Comparison of theoretical and experimental values for plant uptake of pesticide from soil

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Abstract

Pesticides that persist in soils may be taken up by the roots of plants. One way to assess plant uptake is to theoretically predict the extent of plant uptake using a mathematical model. In this study, a model was developed to predict plant uptake of pesticide residues in soils using various parameters, such as pesticide mobility within soil, plant transpiration stream, root–soil transfer rate, plant growth, and pesticide dissipation in either soils or plants. The accuracy of the model was evaluated by comparing the modeled concentrations with measured uptake concentrations of chlorpyrifos (CP) in lettuce, grown on treated soils with concentrations of approximately 10 and 20 mg kg\(^{-1}\) CP. Measured concentrations of CP in lettuce at 21, 30, and 40 d after planting were between the 5\(^{th}\) and 95\(^{th}\) percentiles of model variation. A high correlation coefficient of > 0.97 between modeled and measured concentrations was found. Coefficients of variation of mean factors to residual errors were between 25.3 and 48.2%. Overall, modeling results matched the experimental results well. Therefore, this plant uptake model could be used as an assessment tool to predict the extent of plant uptake of pesticide residues in soils.

Introduction

Many studies have shown that theoretical prediction from mathematical models can assess the extent of plant uptake of hazardous substances, such as heavy metals and organic pollutants persistent in soil [1–6]. However, unlike heavy metals, it is difficult to develop a plant uptake model for organic chemicals, as their dissipation behaviors might be consistent or variable, depending on environmental conditions. Hence, plant uptake models tend to show low availability for organic contaminants. Nevertheless, regulatory authorities in several countries have utilized plant uptake models as a chemical exposure assessment tool, for examples, Soil Screening Levels (SSLs) in USA, Contaminated Land Exposure Assessment (CLEA) in UK, and CSOIL in Netherlands [7–9].

Plant uptake models for pesticides that persist in soils are based mainly on a bioconcentration factor (BCF) that indicates the ratio of pesticide concentrations in the plant and soil [10]. Shone and Wood [11] introduced the root concentration factor (RCF) to demonstrate the relationship between concentrations of pesticides in root and soil solutions. Thereafter, they defined the transpiration stream concentration factor (TSCF) that indicates the ratio of
pesticides transferred from soil solution into the xylem sap of the plant [11,12]. Briggs et al. [13] found a high correlation between RCF or TSCF factors and octanol–water partition coefficients ($K_{ow}$) of pesticides, and these correlations have been mainly considered in studies predicting the dynamic plant uptake of non- and weak-electrolytes as well as acids and bases [3,13].

TSCF has frequently been used in studies of the plant uptake model [14–16]. Charles [17] indicated that TSCF is an important concept for predicting the fraction of pesticides transferred from the roots to various compartments of the plant, such as stems, leaves, and fruits, by the transpiration stream. Moreover, RCF has been occasionally used to simulate the uptake and translocation of pesticides in fruit trees that have very thick fine roots [18]. Unusually, Felizeter et al. [19] used the RCF to model the uptake of acidic chemicals by lettuces, grown under hydroponic conditions.

When modeling the dynamic plant uptake of pesticides persistent in soil, it is necessary to consider adsorption and dissipation interactions between pesticide and soil [20]. These interactions are closely related to the mobility and persistency of pesticides in soil. Strong adsorption of pesticides on soil particles can result in lower soil mobility and reduced plant uptake, and more persistent pesticides may provide opportunities for more consistent uptake by plant roots. Our previous study showed that both the adsorption and dissipation interactions can be used as major parameters to model the residual magnitudes of pesticides in soils [20].

Dissipation of pesticides in plant is also a necessary consideration for modeling the fractions of reduction after uptake by the plant [6]. Dissipation routes of pesticides in plants may include removals by the plant metabolism and reversed translocation from roots to soil. In addition, the dilution of pesticide concentration during plant growth can contribute to dissipating pesticides in plants [21–24]. Juraske et al. [15] modeled the extent of plant uptake using a parameter that described the fraction of pesticide that is reflected back from roots to soil, opposite to the transpiration stream.

The organophosphorus insecticide chlorpyrifos ($O,O$-diethyl $O$-3,5,6-trichloro-2-pyridy phosphorothioate, CP) has been widely used to control various pests occurred in agricultural industry [25]. Although half-lives of CP in soils are variable in the range of 1–100 d [26], the three chlorines in its structure can result in long-term persistence of CP in soils [20,27]. In addition, the low water solubility (1.4 mg L$^{-1}$) and high log $K_{ow}$ (4.7) values of CP may result in insignificant mobility in soil [28–30]. Therefore, it is likely that CP residue in arable soils is exposed to plant uptake, presenting a safety issue with final agricultural products.

This study aimed to develop a plant uptake model of CP from soil using parameters such as pesticide mobility in soil, plant transpiration stream, root–soil transfer rate, plant growth, and pesticide dissipation in either soils or plants. The accuracy of the developed model was statistically assessed by comparing the modeled estimates with measurements obtained from field experiments.

Materials and methods

Development of plant uptake model

Model approach. Pesticides in soils undergo either adsorption on soil particles and organic matters or dissipation by biotic and/or abiotic factors. Pesticide residues that do not undergo adsorption of dissipation in soil are present in soil solution, after which they are absorbed by plant roots with water uptake system. Hereafter, a fraction of absorbed pesticide is reflected back to the soil by the soil–root advection interaction and the resistance of waxy material (Casparian strip), covering the endodermis of the root [17]. Pesticide residues crossing the endodermis are transferred to aerial parts of the plant through xylem vessel. Downward
translocation through the phloem of the plant cannot be included for non-ionic chemicals, such as pesticides, as there is no mechanism that they can retain them in phloem sap [1,31]. Finally, the uptake concentration of pesticides in plants decreases over time with the metabolism and growth of the plant. Taken together, the sequential steps for developing the dynamic plant uptake model in this study were: the dissipation and migration of pesticide in soil; pesticide residues in soil solution; root uptake of pesticide by the transpiration stream; and decrease in pesticide concentration in the plant.

**Concentration in soil solution.** Modeling of plant uptake of pesticide residues in soils was initiated at the point immediately after crops were planted. Based on our previous study [20], the concentration of pesticide in soil at time \( t \) (\( C_e(t) \)) was calculated as the concentration in soil solution (Eq 1):

\[
C_e(t) = C_0 \times (1/2)^{t/T_d}
\]

where \( C_0 \) is the initial pesticide concentration in soil (\( \mu g \) g\(^{-1} \)), \( T \) is the half-life of pesticide in the soil (days), and \( K_d \) is the distribution coefficient of pesticide in soil and soil solution.

**Pesticide concentration in the plant.** The transpiration stream concentration factor (TSCF), which represents the ratio of pesticide concentrations in plant xylem sap and soil solution, is described by Charles [17] and Juraske et al. [15]. TSCF of non-ionic pesticide was estimated using eq 2:

\[
TSCF = 0.756 \times \exp\left[\left(-\log K_{ow} - 2.50\right)^2/2.58\right]
\]

where \( K_{ow} \) is the octanol-water distribution coefficient of pesticide, which is obtained using the eq 3, described by Sabljic et al. [32]:

\[
\log K_{oc} = 0.81 \times \log K_{ow} + 0.1
\]

where \( K_{oc} \) is the partition coefficient between organic carbon and water. TSCF corresponds to the fraction of pesticide absorbed into the plant with the transpiration stream [15]. However, to allow for the fraction reflected back by the endodermis in roots, transfer rate from soil solution to the plant (\( k_{s-p} \)) was specified in eq 4:

\[
k_{s-p} = Q_w \times (1 - TSCF)
\]

where \( Q_w \) is the plant transpiration stream (m\(^3\) day\(^{-1}\)) and the reflected fraction is expressed as ‘1 -TSCF’. A fraction of pesticide absorbed by the plant can be removed by plant. Removal rate of pesticide from the plant (\( k_{rp} \)), based on first order kinetics, was described with eq 5:

\[
k_{rp} = \ln(2)/T_p
\]

where \( T_p \) is the half-life of pesticide in plant. The time-dependent change in plant weight, representing pesticide concentration diluted by plant growth, (\( M_p(t) \)) was calculated using eq 6 (\( t > 0 \)):

\[
\log M_p(t) = I_g \times \exp[k_g \times \log t]
\]

where \( I_g \) and \( k_g \) indicate the logarithmic plant weight at initial time 0, and a plant growth constant, respectively. The final plant uptake model developed in this study was specified by eq 7:

\[
C_p(t) = \frac{C_e(t) \times k_{s-p} \times (1 - \exp[-k_p \times t])}{(k_{rp} \times M_p(t))}
\]
Uptake experiments

Field trials. Three-weeks-old lettuce plug seedlings (Red Lollo Rosa cultivar) were purchased from a commercial plant nursery in Changnyeong, Korea. Uptake experiments were conducted in a greenhouse (2,092 m² in area) on a lettuce farm in Waegwan (WG), Korea, between April 15 and May 25, 2014. Commercial CP (25% of wettable powder), diluted with 5 L of water was sprayed on one experimental plot (n = 3) of 100 × 500 cm using a shoulder-type compression sprayer, equipped with a 1-mm nozzle (KS-10-3, Kwang Sung Co., Daejeon, Korea); no additional pesticides were sprayed throughout the experimental period. Concentrations of CP were approximately 10 and 20 mg kg⁻¹, corresponding to the low (LC) and high concentration (HC) treatments, respectively. Treated soils were homogenized to a depth of 10 cm and aged for 12 h before planting lettuce plug seedlings. The seedlings were planted at intervals of 10 cm. Water was supplied to the seedlings at a rate of 1.7 L/h for 20 min, every 5 d using an overhead sprinkler system. Conditions in the greenhouse were maintained at 23.1 ± 3.13˚C with 60.3 ± 4.74% humidity. A control experimental plot treated with pesticide, but with no plants was prepared at the same time.

Preparation of plant and soil samples. Leaves and roots of lettuce plants (n = 20) were harvested from each experimental plot at 21, 30, and 40 d after the pesticide treatment. Roots were rinsed with running water to remove soil residues and lightly wiped with paper towel. Lettuce weights were measured at each sampling to obtain \( M_p(t) \) values. Lettuce plants were divided into two compartments, roots and leaves that were individually homogenized using a grinder and stored in -20˚C freezer (GC-124HGFP, LG Electronics Inc., Seoul, Korea) prior to pesticide residue analysis.

Soil samples were collected from each experimental plot at 0 (12 h), 7, 14, 21, and 40 d after pesticide treatment, air-dried for 5 d, and passed through a 2-mm sieve. A portion of the samples was used to analyze of soil properties (S1 Table). Water content of soil samples was measured by comparing the change in weight from 24 h of oven-drying, and used to correct the residual concentration of CP in soil.

Pesticide residue analysis. CP residues from soil and lettuce samples, weighing 10 g each, were extracted with 80 mL of acetone. Soil samples were shaken at 200 rpm for 30 min using a shaking incubator (Vision Scientific Co., Ltd., Daejeon, Korea), and lettuce samples were homogenized at 12,000 rpm for 3 min using a grinder (AM-7, Nihonseiki Kaisha Ltd., Tokyo, Japan). Extracts of each sample were filtered through a Büchner funnel lined with filter paper (Whatman No. 2, Buckinghamshire, UK) and transferred into a separatory funnel containing 500 mL water, 50 mL of saturated sodium chloride solution, and 50 mL of methylene chloride. The funnel was shaken vigorously, and the organic solvent fraction was collected after dehydration with anhydrous sodium sulfate. The extraction step was repeated with the remaining residue in the funnel and a further 50 mL of methylene chloride. Organic solvent extracts were combined in one flask and concentrated using a rotary vacuum evaporator (Laborota 4000, Heidolph Instrument GMBH & Co., Schwabach, Germany) at 40˚C. To purify, concentrates were re-dissolved with 10 mL of \( n \)-hexane and loaded into a glass column (16 mm i.d., 30 cm height), packed with 10 g of Florisil. Impurities in samples were removed with 60 mL \( n \)-hexane, and the fraction of pesticide was eluted with 60 mL of ethyl acetate/\( n \)-hexane (95/5, v/v). The final eluate was evaporated, dissolved with 2 mL of acetone, and analyzed using a gas chromatography-mass spectrometer (GC-MS; Shimadzu GC 2010 equipped with a GC-MS QP-2010 Plus, Kyoto, Japan). All analysis of CP residues in soil and lettuce samples were based on a matrix matched calibration (MMC) method. S1 File describes the quality control procedure for the abovementioned analysis, MMC method, and analytical conditions of GC-MS.
The half-life \((T)\) of CP in soil was calculated with the residual data, assuming the first order exponential dissipation (Eq 8):

\[
T = \frac{\ln(2)}{k_{deg}}
\]  

Laboratory experiments

To obtain the \(K_d\) value, adsorption experiments of CP in soil followed procedures described previously [20]. Transpiration stream \((Q_w)\) of lettuce was measured using a potometer, which can record the volume of water uptake in plant. All methods for laboratory experiments are summarized in S1 File. The \(K_d\) and \(Q_w\) values obtained from these experiments were used as parameters in the model.

Assessment of model accuracy

Accuracy of the plant uptake model was assessed by statistically comparing between modeled and measured concentrations. Based on the model assessment method by Juraske et al. [5], we evaluated the mean of the correlation coefficients \((R^2)\) and the error of residuals (ER), known as the standard deviation of the log of residuals between the modeled and measured values.

Estimated CP concentration in leaves

The ratio of CP concentrations in leaves to those in the whole plant \((R_{L/W})\) was calculated using data from the uptake experiments (Eq 9):

\[
R_{L/W} = \frac{C_L}{C_W}
\]  

where \(C_L\) is CP concentration in leaves (mg kg\(^{-1}\)), and \(C_W\) is CP concentration in the whole plant (mg kg\(^{-1}\)). The \(R_{L/W}\) values were used to estimate the CP concentrations in lettuce leaves from the modeled results.

Results and discussion

Quality control

Total ion chromatogram and mass spectrum of CP identified by GC-MS are shown in S1 Fig. The mass spectrum of CP shared > 93% similarities with the mass spectral library data provided by the National Institute of Standards and Technology (NITS). The most intensive target ions (m/z 258 and 314) were used for the selected ion monitoring (SIM) analysis, which were free from fragment ions that appeared in blank soil and lettuce samples. Chromatograms for recovery tests are shown in S2 Fig. Peaks of CP in soil and lettuce that spiked at a concentration of 1.0 mg kg\(^{-1}\) showed clear shape and selectivity. In addition, there were no interfering substances in CP residue analysis. The linearity of calibration curves in the MMC method was acceptable, with correlation coefficients of >0.99. The minimum detectable amount (MDA) and limits of quantitation (LOQs) of CP were 0.1 ng and 0.02 mg kg\(^{-1}\), respectively. Recovery rates of CP that spiked at concentrations of 0.2 and 1.0 mg kg\(^{-1}\) in each sample were satisfactory at 88.1–93.5%, and relative standard deviations (RSD) were <7.2% (S2 Table). Therefore, this method of pesticide residue analysis was able to determine the CP residues from soil and lettuce samples.
Adsorption and dissipation in soil

S3 Fig shows the adsorption behaviors of CP in WG soil. Adsorption of CP in the soil reached equilibrium after 24 h (S3a Fig), and the adsorption isotherm at that time was dependent on the treatment concentrations of CP (S3b Fig), showing a type C curve, as described by Giles et al. [33]. The \( K_d \) values of CP in the soil, calculated from the adsorption isotherm was 82.1 mL g\(^{-1}\) and ranged from 13.4 to 1862 mL g\(^{-1}\), as reported by Moore et al. [34].

Dissipation patterns of CP in soils collected from field experiments are illustrated in S4 Fig. Actual concentrations of CP in soils determined instrumentally at time 0 were 15.2 and 24.9 mg kg\(^{-1}\) in the LC and HC treatments, respectively, which were slightly higher than the nominal concentrations of 10 and 20 mg kg\(^{-1}\) that were expected when treating the pesticide. However, these actual concentrations were sufficient to determine the concentration dependency of pesticide uptake in lettuce. Concentrations of CP in soil at time 0 were used as an initial soil exposure concentration (\( C_0 \)) for plant uptake model. Soil half-lives (\( T \)) of CP in the LC and HC treatments were 17.2 and 7.9 d, respectively, and the dissipation rate was approximately twice as fast in the HC treatment as in the LC treatment. The faster dissipation of CP in the HC treatment implies less opportunity for root uptake of CP. Experimental values of adsorption (\( K_d \)) and dissipation (\( T \)) were used as parameters in the plant uptake model.

Uptake experiment

Measured uptake concentrations of CP in lettuce are shown in Table 1. In the LC treatment, 0.9 mg kg\(^{-1}\) of CP residue was absorbed from soil by lettuce after 21 d of growth, which decreased by 0.2 mg kg\(^{-1}\) (77.8%) after 40 d. Uptake concentration of CP in lettuce in the HC treatment was similar, at 0.8 mg kg\(^{-1}\) after 21 d of growth and declining to 0.1 mg kg\(^{-1}\) (87.5%) at the final sampling time. These results show that the extent of uptake of CP from soil by lettuce was not dependent on the concentration of pesticide in the soil.

Distribution patterns of CP in each compartment of lettuce differed between the LC and HC treatments. In the LC treatment, 50.9–56.1% of the uptake amount was constantly present in leaves throughout the experimental period (Fig 1a). However, the distribution rate in the HC treatment was 56.9% at 21 d of growth period and increased to 86.0% after 40 d of growth (Fig 1b). These results show that the significant amounts of CP can be transferred from roots to edible leaf parts, which may be due to the active growth of leaf parts during the experimental period (S5 Fig). Similar to our findings, Jeon et al. [35] reported the high concentrations (1.36–4.71 mg kg\(^{-1}\)) of boscalid and chlorfenapyr from soil in Korean cabbage leaves. The \( T_p \)
values in the LC and HC treatments were calculated at 8.7 and 6.3 d, respectively, and were used as model parameters to describe the removal rate of CP in lettuce.

Comparison between modeled and measured data

Parameters of the plant uptake model are listed in Table 2. Most of the parameters were obtained from the laboratory experiments, except for half-life values of CP in soil and lettuce. Logarithmical correlation equations between the weight and growth time of lettuce were satisfactory, with correlation coefficients of >0.99, and the $I_g$ and $K_g$ constants in the equations were used as model parameters to calculate $M_p(t)$ values.

Uptake concentrations modeled using these parameters were compared with concentrations measured in the uptake experiments. As shown in Fig 2, the modeled concentration of CP in lettuce was the highest at 3 d of growth for both LC and HC treatments, due to the small weight of lettuce and high residual concentration of CP in soil. Thereafter, they decreased constantly with the increasing weight of lettuce and dissipation of CP in soil. Interestingly, all modeled concentrations in the HC treatment at 40 d were slightly lower than those modeled in the LC treatment, which may be attributed to the faster degradation rate of CP in HC-treated

Table 2. Model parameters used for predicting root uptake of CP by lettuce.

| Input parameter (abbreviation) | Unit      | Value       | Nominal treatment |
|-------------------------------|-----------|-------------|-------------------|
| Initial soil exposure concentration ($C_0$) | mg kg⁻¹ | 15.2        | 24.9              |
| Half-life in soil ($T_s$)     | day       | 17.2        | 7.9               |
| Half-life in plant ($T_p$)    | day       | 8.7         | 8.0               |
| Soil-water distribution coefficient ($K_d$) | mL g⁻¹ | 82.1        | 82.1              |
| Organic carbon-water partition coefficient ($K_{oc}$) | - | 2218.9      | 2218.9            |
| Octanol-water partition coefficient ($K_{ow}$) | - | 1.02×10⁴     | 1.02×10⁴          |
| Transpiration stream ($Q_w$)  | mL day⁻¹ | 46.8        | 46.8              |
| Logarithmical initial plant weight ($I_g$) | g       | 0.3062      | 0.3092            |
| Plant growth constant ($K_g$) | -         | 1.1020      | 1.2031            |

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soils than in LC-treated soils. Similar to the modeled results, the measured uptake concentrations of CP during the uptake experiments were slightly lower in the HC treatment than in the LC treatment.

All measured concentrations were between the 5th and 95th percentiles of model variation. Although we found a large deviation (64%) between modeled and measured concentrations in both treatments at 40 d of growth, the mean value of deviations was acceptable at 27.9%, which is lower than the error value of 36% reported in another study [1]. Therefore, the modeled concentrations corresponded well to the measured concentrations.

Model accuracy assessment
The model accuracy assessment is presented in Fig 3. The correlation between modeled and measured concentrations was high, with $R^2$ values of 0.97 to 0.98, and the ratio between both concentrations was close to an ideal value of 1. Mean ER value between modeled and measured concentrations was 0.18 in the LC treatment, corresponding to a mean factor of -0.70. In contrast, the mean value of ER in the HC treatment was 0.72, corresponding to a mean factor of -1.49. Herein, positive or negative factor values, which represent the mean of the log value of residuals between modeled and measured values, depend on whether the residual value is $>1$ [20]. Coefficients of variation (CV) of factors to ERs ranged from 25.3 to 58.2% and were higher than error values of 19–22.6% reported in other modeling studies [1,15,20].

Variations between modeled and measured concentrations may result from the absence of other influential parameters, such as the desorption and leaching of pesticide in soil, landscape of the field, environmental weather, and contact area between plant roots and soil. In other studies of plant uptake models for hazardous substances, researchers have tried to include such parameters in their models [36–39]. However, there is no complete plant uptake model that satisfies all parameters that are relevant to the plant–soil system.

Estimated concentration in edible leaf parts
Based on the residual data analyzed from the uptake experiments (Table 1), the values of $R_{L/W}$, which is the ratio of concentration in leaves and the whole plant, were calculated (S3 Table).
Fig 3. Comparison between modeled and measured uptake amounts of CP from soil by lettuce (*dotted line* is the ideal relation between both values).

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Fig 4. Correlation between modeled and measured concentrations of CP in leaves of lettuce (*dotted line* is the ideal relation between both values).

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Values of $R_{L/W}$ were between 0.54 and 0.98, and were used to estimate the modeled concentrations of CP in leaves. Fig 4 show that the estimated concentrations of CP in lettuce leaves matched the experimentally-measured concentrations well, with a mean deviation of 27.9%, although the estimated CP concentrations at 40 d of growth deviated from those measured by approximately 64%.

Estimated CP concentrations in leaves were slightly higher in the HC treatment than in the LC treatment, similar to the uptake experiments. Estimated concentrations decreased slightly over time, with faster rates of decreasing CP in leaves estimated in the HC treatment. For both LC and HC treatments, the estimated concentrations ranged from 0.28 to 0.78 mg kg$^{-1}$ during the sampling period of 21–40 d (S3 Table), similar to the concentrations measured in the uptake experiments (0.1–0.8 mg kg$^{-1}$). All estimated and measured concentrations of CP in lettuce leaves exceeded the maximum residue limit (MRL) of 0.1 mg kg$^{-1}$ [40]. Exceeding the MRL demonstrates that cultivating lettuce in soils contaminated with CP at concentrations of >2 mg kg$^{-1}$ may lead to the production of unsafe lettuce at harvest.

In conclusion, our results suggest that the plant uptake model developed in this study could be used as a mathematical assessment tool to predict plant uptake of pesticides that persist in soils. In addition, further studies should be conducted to identify more effective parameters to improve the current low accuracy of plant uptake models.

Supporting information

S1 Fig. Total ion chromatogram (a) and mass spectrum (b) of CP analyzed using GC-MS. (TIF)

S2 Fig. Chromatograms for recovery tests of CP in soil (a, b) and whole plant of lettuce (c, d), spiked at a concentration of 1.0 mg kg$^{-1}$ (a, c—spiked samples; b, d—non-spiked samples). (TIF)

S3 Fig. Adsorption kinetics (a) and isotherms (b) of CP on tested soils (error bar is the standard deviation of the triplicate measurement). (TIF)

S4 Fig. Dissipation behavior of CP in the tested soil (error bar is the standard deviation of the triplicate measurement). (TIF)

S5 Fig. Change in length (a) and weight (b) of lettuce parts during the field experiments. (TIF)

S1 File. Laboratory experiment methods. (DOCX)

S1 Table. Physicochemical properties of the soils used for adsorption tests. (DOCX)

S2 Table. Recoveries of CP in soil and each compartment of lettuce. (DOCX)

S3 Table. Estimated concentrations of CP in leaf parts, calculated using the $R_{L/W}$ values. (DOCX)
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References

1. Juraske R, Vivas CSM, Velásquez AE, Santos GG, Moreno MBB, Gomez JD, et al. Pesticide uptake in potatoes: Model and field experiments. Environ Sci Technol. 2011; 45: 651–657. doi: 10.1021/es102907v PMID: 21141816
2. Kwon CH, Lee YD. Distribution and dissipation of deltamethrin, dimethomorph, imidacloprid, mepanipyrim, and metalaxyl applied to chili peppers. J Korean Soc Appl Biol Chem. 2015; 58: 821–829.
3. Trapp S. Modelling uptake into roots and subsequent translocation of neutral and ionisable organic compounds. Pest Manag Sci. 2000; 56: 767–778.
4. Trapp S, Eggen T. Simulation of the plant uptake of organophosphates and other emerging pollutants for greenhouse experiments and field conditions. Environ Sci Pollut Res. 2013; 20: 4018–4029.
5. Juraske R, Fantke P, Ramírez ACR, González A. Pesticide residue dynamics in passion fruits: Comparing field trial and modeling results. Chemosphere 2012; 89: 850–855. doi: 10.1016/j.chemosphere.2012.05.007 PMID: 22673401
6. Fantke P, Juraske R. Variability of pesticide dissipation half-lives in plants. Environ Sci Technol. 2013; 47: 3548–3562. doi: 10.1021/es303525x PMID: 23521068
7. United States Environmental Protection Agency (U.S. EPA). Technical support document for land application of sewage sludge volume II (EPA 822/R-93-011b). 1992. http://yosemite.epa.gov/water/owrcatalog.nsf/4e73d95b11602f2385256ae1007279fe/38fa4050461763d3985256b0600725389?OpenDocument&CartID=null.
8. Environment Agency (EA). Using science to create a better place: Updated technical background to the CLEA model (Science Report SC050021/SR3). 2008. https://www.gov.uk/government/uploads/system/uploads/attachment_data/file/291014/scho05008bnqw-e-e.pdf.
9. Rijksinstituut voor Volksgezondheid en Milieu (RIVM). Evaluation and revision of the CSOIL parameter set: Proposed parameter set for human exposure modelling and deriving Intervention Values for the first series of compounds (RIVM report 711701021). 2001. http://www.rivm.nl/dsresource?objectid=rivmp:19033&type=org&disposition=inline&ns_nc=1.

10. Trapp S. Fruit tree model for uptake of organic compounds from soil. SAR QSAR Environ Res. 2003; 14: 17–26. doi: 10.1080/1062936021000058755 PMID: 12688413

11. Shone MG, Wood AV. A comparison of the uptake and translocation of some organic herbicides and a systemic fungicide by barley I. Absorption in relation to physico-chemical properties. J Exp Bot. 1974; 25: 390–400.

12. Shone MG, Bartlett BO, Wood AV. A comparison of the uptake and translocation of some organic herbicides and a systemic fungicide by barley II. Relationship between uptake by roots and translocation to shoots. J Exp Bot. 1974; 25: 401–409.

13. Briggs GG, Bromilow RH, Evans AA. Relationship between lipophilicity and root uptake and translocation of non-ionised chemicals by barley. Pestic Sci. 1982; 13: 495–504.

14. Mckone TE, Maddalena RL. Plant uptake of organic pollutants from soil: Bioconcentration estimates based on models and experiments. Environ Toxicol. 2007; 26: 2494–2504.

15. Juraske R, Castells F, Vijay A, Muñoz P, Antón A. Uptake and persistence of pesticides in plants: Measurements and model estimates for imidacloprid after foliar and soil application. J Hazard Mater. 2009; 165: 683–689. doi: 10.1016/j.jhazmat.2008.10.043 PMID: 19036517

16. Rein A, Legind CN, Trapp S. New concepts for dynamic plant uptake models. SAR QSAR Environ Res. 2011; 22: 191–215. doi: 10.1080/1062936X.2010.548829 PMID: 21391147

17. Charles R. Modelling pesticides residues, electronic, scholarly journal. Ph. D. Thesis, École Polytechnique Federale de Lausanne. 2004. https://infoscience.epfl.ch/record/33592/files/EPFL_TH3123.pdf.

18. Hwang Ji, Lee SE, Kim JE. Interpretation and estimation for dynamic mobility of chlorpyrifos in soils containing different organic matters. Environ Geochem Hlth. 2015; 37: 1017–1027.

19. Singh BK, Walker A, Morgan AW, Wright DJ. Effects of soil pH on the biodegradation of chlorpyrifos and isolation of a chlorpyrifos-degrading bacterium. Appl Environ Microbiol. 2003; 69: 5198–5206. doi: 10.1128/AEM.69.9.5198-5206.2003 PMID: 12957902

20. Min ZW, Kim TH, Shin JH, Lee SM, Kim JE. Accelerated effect of ferric salts on degradation of thiophosphate fungicide, tolclofos-methyl by zerovalent iron. J Korean Soc Appl Biol Chem. 2009; 52: 681–687.

21. Macbean C. The pesticide manual. 16th ed. Alton, Hampshire, UK: British Crop Production Council; 2012. pp 199–201.

22. International Union of Pure and Applied Chemistry (IUPAC). Pesticide properties database. http://sitem.herts.ac.uk/aeru/iupac/Reports/154.htm.

23. Kleier DA. Phloem mobility of xenobiotics: I. Mathematical model unifying the weak acid and intermediate permeability theories. Plant Physiol. 1988; 86: 803–810. PMID: 26665992
33. Giles CH, Macewan TH, Nakhwa SN, Smith D. Studies in adsorption part XI. A system of classification of solution adsorption isotherms, and its use in diagnosis of adsorption mechanisms and in measurement of specific surface areas of solids. J Chem Soc. 1960; 111: 3973–3993.

34. Moore MT, Schulz R, Cooper CM, Smith S, Rodgers JH. Mitigation of chlorpyrifos runoff using constructed wetlands. Chemosphere. 2002; 46: 827–835. PMID: 11922063

35. Jeon SO, Hwang JI, Lee SH, Kim JE. Uptake of boscalid and chlorfenapyr residues in soil into Korean cabbage. Kor J Pest Sci. 2014; 18: 314–320.

36. Legind CN, Rein A, Serre J, Brochier V, Haudin C-S, Cambier P, et al. Simultaneous simulations of uptake in plants and leaching to groundwater of cadmium and lead for arable land amended with compost or farmyard manure. PLOS ONE. 2012; 7:

37. Namiki S, Otani T, Seike N. Fate and plant uptake of persistent organic pollutants in soil. Soil Sci Plant Nutr. 2013; 59: 669–679.

38. Carter LJ, Harris E, Williams M, Ryan JJ, Kookana RS, Boxall ABA. Fate and uptake of pharmaceuticals in soil–plant systems. J Agric Food Chem. 2014; 62: 816–825. doi: 10.1021/jf404282y PMID: 24405013

39. Chen Z-X, Ni H-G, Jing X, Chang W-J, Sun J-L, Zeng H. Plant uptake, translocation, and return of polycyclic aromatic hydrocarbons via fine root branch orders in a subtropical forest ecosystem. Chemosphere. 2015; 131: 192–200. doi: 10.1016/j.chemosphere.2015.03.045 PMID: 25863163

40. Ministry of Food and Drug Safety (MFDS). Pesticide residue database. 2015. http://www.foodnara.go.kr/residue/search/list.do?currentPageNo=1&searchType=&searchValue=chlorpyrifos&searchFlag=ALL.