Can the botanical azadirachtin replace phased-out soil insecticides in suppressing the soil insect pest *Diabrotica virgifera virgifera*?

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**Abstract**

**Background:** Due to recent bans on the use of several soil insecticides and insecticidal seed coatings, soil-dwelling insect pests are increasingly difficult to manage. One example is the western corn rootworm (*Diabrotica virgifera virgifera*, Coleoptera: Chrysomelidae), a serious root-feeder of maize (*Zea mays*). We investigated whether the less problematic botanical azadirachtin, widely used against above-ground insects, could become an option for the control of this soil insect pest.

**Methods:** Artificial diet-based bioassays were implemented under standard laboratory conditions to establish dose response curves for the pest larvae. Then, potted-plant experiments were implemented in greenhouse to assess feasibility and efficacy of a novel granular formulation of azadirachtin under more natural conditions and in relation to standard insecticides.

**Results:** Bioassays in three repetitions revealed a 3-day LD50 of 22.3 µg azadirachtin/ml which corresponded to 0.45 µg/neonate of *D. v. virgifera* and a 5-day LD50 of 19.3 µg/ml or 0.39 µg/first to second instar larva. No sublethal effects were observed. The three greenhouse experiments revealed that the currently proposed standard dose of a granular formulation of 38 g azadirachtin/hectare for in-furrow application at sowing is not enough to control *D. v. virgifera* or to prevent root damage. At 10× standard-dose total pest control was achieved as well as the prevention of most root damage. This was better than the efficacy achieved by cypermethrin-based granules and comparable to tefluthrin-granules, or thiamethoxam seed coatings. The ED50 for suppressing larval populations were estimated at 92 g azadirachtin/ha, for preventing heavy root damage 52 g/ha and for preventing general root damage 220 g/ha.

**Conclusions:** There seems clear potential for the development of neem-based botanical soil insecticides for arable crops such as maize. They might become, if doses are increased and more soil insecticides phased out, a promising, safer solution as part of the integrated pest management toolkit against soil insects.

**Keywords:** Integrated pest management, Western corn rootworm, Azadirachtin, *Zea mays*, Biopesticide, Soil insecticide, Biological control

**Introduction**

Corn rootworms are, next to wireworms, grubs and cutworms, serious soil-dwelling insect pests of maize (*Zea mays*). One of the rootworms, the western corn rootworm (*Diabrotica virgifera virgifera*, Coleoptera: Chrysomelidae) is one of the most problematic (Krysan and Miller 1986). Its three larval instars feed almost exclusively on the roots of maize, which becomes apparent when plants lodge (Chiang 1973). *Diabrotica v. virgifera* causes yield losses to large maize production areas of the USA and southern Canada (Kim and Sappington 2005) as well as in Central Europe (Miller et al. 2005).
Affected growers attempt to manage the pest mainly through rotating their fields, thereby interrupting the life cycle of *D. v. virgifera*. Many growers apply granular or fluid soil insecticides, mainly pyrethroids or organophosphates, or use neonicotinoid-coated maize seeds to target the root feeding larvae. North American growers also use transgenic maize expressing different *Bacillus thuringiensis* proteins which are toxic to rootworms (Levine and Oloumi-Sadeghi 1991; Dominguez-Arrizabalaga et al. 2020), but their efficacies under field conditions are variable (Clair et al. 2020; Gassmann et al. 2020). Additionally, broad-spectrum foliar insecticides are occasionally sprayed against the adults using high clearance spraying machinery (Rozen and Ester 2010). Foliar insecticides are often broad spectrum and knock-down contact-pesticides with considerable non-target effects. Also, many soil insecticides and seed coatings are problematic due to their human toxicity and/or serious non-target or other environmental effects.

This has resulted in public concerns and in a ban on the use of neonicotinoid seed-coatings in field crops (Georgiadis et al. 2011), and restrictions in the use chlorpyrifos- and tefluthrin-based soil insecticides in many countries. Only few countries remain, that still have such ingredients on the soil pesticide market, such as the USA. In Hungary, tefluthrin is only registered under special emergency registration. However, tefluthrin will definitely disappear from the European pesticide market (European-Commission 2011) and likely from the entire global market due to its high acute toxicity (World Health Organization 2009). Therefore, options for growers to control *D. v. virgifera*, and also other soil insect pests such as wireworms, grubs and cutworms will be limited, particularly in Europe. Novel pest management solutions and agents are urgently needed, particularly less environmentally-disruptive ones.

For example, neem preparations with their active ingredients of different azadirachtins are widely-used botanical insecticides (Saxena 1989; Scmutterer 1995; Dougoud et al. 2019). Although its modes of action are still somewhat uncertain, they are known to have broad spectrum insecticidal activity as well as some nematocidal, isopodicidal, fungicidal and plant promoter activity (CRC 1989; Doshi et al. 2018, 2020). There are numerous products available in most world regions, mainly against above-ground, soft-bodied insect pests (Dougoud et al. 2019). In some regions, growers also prepare self-made homebrews from leaves or seeds of the tree *Azadirachta indica*, which is of south Asian origin but nowadays widely distributed in the tropics and sub-tropics of many regions (Dougoud et al. 2019). The advantage of neem is that it has a low acute or chronic toxicity to humans and breaks down relatively quickly in the environment (Boeke et al. 2004). Another advantage is that neem is systemic, translaminar as well as of contact mode of activity (Stark and Rangus 1994; Dougoud et al. 2019), allowing its diverse usage. It can directly cause mortality to insects, can inhibit growth and moulting, and even can cause chronic effects such as on insect fertility (Ladd Jr et al. 1984; Al-Sharook et al. 1991; Stark and Rangus 1994; Mehaoua et al. 2013; Merabti et al. 2017).

Astonishingly, there are still limited studies and limited use of neem products against below-ground insect pests (Bhagat 2005). This is surprising because neem leaves, grinded seeds or leftovers from seed processing are occasionally used as a soil amendment against plant parasitic nematodes (Dougoud et al. 2019) as well as a biofertilizer, even in maize (Vageesh et al. 2016). Neem seed extracts also has some fungicidal properties when used as coating of maize seeds (Sowley et al. 2017). Consequently, there exist some experience for in-soil applications of neem, such as leaves or commercial granules (Balaji 2014). Nevertheless, only few products are available for soil application, and knowledge on their effects against soil insects is limited due to their concealment below-ground in or on the roots of crops.

This is particularly true for rootworms (Diabroticina), several of them, as stated above, being serious root-feeding pests of maize. Azadirachtin is known to be toxic to rootworm larvae, such as against *D. speciosa* (Boiça Júnior et al. 2017) or the here-studied *D. v. virgifera* (Xie et al. 1991), and repellent to larvae of cucumber beetles such as *Diabrotica undecimpunctata howardi* (Landis and Gould 1989) or to adults of the closely related *Acalymma vittatum* (Reed et al. 1982). However, further experimentation on the use of azadirachtin against rootworms seem scarce. Only Estes et al.(2018) attempted to examine liquid and granular formulations of neem against *D. v. virgifera* larvae in Illinois, USA, but with inconclusive results. There seems a knowledge gap on how to effectively use neem against rootworms or other soil insect pests, a gap we try to close with the here-presented study.

We aimed at better understanding the pest control potential of azadirachtin using a novel granular neem-based soil insecticide. First, bioassays were conducted to establish the LD$_{50}$ of azadirachtin on the neonates of *D. v. virgifera*. Second, the suggested standard dose of neem granules was compared with lower and higher dosages in relation to standard insecticides in potted-maize plant trials under semi-natural conditions. This set of experimental steps was hoped to allow conclusions on the efficacy and potential feasibility of neem-based granule applications for corn rootworm control. Ultimately, this would help to diversify the currently limited integrated
pest management toolbox against rootworms and potentially other soil insect pests.

Material and methods

Target pest *Diabrotica v. virgifera* LeConte (Coleoptera: Chrysomelidae, western corn rootworm, EPPO code DIABVI) were mass-reared following procedures of George and Ortman (1965), Branson et al. (1975) and Singh and Moore (1999). A non-diapause colony (USDA ARS, Bookings, USA) was used to infest artificial diet-based bioassay with neonates and potted—maize plants in greenhouse experiments with ready-to-hatch eggs (see procedures below). The population is considered susceptible to most insecticides or novel agents as it had not been exposed to any of those, and therefore resistance is considered unlikely (Wright et al. 2000; Magalhaes et al. 2007).

Test agents
An azadirachtin-based granular soil insecticide was tested against *D. v. virgifera* larvae in comparison to a set of commonly used contact and/or systemic insecticides (positive controls) as well as to untreated controls (Tables 1, 2).

In artificial-diet based bioassays against neonates of *D. v. virgifera* in the laboratory, azadirachtin-granules were dissolved in water and 20 µl applied/well/larva (see details below). Those dissolved azadirachtin granules were compared with a commercial fluid azadirachtin formulations and the insecticides tefluthrin, cypermethrin, imidacloprid as well as with untreated control (Table 1).

In potted-maize experiments under greenhouse conditions, azadirachtin-granules were manually applied when the maize seeds were placed into a sowing furrow across the pots (see details below). The azadirachtin-based granular soil insecticide was compared with the soil insecticides tefluthrin and cypermethrin and to thiamethoxam-coated seeds as well as with untreated infested and un-infested control (Table 2).

**Azadirachtin** was tested as per Tables 1 and 2. The main test product was a granule formulation provided by Coromandel International Ltd, India (NeemAzal™ 0.15G similar to Avana™ by Parry America). It contained 0.165% azadirachtin (mainly A with some B, chromatogram-verified, batch 2001-10) as well as 0.35% other neem compounds (Aza-F, Aza-H, Aza-I, salanin,

| Table 1 | Treatment characteristics in artificial diet-based bioassay against neonates of *Diabrotica v. virgifera* under standardised semi-sterile laboratory conditions |
|------------------|------------------|------------------|------------------|------------------|
| **Treatment**    | **Formulation**  | **Dose/ml (a.i.)** | **Dose/cm² (a.i.)** | **Dose/20 µl/0.34 cm² well/larva (a.i.)** | **Experiment number (# plates; # wells)** |
| **Test agent**   |                  |                  |                  |                  |                          |
| Azadirachtin 0.15% (NeemAzaal 0.15G) | Granule | 0.067 mg (0.1 µg) | 0.0038 mg (0.006 µg) | 0.0013 mg (0.002 µg) | 3 (6; 48) |
|                  |                  | 0.67 mg (1 µg)   | 0.038 mg (0.06 µg)  | 0.013 mg (0.02 µg)  | 3 (6; 48) |
|                  |                  | 6.7 mg (10 µg)   | 0.38 mg (0.6 µg)   | 0.13 mg (0.2 µg)   | 3 (6; 48) |
|                  |                  | 66.7 mg (100 µg) | 38 mg (6 µg)       | 1.33 mg (2 µg)     | 3 (6; 48) |
| Azadirachtin 1% (NeemAzaal-T/S 10EC) | Fluid | 0.1 µl (0.1 µg)  | 0.0006 µl (0.006 µg) | 0.0002 µl (0.002 µg) | 3 (6; 48) |
|                  |                  | 0.1 µl (1 µg)    | 0.0006 µl (0.006 µg) | 0.0002 µl (0.002 µg) | 3 (6; 48) |
|                  |                  | 1 µl (10 µg)     | 0.06 µl (0.6 µg)   | 0.02 µl (0.2 µg)   | 1 (3; 24), 2 (6; 48), 3 (6; 48) |
|                  |                  | 3 µl (30 µg)     | 0.18 µl (1.8 µg)   | 0.06 µl (0.6 µg)   | 1 (3; 24) |
|                  |                  | 5 µl (50 µg)     | 0.3 µl (3 µg)      | 0.1 µl (1 µg)      | 1 (3; 24) |
|                  |                  | 10 µl (100 µg)   | 0.6 µl (6 µg)      | 0.2 µl (2 µg)      | 1 (3; 24), 2 (6; 48), 3 (6; 48) |
|                  |                  | 100 µl (1000 µg) | 6 µl (60 µg)       | 2 µl (20 µg)       | 1 (3; 24), 2 (6; 48) |
|                  |                  | 500 µl (5000 µg) | 30 µl (300 µg)     | 10 µl (100 µg)     | 2 (6; 48) |
|                  |                  | 1000 µl (10,000 µg) | 60 µl (600 µg) | 20 µl (200 µg) | 2 (6; 48) |
| Positive controls |                  |                  |                  |                  |                          |
| Imidacloprid 20% (Confidor 200 SL) | Fluid | 0.01 µl (2 µg)  | 0.0006 µl (0.12 µg) | 0.0002 µl (0.04 µg) | 1 (3; 24), 2 (6; 48), 3 (6; 48) |
| Cypermethrin 0.8% (Belem 0.8 MG) | Micro granule | 12.5 mg (100 µg) | 0.73 mg (5.9 µg) | 0.25 mg (2 µg) | 3 (6; 48) |
| Tefluthrin 1.5% (Force 1.5G) | Fine granule | 6.7 mg (100 µg) | 0.38 mg (5.9 µg) | 0.13 mg (2 µg) | 3 (6; 48) |
| Negative controls |                  |                  |                  |                  |                          |
| Untreated-infested | – | – | – | – | 1 (3; 24), 2 (6; 48), 3 (6; 48) |
| Untreated un-infested | – | – | – | – | 1 (3; 24), 2 (6; 48), 3 (6; 48) |

Each diet-filled well infested with one neonate larva

*a Similar to Avana™ by Parry America*
nimbin and fatty acids). The granules were of 2 to 3 mm size (Formulation G of GIFAP code) and of slow release.

As a comparison, a common fluid formulation (NeemAzal™ T/S, Trifolio-M, Germany) was used in artificial diet-based bioassays. It contained 1% azadirachtin A originating from 4% neem seed extract (Azadirachta indica tree), but with unknown concentration of other neem compounds.

Tefluthrin served as a positive control (Tables 1 and 2). The product Force™ 1.5G (Syngenta, Hungary) are fine granules (1 to 2 mm diameter, Formulation FG of GIFAP code) with 1.5% active ingredient.

Cypermethrin served as a positive control (Tables 1 and 2). The product Belem™ 0.8MG (Certis, SBM Development SAS, France) are micro granules (0.8 to 1 mm diameter, Formulation MG of GIFAP code) with 0.8% active ingredient.

Imidacloprid served as a positive control in the artificial diet-based bioassays in the laboratory (Table 1). The product Confidor™ 200SL (17.8% w/w imidacloprid, Bayer Crop Science, Germany) is a soluble concentrate (SL of GIFAP code) with about 20% active ingredient.

Thiamethoxam served as a positive control in the potted-plant greenhouse experiments (Table 2). The product Cruiser™ 350FS (25 to 30% w/w thiamethoxam, Syngenta, Hungary) is a suspension concentrate for seed treatments (FS of GIFAP code) with about 30% active ingredient.

Table 2: Treatment characteristics in potted-maize plant experiments against larvae of Diabrotica v. virgifera under greenhouse conditions

| Treatment | Formulation | Dose/ha (a.i.) | Dose/meter furrow (a.i.) | Dose/10 cm furrow/plant/pot (a.i.) | Dose/cm² (a.i.) | Experiment number (# blocks; # plants) |
|-----------|-------------|---------------|--------------------------|-----------------------------------|----------------|---------------------------------------|
| Test agent | Granule     | 18 kg (27 g)  | 1.3 g (2 mg)             | 130 mg (0.2 mg)                    | 26 mg (40 µg)  | 3 (3; 15)                             |
| Azadirachtin 0.15% (Neem Azaal 0.15G)¹ | 25 kg (38 g) ¹ | 1.9 g (2.8 mg) | 190 mg (0.28 mg)          | 38 mg (56 µg)                      | 1 (3; 15), 2 (4; 20), 3 (3; 15) |
| (limoid)  | 135 kg (200 g) | 10.2 g (15 mg) | 1020 mg (1.5 mg)         | 204 mg (300 µg)                    | 3 (3; 15) |
|          | 250 kg (370 g) | 19 g (28 mg)   | 1900 mg (2.8 mg)         | 380 mg (560 µg)                    | 3 (3; 15) |
|          | 2500 kg (3700 g) | 190 g (280 mg) | 19,000 mg (28 mg)        | 3800 mg (5600 µg)                  | 3 (3; 15) |
| Positive controls | Micro granule | 12 kg (96 g) ² | 0.9 g (7.2 mg)           | 90 mg (0.72 mg)                    | 18 mg (0.14 µg) | 1 (3, 15), 2 (4; 20) |
| Cypermethrin 0.8% (Belem 0.8 MG) (pyrethroid) | 25 kg (200 g) ² | 1.9 g (15 mg)  | 188 mg (1.5 mg)          | 38 mg (0.3 µg)                      | 3 (3; 15) |
| Tefluthrin 1.5% (Force 1.5G) (pyrethroid) | Fine granule | 13.3 kg (200 g) ² | 1 g (15 mg)             | 100 mg (1.5 mg)                    | 20 mg (0.3 µg) | 1 (3, 15), 2 (4, 20, 3 (3; 15) |
| Thiamethoxam 30% (Cruiser 350FS) (neonicotinoid) | Seed coating | 180 ml (63 g/50,000 seeds) ² | 18 µl (6.25 mg/5 seeds) | 3.6 µl (1.25 mg/seed) | 3.6 µl (1.25 mg/seed) | 1 (3, 15), 2 (4, 20) |
| Negative controls | Untreated-infested | – | – | – | – | 1 (3, 15), 2 (4; 20), 3 (3, 15) |
| | Untreated un-infested | – | – | – | – | 1 (3, 15), 2 (4; 20, 3 (3; 15) |

About 2 × 10 cm treatment strips along soil surface in the pots extrapolated to hectare doses of products for 13,300 row meters. 1.5 l soil in pots with maize plants used. Plants of experiment 1 and 3 infested with 50 ready-to-hatch eggs each, and of experiment 2 with 100 eggs. Block numbers reflect the within experiment replicates. Plants numbers reflect the sample size per treatment per experiment.

¹ Standard doses
² Similar to Avana™ by Parry America

Artificial diet-based laboratory bioassays

Experimental setup

To assess lethal doses of azadirachtin on neonates of D. v. virgifera, artificial diet-based bioassays with different dosage were conducted in three replicates under controlled semi-sterile conditions (Table 1). Azadirachtin from a common fluid formulation was compared with a novel granular formulation. The insecticides cypermethrin, tefluthrin and imidacloprid served as positive control. Sterilised tap water or no treatment at all served as negative controls. Each bioassay consisted...
of 3 to 6 polystyrene plates of 96 wells each (07-6096 of Biologix Ltd., USA, or Costar 3917 of Corning Inc., USA). Each well was of 330 µl volume with 5 mm in diameter and 10 mm height, and had a 0.34 cm² surface. Each treatment was applied to 8 wells of each plate per bioassay. Each treatment-dose combination was tested in at least two true replicates.

The larval diet for a bioassay had been prepared 1 day before treatment and infestation. The diet was prepared under semi-sterile conditions following methods of Sutter et al. (1971); Pleau et al. (2002), Moar et al. (2017), Clark and Boland (2016, Genective, pers. comm.). A commercial southern corn rootworm diet was used (Frontier #F9800B, Frontier Scientific Ltd., USA), but maize roots and food colour were added. This diet consists of D(+) sucrose, vitamin-free casein, cellulose, Wesson’s salt mix, methyl paraben fungicide, sorbic acid, cholesterol, raw wheat germ, Vanderzant’s vitamin mix, raw linseed oil, streptomycin sulfate antibiotic, and chlorotetacycline antibiotic. For 100 ml of diet, 13.8 g of the #F9800B diet was grinded and added to 88 ml fluid 60 to 70 °C agar (1.5 g agar CAS 9002-18-0, Chejeter, Japan in deionized water). After blending and cooling to 55 to 60 °C, 0.75 g grinded lyophilized maize roots were added (GLH5939 Pioneer, USA, or Phileaxx RAGT, Hungary) as well as 0.1 g green food colour for better larvae observation (Les Artistes, France). Thereafter, 1.7 to 1.8 ml 10% w/v KOH were added to reach a pH between 6.2 and 6.5. This mix was blended again, and then stirred at 50 to 55 °C. Then, 190 µl diet was pipetted into each 330 µl well filling each to around 2/3rd (repeater pipette P-8, Topscien Co., Ltd, China). Plates with diet were allowed to dry in a laminar flow cabinet during 45 min, and then stored at 3 to 5 °C overnight. The following day, treatments were applied. This is, 20 µl of a treatment were applied to the 0.34 cm² diet surface in each well (10 to 100 µl pipette, Biohit Proline, Finland). Order of treatments were shifted every other plate to avoid edge effects. Plates were dried for 1 to 1.5 h, and then cooled for 1 h in a 3 to 5 °C fridge.

Two weeks prior the bioassays, soil dishes with freshly laid eggs had been removed from D. v. virgifera adult rearing cages to allow sufficient incubation time until egg hatch. Eggs were washed with cool tap water with <0.01% NaOCl through a 300 µm mesh sieve. Around 5000 eggs were added to reach a pH between 6.2 and 6.5. This mix was blended again, and then stirred at 50 to 55 °C. Then, 190 µl diet was pipetted into each 330 µl well filling each to around 2/3rd (repeater pipette P-8, Topscien Co., Ltd, China). Plates with diet were allowed to dry in a laminar flow cabinet during 45 min, and then stored at 3 to 5 °C overnight. The following day, treatments were applied. This is, 20 µl of a treatment were applied to the 0.34 cm² diet surface in each well (10 to 100 µl pipette, Biohit Proline, Finland). Order of treatments were shifted every other plate to avoid edge effects. Plates were dried for 1 to 1.5 h, and then cooled for 1 h in a 3 to 5 °C fridge.

Data assessments and analyses

After 3 and 5 days of incubation, larval mortality, stunting, feeding and contamination were visually assessed through the clear seals using a stereomicroscope (10× magnification, SMZ-B4, Optec, Chongqing, China). Data from a plate were only used when the natural mortality threshold in the untreated control had not been reached, i.e. no more than 3 dead per 8 larvae per column of wells (37.5% threshold). This is in contrast to common practices in bioassays with other insects where the quality acceptance is usually <10% natural background mortality (Dulmage et al. 1990). However, this is rarely achievable with rootworm larvae as the artificial diets known to date remain suboptimal (Hibbard, University of Missouri, 2019, pers. comm.; Huynh et al. 2018).

Stunting was qualitatively assessed as an indicator for sublethal effects in comparison to the size and form of larvae in the untreated control. Feeding was assessed through observing food remains, frass, and diet in the larval gut to assure that diet and a treatment had been ingested. The coefficient of variation (CV) was determined in each bioassay as a measure of data precisions. A CV should ideally be <0.2, and at a CV of >2 further bioassays would be needed (Dulmage et al. 1990). In our experiments, the CVs of 1.2 for bioassay 1, 0.4 for bioassay 2, 0.7 for bioassay 3, and 0.8 for all bioassays, indicated good quality of data (Additional file 1: Table).

Larval data were compared between treatments within each experiment using Chi-Square statistics (because of nominal data type) with an fdr-correction of p-values (Benjamini and Hochberg 1995). To allow across-experiments comparisons, data were standardised to the untreated control data. Distributions of data were investigated using histograms, normal and detrended normal probability Q–Q plots and one-sample
Potted-plant greenhouse experiments

Experimental setup

To assess the efficacy of azadirachtin granules against *D. v. virgifera* larvae under semi-natural conditions, three systematic controlled trials were conducted using infested potted—maize plants in a greenhouse. As positive control served tefluthrin fine granules, cypermethrin microgranules and thiamethoxam—seed coating (Table 2). As negative controls served untreated infested plants as well as untreated uninfested plants. Each treatment was applied into the soil of three to four systematically arranged blocks (=replicates) of five pots. This totalled 15 to 20 data points (=sample size) per treatment per experiment.

In detail, each pot (plastic garden pot, 15 cm inner diameter × 10 cm height, 2 l) was first filled with 1 l sterilized soil. Two maize seeds were added (hybrid Szegedi 386, GK Hungary in experiment 1 and 3, or Futurixx, RAGT, France in experiment 2). Thereafter, 200 ml water were applied to each pot. Treatments were applied either as granules along a 2 cm wide strip across the 10 cm diameter of the pots, or as seed coating. Finally, 1/2 l soil was added burying the treatment and seed 3 cm into the soil leading to a soil surface of 14 cm diameter. The used soil contained 77% sand, 8% loam, 15% clay, 2.8% humus, 1.7% CaCO₃, 0.1% salts, and had a Ph of 7.7 (analysed by Szolnoki Talajvedelmi Laboratorium, Hungary). It had a soil bulk density 0.9 to 1.1 g cm⁻³ and a 7 to 11% soil moisture (γ% = grav.%). An average temperature of 20 ± 5 °C and a relative humidity of 97 ± 3% were recorded 5 cm deep in the soil in the pots as well as 24 ± 4 °C and 44 ± 13% in the air 1 m above the pots using climate data loggers (PeakTech 5185 data logger, Germany). Plants germinated between 4 and 12 days after sowing.

Maize pots were infested with 50 viable ready-to-hatch eggs per plant in experiment 1 and 3 or with 100 eggs in experiment 2. At this point in time, the majority of plants was at 3 leaf stage (height 15 to 20 cm). The eggs were applied in 0.15 to 0.2% aqueous agar with a standard pipette (1 to 5 ml, Eppendorf AG, Germany) in half-portions into two 50 mm deep holes 20 to 30 mm distant from both sides of the plant. A portion of eggs was incubated on moist filter paper at 20 °C in the laboratory to estimate emergence patterns (5 dishes with 10 to 20 eggs each/experiment). They revealed an emergence start 9 ± 5 days after placement. Hatching duration was 10 ± 4 days. Hatching rate was 47 ± 30, 47 ± 23% and 56 ± 19, leading to 24 hatched larvae per pot in experiment 1, 47 larvae in experiment 2, and 28 larvae in experiment 3, respectively. This indicates a medium, but consistent egg quality across experiments, and is comparable to similar studies of Xie et al. (1991).

Data assessments and analyses

Selectivity of the test agents was assessed by recording germination rate, plant phenology and phytotoxicity. Leaf number, plant height and the BBCH growth stage were assessed weekly as well as phytotoxicity according to Anonymous (2009).

At the expected second and early third instar stage, numbers of surviving larvae, root damage and above-ground biomass were assessed. This was 52 ± 18 days after planting and treatment, thus 40 ± 7 days after infestation. Each maize plant was pulled out of the soil, and gently shaken to remove loosely adhering soil particles from roots. Each maize plant was cut 1 cm above roots, and fresh weight, leaf number and plant height were measured. Then, the soil and root of each pot was placed onto a plastic screen for drying out and letting surviving larvae exit and drop onto the wet tissue paper in a tray below following a Berlese approach (Dent and Walton 1998). Larvae and their instars were counted 1, 3, 5 and 7 days later.

The untreated control was aimed to have a minimum level of infestation with 2nd or 3rd instar larvae of 20% to validate the results on agent efficacies. In all experiments, more than 90% of pots of the infested untreated second control yielded larvae. The infested control lead to 6 ± 5 s or third instar larvae/100 applied eggs.

One day after Berlese-placement, the dried roots were removed, gently shaken to remove remaining soil, soaked in water for 5 min, and then washed in 1% NaOCl and then water for 1 min to allow the assessment of root damage. Damage was rated using two scales recommended by EPPO (Anonymous 1999); this is, (1) the non-linear 1.0 to 6.0 traditional IOWA scale (Hills and Peters 1971) which slightly overestimates minor damage; and (2) the linear 0.00 to 3.00 node injury scale (Oleson et al. 2005) which measures only destroyed roots and therefore misses minor damage. To avoid subjective bias on these ratings, root damage was estimated independently by the experimenters, neither of whom knew whether the roots were from a treated or untreated pot.
Distributions of data were investigated using histograms as well as normal and detrended normal probability Q–Q plots (Kinnear and Gray 2000). Equality of variances was assessed using Levene’s test. Influences of treatments on assessed factors were analysed through GLM analyses or through independent samples Kruskal–Wallis H test. Tukey HSD post hoc multiple comparison tests were applied following GLM in case of equal variances, and Games Howell test in case of unequal variances. Logistic regression analyses were applied to assess the dose response of each treatment including the effective dose leading to 50% suppression of the larval populations or root damage prevention (ED50). The mean corrected efficacy of each treatment was calculated relative to the untreated control, this is corrected efficacy \( \% = 100 \times (\text{larvae or damage in control plots} - \text{larvae or damage in treated plots}) / \text{maximum (larvae or damage in control or treated plots)} \) (Toth et al. 2020). As the 1.0 to 6.0 IOWA root damage scale is a non-linear ordinal scale, and a value of 1 equals no damage, the damage data were converted to a 0.0 to 5.0 scale to estimate percent damage prevention across experiments. Results from azadirachtin treatments were validated in relation to the results from the corresponding positive controls of standard insecticides.

**Results**

**Control of neonates in artificial-diet based laboratory bioassays**

Azadirachtin appeared toxic to neonates of *D. v. virgifera* larvae (Figs. 1, 2, Additional file 1: Table). A clear dose–mortality response was observed. The corresponding fit of a logistic regression 3 days after treatment was: \( \text{larval mortality (3d)} = 1 / (1 + \exp(2.22 - 0.71 \times \ln(\text{dose})) \) (Chi-square test for ln of dose: \( p < 0.0001, df = 67; \) McFadden pseudo \( R^2 = 0.70 \), Fig. 1). Accordingly, the 3-day LD50 of azadirachtin was estimated 22.3 µg active ingredient (a.i.)/ml (CI 95% 8.8–56 µg a.i./ml). This corresponds to 1.34 µg a.i./cm² treated surface, and to 0.45 µg a.i./20 µl/larva. The 3-day LD90 was 480 µg a.i./ml (CI 95% 92–2742 µg).

The dose–response did not change much from day 3 until day 5. The 5-day LD50 was 19.3 µg a.i./ml (CI 95% 7.3–51.2 µg) according to the logistic regression fit: \( \text{larval mortality (5d)} = 1 / (1 + \exp(1.99 - 0.67 \times \ln(\text{dose})) \) (Chi-square test for ln of dose: \( p < 0.0001, df = 67; \) McFadden pseudo \( R^2 = 0.70 \), Fig. 1). Accordingly, the 5-day LD50 of azadirachtin was estimated 22.3 µg active ingredient (a.i.)/ml (CI 95% 8.8–56 µg a.i./ml). This corresponds to 1.16 µg a.i./cm² treated surface, and to 0.39 µg a.i./20 µl/larva. The 5-day LD90 was 502 µg a.i./ml (CI 95% 94–2746 µg).

No sublethal effects of azadirachtin such as stunting of larvae were observed (ANOVA for logarithmic model: \( F_{1,41} = 1, p = 0.31, \) adjusted \( R^2 = 0.001)\).

**Control of larvae and prevention of root damage in potted -plant greenhouse experiments**

Azadirachtin treatments at increasing dose reduced the larval survival on the maize roots (relative to control, GLM, \( F_{5,30} = 17.6, p < 0.0001, \) adjusted \( R^2 = 0.73 \)).

Multiple comparison tests revealed efficient control of *D. virgifera* larvae on maize roots by standard doses of tefluthrin and thiamethoxam, but not by standard doses of cypermethrin and azadirachtin (Figs. 3, 4). When increasing doses of azadirachtin, control of larvae became evident. Between 25 and 67% of larvae were killed by a 5× standard-dose (200 g a.i./ha), and 100% control efficacy was reached at a 10× standard-dose (380 g a.i./ha). Doubling the standard dose of cypermethrin did not improve its efficacy in reducing larvae numbers.

The corresponding logistic regression fit of efficacy to different doses of azadirachtin was: \( \text{efficacy in reducing larvae (%) = 1/(1 + \exp(2.203 - 1.14 \times \ln(\text{dose})) \) (Chi-square test for ln of dose: \( p = 0.0027, df = 19; \) McFadden pseudo \( R^2 = 0.43 \), Fig. 3). Accordingly, the ED50 of azadirachtin was 6.9 mg active ingredient/meter of maize furrow (CI 95% 2.6–18.5 mg). This corresponds to approximately 92 g azadirachtin/hectare. The ED90 was 47.7 mg active ingredient/meter of maize (CI 95% 5.2–430 mg).

Reduction of larvae through treatments was only partly reflected in the level of prevention in root damage (Figs. 5, 6). Azadirachtin treatments at increasing dose
improved the level of prevention in root damage (relative to control, GLM, $F_{5,30} = 29.5$ for general root damage, $22.5$ for heavy root damage, adjusted $R^2 = 0.83$ and $0.78$; $p < 0.0001$).

Multiple comparison tests revealed that standard dose of tefluthrin and thiamethoxam consistently prevented the overall as well as heavy root damage (Fig. 4). The standard dose of cypermethrin only inconsistently prevented some of the root damage and the standard doses of azadirachtin was usually not sufficient. When increasing the doses of azadirachtin, prevention of root damage became evident. About 40% of the overall root damage and 67% of the heavy root damage were prevented by a 5× standard-dose (200 g a.i./ha). Root damage was nearly entirely prevented by a 10×-standard-dose (380 g a.i./ha).

The corresponding logistic regression fit of root damage prevention to different doses of azadirachtin was: 

$$\text{efficacy in preventing general root damage (\%) = 1/(1 + e^{2.39 - 0.85 \times \ln(dose)})}$$

(Chi-square test for ln of dose: $p = 0.0075$, df $= 19$; McFadden pseudo $R^2 = 0.55$, Fig. 3). Accordingly, the $ED_{50}$ of azadirachtin was 16.5 mg active ingredient/meter of maize furrow (CI $95\%$ 4.2–65.9 mg).

This corresponds to approximately 220 g azadirachtin/
hectare. The ED90 was 218 mg a.i./meter of maize furrow (CI 95% 9.–5277 mg).

The logistic regression fit of preventing heavy damage to different doses was:

\[
\text{efficacy in preventing heavy root damage (\%)} = \frac{1}{1 + \exp(1.397 - 1.016 \times \ln(\text{dose}))}
\]

(Chi-square test for ln of dose: \( p = 0.011, \text{df} = 19; \text{McFadden pseudo } R^2 = 0.39 \), Fig. 3). The corresponding ED50 of azadirachtin was 3.96 mg active ingredient/meter of maize furrow according to the logistic model fit (CI 95% 1.4–10.8 mg). This corresponds to approximately 52 g a.i./ha. The ED90 was 34.4 mg active ingredient/meter maize furrow (CI 95% 3–392 mg).

Preventing root damage through certain treatments was only little reflected in yield-related parameters. When differences were found between treatments and the untreated control, then their absolute differences were small.

Whilst the standard dose of azadirachtin and up to 2.8 mg a.i./plant/pot (28 mg/m furrow) did not improve biomass of 6 to 10 leaf stage maize, a high dose of 28 mg azadirachtin (280 mg/m) improved biomass (8.3 ± 2.7 g versus 3.9 ± 2 of infested control plants, GLM, \( F_{5,30} = 16 \), adjusted \( R^2 = 0.72; p < 0.0001 \)). The standard dose of tefluthrin also improved biomass in one of three experiments, but no such improvements were detected by cypermethrin or thiamethoxam.

Whilst the standard dose of azadirachtin and up to 1.5 mg a.i./plant/pot (15 mg/m furrow) did not improve height of maize, a high dose of 2.8 mg azadirachtin (28 mg/m) or even 28 mg (280 mg) increased plant height (49 ± 11 cm or 50 ± 11 cm versus 36 ± 15 cm of infested control plants, GLM, \( F_{5,30} = 3.9 \), adjusted \( R^2 = 0.33; p = 0.009 \)). The standard doses of tefluthrin,
cypermethrin or thiamethoxam did not affect plant height in none of the experiments. None of the treatments and applied doses did increase average leaf numbers of 6 to 10 leaf stage maize, except of a small positive effect of thiamethoxam in experiment 2 (9.3 ± 0.9 versus 8.5 ± 0.7 leaves of infested control plants).

Treatments did not affect germination rates (GLM, $F_{12;76} = 1.25; p = 0.27$) and did not lead to any delay in germination ($F = 0.8; p = 0.58$) regardless of azadirachtin or synthetic pesticides and regardless of the different doses being applied. Those treatments also did not cause any phytotoxic effects such as yellowing, chlorosis, necrosis or deformation of leaves or stunting of plants.

Discussion

Our sets of laboratory-bioassays as well as potted-plant experiments confirmed Xie et al. (1991) that the neem plant-derived azadirachtin is toxic to larvae of *Diabrotica v. virgifera*, one of the key pests among rootworms. This is not surprising as azadirachtins are of broad-spectrum activity (Dougoud et al. 2019). Positively, an application of this botanical insecticide as a granule into the sowing furrow can lead to a suppression of the later hatching larvae and to a significant prevention of root damage. Granule applications at sowing are preferred by many maize growers over fluid applications or applications later in the maize growing season (Toepfer et al. 2010). This should be of high interest to industry because of recent bans on the use of a number of soil insecticides and insecticidal seed-coatings due of their either high human toxicity and/or serious non-target effects or other environmental concerns (World Health Organization 2009; European-Commission 2011; Georgiadis et al. 2011).

Consequently, growers in numerous countries are left with few or no management options for soil insect pests in field crops. However, soil insects, such as corn rootworms (*Diabrotica* spp.), cutworms (*Agrotis* spp.), wireworms (*Agriotes* spp.) or grubs (*Melolonthidae*) account for a large proportion of below-ground damage to maize, and the latter two pest groups also to a number of other crops (Toepfer et al. 2014). In Hungary for example, 46% of all the 5000 tons of insecticides sold in 2019 were the granular soil insecticide tefluthrin (Demeter and Lazar
This indicates the high importance of such insecticides for the control of soil insect. However, tefluthrin will probably be retreated from the pesticide markets due to its high toxicity being classified as a WHO-class Ib acutely hazardous ingredient (World Health Organization 2009). Similarly, neonicotinoids have already been phased-out in many countries due to their pollinator toxicity, accumulation in the soil and other environmental effects (Georgiadis et al. 2011). Agri-policies try to address the public concerns with regard to pesticides and try to promote alternative and safer pest management solutions. Examples of such attempts are the directive on “Sustainable Pesticide Use” of the European Union (European Commission 2009) or the “Green Pest Control Policy” of China (Fan 2006; MoA 2011). However, such policies do not necessarily lead to new pest control options, as the discovery and development of novel plant protection agents with new modes of action and low environmental impact is difficult and costly.

We have therefore investigated whether the safe and easily biodegradable botanical insecticide azadirachtin (Boeke et al. 2004), which is widely used against above-ground insects (Saxena 1989), might become an option for the control of soil insect pests. First, we confirmed through laboratory bioassays that the azadirachtins in the here-tested granular formulation were similarly effective as the ones in commonly used fluid formulations. This was necessary because neem products can be variable in their contents of active ingredient(s) as well as in their efficacy for pest control (Stark and Walter 1995; Dougoud et al. 2019). Therefore, the provider of the neem granules had run gas chromatographic analyses prior to our experimentation confirming 0.165% azadirachtin as labelled (mainly A with some B) as well as 0.35% other neem compounds (Aza-F, Aza-H, Aza-L, salanin, nimbins and fatty acids). This is important, because in many neem products, the ingredients are not well assessed and/or declared on the product label (Dougoud et al. 2019). In our bioassays, the dose–response curves of azadirachtin from the granular formulation did not differ from those in common fluid formulations (Fig. 2), indicating a good quality of the test products and correctness of label information.

Our artificial diet-based bioassays on neonates of D. v. virgifera larvae revealed a 3-day LD50 of 22.3 µg azadirachtin/ml which corresponds to 0.45 µg/neonate. Azadirachtin appeared of relatively fast mode of action on D. v. virgifera as the dose–response did not change much from day 3 to day 5. The 5-day LD50 was 19.3 µg/ml which corresponds to 0.39 µg/neonate up to early second instar larva. Also, Xie et al. (1991); Stark and Rangus (1994) and a number of other authors suggested that azadirachtin has contact activity on insects in addition to its widely reported systemic and chronic modes of action. There are a number of LD50 reported for immature stages of several insect groups, such as 77 µg/ml for first instar Ectomyelois spp. (Lepidoptera) over 5 days and 438 µg/ml over 1 day (Mehaoua et al. 2013), 7.6 to 7.7 µg/ml for fourth instars of two Culex spp. (Diptera: Culicidae) with not-reported exposure time (Merabti et al. 2017), and 2.8 µg/ml for young Aphis spp. (Hemiptera: Aphididae) over 7 days (Stark and Rangus 1994). Ladd et al. (1984) reported a low LD50 of only 0.1 µg of topically applied azadirachtin per third instar Popillia japonica (Coleoptera: Scarabaeidae), but over an exposure of 20 days. This suggests that our reported LD50 by 0.45 µg within 3 days or 0.39 µg within 5 days for the much smaller neonates of D. v. virgifera would be much lower when assessing mortality over longer periods. In general, it appears difficult to compare LD50 across studies and insect species due to different experimental setups particularly the involved insect food, exposure periods, different insect weights, and sometimes unclear compositions of azadirachtins and related compounds in the test agents. Despite the often reported LD50 as µg/ml, some of those studies do not report the amount of azadirachtin applied per larva or per insect weight. For example, Xie et al. (1991) reported a 3-day LD50 of only 3.9 (2.5 to 5.9) µg azadirachtin/ml when applying 1.7 ml of the 3.9 µg/ml-solution onto filter paper in a Petri dish with 10 neonates of D. v. virgifera and a maize seedling, thus effectively 6.6 µg. The reported effective dose seems low compared to our data, and reasons are difficult to explain. On one hand, the experimental arrangements of Xie et al. (1991) with filter paper assays in Petri dishes differs to our approach of exposing one larva to azadirachtin on artificial diet, on the other hand their sample size was low (5 Petri dishes only). Qadri and Narsaiah (1978) reported a 1-day LD50 of 1.5 mg azadirachtin/gram body mass of older nymphs of Periplaneta spp. (Blattodea), but there is no such information for other Diabroticina. Our tested D. v. virgifera larvae weighted about 0.42 ± 0.23 mg (across neonates and young second instars). This would roughly correspond to an LD50 of 0.9 to 1.1 mg azadirachtin/gram D. v. virgifera larva, being comparable with the LD50 on Periplaneta. Astonishingly, we did not detect any sub-lethal effects of azadirachtin to D. v. virgifera larvae such as stunting or moulting inhibition. Some authors reported growth inhibitor and deformation effects to insects, such as to Culex larvae (Al-Sharook et al. 1991) or Aphis nymphs (Stark and Rangus 1994). Landis and Gould (1989) reported anti-feeding effects to larvae of Diabrotica undecimpunctata howardi. Although, in our bioassays some larvae of D. v. virgifera moulted to 2nd instar suggesting no inhibitor effects, our assay duration of 5 days might not have been long enough to detect all
possible sublethal effects. Ladd et al. (1984), for example, detected insect growth regulator effects on *Popillia* grubs only within about 20 days after treatment. As for rootworms, however, longer assays are rarely achievable as the known artificial insect diets are suboptimal and contamination rapidly occurs due the non-sterile nature of the available diets (Hibbard, University of Missouri, 2019, pers. comm.; Huynh et al. 2019). Nevertheless, the here-reported LD$_{50}$ levels of azadirachtin of *D. v. virgifera* seem in line with lethal effects to other insect groups, and therefore warranted further investigation.

In a second step, we simulated the efficacy and feasibility of an azadirachtin-based granule application into the soil for corn rootworm control using potted-plant experiments in comparison to standard pesticides. Results showed that standard doses of thiamethoxam-seed coating and tefluthrin-granular soil insecticides applied at maize sowing can well suppress larval populations of *D. v. virgifera* and prevent most root damage (Pilz et al. 2009; Rozen and Ester 2010; Modic et al. 2018; Souza et al. 2020). In contrast, a cypermethrin granular soil insecticide applied at its standard dose at sowing appeared much less effective, and even doubling its dose did not improve efficacy. This confirms variable experiences with cypermethrin-based soil insecticides for rootworm control in field studies (Toth et al. 2020). Unfortunately, also the currently proposed standard dose of 38 g azadirachtin/hectare for granular in-furrow application at sowing appeared not enough to control *D. v. virgifera* or to prevent root damage. However, when increasing doses of azadirachtin, control of larvae and prevention of root damage became evident. At a 5× standard-dose of azadirachtin (200 g/ha), 25 to 67% percent of larvae were killed, about 40% of overall root damage prevented as well as about 67% of heavy root damage. This, as well as the modelled ED$_{50}$ are comparable to or better than the efficacy of most cypermethrin applications. At a 10× standard-dose azadirachtin (380 g/ha), total pest control was achieved as well as the prevention of most root damage. Whether this would be economically feasible is not yet clear. However, such a control efficacy is even better than the efficacies of most applications of tefluthrin granules and comparable to the efficacy of thiamethoxam seed-coatings. This confirms Xie et al. (1991) who applied high doses of azadirachtin as a drench into the sowing furrow in potted-plant trials. This also indicates that both fluid and granular applications into the furrow at maize sowing would lead to control efficacies. As into-furrow applications of agents in maize seem to have little impact on non-targets (limited area treated and below ground) (Babendreier et al. 2015), it is also unlikely that applications of higher concentrations of azadirachtin are environmentally problematic (Boeke et al. 2004). This is also underlined by the biodegradable nature of azadirachtin. Interestingly our experiments showed a slight positive effect of higher doses of neem-granules on plant height and biomass in comparison to similarly effective chemical treatments. Although Xie et al. (1991) did not observe such effects, it may potentially confirm the biofertilizer properties of neem in maize as suggested by Vageesh et al. (2016).

**Conclusion**

In conclusion, there seems clear potential for the development of a neem-based botanical soil insecticide if the required higher concentrations of azadirachtin in the granule, or potentially fluid formulation can be achieved. The currently suggested standard dosage of 38 g azadirachtin/hectare corresponds to 25 kg granules/hectare. Many commercial applicators for fine granules on sowing machines may deliver not more than 20 kg hectare. Therefore, a higher concentration of azadirachtin in less weight of granules would be needed if the application for rootworm control at larger field scale was to become reality. In a next step, larger open field trials using farmer machinery are suggested towards the development of a practical and effective neem-based soil insecticide for corn rootworm control in maize. Those, experiments should clarify whether for example 50, 200, 380 g or more azadirachtin would be needed/hectare under real farming conditions to suppress *D. v. virgifera* larval populations below threshold and to sufficiently prevent root damage to avoid yield losses. If this is achieved, such product(s) may become a promising, safer alternative in the management of rootworms such as *D. v. virgifera* and potentially other soil insect pests. This could become a replacement of some of the banned soil insecticides or insecticidal seed coatings, and will ultimately help diversify the currently limited integrated pest management toolbox.

**Supplementary Information**

The online version contains supplementary material available at https://doi.org/10.1186/s43170-021-00044-9.

**Additional file 1.** Dose-efficacy response of azadirachtin in killing neonates of *Diabrotica v. virgifera* in artificial diet-based bioassays of 96-well plates under standardised semi-sterile laboratory conditions.

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Authors' contributions
All authors have substantially contributed to conception and design, analysis and interpretation of data, as well as to drafting and revising the article. All approved the final version to be published and agreed to be accountable for all aspects of the work. In addition to above, SZT and ST implemented the experiments and collected the data, MSz and ST conducted the statistical analyses, and ST supervised the study and finalised the manuscript. All authors read and approved the final manuscript.

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Availability of data and materials
Meta-data are available in an additional file to the publication. Raw data are deposited in Zenodo (https://doi.org/10.5281/zenodo.4318642). Any other information is available upon request to the corresponding author.

Declarations

Ethics approval and consent to participate
All applicable international, national, and/or institutional guidelines for the care and use of animals were followed. This article does not contain any studies with human participants or vertebrates performed by any of the authors. Informed consent was obtained from all individual participants included in the study.

Consent for publication
All authors agree with this publication.

Competing interests
The authors declared no conflicts of interest.

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