Effect of Organic Residues on Pesticide Behavior in Soils: A Review of Laboratory Research

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Abstract: The management of large volumes of organic residues generated in different livestock, urban, agricultural and industrial activities is a topic of environmental and social interest. The high organic matter content of these residues means that their application as soil organic amendments in agriculture is considered one of the more sustainable options, as it could solve the problem of the accumulation of uncontrolled wastes while improving soil quality and avoiding its irreversible degradation. However, the behavior of pesticides applied to increase crop yields could be modified in the presence of these amendments in the soil. This review article addresses how the adsorption–desorption, dissipation and leaching of pesticides in soils is affected by different organic residues usually applied as organic amendments. Based on the results reported from laboratory studies, the influence on these processes has been evaluated of multiple factors related to organic residues (e.g., origin, nature, composition, rates, and incubation time of the amended soils), pesticides (e.g., with different use, structure, characteristics, and application method), and soils with different physicochemical properties. Future perspectives on this topic are also included for highlighting the need to extend these laboratory studies to field and modelling scale to better assess and predict pesticide fate in amended soil scenarios.

Keywords: soil amendment; organic matter; dissolved organic carbon; agriculture; pesticide fate; dynamics; laboratory experiments

1. Introduction

The use of large quantities of pesticides in today's intensive agricultural systems is a widespread practice for controlling pests, diseases and weeds. This increases the yield per hectare, ensuring the food supply for the world's ever-growing population [1,2], which currently stands at over 7.7 billion people, and is estimated to rise above 9.6 billion by 2050, and reach nearly 11 billion around 2100 [3]. The application of a wide range of pesticides is a regular and required practice in agriculture, as almost 45% of annual food production is lost due to pest infestation or the competition between crops and weeds for soil nutrients [4]. In fact, 3.5 million tons of pesticides are being used, of which 47.5% are herbicides, 29.5% are insecticides, 17.5% are fungicides, and 5.5% are other pesticides [5].

The global pesticide market recorded a value of nearly USD 84.5 billion in 2019, increasing at an annual growth rate of 4.2% since 2015, and it is likely to reach 11.5% with a value of nearly USD 130.7 billion by 2023 [6]. The ten countries consuming the most pesticide in the world are China, USA, Argentina, Thailand, Brazil, Italy, France, Canada, Japan, and India [7].

However, this extensive use of pesticides over recent decades is now of considerable environmental concern because of the release of mobile and/or persistent pollutants into the environment, and the potential accumulation of these toxic substances in soils and/or waters [8–10]. The fate of pesticides and their degradation products determines the contamination of the soil, water and air ecosystems over time. Moreover, if agrochemicals...
remain in the crops, they could finally enter the food chain, posing a threat to human, animal, and plant welfare [11–14].

The contamination of agricultural soils with pesticides could lead to changes in their chemical and biological properties, affecting their quality and causing a negative impact on crop yields [15]. They may impair soil microbial biodiversity and enzymatic activity (a vital indicator of soil tolerance to pollutants), and the associated degradation of soil organic matter (OM) [16,17]. Many reports are available on these negative effects on soil microbial communities [17,18], and on the processes associated with microbial activities [19].

A recent study involving 317 agricultural topsoil samples from the European Union and 76 pesticide residues as target compounds has revealed that 83% of the soils have been contaminated by one or more residues [9]. The contamination of surface and ground waters by pesticides has also been detected in recent years, probably due to deficient pesticide management, and increased by precipitation and/or irrigation that give rise to the runoff or leaching process of these compounds through the soil [20–23]. In fact, the contamination of water by pesticides is increasing in agricultural areas across different countries, and a broad range of pesticide concentrations has been found, in some cases exceeding the limit established for drinking water by European Union (EU) legislation (0.1 µg·L⁻¹) [24–26].

These environmental contamination data highlight the need to roll out strategies to optimize agricultural sustainability by maximizing crop productivity and reducing or preventing soil and water contamination by pesticides. This has been widely addressed in recent years due to the requirement to meet European Community regulations [27]. One of these strategies is based on the in-situ application of organic residues as organic amendments [28]. This method is a common agricultural practice which allows increasing soil OM content, and it can be used to control soil and water contamination by pesticides: (i) promoting the immobilization of pesticides in soil OM, enhancing their subsequent biodegradation, and preventing or reducing their potential mobility into water resources [28–30], and (ii) delivering nutrients to the soil by increasing OM content to promote soil fertility and plant growth and stimulate ecological restoration with concomitant benefits for the health of the soil ecosystem [11]. In addition, organic materials require minimal pre-treatment before their application to the soil because of their biological origin [31].

Large amounts of organic residues are generated from livestock, urban, agricultural and industrial activities, and their management is a topic of environmental and social interest in many countries today due to the problems surrounding their disposal [32,33]. In general, these wastes have a high OM content, and they could be used as organic amendments in agriculture, with this being one of the most sustainable options and with greater environmental advantages. Moreover, numerous organic residues could perform as possible sorbents for pesticides [34–36]. These studies have assessed the effects that organic carbon (OC) from exogenous sources have on the behavior and environmental fate of pesticides in soils due to the affinity of pesticides, which are generally hydrophobic substances, by these organic materials. The OC of the amendments, depending on their nature, composition and content, can modify the main physicochemical processes of pesticides (adsorption-desorption, dissipation and leaching) in soils. These processes determine their efficiency as well as the dissipation or persistence of these compounds in the soil and their effects as potential environmental contaminants of the soil and surface or ground waters [37].

The aim of this review article is to analyze the influence that organic residues applied as soil organic amendments have on the environmental fate of pesticides. It has focused mainly, albeit not exclusively, on research papers published in the last ten years. Special interest has been directed toward the factors that affect the adsorption-desorption, dissipation, and leaching of pesticides in amended soils at laboratory scale. Future perspectives on the joint application of pesticides and organic amendments are also included.
2. Organic Residues as Soil Amendments

2.1. Origin, Characteristics and Impact on Soil Properties

The use of organic residues as soil amendments to improve soil quality and fertility dates back thousands of years. Animal manure and human sewage was already being applied to the soil by Greeks and Romans [38]. Waste products such as crop residues, seashells, farmyard manure and others were previously used to enhance crop growth [39]. The use of organic residues as soil organic amendments is still a widespread and common practice in modern agriculture. In fact, this agronomic practice is on the rise in Europe, the US, and many other countries. Its application improves soil properties, maintaining soil health and productivity, while reducing the disposal of organic wastes into landfills, with the consequent environmental benefits [40]. The restoration and maintenance of soil OM content is one of the main benefits of the application of organic residues to agricultural soil because they add exogenous OM, which contributes greatly to soil fertility and long-term use [41]. OM is the main soil property, as it supports and interrelates the biological, chemical and physical dimensions of soil fertility and health [42]. Furthermore, because OM also contributes to the process of sequestering carbon dioxide from the atmosphere (soil C uptake, and thus climate change mitigation), its improvement has been championed at international forums on food security and climate change [43].

Regarding the benefits that organic amendments have for soil biological properties, they directly stimulate microbial growth and biomass by providing energy and essential nutrients (especially N, P, K, Ca and Mg), or indirectly by promoting plant growth, and consequently the amount of root exudates in the rhizosphere [44]. Moreover, the presence of diverse substrates susceptible to enzymatic hydrolysis within the amendments stimulates soil microbial activities [45]. Soil microbial diversity and composition could also be affected due to a higher availability of nutrients and growth substrates present in the amendments, which increase the number of ecological niches and promote a variety of ecological interactions, such as competition and/or antagonism between organisms [46]. The beneficial effects of organic amendments on the biomass, activity, and diversity of soil organisms have a long-term beneficial impact on soil health [47,48] and also contribute to different ecosystem services (C and nutrient cycling, disease suppression, etc.). However, it is important to stress that microbial responses to the application of organic amendments vary greatly depending on the nature and lability of the amendments’ OM [49].

Organic amendments also influence soil chemical properties in a positive way. Indeed, their favorable effects on soil microbial communities are often linked to changes in soil chemical characteristics [50,51]. Several organic amendments may have a direct effect on soil fertility by providing a wide variety of macro- and micro-nutrients, which support plant and microbial growth [52]. In addition, they may affect soil pH and enhance the cation exchange capacity, thus indirectly influencing nutrient availability, microbial activity and, therefore, soil fertility [53]. Nutrient availability may be influenced by the amendment’s biochemical composition, and in particular by its carbon–nitrogen (C/N) ratio, which may limit soil microbial growth and activity, thereby influencing the rate of OM decomposition and the patterns of nutrient release [49,54].

Soil physical characteristics can also be positively influenced by the application of organic amendments. Accordingly, the addition of exogenous OM directly improves soil structure (better porosity, aggregation and structural stability) [55] and water retention capacity [56], with the associated positive effects previously noted for soil performance and crop productivity. In turn, the stimulation of soil microbial communities through the use of organic amendments may also indirectly improve soil structure, as microbial activity (e.g., through the secretion of exopolysaccharides) and particularly hyphal growth significantly influence soil aggregation and aggregate stability [57]. The increase in soil porosity often reduces soil crusting and bulk density, thus favoring the movement of air and water through the soil matrix, the exploratory capacity of plant root systems, and the development of a suitable environment for soil biological communities [58]. Moreover, organic amendments influence particle size distribution, connectivity and the total surface
area within the soil, increasing the number and types of available niches for biological colonization [49].

The potential positive effects of organic amendments on the soil ecosystem depend on many factors, such as their origin (forestry, farming, urban or industrial wastes, etc.), material stage (solid, semisolid, liquid), subjected or not to treatments (composting, anaerobic digestion, etc.), composition, stability, maturity, and application frequency, method and rate. Moreover, soil type, cropping system, and weather conditions are also important factors. In order to identify the different properties, agronomic potential, and limitations of any given organic amendment for soil and crop health, a thorough characterization of both the organic amendment and the agroecosystem itself needs to be performed before its application [49]. In this sense, however, it is also important to stress that the use of organic residues as a soil organic amendment is not devoid of risk. This agricultural practice may sometimes have unwanted effects on the environment depending on the factors previously cited (nature, origin, dose of application, etc.). Some of the potential negative effects analyzed and determined in the literature are the increase in soil electrical conductivity and its salinization, negative impacts on the sensitivity and resilience of soil bacteria communities, and the release of pollutants into the soil and/or waters (nitrate, heavy metals, antibiotics, polycyclic aromatic hydrocarbons, dioxins, PCBs, etc.) [51,59–63].

2.2. European Legislation on the Use of Organic Residues as Soil Amendments

Economic growth in the EU continues to increase the volume of residues generated. As a result, waste disposal has become a social and environmental concern because it causes the unnecessary loss of materials and energy, environmental damage, and negative effects on health and quality of life. Landfills, for example, occupy space and can pollute air, water and soil, while incineration leads to emissions of air pollutants. The long-term goal of EU waste management policies is to decrease the amount of residues generated, and when their generation is unavoidable, promote them as a resource and achieve higher levels of recycling and safe waste disposal, reducing the negative impacts on the environment and health and ushering in an efficient “Recycling Society” [64].

Two common EU targets for 2030 are recycling 65% of municipal solid waste (MSW) and reducing the corresponding landfill to a maximum of 10% of this figure [65]. In 2018, 5.2 tons of residues were generated per EU inhabitant, and 38.7% of waste in the EU was landfilled and 38.1% was recycled [64]. According to the European Compost Network [66], between around 118 and 138 million tons of bio-residues are generated annually across the EU, of which only about 40% (equivalent to 47.5 million tons/year) is efficiently recycled into high quality compost and digestate. Based on the Status 2019 report [66], a total of 47.5 million tons of bio-residues are treated in 4274 plants, and the predominant treatment process is composting.

Considering that up to 50% of MSW is organic, the bio-residue fraction has a significant role in recycling and developing the circular economy. Most of the MSW generated in the EU is still landfilled (24%) or incinerated (27%), and less than half is recycled (31%) or composted (17%) [64]. According to the data available, there has not been any increase in bio-residue recycling in recent years [67]. Moreover, waste management practices vary widely across EU member countries, with many continuing to send large amounts of MSWs to landfills.

Soil health and quality have been seriously compromised in recent years by constant changes in land use and the depletion of soil OM [68]. Indeed, approximately 45% of Europe’s topsoil (0–30 cm) has a low OM (<3.5%) content [69,70], and the soils in Mediterranean regions are highly susceptible to its loss, with almost 75% recording a low (<2%) or very low (<1%) OM content [71], whereby most of them are considered degraded. The reuse of organic residues as soil amendments in agriculture is an ancient but increasingly popular practice that not only helps to reduce the dependency on agrochemicals, but also constitutes an ecologically, economically, and socially acceptable alternative to landfill
disposal and incineration, contributing at the same time to the objectives of the EU policies of “Zero Waste”, “End-Of-Waste”, and the “Circular Economy Strategy” [46,72].

It should nonetheless be stressed that an organic residue must meet a series of requirements for its potential use as a soil amendment. According to Commission Decision (EU) 2015/2099 of 18 November 2015 establishing the ecological criteria for the award of the EU Ecolabel for growing media, soil improvers and mulch [65], the following definitions apply: “1. Soil amendment means a fertilizer product incorporated into the soil in situ whose function is to maintain, improve or protect the physical or chemical properties, structure or biological activity of the soil, with the exception of limestone amendments, and 2. Organic soil amendment means a soil amendment that contains carbonaceous materials whose main function is to increase the OM content of the soil”. Among the organic residues potentially applicable to soil are those from the following activities: urban (sewage sludge (SS) or MSW), agricultural (crop residues), livestock (manure and slurry), and agro-industrial (wine, beer, sugar and olive production, and mushroom cultivation) [38].

The addition of OM through organic amendments plays a major role in the fate of xenobiotic compounds including pesticides [73,74]. The solid organic matter (SOM) and dissolved organic matter (DOM) of these amendments applied to the soil may modify the physicochemical behavior of pesticides (e.g., adsorption–desorption, persistence, bioavailability, degradation, and mobility), affecting soil quality and surface and ground waters [34,75–79] (Figure 1).

Figure 1. Summary of the organic amendments’ (solid organic matter (SOM) and dissolved organic matter (DOM)) effects on the processes controlling the fate of pesticides in soil.
3. Effect of Organic Residues on the Fate of Pesticides in Soil

3.1. Effect of Organic Residues on the Adsorption-Desorption of Pesticides

The combined application of pesticides and organic residues in soils modifies the former’s physicochemical behavior, mainly through their adsorption-desorption by the amended soils [78,79]. Organic amendments increase soil OC content, and this parameter is the most relevant factor influencing the adsorption process and the affinity of hydrophobic pesticides by soils [80,81]. Adsorption-desorption determines the environmental fate of any organic pollutants in the soil-water environment [82]; it directly or indirectly controls the availability of pesticides to be transported to surface waters by runoff or to groundwaters by leaching, to the air by volatilization, to be degraded/transformed by microbial attack, or be taken up by plants [83]. Thus, the weak adsorption and/or strong desorption of pesticides promotes leaching, run-off, volatilization, biodegradation and even ecotoxicological impacts on non-target organisms, including human beings, while strong adsorption prevents losses of pesticides by such processes [84].

Accordingly, the addition of organic amendments to soil could lead to a greater or lesser degree of pesticide immobilization in the amended soil. This effect has consequences for pesticide degradation, persistence or mobility, enhancing a pesticide’s subsequent chemical, physical, and biological transformation or degradation, decreasing its transport through the soil profile, and consequently reducing groundwater pollution in some cases [28,78]. However, it could also affect the final concentration bioavailable for absorption by the targeted weeds [37]. Therefore, adsorption and desorption processes help to understand how to predict the mobility and availability of pesticides in unamended and amended soils. Numerous references report the ability that organic amendments have to adsorb pesticides [29,85–87].

Adsorption is a physicochemical process in which pesticide molecules are retained on a solid surface (especially by the soil colloidal fraction) within a solution through hydrophobic interactions, van der Waals forces, π-π interaction, and covalent, ionic or hydrogen bonds [88,89]. Soil OM and its more active components, humic acids (HAs) and fulvic acid (FAs), are the principal adsorbents for pesticides, followed by clay colloids and oxyhydroxides of iron and manganese, which interact with pollutants when they reach the soil [90,91]. HA is the OM fraction with the highest reactivity (determined by the number and type of functional groups) and largest surface area. OM’s highly variable composition means it can interact with neutral or ionizable molecules [75]. The pesticide adsorption capacity varies, in general, according to the physicochemical characteristics of the adsorbent and pesticide properties, mainly its water solubility and its hydrophobic, polar, or ionic character [37,92].

The nature and composition of the amendment’s OM vary, with the consequent difficulty in predicting its efficiency for adsorbing pesticides [34,76]. The addition of organic amendments to soil introduces not just SOM but also DOM (Figure 1). The influence of the SOM and DOM content of organic residues on the adsorption of pesticides with different characteristics by amended soils has been frequently studied [34,76,77]. The DOM content in unamended soil is usually very low, but it could become relevant if the organic amendment has a high content in this fraction [93]. DOM is a diverse mixture of complex compounds with different chemical structures and molecular weights that might enhance the formation of multiple interactions with organic pesticides, controlling their distribution in the soil [94]. This is why DOM may modify the movement of pesticides, generally decreasing their adsorption by SOM and increasing their leaching, leading to groundwater contamination [78,95,96], although other authors have indicated that DOM could also be adsorbed by the soil, increasing the adsorption of pesticides and decreasing their leaching [97].

Different processes have been proposed to explain the decreased adsorption of pesticides in the presence of DOM [94,98–100] (Figure 1). These include the competition between DOM and pesticide molecules for the adsorption sites in soil, the saturation of soil adsorption sites by DOM, masking these sites for the adsorption of pesticides, the
co-sorption of pesticides by DOM, and the formation of mobile DOM-pesticides complexes. Some authors have also indicated that DOM has characteristics similar to surfactants with the capacity to decrease surface tension and increase the solubility of pesticides, reducing their adsorption [101]. The extent and nature of DOM-pesticide interactions depend on factors such as pesticide molecular weight and polarity [102]. These relative effects of DOM will be greater for more hydrophobic chemicals, and will be influenced by the concentration, source, size, polarity, and molecular configuration of the organic colloids [95].

Table 1 includes a summary of the main results obtained from the recent literature on the adsorption and/or desorption of pesticides by some organic residues and by soils amended with different organic residues used as amendments. An organic material widely studied for these purposes is biochar (BC). It is an efficient adsorbent and a potential material for soil amendment [30]. It is a carbonaceous and porous product generated from the partial combustion of biomass, and its effects as soil amendment in the adsorption–desorption of pesticides has been assessed from different points of view. The effects of different types of BC, treatments or aging periods, and their different doses or application forms as organic amendments in soils have been reported for pesticides with different characteristics. Parlavecchia et al. [103] have investigated the effect of two types of BC from grape vine pruning residues (BC-G) and spruce wood (BC-S) and two vermicomposts (VC) involving digestates from a mixture of manure and olive mill wastewater (VC-M) and buffalo manure (VC-B) in the sorption-desorption capacity of the fungicide metalaxyl-M. Both types of amendments (BC and VC) have a significant capacity to adsorb the high-water-soluble fungicide. However, BC has recorded a much higher sorption efficiency than VC and lower desorption, which is explained by the composition and structural differences in OM between the two (VC has less aromatic carbon and a higher content of hydrophilic functional groups interacting with polar compounds and solvents than BC). Metalaxyl-M is adsorbed to a similar extent on the two VCs, while a different sorption behavior is observed in BC-G and BC-S due to their different porous structures. Likewise, Wu et al. [89] have assessed the effects of different types of BC from peanuts (BCP), chestnuts (BCC), bamboo (BCB), maize straw (BCM), and rice husk (BCR), and the effects of BCR aging on the sorption, degradation and bioavailability of the herbicide oxyfluorfen in various amended soils. The sorption capacity of the five BC differs significantly due to their physicochemical properties. The sorption capacity of BC for oxyfluorfen is significantly correlated with the specific surface area and elemental composition, but it decreases with longer aging time. BC reduces the bioavailability of oxyfluorfen in amended soils, but a higher bioavailability is recorded with an increase in the aging period of BC. Nevertheless, the sorption capacity of amended soil for oxyfluorfen after six months is still better than the unamended soil, highlighting that BCR is an effective way of reducing the risk of contaminating soil with oxyfluorfen, although it could also diminish the herbicide’s bioavailability and efficacy. Deng et al. [104] have studied the effect that BC obtained from cassava residues at 750 °C (MS750) applied at different rates between 0% and 5% has on the sorption-desorption and mobility of atrazine. The MS750 application significantly enhances the sorption capacity and decreases the sorption reversibility of atrazine in the amended soil compared to the unamended soil, due to the larger surface area and greater aromaticity of MS750 (with favorable sorption domains for organic compounds). Moreover, sorption affinity increases with higher BC application rates, although it is also influenced by solution pH, ambient temperature, and contact time between soil and BC (equilibrium time). The entrapment of atrazine in micropore or pore deformation could lead to desorption hysteresis in BC-amended soils.

Mendes et al. [105] have recently studied the effect that BC from cow bone applied to the topsoil or incorporated into the surface layer has on the sorption-desorption of the herbicides hexazinone, metribuzin, and quinclorac in an unamended soil, pure BC, and BC-amended soil under laboratory conditions. The results indicate low values of $K_f$ adsorption and desorption constants in the unamended soil. BC increases these $K_f$ values, stimulating the retention of all the herbicides in the surface soil. The low C content of BC
has a minimal impact on the total OC of amended soils, with the main changes occurring in pore size (up to 60,000 nm), volume (0.225 cm$^3$ g$^{-1}$), and area (133 m$^2$ g$^{-1}$) to increase herbicide adsorption by the soil. The desorption data for all herbicides were consistent with the values found for adsorption in the unamended soil, where quinclorac was the herbicide with the highest $K_{oc}$ (adsorption) and lowest $K_{oc}$ (desorption). The desorbed amount of herbicides was close to zero after the addition of BC in the soil or in pure BC, confirming the high adsorption potential of BC regardless of the material’s method of application. Similar high sorption has been reported for two weak acid herbicides, aminocyclopyrachlor and mesotrione, in amended soils with the same cow bone char [106]. The comparison of organic residue application at varying rates and with two different particle size groups (0.3–0.6 and 0.15–0.3 mm) indicates that higher BC application rates increases the adsorption and decreases the desorption of both herbicides, regardless of particle size.

The effectivity and rate of application of other organic residues as adsorbents of pesticides has also been reported in different studies. Marín-Benito et al. [107] have studied the effect of large amounts of lignocellulosic residues from forestry and industrial activities on the adsorption–desorption of certain pesticides by soils. The study involves two wood wastes (pine and oak wood) at two different doses (5% and 50%) and various incubation times (0, 5 and 12 months) in two soils with different textures (sandy loam and sandy clay). The effect on the adsorption–desorption of two herbicides and one fungicide (linuron, alachlor, and metalaxyl) has revealed that the application of oak or pine wood to soils increases the adsorption of linuron and metalaxyl by both soils, and of alachlor by the sandy loam soil at a lower dose (5%), while the adsorption of the three pesticides increases under all conditions at the highest dose (50%). The results also indicate the influence of soil type on alachlor desorption and/or its possible bioavailability from wood-soils, but not for linuron and metalaxyl, although this behavior changes with incubation time. The role of the nature of the OC ($K_{oc}$ values) for sorption has been evidenced for alachlor and metalaxyl, but not for linuron. Other residues, such as the SS applied to the soil at various rates (0.1%, 1%, and 10% w·w$^{-1}$), have a non-significant effect on the sorption–desorption of aminocyclopyrachlor and mesotrione [108]. Both herbicides follow a similar adsorption behavior in all treatments, although $K_d$ for mesotrione is $\approx$3.5-fold higher than for aminocyclopyrachlor due to the latter’s higher water solubility. This leads to a higher bioavailability of aminocyclopyrachlor in soil solution for its absorption by weeds and crops.

Fewer studies have addressed the adsorption–desorption of several pesticide-organic residue combinations or the amendment effect on soil physical properties. Duhan et al. [109] have studied the behavior of five herbicides commonly used in sugarcane production (imazapic, atrazine, hexazinone, diuron, and metribuzin) by eleven waste materials (mill muds) and by three soils amended with them at different rates (5–25%, dry weight basis). The authors have observed that all the amendments enhance the adsorption efficiency for four of the five herbicides, depending on the rate of application, especially in the soil with low OC. Even at the lowest application rate, the adsorption of the herbicides increases from two to ten times. Mill muds in soil also reduce the rate and extent of herbicide desorption, especially at a 5% application rate and for mobile herbicides such as metribuzin and atrazine. Marsico et al. [110] have studied the effect that the mucilage extracted from Chia seeds (Salvia hispanica L.) has as a soil amendment on soil physical properties and on the sorption–desorption behavior of four herbicides (MCPA, diuron, clomazone, and terbuthylazine) used in cereal crops. The assessment of the changes in the microstructural characteristics caused by the reactions between the mucilage and soil particles in three soils indicates that mucilage amendment reduces soil porosity due to a decrease in larger pores (radius $> 10$ µm) and a significant increase in finer pores (radius $< 10$ µm), as well as in particle surface. Higher herbicide adsorption has been observed in the amended soils than in the unamended ones. Moreover, herbicide desorption is severely inhibited in the amended soils.
Although many organic amendments have proven to be effective adsorbents of pesticides, only a few studies have evaluated the functional groups involved in the adsorption process [111–114]. Accordingly, Gaonkar et al. [112] have used spectroscopy to characterize the DOM from two organic amendments (mixed waste compost and dried goat manure) and the amended soils, and assess their influence on the sorption of the insecticides dichlorvos and chlorpyrifos. The DOM contained large amounts of highly humified and aromatic molecules. DOM led to a non-significant increase in dichlorvos adsorption (hydrophilic pesticide), due mainly to the additional sites provided by the adsorbing DOM and no interactions between DOM and the insecticide in solution. However, a significant reduction in chlorpyrifos adsorption (hydrophobic pesticide) was observed, probably due to interactions between DOM and the insecticide mostly in solution, and to some extent at the soil/solution interface, increasing the solubilization of chlorpyrifos. This reduction in adsorption depended on the nature and concentration of the DOM, as well as on insecticide properties. In agreement with the adsorption results, chlorpyrifos desorption was significantly increased by the DOM residue. In another recent study, García-Delgado et al. [114] have determined the OC functional groups from four organic amendments (spent mushroom substrate (SMS), GC, manure, and SS) by elemental analysis and $^{13}$C-NMR, and their effects on the adsorption of four herbicides with different structures (triasulfuron, chlorotoluron, flufenacet, and prosulfocarb) by two unamended and amended soils with different textures. The chemical composition and structure of the organic amendments (especially OC content and structural C type), and external factors such as herbicide polarity (hydrophobicity) and soil properties controlled the adsorption process. The adsorption of herbicides was promoted by carbon-rich organic amendments with aliphatic and aromatic structures, while the irreversible adsorption (hysteresis) of herbicides in the amended soils was enhanced by the abundance of O-alkyl and N-alkyl groups of organic amendments.

3.2. Effect of Organic Residues on Pesticide Leaching

Pesticides run off from agricultural soils to surface waters when water exceeds the soil’s infiltration capacity and move to groundwater by leaching or vertical movement in the soil profile. Large macropores act as preferential flow pathways in leaching, prompting the rapid movement of the pollutants through the unsaturated zone [115]. Pesticide runoff and leaching are the main sources of surface and ground water contamination by pesticides, causing environmental problems [116]. The physicochemical properties of the pesticides and soil properties (texture, clay content, OM, and permeability) play a critical role in the leaching process [117].

Pesticides’ leaching potential is generally favored by their high water solubility, long persistence, and low adsorption in soil [118]. Pesticides with an intermediate adsorption rate have a greater tendency to undergo losses through runoff, as weakly adsorbed compounds are more available for leaching through the soil from the surface, while strongly adsorbed ones are less so. Hence, minor leaching is expected in soils with a high OM content because of their greater adsorption [119]. The application of organic amendments to soils increases the soil OM content, but it may also increase the amount of DOM in the aqueous phase and, in general, enhance leaching, as previously explained in Section 3.1 [93] (Figure 1). Nevertheless, the greater or lesser leaching of pesticides in amended soils may not be due solely to the presence of additional SOM or DOM in amended soil, but also to structural changes in soil porosity induced by the higher soil OC content [37].
<table>
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<td>Metalaxyl-M</td>
<td>Silt loam soil (pH 6.70, OC 2.90%)</td>
<td>Biochar from grape vine pruning residues (BC-G) (pH 9.9, OC 75.1%) and spruce wood (BC-S) (pH 9.1, OC 83.8%). Vermicomposts (VC) from manure and olive mill wastewater (VC-M) (pH 7.9, OC 31.6%) and buffalo manure (VC-B) (pH 7.8, OC 36.6%). Biochar/soil: 2% (w w⁻¹)</td>
<td>Sorbent/Solution: 25 mg biochar/5 mL or 3 g soil/8 mL water solution Herbicide concentration: 1–20 mg L⁻¹ Shaken: 24 h, T: 20 °C Analytical determination: HPLC</td>
<td>Metalaxyl sorption order: non-amended soil &lt; soil–VC–M ≤ soil–VC–B &lt; soil–BC–S &lt; soil–BC–G Much higher sorption efficiency by BC than by VC and a lower extent of metalaxyl desorption due to composition and structural differences of the organic matter of BC.</td>
<td>Parlavecchia et al. [103]</td>
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<td>Oxyfluorfen</td>
<td>Loamy clay soil (pH 4.85, OC 0.84%) Sandy loam soil (pH 7.55, OC 0.98%) Clay loam soil (pH 6.59, OC 2.23%)</td>
<td>Biochar from peanut (BCP) (pH, 7.05, C 49.17%), chestnut (BCC) (pH 6.08, C 58.07%), bamboo (BCB) (pH 7.45, C 63.25%), maize straw (BCM) (pH 6.83, C 43.36%), rice hull (BCR) (pH 6.96, C 33.60%) BCR/soil: 0.5%, 1%, or 2% (w w⁻¹)</td>
<td>Sorbent/Solution: 0.1 g biochar/40 mL or 2 g soil/200 mL 0.01 M CaCl₂ Herbicide concentration: 0.05–10 mg L⁻¹ Shaken: 6 days, T: 25 °C Aging time of BCR-soil: 1, 3, 6 months Analytical determination: GC/MS</td>
<td>BC sorption capacities followed the order: BCR &gt; BCB &gt; BCM &gt; BCC &gt; BCP owing to differences in physicochemical properties. BCR sorption capacity decreased with aging time.</td>
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<tr>
<td>Atrazine</td>
<td>Krasnozem soil (pH 7.0, OC 0.89%, clay 28.2%, silt 37.8%)</td>
<td>Biochar from cassava wastes (pH 9.55, C 62.38%) obtained at 750 °C (MS₇₅₀). SSA: 430.4 m²/g, MP: 0.144 m³/g Biochar/soil: 0%, 0.1%, 0.5%, 1%, 3% and 5% (w w⁻¹)</td>
<td>Sorbent/Solution: 0.2–2 g/10 mL 0.01 M CaCl₂ Herbicide concentration: 0.5–20 mg L⁻¹ Shaken: 24 h T: 15, 25, 35 °C, pH: 3, 5, 7, 9 Analytical determination: HPLC</td>
<td>Great sorption capacity for atrazine of MS₇₅₀ in soil due to high surface area and micropore volume. High degrees of aromaticity and hydrophobicity (H/C: 0.02, N + O/C: 0.09) of MS₇₅₀ supplied numerous sorption sites.</td>
<td>Deng et al. [104]</td>
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<tr>
<td>Hexazinone</td>
<td>Sandy loam soil (pH 6.9, OC 0.52%, clay 15.1%, silt 3.3%)</td>
<td>Bone char (BC) (pH 9.72, C 11%) BC/soil: 5% (w w⁻¹) or 60 t ha⁻¹</td>
<td>Sorbent/Solution: 10 g/10 mL 0.01 M CaCl₂ Herbicide concentration: 0.63–3.13 mg L⁻¹, 1.60–8 mg L⁻¹, 0.31–1.56 mg L⁻¹ Shaken: 24 h, T: 20 °C Analytical determination: Liquid scintillation</td>
<td>High sorption of herbicides by BC, regardless of the application form of the material (topsoil or incorporated in the surface layer in leaching columns).</td>
<td>Mendes et al. [105]</td>
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<tr>
<td>Metribuzin</td>
<td>Sandy loam soil (pH 6.9, OC 0.52%, clay 15.1%, silt 3.3%)</td>
<td>Bone char (BC) (pH 9.72, C 11%) BC/soil: 5% (w w⁻¹) or 60 t ha⁻¹</td>
<td>Sorbent/Solution: 10 g/10 mL 0.01 M CaCl₂ Herbicide concentration: 0.63–3.13 mg L⁻¹, 1.60–8 mg L⁻¹, 0.31–1.56 mg L⁻¹ Shaken: 24 h, T: 20 °C Analytical determination: Liquid scintillation</td>
<td>High sorption of herbicides by BC, regardless of the application form of the material (topsoil or incorporated in the surface layer in leaching columns).</td>
<td>Mendes et al. [105]</td>
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Table 1. Cont.

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<thead>
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<tr>
<td>Aminocyclopyrachlor Mesotrione</td>
<td>Clay soil (pH 6.44, OC 2.73%), clay 50.9%, silt 19.6%</td>
<td>Bone Char (BC) (pH 9.72, C 11%)&lt;br&gt;BC/soil: 0%, 1%, 5%, 10%, and 100% (w w&lt;sup&gt;-1&lt;/sup&gt;) or 0, 12, 60, 120, and 1200 t ha&lt;sup&gt;-1&lt;/sup&gt; BC particle size groups: 0.3–0.6 and 0.15–0.3 mm</td>
<td>Sorbent/Solution: 10 g/10 mL 0.01 M CaCl&lt;sub&gt;2&lt;/sub&gt; 0.051 mg L&lt;sup&gt;-1&lt;/sup&gt; (0.32 Bq L&lt;sup&gt;-1&lt;/sup&gt;) aminocyclopyrachlor 5.0 mg L&lt;sup&gt;-1&lt;/sup&gt; (1.13 Bq L&lt;sup&gt;-1&lt;/sup&gt;) mesotrione Shaken: 24 h, T: 20 °C Analytical determination: Liquid scintillation</td>
<td>Higher BC rates (regardless of the particle size) increased both herbicides adsorption and decreased their desorption.</td>
<td>Mendes et al. [106]</td>
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<tr>
<td>Linuron</td>
<td>Sandy loam soil (pH 6.3, OC 0.51%, clay 11.8%, silt 13.6%), Sandy clay soil (pH 6.9, OC 1.04%, clay 38.1%, silt 5.8%)</td>
<td>Pine Wood (OC 41.6%, DOM 1.62%, lignin 24.4%), oak wood (OC 38.5%, DOM 6.86%, lignin 18.2%) Wood/soil: 5% and 50% (w w&lt;sup&gt;-1&lt;/sup&gt;) or 1.2, 12, and 120 t C ha&lt;sup&gt;-1&lt;/sup&gt;</td>
<td>Sorbent/Solution: 5 g/10 mL water solution 1–25 mg L&lt;sup&gt;-1&lt;/sup&gt; (100 kBq L&lt;sup&gt;-1&lt;/sup&gt;) Shaken: 24 h, T: 20 °C Incubation times: 0, 5 and 12 months Analytical determination: Liquid scintillation</td>
<td>Pesticide adsorption increased with high wood dose but OC nature was not relevant. Adsorption did not change after incubation times. The adsorption irreversibility decreased in presence of wood for alachlor and increased that of linuron and metalaxyl.</td>
<td>Marin–Benito et al. [107]</td>
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<tr>
<td>Imazapic</td>
<td>Red Ferrusol (pH 7.1, OC 2.1%, clay 41%, silt 23%), Atrazine (pH 7.0, OC 2.3%, clay 30%, silt 22%), Hexazinone (pH 5.7, OC 0.9%, clay 30%, silt 22%), Diuron (pH 6.5, OC 3.5%, clay 22%, silt 8%), Metribuzin (pH 6.5, OC 3.5%, clay 22%, silt 8%)</td>
<td>Eleven mill muds/ash from different sugar mills (pH 6.04–7.26, OC 27.7–37.8%) Mill muds/soil: 5–25% (w w&lt;sup&gt;-1&lt;/sup&gt;)</td>
<td>Sorbent/Solution: 1 g/5 mL 0.01 M CaCl&lt;sub&gt;2&lt;/sub&gt; 0.5 mg L&lt;sup&gt;-1&lt;/sup&gt; Shaken: 24 h, T: 25 °C Analytical determination: Q-TOF</td>
<td>Sorption order: diuron &gt; atrazine = metribuzin &gt; hexazinone = imazapic (consistent with herbicide properties). Mill muds at 5% dose increased herbicide retention up to tenfold. Amendments reduced desorption of mobile herbicides in low OC soils.</td>
<td>Duhan et al. [109]</td>
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<tr>
<td>MCPA</td>
<td>Sandy loam soil (pH 7.93, OC 0.54%, clay 6.7%, silt 16.8%)</td>
<td>Mucilage extracted from chia seeds (Salvia hispanica L.)</td>
<td>Sorbent/Solution: 0.5 g unamended or amended soil/8 mL water solution</td>
<td>Soil porosity decreased by mucilage amendment. Sorption of herbicides increased in amended soils (sandy–loam &lt; loam &lt; clay–loam). Diuron recorded the highest K$_d$ value and desorption was observed only for terbuthylazine.</td>
<td>Marsico et al. [110]</td>
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<tr>
<td>Diuron</td>
<td>Loam soil (pH 6.77, OC 1.77%, clay 22.1%, silt 34.2%)</td>
<td>Organic residue/soil: 10% (w w$^{-1}$)</td>
<td>Herbicide concentration: 1 mg L$^{-1}$</td>
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<tr>
<td>Clomazone</td>
<td>Clay loam soil (pH 8.14, OC 1.38%, clay 31.1%, silt 26.8%)</td>
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<td>Shaken: 24 h, T: 20 $^\circ$C</td>
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<tr>
<td>Terbuthylazine</td>
<td>Sandy loam soil (pH 7.36, OC 1.20%, clay 17%, silt 25%)</td>
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<td>Analytical determination: HPLC</td>
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<tr>
<td>Dichlorvos</td>
<td>Sandy soil (pH 8.52, OC 0.7%, clay + silt 9.3%)</td>
<td>Compost (C) from mixed wastes (pH 6.61, OC 29.5%, DOM 354 mg L$^{-1}$), and dried goat organic manure (OM) (pH 8.67, OC 14.4%, DOM 620 mg L$^{-1}$)</td>
<td>Sorbent/Solution: 5 g soil/100 mL in C-DOM or 0.01 M CaCl$_2$ Herbicide concentration: 0.1–10 mg L$^{-1}$ (chlorpyrifos) 0.25–100 mg L$^{-1}$ (dichlorvos)</td>
<td>C–and OM–DOM increased dichlorvos sorption (S &lt; S–OM–DOM &lt; S–C–DOM) and decreased chlorpyrifos sorption (S &gt; S–C–DOM &gt; S–OM–DOM). Humified and aromatic nature of DOM determines the interactions with pesticides with different hydrophobic character.</td>
<td>Gaonkar et al. [112]</td>
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<td>Chlorpyrifos</td>
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<td>Organic residues/soil: 2.5 and 5% (w w$^{-1}$)</td>
<td>Shaken: 24 h, T: 25 $^\circ$C</td>
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<tr>
<td>Triasulfuron</td>
<td>Sandy loam soil (pH 7.36, OC 1.20%, clay 17%, silt 25%)</td>
<td>Spent mushroom substrate (pH 7.9, C 26.4%, DOM 1.29%), green compost (pH 7.2, C 23.6%, DOM 0.69%), manure (C 18.5%, DOM 1.32%), sewage sludge (pH 7.6, C 28.9%, DOM 1.18%) Organic residues/soils: 10% (w w$^{-1}$)</td>
<td>Sorbent/Solution: 5 g soil or 0.1 g organic residues/10 mL 0.01 M CaCl$_2$ Herbicide concentration: 1–25 mg L$^{-1}$ (TSF, CTL, FNC) 0.25–10 mg L$^{-1}$ (100 Bq mL$^{-1}$) (PSC)</td>
<td>Highest adsorption for prosulfocarb (lowest water solubility and highest K$_{ow}$) in all materials. Aliphatic and aromatic structures optimize adsorption and O-alkyl and N-alkyl groups enhance desorption hysteresis.</td>
<td>Garcia–Delgado et al. [114]</td>
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<tr>
<td>Prosulfocarb</td>
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<td>Shaken: 24 h, T: 20 $^\circ$C</td>
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<tr>
<td>Chlorotoluron</td>
<td>Loamy sand soil (pH 7.61, OC 0.9%, clay 13%, silt 6%)</td>
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<td>Analytical determination: HPLC/MS and Liquid scintillation</td>
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<td>Flufenacet</td>
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Table 2 includes a summary of the main results obtained from recent literature that has evaluated pesticide mobility in amended soil, including the effects of DOM extracted from organic residues. These studies have been reported for compounds with different properties, and they have generally been conducted under laboratory conditions. These laboratory assays are usually based on two types of soil columns: undisturbed soil cores taken directly in the field to preserve the soil macrostructure within the soil profile [120,121], or packed soil columns [78,96]. The effect of organic amendments on the leaching of pesticides is assessed in both types of columns after washing with a continuous flow of water or aqueous solution of CaCl$_2$ (saturated flow) or a discontinuous flow (saturated/non-saturated flow). Several pesticide leaching studies under different flow regimes have been conducted with columns of soils amended with a wide variety of organic residues, such as olive mill waste [122,123], SMS [96], sheep manure, spent coffee grounds, composted pine bark, and coir [124], BC [125–127], manure and fly ash [128,129], winery vermicompost [130], agro-industrial and composted organic wastes [131], or GC [132]. In general, these studies have revealed a decrease in pesticide leaching because of enhanced pesticide sorption due to the influence of the organic amendments in soils.

The effects of BC as a soil amendment for reducing the movement of pesticides to surface and ground waters has been evaluated together with the effects of increasing pesticide sorption, as previously indicated. Delwiche et al. [133] and Deng et al. [104] have reported a sharp decrease in atrazine leaching after amending the soil with BC from pine chip and from cassava residues, respectively. The presence of more macropores in the BC structure was indicated as being responsible for this decreased atrazine leaching, while pore deformation in BC decreased pesticide leaching in homogenized soil [134]. Tang et al. [135] have reported that BC from bamboo greatly reduces organochlorine pesticide leaching (by up to 65%), and Wang et al. [136] have reported that 60% of sulfamethoxazole is lost/leached in unamended soil and reduced to 2–14% after applying BC. Mendes et al. [105,106] have recently studied the high potential of BC from cow bone to adsorb and reduce/minimize the desorption of the herbicides hexazinone, metribuzin, quinclorac, aminocyclopyrachlor, and mesotrione. An efficient decrease in their leaching in BC-amended soils has been identified even at relatively low application rates.

The simultaneous effect of other organic residues on different processes affecting pesticide dynamics in amended soils, including leaching, has also been also reported. Pérez-Lucas et al. [137] have assessed the effect of two different organic residues (composted sheep manure (EC) and coir (CR)) on the sorption, persistence, and mobility of one herbicide (alachlor (AL)) and two insecticides (chlorfenvinphos (CF) and chlorpyrifos (CP)) with different physicochemical properties. The incorporation of organic wastes in the soil significantly increases the sorption of all the pesticides (especially CF) and decreases their degradation. Leaching experiments conducted in disturbed soil columns have shown a rapid leaching for AL through the unamended soils and a lower retention than for organophosphate insecticides. CF and CP record a significantly lower leaching rate than AL (especially CP) and lower concentrations in leachates. There is a clear decrease in the amount recovered in leachates in the amended soils, except for CP, whose recoveries hardly change. By contrast, Jiang et al. [84] have investigated the effects of sugarcane bagasse compost (SBC) and chicken manure compost (CMC) on the adsorption–desorption, leaching and bioavailability in soil of the fungicide penconazole under laboratory conditions. The autoclave-treated SBC or CMC applied at 2.5% and 5.0% (w w$^{-1}$) promoted the adsorption capacity of soils for penconazole, whereas desorption was radically reduced, with the effect being enhanced by the higher amount of organic amendment (Table 1). Furthermore, column leaching experiments indicated that SBC or CMC limited the transport of penconazole through the soil columns, decreasing its concentration in the soil leachate. These results have revealed major changes in bioavailability experiments under SBC or CMC application, showing that the organic amendments influence the uptake and translocation of penconazole in plants.
The effects of unamended and amended soil properties on the leaching of certain pesticides, such as clothianidin, have been reported. Samarendra–Singh et al. [138] have studied this insecticide’s leaching potential through packed columns with or without farmyard manure (FYM) in a clay loam and a sandy loam soil. The results indicate that the insecticide has less potential to leach in soils with a higher percentage of clay and OC. Likewise, clothianidin may sink to a lower soil profile in the highly porous sandy loam soil under high rainfall conditions, and may contaminate groundwater, so the use of FYM was a good option to reduce its leaching.

Some leaching studies have been conducted under dynamic conditions to explore, together with pesticide characteristics, the changes in their mobility and the potential contamination of groundwater in a more realistic way [96]. Experiments performed under saturated flow conditions simulate the worst-case scenario [125,129,139], while experiments performed under conditions of saturated-unsaturated flow, such as precipitation and irrigation events, simulate real field conditions [123,124]. In general, mobility studies in amended soils have been performed under saturated or unsaturated flow regimes, with fewer studies under both flow conditions or under different states of aging pesticides in amended soils in spite of their importance for determining pesticides’ impact on mobility [96,120,132]. Álvarez-Martín et al. [96] have reported the leaching of two fungicides, tebuconazole (non-polar) and cymoxanil (polar), in packed soil columns, evaluating the influence of SMS at different rates (5% and 50% w w$^{-1}$) as an amendment, and saturated and saturated-unsaturated water flows as leaching regimes. SMS decreased the leaching of tebuconazole in the amended soil under both leaching regimes, while it did not do so for cymoxanil under saturated flow. However, the leached amounts of both fungicides decreased when a saturated-unsaturated flow was applied in SMS-amended soils. A significant decrease in leaching was also observed after fungicide incubation in the column, especially in soil + SMS 50% when both flows were applied. A high SMS dose decreases fungicide leaching but may increase its adsorption in a non-extractable form over time, reducing its bioavailability (non-polar fungicide) or mineralization (polar fungicide). Similarly, Barba et al. [132] have assessed the effects of GC in the leaching of the herbicide prosulfocarb through packed soil columns, with saturated or saturated-unsaturated flows as irrigation regimes and different herbicide incubation times after its application. The amount of herbicide retained in the column was higher in the amended soil under saturated flow than in the unamended one, indicating a stronger interaction between the herbicide and the amended soil. However, the total amounts retained under saturated-unsaturated flow were similar in all the treatments. The incubation time did not significantly affect herbicide retention, but it considerably increased the mineralized amount under saturated flow.

Other investigations in the literature have focused on DOM’s impact on pollutant solubilization. It is often the case that not only the quantity but also the quality of the DOM is a principal factor influencing the potential enhancement of a pesticide’s solubility, besides its physicochemical characteristics [91,140]. Chabauty et al. [141] have studied the effects of DOM from the top soil on the leaching of two pesticides, isoproturon and epoxiconazole, and two pharmaceutical compounds, ibuprofen and sulfamethoxazole, in cultivated soils receiving repeated application of combined compost consisting of green wastes and sewage sludge (SGW). Percolation experiments were performed in undisturbed soil columns with and without DOM from two soils to characterize DOM’s transport and dynamics in the Bt horizons, as well as its effect on water transport and the interactions between organic pollutants and deeper soil horizons. The illuviated Bt layer may act alternatively as a DOM source and sink DOM according to the quality of the percolating solutions. DOM significantly increases the mobility of all the organic contaminants, but the effects fluctuate according to the molecules’ hydrophobic nature and ionic character.
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<tr>
<td><strong>Alachlor</strong></td>
<td>Hypercalcic calcisol (pH 7.9, OC 0.9%, clay 29.1%, silt 33.4%)</td>
<td>Composted sheep manure (EC) (pH 8.3, OC 264.9 g kg(^{-1})) and Coir (CR) (pH 7.5, OC 442 g kg(^{-1})) Organic residues/soils 1%</td>
<td>Packed columns (5 cm i.d. × 30 cm length) of S, S + EC and S + CR. Saturation with CaCl(_2) solution (0.01 M) at maximal WHC. Drainage for 24 h. Determination of PVs (mL) of soil columns. Application of pesticide (1 mL). Leaching volume 750 mL of CaCl(_2) solution (0.01 M) for 10 days. Leached volume/day: 50 mL. Analytical determination: GC/MS</td>
<td>Highest leaching for alachlor. Chlorfenvinphos and chlorpyrifos had low leachability through soil columns (related with their low water solubility). Both compounds were recovered in higher proportions from the soil column than alachlor.</td>
<td>Pérez–Lucas et al. [137]</td>
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<td><strong>Chlorfenvinphos</strong></td>
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<td><strong>Chlorpyrifos</strong></td>
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<td>C 150 µg mL(^{-1})</td>
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<td><strong>Penconazole</strong></td>
<td>Sandy soil (pH 6.20, OC 0.56%, clay 14.92%, silt 33.44%)</td>
<td>Sugarcane bagasse compost (SBC) (pH 6.32, OC 56.16%), chicken manure compost (CMC) (pH 6.27, OC 27.41%). Organic residues/soil: 2.5% and 5.0% (w w(^{-1}))</td>
<td>Glass columns (4.8 cm i.d. × 32 cm length). Pre-saturation with CaCl(_2) solution (0.01 M) (500 mL) for 16 h. Application of pesticide (1 mL). Leaching volume 2500 mL of CaCl(_2) solution (0.01 M) for 50 h. Leached volume 50 mL fractions. Extraction of pesticide from soil column (each 5 cm). Analytical determination: UPLC/TUV</td>
<td>SBC and CMC reduced penconazole leaching by decreased soil porosity and increased adsorption by amended soils. Inhibition of leaching by CMC was lower than by SBC (due to differences in sorption capacity). On the contrary, its content was higher in SBC-soil than in CMC-soil columns.</td>
<td>Jiang et al. [84]</td>
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<td><strong>Clothianidin</strong></td>
<td>Clay loam S1 (pH 5.06, OC 0.95%, clay 30.4%, silt 36.5%) Sandy loam S2 (pH 8.41, OC 0.29%, clay 10.4%, silt 18.1%)</td>
<td>Farm yard manure (FYM) (pH 6.6, OC 23.7%)</td>
<td>Packed soil columns (2.1 cm i.d. × 50 cm length) (200 g S or S + FYM). Pre-saturation overnight in water. Leaching flow: 400 mL of water (1156 mm of rainfall) as continuous flow, or amounts of 20, 40, 80 and 160 mL of water (51.92, 103.85, 207.71 and 415.42 mm of rainfall) as discontinuous flow). Extraction of pesticide from column soil (each 5 cm). Analytical determination: HPLC/PDA</td>
<td>Clothianidin leaching was minimized in S1 compared to S2 after FYM application. Both soils concentrated maximum residue with or without FYM in 0–20 cm soil depth. Clothianidin did not leach under different and discontinuous flow conditions.</td>
<td>Samarendra–Singh et al. [138]</td>
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<td>14C- Tebuconazole 14C -Cymoxanil C 1 mg and 10 kBq mL⁻¹</td>
<td>Sandy clay loam soil (pH 7.52, OC 0.67%, clay 21.1%, silt 11.9%)</td>
<td>Spent mushroom substrate (SMS) (pH 6.97, OC 24.5%, DOM 1.91%)</td>
<td>Pack soil columns (3 cm i.d. × 25 cm length) (100 g S, S + SMS5 and S + SMS50). Chloride as an ion tracer. Leaching flow: 500 mL of CaCl₂ solution (0.01 M) (12 PV). Washing flow regimes: saturated (continuously pumped for ≈8 h) and saturated–non saturated (20 days, 25 mL/day). Columns nonincubated and incubated over 30 days. Analytical determination: Liquid scintillation</td>
<td>Amendments decreased leaching of tebuconazole under different flow conditions, and decreased leaching of cymoxanil under saturated-unsaturated flow. Ageing favored retention decreases tebuconazole leaching or cymoxanil mineralization.</td>
<td>Álvarez–Martínez et al. [96]</td>
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<tr>
<td>14C-Prosulfocarb C 1 mg and 10 kBq mL⁻¹ (2.5 times the agronomic dose)</td>
<td>Sandy clay loam soil (pH 7.35, OC 1.30, clay 17%, silt 25%)</td>
<td>Green compost (GC) (pH 7.20, OC 24.1%, DOM 0.703%)</td>
<td>Pack soil columns (3 cm i.d. × 25 cm length) (100 g S and S + GC). Chloride as an ion tracer. Leaching flow: 500 mL of CaCl₂ (0.01 M) (12 PV) under saturated flow and under saturated–unsaturated flow (20 mL/ day). Columns nonincubated and incubated over 28 days. Analytical determination: Liquid scintillation</td>
<td>Leached amounts decreased in S and S + GC columns after incubation. Retained amounts were lower in S than in S + GC columns under saturated flow. Prosulfocarb was retained in the first segment of columns under all conditions. Herbicide incubation increased the mineralized amount under saturated flow.</td>
<td>Barba et al. [132]</td>
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<td>Isoproturon C 47–59 µg L⁻¹</td>
<td>Loamy soil Ap (0–28 cm)</td>
<td>Combined compost of sewage sludges and green wastes (SWG) (pH 6.9, OC 15.8 g kg⁻¹)</td>
<td>Undisturbed soil cores (14 cm i.d. × 30 cm length) with 5300 cm³ volume. Bromide as ion tracer. Leaching experiments with synthetic water, DOM of soil or soil + SWG. Unsaturated steady–state flow regime of two consecutive rainfalls of 1.76 mm h⁻¹ intensity and separated by a 1–week flow interruption on triplicated cores for 28 days. Analytical determination: UHPLC/MS/MS</td>
<td>DOM increased mobility of Br- and all pollutants. The mobility increase was greater for more hydrophobic compounds (epoxiconazole and ibuprofen). DOM can also enhance the transport of anionic molecules but for these compounds also depend on their affinity for the soil matrix including soil solution composition and its pH.</td>
<td>Chabauty et al. [141]</td>
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3.3. Effect of Organic Residues on Pesticide Dissipation

Pesticide degradation refers to its breakdown into environmentally stable and compatible substances, and together with adsorption it is the main process that determines its fate in soil and water environments [142]. Data on the degradation kinetics of pesticides are necessary for characterizing their concentration over time and evaluating their persistence when applied in the field, as well as their potential threat to both target and non-target species.

Pesticide degradation in soils is informed by abiotic and biotic factors. Abiotic degradation involves hydrolysis and photodegradation, while biotic degradation involves the reactions of microorganisms (mainly bacteria and fungi). Microbial degradation is generally considered to be mainly responsible for pesticide breakdown, resulting in less toxic compounds and affecting even the most persistent pesticides [143–146]. The level of biodegradation can fluctuate from minor changes, without significantly compromising a pesticide’s chemical and toxicological properties, to the compound’s complete mineralization. In general, soil microorganisms use pesticides as a direct source of energy and nutrients [75], or through co-metabolism [147]. The main factors affecting pesticide degradation include different environmental aspects (soil type, soil OM, irrigation, weather -mainly temperature and precipitation), pesticide formulation (individual or combined compounds), and application method (single or repeated application) [148].

The control of the pesticide adsorption–desorption processes in soil by SOM and DOM from the aforementioned organic amendments may affect their dissipation (Figure 1). In general, SOM favors the adsorption of pesticides, and this process controls their degradation, as it determines their availability in the soil. Adsorption decreases their bioavailability for degradation by soil microorganisms and the rate at which they are degraded [75,83,149]. Organic amendments could therefore modify the persistence and dissipation of pesticides in soils by enhancing their adsorption as a consequence of increasing soil OC [77–79]. In some cases, organic materials have decreased the half-life of a pesticide, while in others there has been an increase or even no effect at all [79,150,151]. In turn, DOM may also affect the degradation and mineralization kinetics of pesticides by (i) decreasing pesticide adsorption and enhancing its bioavailability or by contrast, (ii) increasing the adsorption of the compounds and decreasing their bioavailability according to some of the different processes explained in Section 3.1 [94,101,140,152]. DOM’s influence in pesticide biodegradation depends on its source, nature and concentration, and therefore on soil type and pesticide characteristics [153]. Simultaneously, OM, nutrients, and sometimes the microorganisms in the amendment may modify/increase the activity of soil microbial communities, and the stimulation of microbial activity may increase pesticide degradation [154–156]. In fact, most of the organic residues allow the growth of fungi, which produce extracellular ligninolytic enzymes that promote pesticide degradation [157].

Table 3 includes a summary of the main results reported in recent experiments carried out under laboratory conditions, evaluating the effect that the application of organic residues has on the bioavailability, degradation, dissipation and/or persistence of pesticides in amended soils. In general, different factors have been considered, such as soil amendments of different nature, origin, rates or incubation times, and pesticides with different characteristics and behavior.

BC’s effect on the dissipation of pesticides has been widely investigated, together with its impact on the adsorption–desorption and leaching processes. Yavari et al. [146] have studied the effects that two BCs obtained from the empty fruit bunch (EFB) of oil palm and rice husk (RH) have on the photolysis and biodegradation of the herbicides imazapic, imazapyr, and their combination (OnDuty®). Polar herbicides are resistant to hydrolysis degradation, and photolysis rates are reduced significantly in the presence of BC. EFB-BC had a greater effect on the increase in DT_{50} due to its chemical composition and surface functional groups. In contrast to photolysis, the biodegradation of imidazolinones is significantly accelerated in the presence of both BCs, although RH-BC has a greater impact. The effects of other BCs derived from rice straw (RS), corn straw (CS), chicken
manure (CM) and tire rubber (TR) on the dissipation of the fungicide cyazofamid and its metabolite CCIM (4-chloro-5-p-tolylimidazole-2-carbonitrile) have been assessed by Tang et al. [158]. Cyazofamid dissipation is either accelerated or depressed, depending on BC type, application rate, and soil moisture. Adsorption, hydrolysis and microbial degradation are all involved in cyazofamid dissipation. CM- and CS-BC enhance fungicide dissipation through the biodegradation related to changes in the microbial community. The residual amounts of metabolite CCIM increase by 8–15 times after BC application, regardless of BC type and cyazofamid dissipation. Tang et al. [158] provided new insights into BC application for soil remediation and highlighted the potential risk of the build-up of metabolites in BC-amended soil, especially when the metabolites have higher toxicities than the parent compound.

The impact on pesticide degradation of other organic amendments applied to soils has been investigated for the herbicide imazapic under various environmental conditions, including temperature, soil moisture, and soil pH [159]. CM and biogas slurry (BS) have been applied at different rates, and herbicide degradation has been studied at different temperatures (15, 25 and 35 °C), soil moisture content (15%, 40%, 60% and 90%), and soil pH (6.0, 7.0 and 8.0). The imazapic degradation rate increases with higher temperature, soil pH, and soil moisture, and decreases with OM content. A long residual period of herbicide has been observed, and microbial degradation has been proposed as the primary degradation mechanism (high temperatures and moist soil conditions favor microbial activity and accelerate imazapic degradation). Microbial degradation has been correlated with soil properties and adsorption. The combined effect of temperature and organic amendments on the degradation of chlorotoluron and flufenacet has also been studied by Marín-Benito et al. [160]. They have assessed herbicide degradation kinetics at two different temperatures, 6 and 16 °C, in an unamended soil and one amended with SMS and GC, and the formation of the main metabolites of both herbicides over degradation time. The degradation of flufenacet, the more hydrophobic herbicide, was slower than chlorotoluron in all the treatments. Nonetheless, the addition of SMS and GC to the soil decreases their degradation rates because of their higher sorption and lower bioavailability. This impact depends on the herbicide and incubation temperature. Faster degradation has been confirmed for both herbicides at 16 °C, possibly due to increased microbiological activity with the higher temperature, and a significant presence of metabolites has been observed in all the soil treatments. The effect of GC as soil amendment on the dissipation of the herbicide prosulfocarb (14C-labeled) has also been assessed by Barba et al. [161], determining the dissipation mechanisms and the total mass balance of herbicide applied at two rates (4 and 10 mg kg⁻¹). The herbicide records higher bioavailability and faster dissipation in the unamended soil than in the amended one due to its lower sorption, irrespective of the dose applied. A temporal decrease in extractable amounts has been observed in all the soil treatments, revealing herbicide aging over time. Prosulfocarb mineralization and non-extractable residues record a slow increase over incubation time. At the end of the dissipation period, the total 14C mass balance is approximately 60–70% under all conditions, indicating that part of the herbicide has probably been lost through volatilization. Hence, GC influences herbicide bioavailability by decreasing its biodegradation.

As regards the impact that the pesticide application method (single or repeated) has on pesticide dissipation in the soil, it has been frequently reported in unamended soils with significant conclusions. On the one hand, the repeated application of pesticides to the same soil over a long period is a common agricultural practice, and it can influence pesticide dissipation, accelerating biodegradation by adapted soil microorganisms, which may reduce or even eliminate the pesticide’s efficacy for controlling the target pest, disease or weed [162,163]. On the other hand, pesticide residues could remain in the soil after repeated applications, affecting soil microbial function and structure. In some cases, repeated application may have a detrimental effect on the microbial community by degrading the pesticide, significantly compromising its performance [163,164]. In amended soils, however, laboratory studies on pesticide dissipation have been conducted mainly with a
single application of just one compound [29,165,166], while only a handful of studies have reported the effects that repeated and/or combined pesticide applications have on their dissipation in amended soils. Pose-Juan et al. [167] have studied the effects of two repeated applications of mesotrione, pethoxamid, and triasulfuron on their dissipation rates in unamended soil and soil amended with GC or SS. Repeated herbicide application in soils has prompted dissipation effects that differ to those observed with only a single application. For mesotrione, this repeated application has an inhibitory effect on its dissipation rate in all the soils studied, and the herbicide could become more persistent or increase its accumulation in amended soils. However, repeated applications of pethoxamid and triasulfuron accelerate their dissipation in amended soils. The formation of bound residues over time has been suggested as the process responsible for an apparent dissipation, decreasing the bioavailability of the compound to be degraded. The dissipation mechanism of these herbicides has been supported by the adsorption constants determined, indicating the highest adsorption by amended soils (Table 1).

4. Conclusions

This review evaluates how the dynamics of pesticides in soil is modified by the application of organic residues as soil amendments. By focusing on the main processes (adsorption–desorption, dissipation and leaching) that govern the fate of pesticides in soils, the review highlights the complexity of extrapolating the behavior from some pesticides to others in the presence of different organic residues, even in the same soil. Multiple reasons intrinsically related to soil, pesticide and organic residue properties and composition limit this extrapolation. However, all these factors and the combinations tested at laboratory scale constitute a major limitation. Working at laboratory scale represents the platform, and first step of any subsequent successful research because it involves doing so under controlled conditions oriented towards target variables. This allows analyzing the impact of the application of organic amendments to soils on an individual basis, influencing the environmental fate of pesticides and testing different variables to optimize the coexistence of both agricultural practices.

5. Future Perspectives on the Application of Pesticides and Organic Amendments in Soils

Once the influence of organic amendments on an individual pesticide has been studied and documented by laboratory assays, the next step should be to extend these studies to field scale under real conditions. This will allow considering additional factors such as the undisturbed soil structure, precipitation, temperature, irrigation, or the presence of crops affecting the dynamics of pesticides through the uptake process or simply controlling for water dynamics. A holistic view jointly evaluating the different pesticide processes and their interactions affected by the application of the organic amendments will contribute to a more complete and more effective interpretation of the environmental fate of pesticides in amended soils. Despite the potential relevance of the results obtained in this kind of study, only a few assays combine pesticides-organic amendments at field scale [149,168–170].

Finally, the third step prior to the use of organic residues as soil amendments from a beneficial agronomic point of view that avoids compromising the sustainability of soil and water resources would be to simulate the behavior of pesticides in amended soil scenarios. There is an important gap in this research field, as evidenced by the scarce studies reported in the literature, especially using the pesticide fate models usually applied in a regulatory context [171–173].

The optimization of the use of pesticides together with organic amendments in soil therefore calls for field studies to assess pesticide behavior under different amended soil scenarios, together with models for simulating and predicting pesticides’ environmental fate as future perspectives.
Table 3. Dissipation of pesticides in amended soils.

<table>
<thead>
<tr>
<th>Pesticide</th>
<th>Soil Characteristics</th>
<th>Organic Amendment/Dose</th>
<th>Experimental Design</th>
<th>Results</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Imazapic</td>
<td>Clay loam soil (pH 6.36, OC 0.99%, clay 37.9%, silt 21.58%)</td>
<td>Biochar from fruit bunch of oil palm (EFB) (pH 6.13, C 58.60%) and rice husk (RH) (pH 6.32, C 48.26%)</td>
<td>Hydrolysis: Solution/herbicide concentration 50 mL/10 mg L(^{-1}), pH 3, 7 and 9 Photodegradation: Soil/herbicide 30 g/0.2 µg g(^{-1}). Biodegradation: Soil/herbicide 10 g/0.2 µg g(^{-1}). T: 25 °C. Relative humidity: 85%. Time up to 70 days. Analytical determination: HPLC</td>
<td>All herbicides were resistant to hydrolysis degradation. Photolysis rates of herbicides were reduced by use of biochar, particularly EFB. Biodegradation of herbicides accelerated significantly by the use of biochars.</td>
<td>Yavari et al. [146]</td>
</tr>
<tr>
<td>Imazapyr mixture</td>
<td></td>
<td>Biochar from fruit bunch of oil palm (EFB) (pH 6.13, C 58.60%) and rice husk (RH) (pH 6.32, C 48.26%)</td>
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<tr>
<td></td>
<td>(Onduty\textsuperscript{®})</td>
<td>Biochar/soil: 1.0% (w w(^{-1}))</td>
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<tr>
<td>Cyazofamid</td>
<td>Silty soil (pH 7.14, OC 0.71%, clay 8.25%, silt 85.55%)</td>
<td>Biochars from rice straw (RS) (pH 9.87, C 36.58%), corn straw (CS) (pH 9.97, C 57.33%), chicken manure (CM) (pH 8.16, C 27.73%) and tire rubber (TR) (pH 8.82, C 74.60%)</td>
<td>Soil samples: 20.6 g Pesticide applied: 2.5 mg/kg of dry soil Soil moisture content: 40% WHC Incubation T: 25 °C Sampling times up to 40 days Analytical determination: HPLC/MS/MS</td>
<td>Cyazofamid dissipation order: CS &gt; RS &gt; CM TR depressed cyazofamid dissipation. Adsorption, hydrolysis and microbial degradation were all involved in its dissipation. CM and CS enhanced the cyazofamid dissipation by biodegradation. CCIM residual increased by 8–15 times after biochar application, regardless of biochar type.</td>
<td>Tang et al. [158]</td>
</tr>
<tr>
<td>Metabolite CCIM</td>
<td></td>
<td>Biochar/soil: 3% (w w(^{-1}))</td>
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<tr>
<td>(4-chloro-5-p-tolylimidazole-2-carbonitrile)</td>
<td></td>
<td>Biochars from rice straw (RS) (pH 9.87, C 36.58%), corn straw (CS) (pH 9.97, C 57.33%), chicken manure (CM) (pH 8.16, C 27.73%) and tire rubber (TR) (pH 8.82, C 74.60%)</td>
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<td>Tang et al. [158]</td>
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<tr>
<td>Imazapic</td>
<td>Silty soil (pH 8.1, OM 0.55%, clay 9.1%, silt 69.1%)</td>
<td>Chicken manure (pH 7.1, OM 21.5%), biogas slurry (pH 7.0, OM 20.1%) CM/soil: 2.1–16%; BS/soil: 3.6–0.9%</td>
<td>Soil samples: 1000 g Pesticide applied: 20 mL (50 mg L(^{-1})) Incubation T: 15, 25 and 35 °C Soil moisture contents: 15%, 40%, 60%, and 90% pH values: 6.0, 7.0, and 8.0 Sampling times up to 150 days Analytical determination: HPLC/MS/MS</td>
<td>Imazapic degradation rate increased with temperature, soil pH, and soil moisture, and it decreased with OM content. Biogas slurry accelerated imazapic degradation (significant microbial contribution to its degradation).</td>
<td>Su et al. [159]</td>
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<td></td>
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<td>Biochar/soil: 3% (w w(^{-1}))</td>
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<tr>
<td>Chlorotoluron</td>
<td>Sandy loam soil (pH 6.34, OC 0.77, clay 14.9%, silt 4.7%)</td>
<td>Spent mushroom substrate (SMS) (pH 7.9, OM 59.4%, DOM 0.8%) Green compost (GC) (pH 7.2, OM 46.0%, DOM 0.7%). SMS and GC/soil: 140 and 85 t ha(^{-1})</td>
<td>Soil samples: 600 g Pesticide applied: 14 mg (chlorotoluron) and 5.5 mg (flufenacet)/kg of dry soil Soil moisture content: 40% WHC Incubation T: 6 and 16 °C Sampling times up to 67 or 273 days Analytical determination: HPLC/MS</td>
<td>Flufenacet degradation was slower than that of chlorotoluron. Amendments increased DT\textsubscript{50} values for both herbicides incubated at both temperatures, especially at 16 °C due to the higher microbiological activity.</td>
<td>Marin–Benito et al. [160]</td>
</tr>
<tr>
<td>Pesticide</td>
<td>Soil Characteristics</td>
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<tr>
<td>14C- Prosulfocarb</td>
<td>Sandy clay loam soil (pH 7.35, OC 1.30, clay 17%, silt 25%)</td>
<td>Green compost (GC) (pH 7.20, OC 24.1%, DOM 0.703%) GC/soil: 20% w w⁻¹ (180 t ha⁻¹)</td>
<td>Soil samples: 500–700 g Pesticide applied: 4 and 10 mg kg⁻¹ of dry soil and 100 Bq g⁻¹ Soil moisture content: 40% WHC Incubation T: 20 °C Sampling times up to 50 days Analytical determination: HPLC/MS and Liquid scintillation</td>
<td>Highest DT₅₀ values in amended soil. They increased with the herbicide concentration in unamended soil but decreased in amended soil. Lost through volatilization of herbicide was consistent with the total ¹⁴C mass balance close to 70% at the end of dissipation period.</td>
<td>Barba et al. [161]</td>
</tr>
<tr>
<td>Mesotrione</td>
<td>Sandy loam soil (pH 6.3, OC 0.49%, clay 10.7%, silt 5.9%)</td>
<td>Sewage sludge (SS) (pH 7.08, OC 8.06%, DOM 0.102%) Green compost (GC) (pH 6.73, OC 27.0%, DOM 2.17%) SS or GC/soil: 50 t ha⁻¹</td>
<td>Soil samples: 800 g Pesticide applied: 2 mg kg⁻¹ of dry soil Soil moisture content: 40% WHC Incubation T: 20 °C Sampling times up to 99, 43 and 144 days Analytical determination: HPLC/MS</td>
<td>Repeated application of pesticides decreased (mesotrione) or increased (pethoxamide) its dissipation rate in all treatments. For triasulfuron, it increased only in amended soils. Highest DT₅₀ values for pethoxamid and triasulfuron in S + GC, and for mesotriona in S + SS.</td>
<td>Pose et al. [167]</td>
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</table>
Author Contributions: Conceptualization, M.J.C., M.J.S.-M., M.S.R.-C. and J.M.M.-B.; writing—original draft preparation, M.J.C., M.J.S.-M. and J.M.M.-B.; writing—review and editing, M.J.S.-M., M.S.R.-C. and J.M.M.-B.; supervision, M.J.S.-M., M.S.R.-C. and J.M.M.-B.; project administration, J.M.M.-B.; funding acquisition, J.M.M.-B. All authors have read and agreed to the published version of the manuscript.

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