterized by a high solubility, in the order of 6 to 8 g/l at ambient conditions, resulting in significant infiltration and consequent mobility in soil and subsoil (Oliveira Jr et al., 2001; Sakaliene et al., 2007) and relatively high volatility, with consequent drift problems and potential risks for users (Ding et al., 2019; Egan and Mortensen, 2012).

Natural materials were tested in this study for the development of the nano-formulation, namely mineral carriers (montmorillonite clays and a zeolite), and carboxymethyl cellulose (CMC) as a coating agent to minimize the loss of herbicide and modulate its release over time. It is known that mineral particulate matters such as silica and clay have a good affinity with a broad set of polar or ionizable pesticides, and have been

successfully proposed as carriers in the development of nano-formulations. In most cases, natural particles are chemically modified to improve their affinity to the pesticidal molecules, such as in the case of organoclays (Cabrera et al., 2016; Cornejo et al., 2008; Hermosin et al., 2001). Synthetic hollow silica particles have also been successfully employed (Bueno and Ghoshal, 2020; Gao et al., 2020; Nuruzzaman et al., 2020). Less frequently, unmodified mineral particles have been directly used, e.g. montmorillonite for hexazinone (Celis et al., 2002), simazine and 2,4-D (Cox et al., 2000). The latter approach, even if often slightly less performing in term of loading capacity, was preferred in this work due to its higher sustainability and environmental compatibility, easier implementation and reduced cost.

Polymers, and in particular biopolymers, are currently used in the form of capsules (Petosa et al., 2017), polymeric microbeads and as a coating of solid carriers to control the release of encapsulated pesticides and fertilizers (Joshi et al., 2020; Shasha et al., 1976), protect labile compounds (Elhaj Baddar et al., 2020), and prevent excessive volatilization (Azeem et al., 2014; Dimkpa et al., 2020). Alginate is by far the most applied biopolymer (Kenawy and Sakran, 1996), even though carboxymethyl cellulose (Kök et al., 1999), lignin (Behin and Sadeghi, 2016), starches (da Costa et al., 2019), bio-based plastics (Riggi et al., 2011) and others have been employed.

In this work the efficacy and efficiency of four different natural mineral carriers, as well as the opportunity of using a bio-polymeric coating to control the dicamba release were evaluated through different tests. The best performing formulation was identified based on a balance between technical constraints and potential environmental effects. The ideal nano-formulation should be the one showing the highest loading capacity (evaluated in terms of mass of dicamba adsorbed per mass of carrier), the highest colloidal stability in a broad range of hydrochemical conditions (to guarantee that the formulation can be diluted in water without an abrupt aggregation and/or sedimentation of the carriers, which would hinder its use), the lowest release of dicamba after dilution, the lowest loss of dicamba due to volatilization (which represents one of the main environmental problems of this AI), the lowest mobility in the porous medium (thus limiting as much as possible the potential spreading of dicamba, if infiltrated with rain or intense irrigation), and obviously the

most efficient in controlling weeds. Clearly not all these aspects – sometimes contrasting – can be maximized, and the optimal nano-formulation was identified as the most reasonable compromise.

2. MATERIALS AND METHODS

2.1. Materials

Four carriers, all provided in the form of dry powders, were tested: a K10 Na-montmorillonite (Sigma Aldrich, labelled K10 in the following), a generic Na-montmorillonite (Indigo Herbs, UK, labelled Na-M2), a Ca-montmorillonite (PraNaturals, UK, labelled Ca-M), and a zeolite (NaturaForte, Germany, labelled ZEO). A food-grade, low-molecular-weight carboxymethyl cellulose (WALOCCEL® CRT30GA, Dow Chemical Company Ltd, US, labelled CMC) was tested as carrier coating.

Structure, shape and elemental composition of the carriers were investigated using SEM-EDS microscopy (JEOL, Japan) directly on the dry powders. The carrier particle size was determined using a disk centrifuge (DC24000 UHR, CPS, US). To this aim, the samples were dispersed in deionized water (DIw), allowed to hydrate, sonicated for 25 minutes prior measurement and analysed at a disc rate of 3000 rpm in the range 70-0.1 microns. The zeta potential of the four carriers dispersed in DIw (following the same preparation protocol) with addition of NaCl (1 mM to 100 mM) or CaCl₂ (0 to 1 mM) were determined with electrophoretic measurements using dynamic light scattering DLS (Zetasizer Nano ZSP, Malvern Instrument, UK).

Dicamba was provided by Alfa Chemistry (US). A commercial herbicide (Mondak 21 S, Syngenta, Italy) containing dicamba (not nano-formulated) at a nominal concentration of 243.8 g/l was used as a comparison in volatilization, transport and efficacy tests.

The tap water (TAPw) used for release and transport tests was drawn from the municipal water supply network and chemically analysed for salts, pH, EC and TDS (data reported in Supporting Information, Table S1).

A medium silica sand (Dorsilit 8, Dorfner, Germany; d_{10} , d_{50} and d_{90} equal respectively to 0.415, 0.45 and 0.5 mm), was used for volatilization and column transport tests. A coarse sand (Dorsilit 5G, Dorfner, Germany; d_{10} , d_{50} and d_{90} equal respectively to 1.12, 1.58 and 1.9 mm) and a sandy loam soil (collected at DISAFA - University of Torino, d_{10} , d_{50} and d_{90} equal respectively to 0.04, 0.065 and 0.148 mm) were used for volatilization tests. For all sands, the granulometric curve (data reported in Supporting Information, Figure S1) was determined via dry sieving, while real soil undergo to wet sieving.

2.2.Methods

2.2.1. Nanopesticide preparation

The carriers were loaded with dicamba using direct adsorption. Briefly, the carrier was dispersed in a dicamba solution (in the range 0.125 g/L to 7 g/L). The suspensions were continuously mixed with a magnetic stirrer in a closed vessel with reduced headspace to minimize volatilization. Contact times of 24 h and 2 h were used. After loading, the suspensions were centrifuged for 20 min at 7000 rpm (5259 rcf) and the precipitate was collected and stored as stock nano-formulation.

Adsorption isotherms were determined for all tested carriers and loading conditions. After centrifugation, the supernatant was filtered with a 0.45 micron syringe PTFE filter (Thermo Fisher Scientific, Waltham, USA) and the dicamba concentration in the supernatant ($C_{w,eq}$ [$M L^{-3}$]) was evaluated via Uv-vis spectrophotometry (Specord S600, Analytik Jena, Germany) at a wavelength of 280.5 nm (the calibration curve is provided in Supporting Information, Figure S2). The lowest dicamba detection limit was 50 mg/l. The adsorbed concentration, or adsorption capacity, expressed as dicamba mass adsorbed per unit mass of carrier, $C_{s,eq}$ [$M M^{-1}$], was determined from $C_{w,eq}$ measurements via mass balance.

In case of polymer-coated formulations, the CMC was added to the batch to reach a final polymer concentration of 0.5 g/l. Polymer adsorption onto the loaded carrier was promoted via stirring for 2h.

2.2.2. Release tests

Release tests were conducted on all uncoated nano-formulations and on K10 coated formulations. All tested stock samples were obtained via loading of the carrier in a 7 g/L dicamba solution and with a contact time of 24 h, following the procedure described above.

The stock nano-formulation without any pre-treatment was dispersed in DIw or TAPw, depending on the tested condition, at a concentration of 5 g/L (1g of nano-formulation in 200 ml). The suspension was maintained in agitation with a magnetic stirrer at 200 rpm in a beaker sealed with parafilm, with reduced headspace to minimize volatilization, for at least 6 hours. Release tests in DIw and TAPw from loaded K10 montmorillonite, both uncoated and coated with CMC, were prolonged to 30 h. A sample was periodically collected, filtered with a PTFE syringe filter to remove the carrier and analysed via spectrophotometry to determine the concentration of dicamba released from the nano-formulation.

The fraction of herbicide retained on the carrier at a given time, $R(t)$, was evaluated as:

$$R(t) = 1 - \frac{C_w(t) \cdot V_w(t)}{C_{s,eq} \cdot M_{s,c}} \quad (\text{eq. 1})$$

where $C_w(t)$ is the dicamba concentration measured in the water sample collected at time t [M L^{-3}], $V_w(t)$ is the dispersion volume at time t [L^3], $C_{s,eq}$ is the adsorbed concentration at the beginning of the test, and $M_{s,c}$ is the dry mass of carrier present in the system at the beginning of the test [M].

2.2.3. Volatilization tests

Volatilization tests were performed for uncoated and CMC-coated K10 nano-formulations (labelled respectively K10 and K10-CMC) and compared with a pure dicamba solution and with the commercial product. All samples were diluted in DIw in order to reach the same concentration of the AI (3 g/l).

In pre-screening tests, direct volatilization from the solution/suspension was assessed. Each sample was exposed to ambient air on a Petri dish (Mueller, 2015; Strachan et al., 2010), which was periodically weighted for 30 hours. Tests were run at least in duplicate, at room temperature of $+22.9 \pm 0.3$ °C and relative humidity of 55.6 ± 4.4 %. After 30 hours the remaining mass of dicamba was determined via solvent extraction method: acetonitrile and water at a mass ratio of 7:3 were added and kept in agitation for 24h in a closed vial with minimal headspace; the solution/suspension was then filtered using a PTFE syringe filter and analysed with UV-vis spectrophotometry. Two blank sets of tests containing only K10 and one with only water were run to check possible releases from sand and carriers that may interfere with the UV-vis analysis (no interference was observed).

Volatilization from dicamba-sprayed soil was then assessed in a second set of tests. The coarse and medium sands and the sandy loam soil were air-dried; the soil was sieved to remove particles and conglomerates larger than 2mm. 15 g of soils were added to each Petri dish, sprayed and treated similarly to the pre-screening tests. Blank tests that included soil only (no spray), soil sprayed with water only and soil sprayed with K10 suspension (no dicamba) were run in parallel.

2.2.4. Column transport tests

Column transport tests in saturated conditions, aimed at mimicking carrier and pesticide transport in groundwater, were performed for K10 without pesticide loading (with or without polymeric coating), pesticide-loaded K10 (with or without coating), pure dicamba solution and the commercial product. A Plexiglas column with adjustable ends and internal diameter of 1.6 cm was wet-packed with 36.5 g of Dorsilit 8 sand to an average length of 11.61 (± 0.15) cm. The sand was hydrated and degassed prior packing to remove residual air microbubbles. A polypropylene filter with 120 μm mesh was placed at the top and bottom of the column to avoid sand from entering inlet and outlet tubing. The solutions/suspensions were injected in saturated conditions with a peristaltic pump (ISMATEC REGLO Analog MS-4/8, Cole-Parmer, Germany) at a constant flow rate of $1.46 \cdot 10^{-8}$ m³/s, corresponding to a Darcy velocity of $7.26 \cdot 10^{-5}$ m/s. Inflow and outflow

concentration of solutes and suspensions was monitored via optical density measurements using the UV-vis spectrophotometer equipped with front-through quartz cells with 5 mm lightpath (Hellma, Germany). Monitoring wavelengths of 198.5 nm, 280.5 nm and 350 nm were adopted for NaCl, dicamba and carriers, respectively (calibration curves in Supporting Information, Figures S2 and S3). For tests involving the nano-formulations, outflow samples were also manually collected, filtered and analysed to reconstruct the breakthrough curve of free dicamba.

The nano-formulations were injected at a carrier concentration of 0.9 g/l, corresponding to a dicamba concentration of approximately 50 mg/l (the actual concentration slightly varied based on the specific tested formulation). This concentration corresponds to approximately 10% of the typical recommended concentration for dicamba-based products when applied on weeds: after field applications the most commonly detected dicamba concentration in leaching water is 3% to 10% of the applied one (Sakaliene et al., 2007; Tindall and Vencill, 1995), and consequently the most severe expected scenario for groundwater contamination was adopted.

Preliminary transport tests of K10 alone (without dicamba loading), with and without CMC coating, followed this protocol:

1. Pre-equilibration with DIw for at least 5 pore volumes (PVs)
2. Tracer injection (NaCl 30 mM) for 5 PVs
3. Flushing with DIw for 5 PVs
4. Injection of K10 dispersed in DIw for 5 PVs
5. Post-flushing with DIw for at least 5 PVs

The injection protocol for pesticide transport tests (including dicamba solution, nano-formulations and commercial herbicide) included the following steps:

1. Pre-equilibration with DIw for at least 5 PVs
2. Pre-flushing with background electrolyte solution (NaCl 30 mM) or TAPw (depending on the specific test) for 5 PVs

3. Injection of the dicamba solution/nano-formulation (dispersed in NaCl 30 mM solution or in TAPw) for 5 PVs
4. Post-flushing with NaCl 30 mM solution or TAPw for 5 PVs
5. Second post-flushing with DIw for at least 5 PVs

The porosity and the dispersivity coefficient of the sand-packed columns were determined for each transport tests by least-squares fitting the NaCl breakthrough curve (i.e. steps 1 and 2 of the injection protocols) to the classic advection-dispersion partial differential equation for conservative solutes:

$$\varepsilon \frac{\partial C_t}{\partial t} = -q \frac{\partial C_t}{\partial x} + \alpha_x \frac{\partial^2 C_t}{\partial x^2} \quad (\text{eq. 2})$$

where C_t is the tracer concentration [$M L^{-3}$], q is Darcy velocity [$L T^{-1}$], ε is the effective porosity [-] and α_x is the dispersivity coefficient [L] of the porous medium.

The transport of the carriers was modelled using the modified advection-dispersion-deposition equation usually adopted to describe colloid transport in saturated porous media:

$$\begin{cases} \varepsilon \frac{\partial C}{\partial t} - \sum_i \rho_b \frac{\partial S_i}{\partial t} = -q \frac{\partial C}{\partial x} + \alpha_x \frac{\partial^2 C}{\partial x^2} \\ \rho_b \frac{\partial S_i}{\partial t} = \varepsilon k_{a,i} (1 + A_i S_i^{\beta_i}) C - \rho_b k_{d,i} S_i \end{cases} \quad (\text{eq.3})$$

where C is the carrier concentration in water [$M L^{-3}$], S_i is the concentration of carrier particles retained on the solid grains due to the i -th retention mechanism [$M M^{-1}$], ρ_b is the bulk density of the porous medium [$M L^{-3}$], $k_{a,i}$ is the carrier attachment rate due to the i -th retention mechanism [T^{-1}], $k_{d,i}$ is the corresponding detachment rate [T^{-1}], A_i and β_i are empirical coefficients specific to the deposition mechanism [-]. The first equation refers to particle transport in water and the second one describes the deposition mechanism(s). The parenthesis in the second equation is the generic formulation for particle retention mechanisms proposed by Tosco and Sethi (Tosco and Sethi, 2010). For linear attachment, $A_i = \beta_i = 1$; for blocking, $\beta_i = 1$ and $A_i = -1/S_{\max,i} < 0$ (where $S_{\max,i}$ is the maximum concentration of particles retainable on the solid matrix due to the i -th retention mechanism); for ripening, $A_i > 0$ and $\beta_i > 0$. In case of irreversible deposition, the second term vanishes being $k_{d,i} = 0$.

In this work the following two-site deposition model was adopted:

$$\begin{cases} \varepsilon \frac{\partial C}{\partial t} - \rho_b \frac{\partial S_1}{\partial t} - \rho_b \frac{\partial S_2}{\partial t} = -q \frac{\partial C}{\partial x} + \alpha_x \frac{\partial^2 C}{\partial x^2} \\ \rho_b \frac{\partial S_1}{\partial t} = \varepsilon k_{a,1} C \\ \rho_b \frac{\partial S_2}{\partial t} = \varepsilon k_{a,2} (1 + A_2 S_2^{\beta_2}) C \end{cases} \quad (\text{eq. 4})$$

The breakthrough curves of tracer and particles were inverse-fitted to the respective transport equations using the software MNMs 2018 (Micro-and Nanoparticle transport, filtration and clogging Model - Suite) (Bianco et al., 2016; Mondino et al., 2020; Velimirovic et al., 2020). A porosity of 0.31 (± 0.02) and dispersivity of $1.84 \cdot 10^{-4} (\pm 3 \cdot 10^{-7})$ m were obtained from tracer tests, and were assumed valid also for particle transport, since no evidence of particle early breakthrough was observed.

Dicamba adsorption/desorption onto/from the carriers was not modeled since all nano-formulations were prepared for injection a few hours in advance, thus allowing the complete release of the pesticide before injection into the column.

2.2.5. Weed control efficacy tests

Weed control efficacy of the commercial dicamba, K10 and K10-CMC was evaluated in greenhouse on *Solanum nigrum* and *Amaranthus retroflexus*, two dicamba sensitive weeds, grown in pot filled with commercial potting mix. Each pot contained 5 seeds of a single weed species and 4 replicate pots were prepared for each combination of herbicide formulation, dose and species. The experiment was repeated twice, in May and August 2019. When weeds reached a two to three leaf stage, they were sprayed with the following equivalent dose of dicamba: 0 (untreated control), 146.3, 195.0, 219.4, 243.8, 268.2 and 292.6 g AI ha⁻¹, corresponding to the following volume of commercial dicamba: 0, 0.6, 0.8, 0.9, 1 (field rate), 1.1. and 1.2 L ha⁻¹. Seedrape oil (Codacide, Corteva) used at 1 L ha⁻¹ as an adjuvant was added to all the spray solutions. Herbicide treatment was performed using a cabinet sprayer equipped with a single flat fan nozzle (Teejeet AI11002-VS), calibrated to deliver 300 L ha⁻¹ at a pressure of 203 kPa. After treatment, pots were randomly

arranged in greenhouse benches until the study ended. Average greenhouse mean, maximum and minimum air temperature was 24.03, 37.0, and 15.1°C, respectively, in the May experiment and 25.0, 18.2, and 38.9°C, respectively, in the August experiment. Average air humidity was 68.5 and 74.5% in May and August experiment, respectively. Natural light was supplemented with metal-halide lamps to obtain a photosynthetic photon flux density of about 300 $\mu\text{mol s}^{-1} \text{m}^{-2}$ with 16 h photoperiod. At 21 days after treatment the fresh biomass of treated plants was weighted by cutting them just above the soil level. Treatment efficacy was expressed as percentage of fresh aboveground weight relative to the untreated control (relative weight %). Values of relative weight may range from 0% (complete plant desiccation) to 100% (fresh weight of untreated plants).

Data of percentage of relative weight were first analysed to check whether they could be described using a single model (the three-parameter log-logistic regression model in Eq. 5) after pooling the data of May and August experiments. The data were first fit as a pooled data set and then as two separate data sets. ANOVA was conducted to verify the presence of significant differences between the two analyses ($P \leq 0.05$). Given that no differences were found, the relative weight data of the two experiments were analysed as a single data set.

A dose-response curve was built for each herbicide formulation, separately per weed species. The percentage of relative weight of treated plants was fit against herbicide rates according to a three-parameter log-logistic regression model (Equation 5):

$$Y = \frac{d}{1 + \exp\{b[\log(x) - \log(e)]\}} \quad (\text{eq. 5})$$

where Y is the relative weight of treated plants, x is the herbicide rate expressed in g AI ha^{-1} , d is the upper limit, and b is the relative slope at the point of inflection e . Model fitting was performed using the *drm* function of the DRC add-on package of the open-source program R (Fogliatto et al., 2021; Team, 2019). The effective herbicide dose required to reduce plant relative weight by 50% (ED_{50}) and 90% (ED_{90}) compared with the values observed at 0 g AI ha^{-1} were calculated from the fitted model using the *ED* function of the

DRC package. Within each species and for a same ED level (ED_{50} and ED_{90}), ED values were compared using the function *EDcomp* of the DRC add-on package.

3. RESULTS AND DISCUSSION

3.1. Carrier characterization

Information on particle size, zeta potential and fraction of retained herbicide after dilution was used to identify the best performing carrier and coating, to be further assessed in the following steps of the study.

The particle size distribution was measured for the carriers dispersed in deionized water using a disk centrifuge (Supporting Information, Figure S4). Table 1 reports the D_{10} , D_{50} and D_{90} values obtained for all samples from the cumulated particle size distribution. D_{50} is in the range 0.5 - 2.5 μm for all samples. The particles are broadly distributed, with uniformity coefficient above 4 in all cases, and show a dominant fraction in the micrometer range, and a secondary fraction in the nanometer range. In particular, a wide peak in the range 2 - 4 μm was observed for all materials except the zeolite (ZEO), which showed a wider peak in the range 3 - 10 μm . Only the Na-M2 sample showed a clear second peak, around 200 nm.

The zeta potential (Supporting Information, Figure S5) of the carriers was negative in all explored conditions, as expected for clays and zeolites. More negative values were measured for particles dispersed in DIw; when particles were dispersed in NaCl or CaCl₂ solutions at a high salinity, the zeta potential approached zero, suggesting lower colloidal stability. It is worth to highlight that Ca-M particles showed, in all solutions, zeta potential values closer to neutrality compared to the other particles, suggesting that they may be more prone to aggregation. This was confirmed by a general tendency of Ca-M to sediment faster when dispersed in solutions other than DIw (including tap water). Such a fast sedimentation rate, visible even by eye, cannot be attributed to a difference in size of primary particles nor to a higher density. This behaviour compromised the use of Ca-M in some tests, as discussed in the following paragraphs.

3.2. Adsorption and release tests

The capability of the four candidate carriers to adsorb the pesticide was evaluated first based on adsorption isotherms. To this aim, carriers were loaded with dicamba, in the absence of polymeric coating, in a broad range of herbicide concentration (0.125 g/L to 7 g/L). The upper concentration limit was selected close to the dicamba solubility (8 g/L at pH 1.9).

A contact time of 24 h was first selected based on the literature (Azejjel et al., 2009; Carrizosa et al., 2001). The adsorption isotherms (Figure 1) showed a linear relationship between adsorbed and dissolved herbicide concentration (respectively $C_{s,eq}$ and $C_{w,eq}$) in the explored concentration range. Thus, experimental data were fitted with a linear isotherm, $C_{s,eq} = K_d \cdot C_{w,eq}$. All carriers reached the highest loading capacity for the highest tested concentration. Comparing carriers, the calcium montmorillonite (Ca-M) and one of the sodium montmorillonites (Na-M2) reached the highest loading capacities, corresponding to adsorbed concentrations of approximately 120 mg of dicamba per gram of carrier in both cases. The fitted partition coefficients are, respectively, 0.0251 L/g ($R^2 = 0.9759$) and 0.0216 L/g ($R^2 = 0.9966$). The K10 sodium montmorillonite reached a maximum of 80 mg/g, with $K_d = 0.0124$ L/g ($R^2 = 0.9612$); the lowest loading capacity was registered for the zeolite (ZEO), which showed a maximum retained dicamba mass of 62 mg/g, with $K_d = 0.0122$ L/g ($R^2 = 0.9893$).

A shorter contact time (2 h) was also tested in view of a possible optimization of the loading procedure. However, the results were not univocal (Supporting Information, Figure S7), showing for Ca-M a loading capacity comparable to the one obtained in 24 h, a slightly higher $C_{s,eq}$ for K10, and lower for Na-M2 and ZEO. The results were only partly reproducible, thus suggesting that for at least some carriers a contact time of 2 hours does not guarantee equilibrium between phases. In particular, the structure of zeolite, characterized by cages and channels between the structural tetrahedral that form the primary porosity, is likely responsible for the lowest performance of the ZEO sample at short contact time (Rhodes, 2010; Stocker et al., 2017).

Based on adsorption isotherms, a contact time of 24 h was therefore adopted for the following steps of the study, for all carriers.

Release tests were carried out diluting the nano-formulations in DIw or TAPw, the latter to simulate conditions similar to real pesticide application in the field. The nano-formulation concentration after dilution (5 g/l) corresponded to a dicamba concentration of 0.42, 0.58, 0.55 and 0.34 g/l, respectively for K10, Na-M2, Ca-M, ZEO (compare to adsorption isotherms). This dilution ratio was selected appropriately to guarantee a dicamba concentration in the range of recommended application rate of the commercial products (namely 0.12-1.46 g/l of dicamba, depending on the specific crop).

Release tests in DIw (Figure 2a) were used as a screening for the selection of the best candidate carrier, in combination with adsorption and characterization results. When the uncoated nano-formulations were diluted in DIw, Na-M2 showed the lowest percentage of retained pesticide, both on the short term (less than 1 hour) and on a longer time frame (60 hours and later). The highest release was observed in less than 1 hour, with a subsequent partial re-adsorption on later stages. A final percentage of approximately 30% of dicamba retained on the Na-M2 carrier after dilution was registered for the uncoated carriers, and similar results were obtained also for the CMC-coated carrier (data not reported). This result indicates that Na-M2, even if capable to perform better than other carriers in terms of adsorption capacity, is not a suitable candidate for the dicamba nano-formulation. Even if a high release in a short time could be desirable for applications where an initial higher herbicide quantity is required, in light of the high dicamba volatility a Na-M2 based nano-formulation would not be the optimal choice because of the high risk of pesticide losses in air.

Ca-M showed the highest percentage of retained pesticide after dilution in DIw (approximately 80%). However, a peculiar behaviour was observed for this carrier: a fast, significant release was registered on a short time frame (1 h), leading to a retained percentage of 65%. In a longer time frame, part of the pesticide was re-adsorbed, leading to a final percentage of 80% retained dicamba approximately two hours after dilution. Based on high retention only, Ca-M would have been the best candidate carrier for the development of the nano-formulated dicamba. However, the high variability over time of the retained dicamba may lead to a partly unpredictable behaviour at the field scale: it is not possible to assume a priori how much time

would it take for a farmer to start the field application of the pesticide after its dilution. If this time is shorter than one hour, it is possible that a high fraction of AI is dissolved in water, and therefore prone to volatilization and/or free infiltration in the subsoil. Moreover, its relatively poor colloidal stability, highly sensitive to salt content even in the presence of a polymeric coating, weakens its suitability for real-scale applications, where ionic strength and salt composition of natural or tap water used for pesticide dilution is not under control and highly variable. In light of these considerations, Ca-M was not further considered in the development of the dicamba nano-formulation.

ZEO and K10 showed similar behaviours, with approximately 55% of dicamba remaining adsorbed on the carrier 6 hours after dilutions, therefore resulting suitable candidate carriers. However, due to the higher adsorption capacity of K10 compared to the zeolite, K10 was selected as the best candidate for the next steps of the study. It is also worth to mention that the K10 carrier showed the most constant trend in dicamba release, with an initial fast release in the first 10 minutes, followed by a fast stabilization.

The polymeric CMC coating helped reducing the dicamba release from K10: CMC-coated particles diluted in DIw (Figure 2b) showed a fast (less than 10 mins) stabilization on a plateau corresponding to approximately 80% of retained dicamba, comparable with uncoated Ca-M, with an overall increase of retained AI of approximately 25% with respect to the uncoated K10. The coating is expected to primarily act by hindering the herbicide desorption, increasing the diffusive path toward the bulk fluid (Rashidzadeh et al., 2017). Conversely, when the nano-formulations were diluted in TAPw, the effect of the CMC coating on the release was reduced (Figure 2b), particularly on the long term: the released dicamba was 28% after 6h from both formulations, with a slightly higher release for the coated formulation in the first 3 hours. The major elements expected to influence the release behaviour are pH and salt content. The pH of the formulations diluted in tap water was 8.4 and, since dicamba has a very low pK_a ($pK_a=1.95$), in this alkaline environment the herbicide is present primarily as a deprotonated species; the presence of ionic species in water, such as HCO_3^- and PO_3 , could inhibit the release from the clay surface when coating is not present.

3.3. Volatilization tests

Pre-screening volatilization tests were performed exposing to ambient air the diluted K10-based nano-formulations (K10 and K10-CMC), pure dicamba solution and diluted commercial product on Petri dishes, without soil. The results (Supporting Information, Figure S8) showed a marked volatilization of pure dicamba: after 24 hours about 30% of the AI was lost via volatilization, indicating a moderate attitude of the AI to pass in air phase. Volatilization from the commercial product was negligible, with 99.7% of the pesticide mass remaining after 24 hours, likely thanks to the co-formulants, aimed, among other purposes, at increasing wettability and adhesive performance of the formulated dicamba, thus reducing water-air mass exchange. The uncoated K10 formulation, despite the high amount of AI released in water, limited the volatilization losses to 7.6% of the initial dicamba mass. Interestingly, a simple mass balance suggests that the presence of the K10 particles in the solution inhibits also the volatilization of the freely dissolved dicamba: for uncoated nano-formulations diluted in DIw, approximately 45% of the dicamba initially adsorbed on the carrier is expected to be released after dispersion in DIw (based on release tests), 30% of which should in principle be lost via volatilization (based on pure dicamba volatilization results). However, this would correspond to an overall volatilization of 13.5% of the initial dicamba mass, while only 7.6% was observed here. This could be linked to the specific experimental configuration (i.e. concentrated formulation applied on a Petri dish): the rapid water evaporation (expected to occur the first 1h of the test) may have left the AI in contact with the air phase, with higher mass exchange in the first phase of the test; later on, tortuous paths can have formed through the clay film over the Petri dish while water was evaporating, with an overall reduced contact between dissolved dicamba and air. However, more interestingly, it is also possible that, during water evaporation, part of the dissolved dicamba re-adsorbed onto the clay particles due to the altered equilibrium between the phases (Sciumbato et al., 2004; Strachan et al., 2010). At this stage it is not possible to discriminate between the two processes and, likely, the observed results are obtained as a combination of both. For the CMC-coated formulation a similar trend was observed, with even more limited volatilization,

resulting in an overall loss of 4.6% of the initial dicamba mass after 30 hours, thus confirming the usefulness of the polymeric coating also to prevent volatilization (Rashidzadeh et al., 2017).

The dicamba formulations applied to soils evidenced similar volatilization trends (Figure 3). As a general rule, dicamba alone showed the highest losses: in medium and coarse sand the volatilization was very similar, with approximately 26% of AI losses in 24h; in sandy loam, the volatilization rate was lower (14%). Also for the commercial formulation and for the two nano-formulations, the volatilization was more pronounced in the sand samples and reduced in the sandy loam, suggesting a partial affinity of the compound to the fine fraction of the soil. Contrary to the preliminary volatilization tests performed assessing direct volatilization from the solution, when applied to soils the commercial formulation and the CMC-coated nano-formulation showed comparable volatilization rates. In coarse sand and sandy loam the AI loss is respectively close to 9% and 4.5%, while in medium sand the coated nano-formulation performed significantly better than the commercial dicamba-based product, showing a loss of 5% versus 9.6%. It is finally worth to highlight that in all cases, but particularly in the two sand samples, the presence of the CMC coating significantly reduced the dicamba volatilization (from 15.6% to 5% in medium sand, from 17.2% to 7.4% in coarse sand, and from 5.4% to 5% in sandy loam), thus confirming the key role of the polymeric coating in controlling the release of dicamba from the montmorillonite carrier.

3.4. Transport in the porous medium

Preliminary transport tests were performed injecting the K10 carrier with and without the polymeric coating in sand-packed columns to assess the potential role of the polymeric shell in modifying particle transport. In DIW the presence of the CMC coating is not expected to play a significant role in particle transport. The zeta potential of both coated and uncoated particles is strongly negative (-25.2 ± 0.46 mV for uncoated K10, -38.4 ± 0.89 mV for CMC coated K10). To help understanding transport-controlling processes, DLVO interactions were estimated (Elimelech, 1995). Both particle-particle and particle-sand DLVO interaction profiles are repulsive without secondary minima, which could suggest mild aggregation and/or deposition

(Supporting Information, Figures S9a and S10a). Coherently, the particle breakthrough curves (BTCs) obtained from column transport tests (Figure 4) show a non-negligible mobility of both coated and uncoated carrier. In both cases the breakthrough concentration approached 60% of the injected one, showing minimal influence of the coating. The mass balance (Table 2) indicates a slightly higher mobility for the CMC-coated particles, coherently with slightly more repulsive particle-particle and particle-collection DLVO profiles: 67.22% of the injected mass of coated K10 was eluted at the end of the test, while 63.09% was recovered at column outlet for bare K10. Based on the measured zeta potential values and the clearly repulsive DLVO interaction it is possible to attribute the carrier retention in the porous medium mainly to physical mechanisms, above all to mechanical filtration: the K10 particles, even if stably dispersed in the injected suspension, are sufficiently large to partly interact with the porous medium; the ratio of K10 d_{90} (3.746 μm) to sand d_{10} (370 μm) approaches the critical ratio of 1% above which a partial mechanical filtration of particles can be observed (Luna et al., 2015; Xu et al., 2006). The experimental breakthrough curves were modelled using a dual deposition site attachment/detachment model (eq. 4) with a linear irreversible deposition site (representing mechanical filtration) and a linear reversible deposition site (representing physical-chemical interactions, obtained imposing $A_2 = 0$ in the model equation). They showed a good agreement between simulated and measured BTCs. The fitted coefficients are reported in the first two columns of Table 2. The first interaction site is dominant, thus reflecting the dominance of mechanical filtration as retention mechanism. The particle retention observed in these tests can be assumed to be the minimum to be reasonably expected in all experimental conditions, due to the clearly repulsive interactions.

When comparing the different dicamba-based formulations dispersed in NaCl 30 mM solutions (Figure 5), a significant discrepancy between nano-formulated and not nano-formulated suspensions is observed. Pure dicamba and the commercial product, as it can be denoted from their breakthrough curves, were transported similarly to a trace. They reached $C/C_0 = 1$ with no evident delay (thus indicating the absence of any relevant adsorption phenomenon onto the silica sand) and all injected mass is recovered at column outlet at the end of the test (recovered mass of 99.35% for pure dicamba and 98.44% for the commercial product).

The K10-formulated dicamba, both in presence and absence of coating, showed a remarkably limited mobility compared to pure and commercial dicamba. During the injection and subsequent flushing at constant NaCl concentration (i.e. in PVs 1 to 10), the uncoated formulation did not show any appreciable breakthrough, while the CMC-coated formulation reached a maximum outflow concentration equal to 15.3% of the injected one (Figure 5, respectively red and green curves). In terms of mass balance, this corresponds to 1.74% of injected uncoated K10 nano-formulation reaching the column outflow and 14.45% of the coated one in the first PVs. The breakthrough curves of the two carriers were successfully fitted with eq. 4 (Supporting Information, Figure S11). In this case the fitted attachment coefficients for site 1 (linear irreversible attachment) are approximately one order of magnitude higher than those obtained for the carriers in DIw, both for coated and bare K10, indicating that a stronger physical retention was occurring for the nano-formulations dispersed in the NaCl solution compared to the carriers dispersed in DIw. As for the second interaction site, for the CMC-coated K10, a linear reversible interaction ($A_2 = 0$) was adequate to represent the physical-chemical interaction, while for the bare particles a ripening mechanism ($A_2 > 0$) correctly described the particle physical-chemical interactions occurring in the porous medium. This is in agreement with the predicted DLVO interaction profiles (Supporting Information, Figures S9b and S10b): for the CMC-coated nano-formulation, a weakly repulsive profile is obtained for particle-particle interaction, with a shallow secondary attractive minimum and a very limited energy repulsive barrier (~ 0.2 KT), suggesting that aggregation may partly occur in the suspension. Conversely, for the uncoated K10 formulation, an entirely attractive profile is obtained, indicating that particles are attracting each other and agglomerating. As for particle-collector interaction, on the contrary, repulsive profiles are observed, even though the repulsive energy barrier against deposition is of limited extent ($\sim 2-4$ KT). In terms of particle transport, this suggests that, for the uncoated formulation, particles are partly aggregated, thus prone to a more pronounced mechanical filtration (represented by the first interaction site) with respect to the carrier in DIw, and deposited particles are attracting suspended ones, thus resulting in a strong ripening (reflected by the second interaction site). For the coated nano-formulation, the CMC shell prevents excessive aggregation,

mechanical filtration is more limited and ripening does not occur, at least on the time scale and travel path explored in these tests.

It is worth to recall that, when the nano-formulations are diluted in water, a fraction of dicamba is released and therefore is present as freely dissolved compound in the injected suspensions; in these tests this fraction corresponded approximately to 25% of the total dicamba for uncoated K10 and 20% for CMC-coated K10 (compared with release tests). The breakthrough curves for the free fraction of dicamba (Supporting Information, Figure S12) show that, for both nano-formulations, the free compound is transported through the sand-packed column similarly to a tracer, with no evidence of any retention. Based on this additional information, it is possible to draw a mass balance for the total dicamba (i.e. both freely dissolved and embedded in the carrier), which indicates that, at the end of the first 10 PVs, for the uncoated carrier 10% of the injected dicamba has been eluted from the column, and 20.73% has been eluted for the CMC-coated carrier. This finding suggests that, even though the polymeric coating helps preventing excessive release and volatilization of the AI from the carrier, it slightly enhances mobility compared to bare K10 particles. However, in the second part of the transport tests (PVs 10 to 20), when the columns were flushed with DIw, the abrupt change in ionic strength generated a markedly different response of bare and CMC-coated carriers. The uncoated nano-formulation was strongly mobilized by the step change in ionic strength, resulting in the remobilization of most formulation previously retained in the column (79.68% of the K10 carrier and 88.43% of the total injected dicamba was eluted at the end of the test). Conversely, the mobilization of the CMC-coated nano-formulation was minimal, resulting in an overall elution of 15.35% of the injected carrier, corresponding to 24.52% of the total injected dicamba. This last finding remarkably suggests that, even though the CMC coating imparts a slightly higher mobility to the nano-formulation, it also reduces the effects of changes in ionic strength. In particular, it decreases the possibility of the retained nano-formulation re-mobilization in case the salt concentration is reduced, for example, due to intense infiltration of rain.

3.5. Weed control efficacy tests

The dose-response curve highlighted a relative weight reduction of the treated weeds at increasing herbicide rates (Figure 6 and Table S2 in Supporting Information). All the herbicide formulations showed higher efficacy on *S. nigrum* than against *A. retroflexus*. The efficacy was consistent between herbicides in the two species, highlighting a higher relative weight reduction with K10-CMC, followed by commercial dicamba and K10. However, significant differences in ED values were found in *S. nigrum* only, with K10 less effective than the other two herbicide formulations. In particular the herbicide rate able to reduce weed biomass in *S. nigrum* by 50% was 1.5 g AI/ha and 5.8 g AI/ha with K10-CMC and commercial, respectively.

At 90% weight reduction (corresponding to a relative weight of 10% in comparison to control) all the herbicides acted similarly in the case of *A. retroflexus*, in which values of herbicide rates higher than 100 g AI/ha were necessary to obtain a similar efficacy level. In the case of *S. nigrum*, 90% weight reduction was obtained at 53.1 g AI/ha and 82.5 g AI/ha with K10-CMC and commercial, respectively, values much smaller than that required for K10 (156.3 g AI/ha).

4. CONCLUSIONS

In this work a novel approach based on the use of natural clays to reduce the environmental mobility of dicamba was proposed. Four candidate carriers (namely, two Na-montmorillonites, a Ca-montmorillonite and a zeolite) were tested, and all proved to be effective in adsorbing the herbicide. However, not all of them guaranteed easy applicability or reduced release in water when the formulations were diluted (i.e. mimicking the preparation of the product prior field application). A key role was played by the polymeric coating, formed by carboxymethyl cellulose, a food-grade biodegradable polymer used in a broad range of applications, from food industry to enhanced oil recovery to pharmaceuticals. The best performing formulation, namely CMC-coated Na-montmorillonite K10, was identified as the most advantageous compromise between technical constraints and potential environmental effects. From the technical point of view, it showed a good control

of dicamba release after dilution and good colloidal stability, which allows its application using conventional pesticide spraying equipment. Its efficacy in the greenhouse tests against target weeds was similar or sometimes higher than that of commercial dicamba formulation, and showed promising prospects for improvements, which will be the focus of upcoming work.

Compared to other approaches previously proposed in the literature, based on organoclays and, more in general, of chemically modified adsorbing carriers, in this work the focus was on the use of natural unmodified materials as carriers, and on the development of a simple and low-impact preparation method, applicable at room temperature and pressure without addition of chemicals. The loading capacity obtained in this study is comparable or lower than those reported in the literature for organoclays (e.g. (Carrizosa et al., 2001)), but the CMC-coated K10 showed a good control of the major environmental criticalities of dicamba, namely volatilization and mobility in the subsoil. In particular, the CMC-coated K10 allowed a control of volatilization losses comparable to a commercial dicamba-based product, even in the absence of specific co-formulants, which are present in the commercial formulation but were not included in the nano-formulations.

Concerning the mobility in the subsoil, in this work a preliminary assessment was performed focusing on the saturated zone, a potential major route for dicamba migration in the subsoil. The remarkable mobility of pure and commercially formulated dicamba observed in transport tests, both in synthetic and real water indicates that, in case dicamba reaches an aquifer system as a free compound, it is expected to be highly mobile without any significant attenuation (except obviously for degradation processes, which are not appreciable on the short time scale of the experiments performed in this study). Conversely, the use of a mineral carrier significantly reduced the potential mobility, in all explored conditions. Also in this case, the polymeric coating played a key role. Even though it imparted a slightly higher mobility to the nano-formulations, compared to bare carriers, it also significantly reduced the risk of re-mobilization when abrupt hydrochemical perturbations were applied, namely, when the columns were flushed with deionized water. Even if this condition is clearly unrealistic in a field-scale scenario, it has been adopted here as an extremized simulation of intense rain events: in this case the precipitation, characterized by a significantly lower salinity than

groundwater, may infiltrate and, particularly for shallow aquifer systems, can significantly reduce the local groundwater salinity, with the risk of local re-mobilization of the nano-formulation. Clearly an in-depth study of the potential mobility of the new nano-formulations in the subsoil requires the assessment of a broader range of experimental conditions, and a detailed investigation focused on the top soil, which is beyond the scope of this paper. More in general, deeper investigation is needed on several other aspects touched in this work, including for example a further optimization of the preparation procedure, and a more detailed evaluation of the nano-formulation efficacy toward target weeds. However, the authors believe that even the preliminary results presented here already provide a first insight on the potentialities of natural clays as a low-impact solution to reduce environmental drawbacks of critical agrochemicals.

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Conflicts of interest

There are no conflicts to declare

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Figures

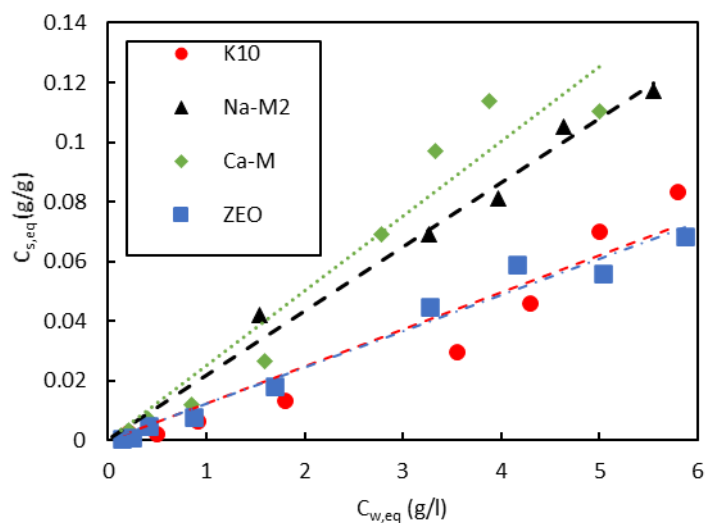


Figure 1 Adsorption isotherms for the four candidate carriers (K10, Na-M2, Ca-M, ZEO) obtained for a contact time of 24 hours. Experimental data (dots) and least-squares fitted linear isotherms (lines).

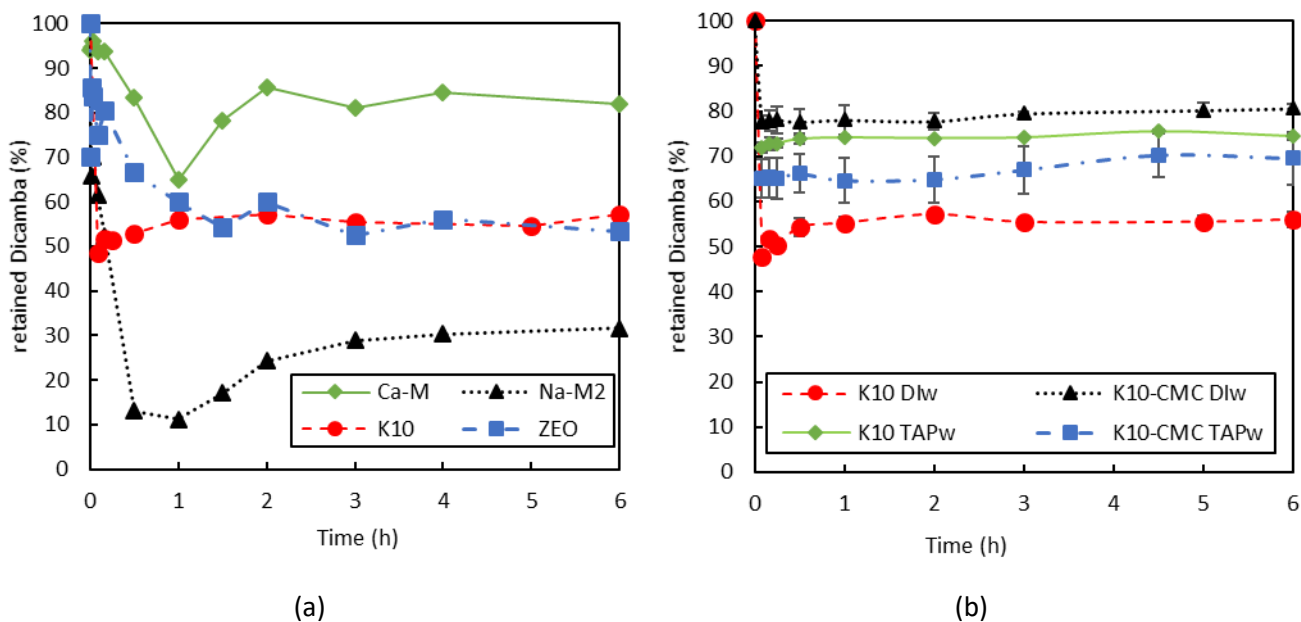


Figure 2 Release tests: retained dicamba $R(t)$ expressed as a percentage for the four carriers (ZEO, Na-M2, Ca-M, ZEO), after dilution in deionized water (DIw), without CMC coating (a), and on K10, after dilution in deionized water (DIw) or in tap water (TAPw), with and without CMC coating (b)

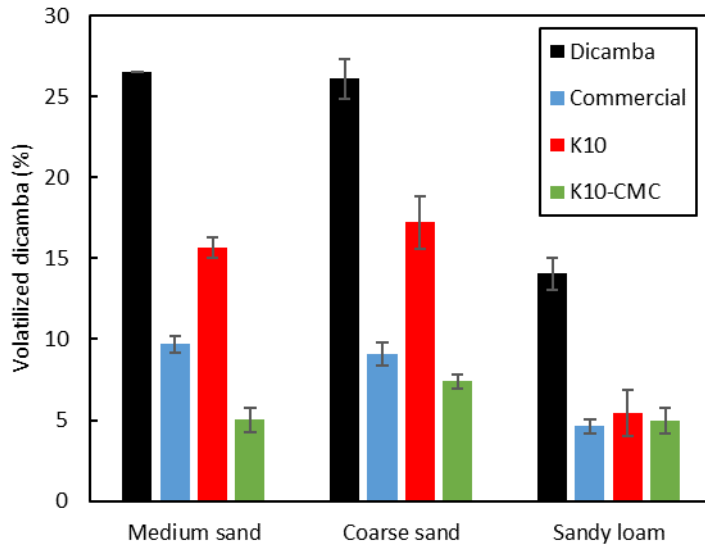


Figure 3 Volatilization from soils: dicamba loss 24 h after application to different soils, reported as percentage of volatilized mass with respect to applied mass

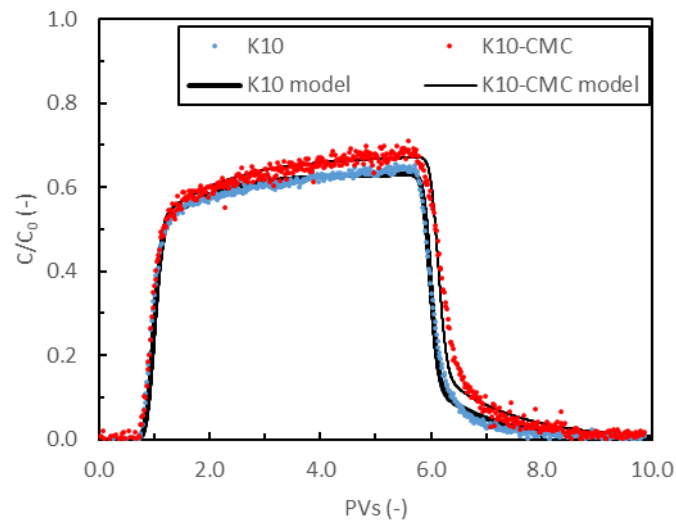


Figure 4: Breakthrough curves for uncoated (blue) and CMC-coated (red) K10 carrier, without dicamba. Experimental data (coloured dots) and least-squares model curves (black lines) are reported.

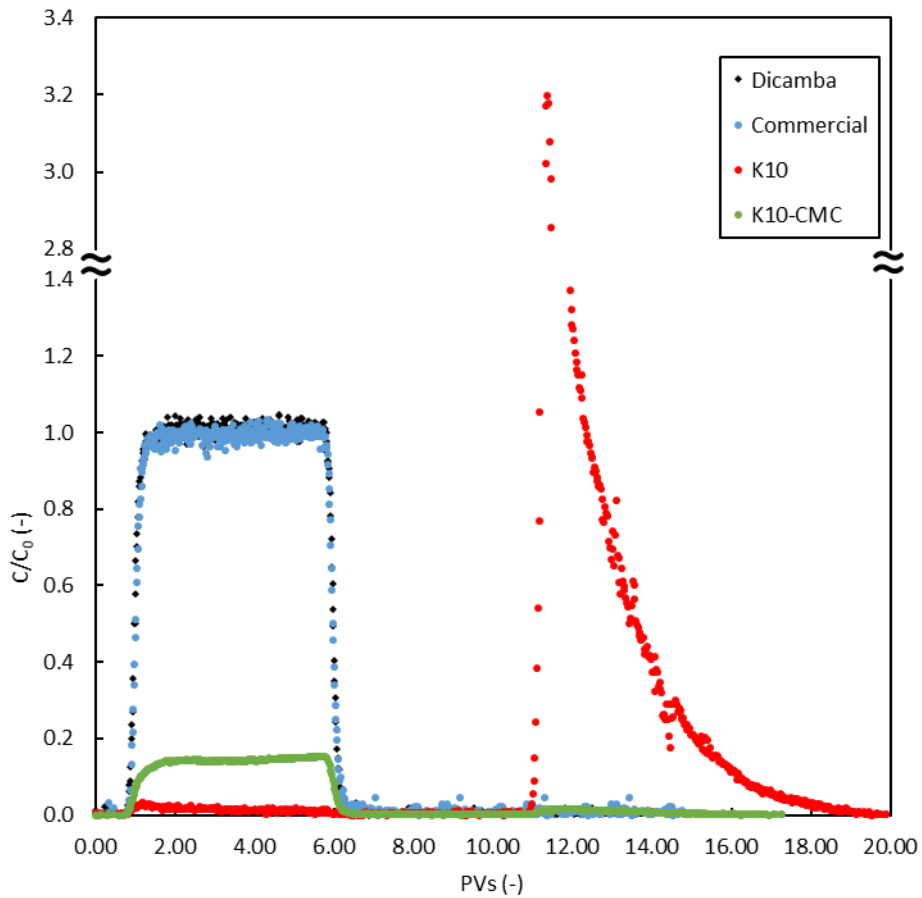


Figure 5 Breakthrough curves for non-formulated dicamba (black), commercial formulation (blue), uncoated (red) and CMC-coated (green) nano-formulations. All formulations were dispersed in NaCl 30 mM solution (injection: pore volumes 0 to 5); the first flushing (pore volumes 5 to 10) was performed injecting pesticide-free 30 mM NaCl solution; the second flushing (pore volumes 10 to 20) was performed with DIw.

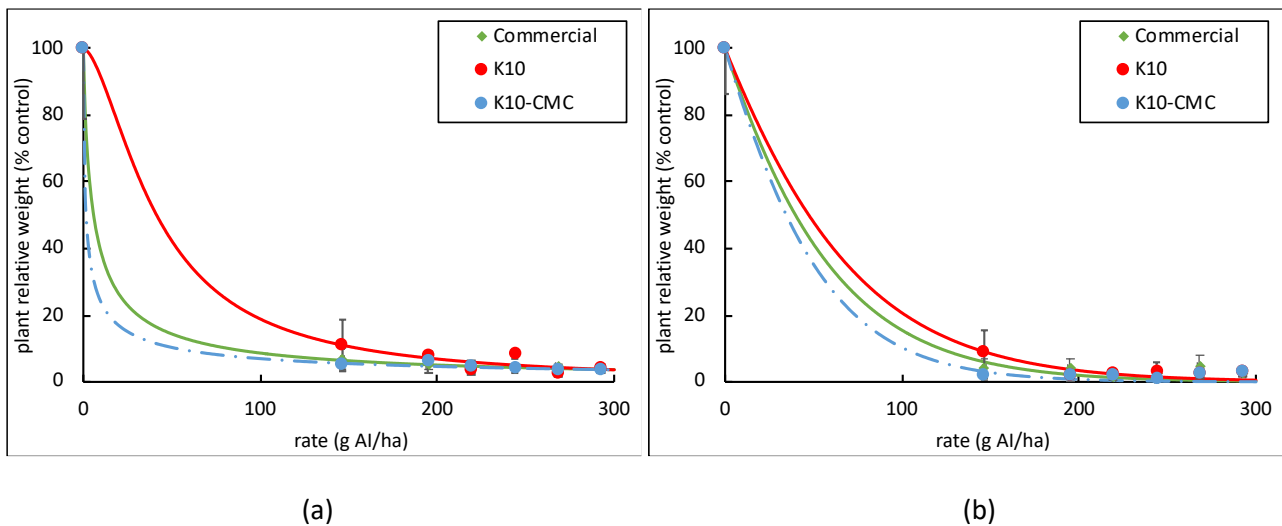


Figure 6 Dose-response curves between *Solanum nigrum* (a) and *Amaranthus retroflexus* (b) plant relative weight and herbicide dose of K10-CMC, commercial dicamba and K10 formulations.

Tables

Table 1 Size and zeta potential in DIW and their variation in NaCl different ionic strength

	Particle size distribution (μm)				Zeta potential (mV)		
	D ₁₀	D ₅₀	D ₉₀	U=D ₆₀ /D ₁₀	in DIW	in NaCl 100 mM	in CaCl ₂ 2mM
K10	0.287	1.561	3.746	6.49	-18.9 ± 2.0	-14.2 ± 0.7	-6.2 ± 0.5
Na-M2	0.154	0.570	2.834	6.06	-26.5 ± 2.6	-14.6 ± 2.8	-13.2 ± 1.1
Ca-M	0.180	0.656	2.262	4.93	-14.3 ± 0.9	-10.2 ± 0.1	-8.0 ± 0.6
ZEO	0.674	2.525	2.525	4.39	-18.2 ± 0.5	-8.7 ± 0.3	-12.3 ± 1.0

Table 2 Mass balance and fitted model parameters for the transport of carriers and formulations in the porous medium

			K10 carrier only		Dicamba formulations			
			K10	K10-CMC	Dicamba	Commercial	K10	K10-CMC
Mass balance	Carrier	Eluted mass @10 PVs	63.09%	67.22%	n.a.	n.a.	1.74%	14.45%
		Eluted mass @20 PVs	n.a.	n.a.	n.a.	n.a.	79.68%	15.35%
	Free dicamba	Eluted mass @10 PVs	n.a.	n.a.	99.35%	98.44%	99.77%	100.00%
	Total dicamba	Eluted mass @10 PVs	n.a.	n.a.	99.35%	98.44%	10.06%	20.73%
		Eluted mass @20 PVs	n.a.	n.a.	n.a.	n.a.	88.43%	24.52%
Modeling	Site 1	k _{a,1} (1/s)	6.99·10 ⁻⁴	6.32·10 ⁻⁴	n.a.	n.a.	4.00·10 ⁻³	4.48·10 ⁻³
	Site 2	k _{a,2} (1/s)	3.53·10 ⁻⁴	4.32·10 ⁻⁴	n.a.	n.a.	6.40·10 ⁻³	9.70·10 ⁻⁴
		k _{d,2} (1/s)	1.68·10 ⁻³	1.38·10 ⁻³	n.a.	n.a.	2.68·10 ⁻⁵	6.07·10 ⁻³
		A ₂ (-) (*)	0	0	n.a.	n.a.	998.1	0
		β ₂ (-) (**)	1	1	n.a.	n.a.	1	1
	Fitting	R ²	0.9963	0.9937	n.a.	n.a.	0.8384	0.9956

(*) fitted only for dicamba nano-formulations

(**) not fitted