Difficulties in Potato Pest Control: The Case of Pyrethroids on Colorado Potato Beetle

Imola Molnar and Elena Rakosy-Tican *

Abstract: Colorado potato beetle (Leptinotarsa decemlineata, CPB) is considered one of the most persistent crop pests because it is highly adaptable and can rapidly develop insecticide resistance. Nowadays, this beetle is resistant to over 54 different insecticides. In the absence of competitive alternatives, the answer of farmers to the high degree of pest adaptability is to increase the number of pesticide treatments or apply chemicals with different modes of action. Such a strategy increases the risk of intoxication in non-target organisms and leads to environmental pollution, augmenting the carbon footprint. Furthermore, these strategies are also unsustainable and inefficient for pest management in the long-term. The time has thus come to reform existing agriculture practices, for which the implementation of an integrated pest management strategy would be a more feasible tool. Applying a sustainable pest management strategy is indispensable for a better understanding of the status and mechanisms of insecticide resistance. Effective pest management requires monitoring the resistance of pests and developing a well-programmed pesticide treatment to simultaneously reduce the insecticide selection pressure and environmental pollution. In this context, here we present a review on the difficulties of potato pest control using as a case study the resistance of CPB to pyrethroids.

Keywords: Colorado potato beetle; insecticide resistance; potato; pyrethroids

1. Introduction

Pesticides are natural or synthetic substances used to kill, deter or control undesirable organisms. The pesticides family includes herbicides, insecticides, fungicides and disinfectants for preventing the spread of bacteria or viruses [1]. The toxicity of pesticides is determined not only by the concentration of the substances but also by the exposure time [2]. The irresponsible use of these chemicals substantially increases our environmental footprint. Generally, small residues of synthetic pesticides can persist for a long time in the environment (soil, fresh or sea water, on the surface of vegetables or even inside the crops and derived products), which also increases the possibility of poisoning non-target organisms, including humans. Nevertheless, global food production will need to increase in order to feed the increasing population, which will drive an intensification of agricultural production and lead to an increased demand for agrochemicals [3].

Despite the extensive use of pesticides with various active ingredients, an estimated 34% of all potato crops are lost annually to insect pests. Major pathogens (Phytophthora infestans, Alternaria solani, Thanatephorus cucumeris), viruses (potato leafroll virus, potato virus Y) and animal pests (potato cyst nematodes, Colorado beetle, Phthorimaea operculella) are widely distributed, which can wreak havoc on potato fields if left uncontrolled. Without crop protection almost 75% of potato production would be lost [4,5].
2. A Brief Historical Overview of Insecticide Use

Botanical preparations (i.e., *Dalmatian pyrethrum* flowers, elderberry flowers, tobacco infusions) were the first substances used as insecticides by ancient people. The next major stage was the use of inorganic insecticides such as arsenic and boric acid, which offered only marginal control of insects. Starting in the middle of the 19th century, pest control with insecticides gained wider use. For example, Paris green (mixed copper acetoarsenite) was successfully used to control the Colorado potato beetle, *Leptinotarsa decemlineata* (Say), (CPB in many countries around the world. A step forward in the early 20th century was the discovery of synthetic organic insecticides, which opened new horizons in pest control [6]. In 1939, the insecticidal activity of dichlorodiphenyltrichloroethane (DDT) was discovered, and this substance led to the development of chlorinated compounds like lindane and hexachlorocyclohexane. These chlorinated insecticides dominated the market for nearly three decades, when another promising insecticide group was developed, the organophosphates (parathion, malathion and diazinon) [7]. In the middle of the 20th century, synthetic carbamates were introduced. The most widely used carbamates—carbaryl and aldicarb—are derivatives of physostigmine obtained from Calabar bean [8]. The success of pyrethroids started in the 1970s when permethrin and deltamethrin were discovered [9]. The main advantage of pyrethroids compared to other synthetic insecticides was the requirement of lower doses to achieve the expected potent effect. In the last decades, neonicotinoids (i.e., imidacloprid and dinotefuran) were developed, which are systemic insecticides and therefore used to treat seeds to provide plant protection against pests from germination until full development [9–11].

3. Types of Insecticides Affecting Different Membrane Channels

Nowadays, different classes of insecticides are available on the market, which have a wide range of mechanisms of action. This section only includes those insecticides that act directly or indirectly on different membrane channels. Understanding the various actions of insecticides on ion channels will be important when the cellular mechanisms and effects of pyrethroids are discussed. The other types of insecticides are equally important, but they are not closely related to the current topic. The newest information about the existing insecticide classes and their mode of action was recently published by the Insecticide Resistance Committee (IRAC) [12].

Ryanodine and synthetic diamides are ligands of ryanodine receptors (RyRs), which are members of the ligand-gated calcium channel family and are mainly located on the sarcoplasmic reticulum of muscle cells and the endoplasmic reticulum of neuronal cells [13,14]. Ryanoids and synthetic analogs induce cessation of feeding, lethargy, paralysis, and death. The molecular background of these symptoms is based on the prolonged activation of RyRs by ryanoids, which leads to an elevated level of cytoplasmic calcium. The increased concentration of Ca$^{2+}$ in muscle cells induce sustained skeletal muscle contraction and paralysis, while in neurons induce cytotoxicity and altered neuronal activity (Figure 1) [13,15].

Neonicotinoids (e.g., imidacloprid, dinotefuran) represent a large group of insecticides that act selectively on the nicotine receptors of acetylcholine (nAChRs), which play a crucial role in synaptic transmission in the central nervous system of insects [7]. The nAChRs are activated through the binding of acetylcholine (ACh) neurotransmitters and are deactivated when the acetylcholinesterase enzyme (AChE) rapidly hydrolyzes the ester linkage in acetylcholine. The same activation mechanism of nAChRs also occurs in the presence of neonicotinoids. Only the deactivation process is missing, because AChE cannot break down and inactivate these types of insecticides. This persistent activation leads to an overstimulation of the cholinergic synapses and results in hyperexcitation, convulsions, paralysis, and finally insect death (Figure 1) [16,17]. In the last decades, neonicotinoids were widely used chemicals to control undesirable organisms. Some studies pointed out that neonicotinoids can accumulate in the environment, which increases the risk of long-term intoxication [18]. In addition, it was confirmed that this group of insecticides represents a danger for honey and wild bees. Therefore a ban on outdoor use of several
neonicotinoids (clothianidin, imidacloprid, thiamethoxam (2013) and thiacloprid (2020)) was implemented in European countries [17,19].

Figure 1. Molecular actions of different classes of insecticides on ion channels. In normal conditions, the alternation of activation and inhibition of ion channels are essential in neurotransmission. Any external factor which upsets this balance prevent the formation of normal membrane potentials, which leads to unconventional neurotransmission (e.g., hyperexcitation of cells). The induced irregular neurotransmission may produce different symptoms (e.g., convulsions, paralysis) and even lead to death. (Abbreviations: GABA = γ-aminobutyric acid, Ach = acetylcholine, AChE = acetylcholinesterase, RyR = Ryanodine receptor, VGSC = voltage-gated sodium channel, GABAR = GABA receptor, nAChR = nicotinic acetylcholine receptor, Na = sodium ion, Cl = chloride ion, K = potassium ion, Ca = calcium ion).

The organophosphates (OPs) and carbamates are inhibitors of the acetylcholinesterase enzyme (AChE), which increase synaptic concentrations of acetylcholine leading to hyperexcitation of the CNS (Figure 1). The signs of intoxication of organophosphates include restlessness, hyperexcitability, tremors, convulsions, and paralysis. In mammals there is also a significant involvement of peripheral cholinergic systems in OP and carbamate poisoning, leading to the so-called Salivation, Lachrymation, Urination, Defecation, Gastrointestinal upset and Emesis (SLUDGE) syndrome [20].

The insecticides 4,4′-diisothiocyanato-2,2′-stilbenedisulfonic acid (DIDS), anthracene-9-carboxylic acid, 5-nitro-2-(3-phenylpropylamino)benzoic acid (NPPB) and indanyloxyacetic acid (IAA-94) inhibit the current of the voltage-gated chloride channel (VGCC). These compounds function as dietary insecticides and present dose-dependent effects with low mammalian toxicity rates [21].
The lindane, endosulfan, isoxazoline, and metadiamide insecticides inhibit the proper functioning of the γ-aminobutyric acid (GABA) receptors, therefore, affect Cl− influx (Figure 1) [22]. Only lindane and endosulfan are still in use, but they are also gradually being phased out due to their high mammalian toxicity and especially, their high dermal toxicity.

Both types of chloride channel blockers (VGCC-, GABA receptor-blockers) induce CNS hyperexcitability and convulsions not only in insects, but also in mammals [21,22].

All presented substances alter normal neurotransmission by modifying the functions of different ion channels. These insecticides often overstimulate the firing of excitatory neurons, which lead to hyperexcitation, or they achieve the same effect in a completely different way by blocking the inhibitory neurotransmission (Figure 1).

3.1. Pyrethroids

Pyrethroids are synthetic structural derivatives of insecticidal pyrethrins found in the pyrethrum extract of Chrysanthemum species and can be characterized by having a high degree of lipophilicity [23]. Pyrethroids belong to the fourth group of insecticides according to the WHO classification and include 42 substances [24]. Natural pyrethrins consist of six compounds: pyrethrin I and II, jasmolin I and II, and cinerin I and II. Pyrethrum extract is the most effective natural insecticide, but it is unstable in light and air, which decreases its effectiveness in crop protection. During the development of synthetic pyrethroids, the main goal was to modify the structure of pyrethrins to become more stable in lightning conditions, but do not decrease the potency and rapidity of insecticidal activity and/or have relatively low mammalian toxicity [25]. Pyrethroids can be classified into Type I or Type II groups, depending on the alcohol substituent in the molecule. The members of Type I group contain descyano-3-phenoxybenzyl or other alcohol chain moieties. The older Type I compounds without phenoxybenzyl alcohol (e.g., bioallethrin, tetramethrin) are relatively unstable in outdoor conditions. Therefore they were mainly used as household disinsectants for flying or crawling insects. The introduction of phenoxybenzyl (e.g., permethrin) or other halogenated alcohols (e.g., tefluthrin) improved the chemical stability of these compounds, which opened the possibility of applying pyrethroids in the field. The Type II pyrethroids contain α-cyano-3-phenoxybenzyl alcohol, which yields a 3-6-fold increase in its insecticidal performance compared to the cognate non-cyano analogs (Figure 2) [26].

![Figure 2](image_url)

**Figure 2.** Chemical structure of pyrethroid insecticides (From left to right: pyrethrin I, permethrin, cypermethrin). The introduction of phenoxybenzyl alcohol moiety into the chemical structure of natural pyrethrins increases the stability of the Type I pyrethroids. All Type II pyrethroids have a cyanide group.

The chemical structure of pyrethroids is highly flexible with many attainable geometries. Therefore, they can be used as mixtures of active isomers (e.g., allethrin) or as single isomers (deltamethrin and esfenvalerate) [27].
3.2. Toxication Symptoms of Pyrethroids

The new generation of synthetic pyrethroids was selected to be less sensitive to photodegradation, but despite this property, in general, pyrethroids are still highly photolabile which greatly reduces their potential for mammalian poisoning. They proved to be 2250 times more toxic to insects than vertebrates due to the smaller size of insects, lower body temperature and more sensitive sodium channels [28]. A single high dose of Type I pyrethroids could cause hyperexcitability and convulsions in insects and whole-body tremor, furthermore aggression, convulsive twitching, named as T syndrome in mammals. Whereas Type II pyrethroids induce ataxia and incoordination in insects, while in mammals, they produce choreoathetosis, salivation, clonic seizures named as CS syndrome [11].

Pyrethroids may enter the body through skin contact, inhalation and food or water ingestion [29]. Furthermore, pyrethroids are toxic to aquatic animals, e.g., shellfish (crayfish, lobster), and fish (Danio rerio Hamilton-Buchanan, carp and rainbow trout), affecting not only the ion channels in neuronal membranes, but also in the mitochondrial membranes [30–34]. All pyrethroids are rapidly metabolized in mammals and their metabolites are almost completely excreted within several days, except for the cyano moiety. Therefore, tissue residues in mammals are relatively low [10,35]. The major metabolic reactions of pyrethroids are the following: oxidation of methyl groups and aromatic rings; hydrolysis of the ester linkage; and several types of conjugation reactions, which are mediated by carboxylesterases and members of cytochrome P450 [36].

Even though pyrethroids are relatively safe insecticides rapidly metabolized in mammals, chronic exposure or a high single dosage of pyrethroids can result in intoxication with different severity of symptoms, or even death [37]. After the consumption of different types of food: pasta, rice, whole meal bread, and fruits from sprayed areas, the metabolites of pyrethroids were detected in the urine [38,39]. In mammals, age-related sensitivity was observed to pyrethroids, for example: deltamethrin was more toxic to young rats than to adults, or cypermethrin proved to be more toxic in younger individuals [40]. Furthermore, due to the lipophilicity feature of pyrethroids, they can easily pass through the blood-brain barrier and at high concentrations they can also have neurotoxic effects [41,42]. Developmental pyrethroid exposure results in alterations of the nervous system structure and functions, which manifests in deficits of motor coordination, learning, and memory performance. Long-term exposure of children to permethrin caused an increase in the number of permethrin metabolites in the urine, as well as behavioral changes, in particular an increase of aggressive behaviors and sleep disorders [43]. Furthermore, children were characterized by impaired memory, poorer verbal abilities and decreased intelligence scores [44]. Many other side effects of permethrin exposure were observed: household usage of permethrin is often associated with allergies and asthma, especially in children. Skolarczyk et al. [45] demonstrated an immunosuppressive effect of pyrethroids in the case of children. Furthermore, permethrin metabolites were detected in breast milk from women in Brazil, Colombia and Spain [46]. Daily exposure to pyrethroids leads to neuronal cell loss in various brain regions, which is a result of increased production of reactive oxygen species, increased expression of pro-apoptotic genes: p53 and Bax, and decreased expression of anti-apoptotic Bcl-2 [40,47]. Nonetheless, the U.S. Centers for Disease Control recommends the use of pyrethroid repellents as safe substances to protect pregnant women against Zika infection [48,49].
4. Mechanism of Action of Pyrethroids

4.1. Pyrethroids Effects on Voltage-Gated Sodium Channels (VGSCs)

Pyrethroids are predominately absorbed through the insect cuticle, and after reaching the nervous system, disrupt the membrane potential and neurotransmission, causing knockdown and subsequently death. Under normal conditions, nerve impulses (action potential) consist of arranged ion movements through specific ion channels. The action potential has three distinct phases: in the initial step, nerve impulses open the VGSCs and Na\(^+\) ions flow into the cells leading to membrane depolarization (positive wave), followed by a repolarization wave from the efflux of K\(^+\) ions when the value of the membrane potential becomes negative again; finally comes the restoration of the resting potential by the active transport of Na\(^+\)/K\(^+\) pump. The generated action potential moves towards the axon until reaching the nerve terminal, which induces different neurotransmitters [50].

VGSCs consist of two subunits: \(\alpha\) subunit, which forms the pore and an auxiliary subunit. The \(\alpha\) subunit contains six transmembrane segments (S1–S6), where S4, S5 and S6 segments form the channel pore and confer selectivity for Na\(^+\). The S6 segment is involved in the activation gating of the channel allowing Na\(^+\) inflow. During the resting potential and repolarization phase of an action potential, the pores of VGSCs are closed [51,52]. In insects, a single gene encodes the VGSC \(\alpha\) subunit, while mammals have nine distinct pore-forming \(\alpha\) subunit genes. Sodium channel isoforms expressed in various mammalian tissues exhibit different sensitivity to pyrethroids. Mutations in the sodium channel could induce pyrethroid resistance. This phenomenon was observed in case of several model species and insect pests (Drosophila melanogaster, Xenopus laevis and Colorado potato beetle). The majority of resistance-associated mutations are found in the intracellular linkers between S4, S5 or within transmembrane helices S5 and S6. Genetic mapping of pyrethroid resistance genes in insects clarified the pyrethroid-binding site on VGSC that was localized between the transmembrane helices S4 and S5 segments [53].

Pyrethroids delay VGSC inactivation resulting in a prolonged Na\(^+\) current, which leads to hyperexcitation of neurons. In general, Type II compounds delay the inactivation of sodium channels substantially longer than Type I compounds do, which induce multiple spike discharges in neurons [54]. Type II pyrethroids depolarize the ax on membrane potential, which reduces the amplitude of the action potential and eventually leads to the loss of electrical excitability. All these effects occur mainly because pyrethroids extend the duration of the Na\(^+\) flowing through sodium channels by slowing down or preventing their inactivation [54]. Moreover, the prolonged Na\(^+\) flow increases the release of neurotransmitters from presynaptic nerve terminals. The different effects of Type I and Type II pyrethroids are due to the variance in the duration of modified sodium currents: Type I compounds produce long action potentials (burst discharges) as a result of a single stimulus with little or no effect on the resting potential, while Type II compounds do not induce repetitive discharges, they cause a use-dependent blockage of the action potential coupled with depolarization of the resting potential [11].

Pyrethroid insecticides primarily target VGSCs, but also affect the function of other voltage-gated channels such as calcium and chloride channels and the binding of pyrethroids to GABA receptors was also observed. The effects of pyrethroids on these alternative targets have been demonstrated mainly in mammals, but recent studies showed that these membrane proteins are also targeted by pyrethroids in insects [53,55–57].

4.2. Alternative Targets of Pyrethroids

Several studies have demonstrated that pyrethroids influence neuronal voltage-gated calcium channels (VGCCs) in the same concentration range observed in the case of neuronal VGSCs (Figure 1). Allethrin (Type I) blocked recombinant rat L-, P/Q- and T-type VGCCs at a concentration like that required for the VGSC channels in human embryonic kidney cells [58–60]. Direct effects of pyrethroids on VGCCs explain the enhanced neurotransmitter release in the CNS, resulting in CS syndrome of intoxication [61]. In insects, pyrethroids also influence the function of VGCCs. Wang et al. [53] demonstrated that deltamethrin
might act on T-type VGCCs in brain cells of honeybees and result in behavioral abnormalities including swarming, feeding, learning, and acquisition. VGCCs are also alternative targets for Type II pyrethroids. Deltamethrin induces concentration-dependent blockage of chloride channels in muscle cells, as well as in peripheral and CNS neurons. Pyrethroids inhibit the Cl\(^-\) permeability leading to increased muscle membrane resistance and salivation, which are symptoms of CS syndrome (Figure 1) \[62,63\]. Jenson \[56\] proved for the first time that Type II pyrethroid block chloride channels in Spodoptera frugiperda and Anopheles gambiae insect cells. Moreover, Type II pyrethroids bind to and block GABA receptors in mammalian brain preparations. However, higher concentrations are needed than in case of sodium channels. The blockage of GABA receptors is an indirect neuroexcitatory effect involving the removal of inhibitory neuronal input, leading to convulsions \[11,53\]. Furthermore, pyrethroids could directly act on GABA receptors and reduce GABA-mediated responses in insects. The blocking effect of deltamethrin on GABA receptors was observed in Anopheles gambiae and Apis mellifera (Figure 1) \[55,64\].

5. Insecticide Resistance—The Case of Colorado Potato Beetle

The CPB has been considered as a model for pest mismanagement, because it is highly adaptable and can rapidly develop insecticide resistance. Due to its feeding, mating habits and great adaptability, CPB has developed this “superpower”. CPB is the most devastating defoliator of cultivated potatoes, but also may cause significant damage to tomatoes and eggplants \[65,66\]. Nowadays, the CPB distribution area is around 16 million km\(^2\) in North America, Europe and Asia, which is extremely large compared to its initial estimated 5000 km\(^2\) living territory \[67,68\].

In continental climates, with colder winters, CPB adults enter diapause, which is induced by a short-day photoperiod, falling temperature, decrease of forage quality (yellowed leaves) and pass the wintertime in the soil \[69\]. In these areas, CPB generally brings up two generations per year, but in mild climate zones, three or even four generations may grow up \[70\]. During the spring or early summer, adult beetles colonize potato fields, they are attracted to volatile compounds of the Solanum species and immediately start to reproduce \[71,72\]. CPBs have high fecundity: female beetles may lay up to 800 eggs in their life \[67\]. The life cycle of CPB involves seven life stages (egg-4 larval stages-pupae-adult), larvae undergo metamorphosis to achieve adult form. Pupation occurs in the soil, and adults emerge one week later, depending on temperature \[68\]. Larvae and adults consume potato leaves and are especially voracious. Due to their mating and destructive feeding habits, they can completely destroy potato fields, if left uncontrolled \[68\]. Despite the relatively young evolutionary age of CPB, they possess a high intraspecific polymorphism, which provides a broad range of ecological plasticity and adaptability to different biotic and abiotic factors \[68,73\]. Transposable elements, which comprise at least 17% of the CPB genome and high levels of nucleotide diversity in this pest population, facilitate rapid evolutionary changes \[74\].

Defoliation of potato plants by the CPB can significantly decrease tuber production \[75\]. A loss of more than 75% of the foliage can cause a total crop loss \[76\]. Several important potato producing countries, such as Russia, Ukraine, Poland and others in Eastern Europe, have reported high yield losses due to the CPB invasion \[77\].

For over 125 years management of CPB has been mainly based on chemical control. In Europe, potato crops are treated with insecticides against CPB 3–4 times per year to achieve the expected yield. Increasing the dosage provides only short-term relief but greatly contributes to the development of resistance. Nowadays, the control of CPB has become difficult due to the rapid evolution of its resistance to most insecticides through both, cross-resistance and multiple resistance to different compounds. Unsuccessful sub-lethal pyrethroid control of CPB leads to undesired side effects: it has a positive fitness effect over generations in CPB, by increasing the survival rate of the insect and reaching higher body mass, which is known to facilitate the population growth and invasion success \[78\]. Moreover, the development of resistance to insecticides significantly increases
potato production costs, it contaminates the environment and disturbs the equilibrium of agro-ecosystems [70,79].

High predisposition to resistance development in CPB is caused by several distinct factors: co-evolution of the CPB and Solanaceae species promoted the development of the enhanced detoxifying ability of beetles to tolerate high concentrations of toxic glycoalkaloids which is a specific repellent chemical of the Solanaceae species. Similarly, the high fecundity rate increases the appearance of random mutations in CPB populations, while a large number of laid eggs by a resistant beetle also ensures successful integration of the new trait. Furthermore, the narrow host range of beetles reduces the possibility of the existence of undisturbed refugees, where susceptible genotypes may escape exposure to chemicals. Last but not least, excessive pest control with different pesticides has been a major concern due to the repeated development of resistance in the CPB population [79,80].

Known mechanisms of CPB resistance to insecticides include mutations in genes encoding target receptors or enzymes (e.g., VGSCs, AChE), enhanced metabolism involving esterases, carboxylesterases, and monooxygenases, as well as reduced insecticide penetration and increased excretion (Figure 3) [67].

![Figure 3. CPB resistance mechanisms to the organophosphate, carbamate, neonicotinoid and pyrethroid insecticides. Black arrows indicate the mode of action of different insecticides. Red arrows indicate the developed CPB resistance mechanisms, which prevent the harmful effects of insecticides (red X).](image_url)

Resistance mechanisms may be highly diverse, even within a relatively narrow geographical area and between the life stages of the beetle [66,81]. The first recorded case of the resistance of this pest to a synthetic organic compound was the failure of DDT in 1952. Since then, CPB has developed resistance to over 54 different insecticides including organophosphates, pyrethroids, neonicotinoids and others. [79,82,83]. Over 300 documented cases of insecticide resistance are listed in the literature coming from Asia, Europe, and North America [84].

In 1965, an endosulfan (organochlorine insecticide)-resistant CPB population was recorded for the first time [5]. The CPB resistance mechanism involved in defense against endosulfan is mainly metabolic, by increasing the enzymatic activity and increasing the number of enzymes: esterases, monooxygenases, and glutathione-S-transferase (Figure 3) [85].
The Colorado potato beetle’s resistance to azinphosmethyl (an organophosphate) and carbofuran (a carbamate) were observed as early as the late 1970s [5]. This resistance is associated with a point mutation in AChE, a substitution of adenine with guanine at position 980 in the AChE gene, which leads to the change of serine to glycine (S291G) [86]. Furthermore, two other AChE mutations were identified in CPB, which were associated with resistance to these insecticides, R30K (replacement of the amino acid arginine with lysine) and I392T (replacement of isoleucine with threonine) [87]. The S291G mutation leads to conformational changes at the α-helix level in AChE that affect the binding sites for the insecticides (Figure 3) [86,88].

The intensive use of pyrethroids in controlling pests and disease vectors has been one of the main factors contributing to the development of pyrethroid-resistance among insects, which nowadays has grown into a global problem. One of the major mechanisms of pyrethroid-resistance, which reduces neuronal sensitivity to this class of insecticides, is known as knockdown resistance (kdr) of VGSCs. This mutation was initially found in *Musca domestica* [89], but soon in many other insects such as *Blatella germanica* [90,91], *Anopheles gambiae* [92] and *Myzus persicae* [93]. Nowadays, the kdr mutation has been documented globally in most arthropod pests and disease vectors [51]. The kdr mutation consists of substituting leucine for phenylalanine at position 1014 (L1014F), which confers resistance by reducing the binding of pyrethroids to sodium channels.

The kdr mutation has been identified as the main mechanism of resistance to pyrethroid insecticides in CPB (Figure 3) [82]. The kdr mutation shows an incompletely recessive inheritance in the CPB population [78,82]. In Europe, pyrethroid resistance due to kdr was detected in CPB field populations in Bulgaria, Czech Republic, Germany, and Serbia [78,81,86,94,95]. In addition to L1014F, two more target site mutations (T929I, T929N) were described in the sodium channel, which confer a super-kdr-like phenotype in the Bulgarian CPB population (Figure 4) [95]. Co-occurrence of more than one resistance-associated mutation drastically reduces the VGSCs sensitivity to pyrethroids, compared to single mutations.

![Figure 4](image-url)  
*Figure 4.* Identified mutations in the voltage-gated sodium channel in Colorado potato beetle populations that are associated with pyrethroid resistance (adapted from Rinkevich et al. [51]). The sodium channel protein contains four homologous repeats (I–IV), each part has six transmembrane segments (S1–S6). The red circle mutations were identified in pyrethroid resistant CPB: L1014F, T929I and T929N.

Oxidative metabolism participation in pyrethroid resistance is also well documented in the CPB population [94,96]. Resistant beetles detoxify the toxin (insecticide) faster than susceptible ones, or quickly rid their bodies of the toxic molecules. Resistant populations possess higher levels or more efficient forms of these detoxifying enzymes [67,94].
In the USA, several CPB populations are already resistant to pyrethroids, while in Europe, even though the pest control with pyrethroids is still effective, reduced susceptibility of pests to this chemical has also been observed [81,86,94].

Since pyrethroid resistance evolved in CPB populations, neonicotinoid insecticides have been the cornerstone of CPB pest management in cultivated potatoes. However, the widespread use of neonicotinoids in the last decades has now led to the development of neonicotinoid-resistant CPB populations [82]. Nicotinoid resistance of CPB is due to overexpression of genes encoding detoxifying enzymes: cytochrome P450, glutathione S-transferase, UDP-glycosyltransferase, and xenobiotic transporters, i.e., ATP-binding cassette (ABC) transporter, which participates in the excretion process of metabolized insecticides [97,98]. Furthermore, other mechanisms are involved in CPB resistance to neonicotinoids: mutations in the ACh receptor [99], reduced absorption and penetration of insecticides, increased activity of carboxylesterases and monoxygenases (Figure 3) [79].

The increased number of resistant CPB populations has a substantial impact on potato cultivation, undermining the effective production of this crop every year. With the persistent resistance occurrence in the CPB populations and the large impact of this pest on the potato industry, research in these areas could help in the development of better pest management strategies (integrated pest management—IPM), which have to focus on: (1) understanding the status and mechanisms of insecticide resistance by monitoring the resistance of local CPB populations; (2) overcoming or delaying resistance to existing insecticides by using well-established insecticide treatment plans (well-programmed pesticide treatment, combination of effective pesticides by omitting chemicals to which local CPB population is resistant) and (3) preventing the development of resistance to new pesticides by reducing the insecticide selection pressure with parallel application of other strategies than the chemical control (e.g., crop rotation, plantation of sensitive trap plants or resistant potato cultivars, treatments with bio-insecticides).

6. The Numbers behind Potato Cultivation and Insecticide Use

Chemical pest control was already preferred in the last century and it is still the predominant type of pest control. In case of CPB, insecticides are the most important control measure on commercial potato farms. However, recent concern about the harmful effect of chemical pesticides on growers, consumers and the environment has encouraged scientists to consider alternative, safer and more effective control agents [65,66]. Furthermore, in 2009 the European Commission of the European Parliament proposed a plan to transform the European Union’s agricultural system to a more sustainable and safer from health perspectives [100]. According to the plan, the main task is to reduce the risks and impacts of the excessive use of pesticides (the target is a reduction of 50% in the next decade), in addition, promote the use of alternative techniques (e.g., biological, mechanical control) and compliance with the principles of integrated pest management (IPM) [100]. Nonetheless, insecticide sales significantly increased in recent years in around 60% of the member states of the European countries (Figure 5) [101].

Germany, Italy and Spain are the top insecticide users from the European countries, while France, Ukraine and Romania use most of pyrethroids for pest control (Figure 6) [102]. France is the list leader in insecticides usage for potato production, followed by Netherland and Romania (Figure 7a) [102]. The first two countries are the main potato producers in terms of yield, while Romania occupies only 32nd place in the list based on statistics from the 2017–2019 period (Figure 7b) [102].
Figure 5. Percentage changes in insecticide sales by European countries from 2014–2016 vs. 2017–2019 (the presented graphic is based on data obtained from EUROSTAT database [101]).

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Figure 6. The European countries that used the most insecticides (total insecticide use in tonnes), between 2017 and 2019. The smaller right-sided graphic describes exclusively the pyrethroid use in these countries (the presented graphics are based on data obtained from FAOSTAT database [102]).
France is the list leader in insecticides usage for potato production, followed by the Netherlands and Romania (Figure 7a) [102]. The first two countries are the main potato producers in terms of yield, while Romania occupies only 32nd place in the list based on statistics from the 2017–2019 period (Figure 7b) [102].

Figure 7. (a). Percentage distribution of insecticide use for potato cultivation in European countries (statistics for the last 10 years) (b). Potato yield by countries between 2017 and 2019 (tonnes/ha) (the presented graphics are based on data obtained from FAOSTAT database [102]).

7. Perspectives to Reduce Insecticide Use

Nowadays, there is an increasing awareness of the side effects of insecticides and researchers are looking for a more eco-friendly and sustainable solution to combat insect pests such as CPB. Strategies that meet the abovementioned criteria could be indirect and direct alternative control methods. Indirect alternative control methods are crop rotation [103], straw mulch [104], trap cropping [105] or selection of plant varieties [106]. Among all of them, crop rotation and selection of plant varieties are the most powerful in CPB control, but crop rotation alone is not effective enough, only slowing down CPB population buildups. On the other hand, there are only a few cultivars, which have some
degree of resistance to CPB (e.g., cv. Dakota Diamond) [107,108]. Direct alternative methods to control CPB include behavioral interference methods [109], physical and mechanical control: bug vacuum [110], thermal pest control [111], use of natural enemies [112,113], use of the plant extracts and botanical insecticides [108]. Natural compounds and/or natural enemies are especially used in organic agriculture, but there is a growing interest in conventional agriculture. In Europe, there are only a few natural antagonists or parasitoids of the CPB beetle, and only a few studies are investigating the antibiotic effects of these bioinsecticides on CPB, therefore the beetle-killing and or beetle-suppressing efficacy of these compounds needs to be further studied [114,115]. *Pterostichus melanarius* Illiger [11,116] and *Chrysoperla carnea* Stephens [109] were found feeding on *L. decemlineata*. Two *Tachinidae* parasitic flies, *Myiopharus aberrans* L. and *M. doryphorae* Riley were found to parasitize *L. decemlineata* with relative success [117,118]. The disadvantages of using predators are multilayered. First of all, the newly introduced predator species can have an unknown effect on the global ecosystem, furthermore, their maintenance and production in adequate quantities are expensive and last but not least for releasing the predators, the purchase of additional equipment (mechanical distributors) is required [119]. Additionally some field studies suggest that natural enemies have a minimal effect [120–122].

Several species of *Bacillus*, including *B. pumilus*, *B. cereus*, *B. megaterium* de Bary [123], and *B. thuringiensis* Berliner (Bt) [113,124,125], were found to increase *L. decemlineata* mortality. *Bt* toxin produced by these microorganisms is only effective if it is ingested. Field experiments demonstrate that spraying *Bt* is most successful against newly hatched CPB larvae, so it should be used in the fields around this time [126,127].

Furthermore, biotechnological tools can be used, such as somatic hybridization, RNA interference (RNAi) [5,128], or CRISPR-Cas [129], to induce host plant resistance against CPB or CPB control. Among the wild potato relatives, several are described as resistant species for CPB attack through different mechanisms like special glycoalkaloids such as leptine I and II synthesized mainly in the leaves of *Solanum chacoense* or the tomatine present in some wild potato species and glandular trichomes mainly with repellent effect on the beetle. Some of the wild relatives of potato, which proved resistant are: *S. bertaulthii*, *S. tarijense*, *S. neocardenasii*, *S. chacoense* and *S. bulbocastanum* [77,130–132]. The differences in ploidy levels between cultivated potato (4x) and wild *Solanum* species (mainly 2x) make it difficult to transfer the valuable resistance genes into cultivated potato germplasm using classical hybridization. One way for transferring multiple resistance genes from wild *Solanum* species is somatic hybridization via protoplast electrofusion. The advantages of this biotechnological approach are: (i) hybridization of sexually incompatible but related species that is impossible or difficult to realize using classical methods, (ii) the transfer of polygenes involved in resistance and/or multiple resistance traits, (iii) the somatic hybrid plants obtained through this technology are not considered as GMO in Europe (directive 2001/18/EC- annex 1B) [133]. Several studies demonstrate the possibility of transfer of the CPB resistance genes from wild relatives (*S. tarnii*, *S. cardiophyllum* and *S. chacoense*) to somatic hybrids and back-cross progenies (BCs) [134–137]. During the development and selection of new CPB resistant host plants, there is the chance to learn more about action-mechanisms of different metabolites involved in the plant defense. This could contribute to the discovery of novel, eco-friendly ways to combat this voracious pest. For example, the application of molecular markers for the detection of leptins [136] and the discovery of new quantitative trait locus (QTL) markers, which correlate with glandular trichomes excretory functions, would pave the way for marker-assisted introgression breeding. It is likely that there are still other factors which are involved in CPB resistance. However, these action-mechanisms are yet to be discovered and hopefully can also be used in the development of host plant resistance.

Moreover, it would be necessary to establish a uniform statistical follow-up system in all European countries in order to correctly estimate the situation of pesticide use, to accurately monitor the risk of pesticides. According to the present situation, the countries should report statistics on the agricultural use of pesticides for selected crops after every
5 years and these reports are not uniform, detailed and unfortunately appropriately upgraded. The EU’s executive organization aims to regulate pesticides statistics until 2023 in order to overcome such data gaps [100].

Furthermore, a better communication with farmers would improve the success of the implementation of integrated pest management (IPM) strategies. In many countries, such as Romania, farmers have little knowledge about biological and environmental issues concerning CPB. They have only one tool in their pocket to protect their potato plantations against this voracious pest, which is nothing else than the insecticides. Therefore, there is an urgent need to develop a local specific IPM strategy and to make it available and well-explained to farmers in order to improve the integration in cultural practices of the new, more eco-friendly strategies. For example: by using a well-programmed manner to apply only the effective insecticides, which could be selected based on monitoring of insecticide resistance of local CPB populations, the number of insecticide treatments could be substantially reduced. Other strategies would be the plantation of resistant potato cultivars or very sensitive genotypes as trap plants and the application of bio-insecticides; moreover, the use of natural enemies should reduce the monopoly of chemical insecticides.

8. Conclusions

Insects have a huge negative impact on potato and other crop yield and there is an urgent need for alternative, innovative but also scientifically sound strategies to combat them in a more environmental-friendly way, keeping in mind the permanent reduction of carbon footprint and development of sustainable methods to fight pests.

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