Recent advances in engineering crop plants for resistance to insect pests

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Abstract

Background: While the rapidly increasing global population has led to a dramatically increased demand for the agricultural production, there have been heavy economic losses owing to various pest attacks on different food crops. The advancement of various biotechnological techniques have come as a boon in addressing the global concern and leads to the development of novel varieties that have proven to be highly economical, pesticide resistant and environmentally safe.

Main body: The present review was aimed to update the recent developments that have taken place in the field of crop production. Major focus was laid predominantly on such genes that have demonstrated positive effects and proved to be of commercial success at the market primarily due to the development of pest-resistant transgenic food crops with expression of Bacillus thuringiensis toxins. This technology has been effective against a wide range of pests including coleopterans, lepidopterans, hemipterans, dipterans, strongylida (nematodes) and rhabditida. In similar lines various plant derived toxic proteins were also discussed along with different genes that code for insect resistant proteins such as δ-endotoxins and secreted toxins. This article also helps in understanding the structural features of the genes that are endowed with insect resistance followed by their mechanism of action on pests. Further the role of secondary metabolites in controlling the pests was addressed. The Pros and Cons of existing tools of insect pest management were demonstrated.

Conclusions: Novel technologies are necessary in crop improvement to progress the pace of the breeding programs, to confer insect resistance in crop plants. Therefore, the future aim of crop biotechnology is to engineer a sustainable, multi-mechanistic resistance to insect pests considering the diversity of plant responses to insect attack.

Keywords: Genetic engineering, Insect resistance, Toxins, CRISPR/Cas9, RNAi, Phytophagous, Stacked traits

Background

Genetic engineering is a deliberate process of making changes to the characteristics of an organism by changing its genetic material. Genetic engineering in crop plants mainly offers two advantages i.e., (1) combining several individual, commercially useful genes to form gene cassettes and (2) reducing the time to introgression of these genes into a single genetic background. Since the “first report of genetically modified plants appeared in 1984 (Horsch et al. 1984), there has been a very rapid progress directed at using this novel technology for the practical ends of crop improvement. Protection of crop plants from insect pests was quickly seized upon as a major goal of plant genetic engineering (National Council 2000). The potential size of this market attracted major attention of a number of commercial organizations and the potential economic importance of this sector of biotechnology is "finally becoming more widely recognized (Burke and Thomas 1997). The practical application of plant genetic engineering involves two equally important technologies; cellular and molecular biology. The list of crop species

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which are amenable to genetic engineering has grown steadily and now includes majority of the crop species and many minor previously orphan crops. Concurrently, the list of useful genes for introduction into transgenic crops has not grown at a similar pace, although several different genes which might be useful for crop protection have been proposed. In this review, concentration was laid primarily on those genes which demonstrated effects in transgenic plants and their commercial success at the market. By far the greatest research effort in developing pest-resistant transgenic crops has gone into expression of Bacillus thuringiensis (Bt) toxins in plants (Flores et al. 2005).

**Main text**

**Phytophagous insects**

Insects belonging to the class Insecta are the most diverse group of arthropods and two thirds of them are phytophagous i.e. feed on living plant parts. Majority of the insect pests feed on one or few related plant species (monophagous) and few insects feed on multiple hosts (polyphagous). Intensity of crop damage by an insect depends upon the type of mouthparts which in turn decides the method of feeding. The insect feeding modes are broadly grouped into two types i.e., sucking/houstellate type, biting and chewing type and accordingly their mouthparts are evolved to suit their feeding style. Insects with sucking type of mouthparts causes substantial harm to the crop plants by damaging the vascular system of the plant (phloem and xylem), impairing transportation of water, minerals and food. In addition to that, they also cause additional damage to crop plants by transporting disease causing pathogens such as bacteria, fungi and viruses (Yadeta and Thomma 2013). Aphids, whiteflies, leafhoppers, thrips and honeybees are the examples for piercing and sucking type of mouth parts. On the contrary, insects with biting and chewing type of mouth parts cause mechanical damages to plant parts as they chew plant tissues such as root, stem, leaf, flowers and fruits affecting water and mineral acquisition photosynthesis and seed set. Grasshoppers, beetles, moths are the examples for insects with chewing type of mouth parts (Douglas 2018). More than 500,000 insect species belonging to majorly 8 out of 36 insect orders are known to be phytophagous (Bensoussan et al. 2016). Phytophagy is very common in the insect pests belonging to Lepidoptera (moths and flies), Coleoptera (leaf beetles and weevils), Hemiptera (Aphids, Planthoppers, sucking bugs), Diptera (gall flies, leaf miners, fruit flies), Hymenoptera (sawflies, wasps), Thysanoptera (thrips), Orthoptera (locusts, grasshoppers, crickets) and Phasmatodea (leaf, stick insects) orders (Chapman 2009).

**Yield losses due to insect pest**

Annually, global yield losses due to insect pests were estimated to be 18–20% of the annual crop production estimated at a value of more than US$470 billion (Sharma et al. 2017) and 30–35% in Indian agriculture which costs around US $36 billion, demonstrating adverse influence on the agricultural market, food security, and farmers profits. During the twenty-first century, these losses declined to 17.5 from 23.3% from the early 2000s, a positive indicator which can be largely attributed to the increase in the use of transgenic crops (Dhaliwal et al. 2010). Apart from causing significant yield losses by direct damage, insect pests can also cause indirect damage by transmitting disease causing pathogens. Sustainable increase of global food production without causing ecological damage has gained importance and many developed countries started investing in the development of eco-friendly pest management tactics during the 1980s itself (Swanton and Weise 1991), particularly important to meet the increased food supply of the burgeoning human population by 2050. This can be achieved by increasing the automation of agriculture concurrently utilizing biotechnological innovations. By 2050, growing-season temperatures will probably go beyond those documented in the past century and may significantly eases crop yield. On the other hand, models assessing the effects of climate warming on crop yields seldom consider impacts on insect pests, despite the damages that result directly from pest infestations and indirectly from pesticides applied to reduce pest damage (Rosenzweig et al. 2014). It’s predicted that the insect pests are expected to differ in their response to increasing temperatures geographically and among crops (Lehmann et al. 2020).

Interaction between the temperature, physiology and demography of insects to project the upcoming impact of insects on crop production was well established by Jamieson et al. (2012) which helps to understand the complex climate-mediated effects on plant–insect and multitrophic level interactions as well as the roles of plant eco-physiological processes in driving both bottom-up and top-down controls. It was also estimated that pest-related changes in yields of the principal grain crops such as maize, rice and wheat, which collectively account for 42% of direct calories taken by humans globally (Deutsch et al. 2018). The information regarding yield losses due to insect pests on crops like rice is very limited and also demonstrated the urge for the development of pest management practices. Herbivorous insects destroy nearly one fifth of the crop production on the planet annually (Oerke 2006). The most important potential reason for proliferation of pests is the creating man-made habitats, i.e., agro ecosystems that fulfill human
food requirements, where crops are selected for their huge size, high yield, nutritious value, and gathered in a confined area (Rembialkowska 2007). This does not just suffice human demand but provides a highly conducive environment for herbivorous insects at the same time. Artificial crop selection methods made it possible to meet the demands of human consumption, but on the flip side it was subjected to be conducive for infestation of insect pests. For instance, infestation by kahpra beetle (Trogoderma granarium Everts) larvae in grain kernels has negative effect on mineral composition, existing carbohydrates, protein, starch digestibility and bioavailability. The spoilage of kernels is more with the contamination of T. granarium body parts than consumption. Consumption of some of these contaminants in food may cause serious health issues (Athanassiou et al. 2019). Quantitative information on yield losses due to pests is vital to the development of sound pest management practices. It is therefore surprising to see how limited our information is today in the case of rice, the most important food crop worldwide.

### Pros and cons of existing tools of insect pest management

#### Physical methods

Farmers follow several cultural, mechanical, chemical, biological, botanical, genetic and regulatory practices to control the insect pests. In physical method, the insect pests were dislodged by spraying the plant with the water which knocks off aphids and mites, and however this is mostly in the case of household pests. The larvae from the bagworm can be picked off from the infested plant (Fields and Muir 2018). Well-known methods like traps are used to remove apple maggots, corn borers, fruit flies, bag worms, corn earworms, and peach tree borers through physical and/or chemical-based traps (Wojtowski 2010). Plant replacement technique was used to reduce the circulation and reduced many disease problems (Banks and Fields 1995). The two grain pests of controlling insect pests include living organisms that can kill the pest and the second incorporation of naturally occurring biochemicals that are harmful to the pest yet often are harmless to the consumers and the ecosystem. The insect pests are attacked by natural predators that are advantageous to the landscape. These beneficial insects often exist in the landscape naturally, but they also can be introduced. “Beneficials” may be predators or parasitoids. One common example of a beneficial predator is the lady beetles. Both the larvae and adult lady beetles prey on aphids and other soft-bodied insects. Other predators include lacewings, spined soldier bugs, flower flies, and spiders. Parasitoids live on and often kill their host. Some parasitic wasps use caterpillars, white-flies, aphids, and soft scales as hosts. An example of an organism that possess a naturally occurring biochemical is the bacterium Bt. Bt produces insecticidal proteins that are poisonous to specific group of insects, yet harmless to other organisms. Bt can be sprayed on plants and when the sensitive/susceptible insect pest feeds on the culture sprayed on the plant, protein gets ingested into the larvae and the larvae get killed (Mishra et al. 2017).

#### Chemical methods

Conventional chemicals are the primary go-to method of control and are often the most effective means of control. However, it should not be looked upon as the only method of control, as it can create selection pressure leading to resistance problems in pests which severely affects the sustainability of chemical methods. To use these chemicals to the fullest and have the greatest effect, chemicals need to be applied on a specific part of the plant when the pest is most vulnerable. Always chemical controls are to be applied according to instructions on the label. In many cases, environmentally safe pesticides such as horticultural oil or insecticidal soap are effective choices (Oerke 2006). However, applications must be timed carefully to have the greatest effect on the pest insect population. Because they have no residual activity after they have dried, soaps and oils are usually the option as they are the least disruptive to populations of beneficial organisms (Khan et al. 2008). The label on every pesticide formulation displays a warning sign like caution, warning, danger which indicates the level of toxicity of the chemical. These signs help to choose the least hazardous material among the effective alternatives. For the most landscape pests, it’s needed to consider pesticides in only the first two categories. Some pesticide formulations can be applied only by applicators with special training and who are certified by the state’s department of agriculture.

#### Biological methods

Biological methods of pest control can be an additional pest control tool especially when chemical methods of eradicating the insect pest have an unconstructive effect on the environment. In order to minimize the adverse effects on the environment, Sterile Insect Technique (SIT), which involves mass-rearing and releases of sterile insects is used for pest control. Sterile males must compete with wild males and reduce reproduction by wild females; in some cases the goal is eradication and in other programs it is to suppress pests. Ultimately, the goal is to reduce crop damage or transmission of insect-vectored diseases (Hoy 2003). SIT programs were highly effective in eradication of the screwworm (Cochliomyia hominivorax Coquerel) in North and Central Americas, and the Mediterranean fruit fly (Ceratitis capitata Wiedemann)
in Florida and other locations (Alphey 2016). Large numbers of insects were sterilized by irradiation at unavoidable cost (Hoy 2003). In most of the cases, males were released involving sex-separation which is a complicated process. Both the difficulties of cost and sex separation can be prevented by using engineered strains carrying a dominant, repressible, lethal gene or genetic system (Alphey 2002). The chemical pesticides are substituted by the microbial insecticides, their usage is constrained to kill a narrow spectrum of insect species. Thus, making Bt, a soil-borne bacteria, as a right choice, since long back to control the insect pest to overcome effects of synthetic chemical insecticides (Gupta and Dikshit 2010).

**Plant toxic proteins with insecticidal properties**

**Lectins**

Lectins are proteins extensively found in nature produced by plants and other organisms including mammals. These are of non-immune origin that possesses one non-catalytic domain that specifically and reversibly binds to mono- or oligosaccharide (Macedo et al. 2015). These are multivalent so that agglutinate cells bind to the brush-border membrane of the insect’s intestinal epithelial cells or, in the case of chitin-binding lectins, to the peritrophic membrane. The other way of toxicity is to bind to the two glycosylated digestive enzymes, particularly effective against the sap sucking: Hemiptera, Coleoptera, Lepidoptera and Homoptera (Chougule and Bonning 2012). Though activity of the lectin GNA (Galanthus nivalis agglutinin; GNA) from the snowdrop plant (Galanthus nivalis) against aphids, sap-sucking insects was demonstrated, harmful effects of GNA present in transgenic potatoes given raw as feed to rats showed an evidence to screen lectins more consistently in future (Ewen and Pusztaï 1999; Macedo et al. 2015).

**Ribosome-inactivating proteins**

Ribosome-inactivating proteins (RIPs) are capable of enhancing plant resistance to insect pests such as lepidopterans (Dowd et al. 2006), coleopterans (Kumar et al. 1993) and dipterans (Shahidi-Noghabi et al. 2008). The insecticidal activity of the RIPs can be demonstrated by feeding the insect pests with artificial diet incorporated by RIPs. For example, the feeding of tobacco aphid (Myzus nicotianae) Blackman on leaves from transgenic tobacco plants overexpressing SNA-I retarded the growth and reduced the fecundity and adult survival (Shahidi-Noghabi et al. 2008). In addition, an artificial diet added with a different type-I RIPs also reduced the fecundity and survival of velvet bean caterpillar (Anticarsia gemmatalis) Hübnner, beet armyworm Spodoptera exigua (Hübnner) and fall armyworm (Spodoptera frugiperda J.E. Smith) (Bertholdo-Vargas et al. 2009). Recent studies have suggested that type-I and type-II RIPs from apple (Malus domestica Borkh.) have a strong aphicidal activity. The nymphal survival of aphids (M. nicotianae) in apples was greatly reduced by the artificial diet included with the purified recombinant proteins for type-I RIPs and type-II RIPs (Bolognesi et al. 2016).

**Bacillus thuringiensis**

*Bt* is a Gram-positive spore-forming bacterium that produces parasporal crystal proteins encoded by the *cry* or *cyt* genes, toxic to many insect species. The 30% dry weight of the spore is represented by the parasporal crystal proteins (Mishra et al. 2017). *Bt* is more or less exclusively active against larval stages of different insect orders and kills the insect by disruption of the midgut tissue followed by septicaemia (Raymond et al. 2010). It consists of one or more pro-toxin species weighing about 160 kDa. The proteolyzed pro-toxins are converted into peptides, weigh up to 55 kDa to 70 kDa are predominantly toxic to lepidopteran, dipteran, or coleopteran insects (Pardo-Lopez et al. 2013).

The first δ-endotoxin coding gene was sequenced in 1985 (Schnepf and Whiteley 1985). Till date, 806 *cry* genes that code for Cry proteins were identified, which are usually located on the large plasmid (Palma et al. 2014). Most of the Cry proteins possess insecticidal activity and are exploited for the control of insect pests primarily in agriculture. Some of the recent studies have reported unique Cry proteins which are non-insecticidal, but have the ability to kill human cancer cells (Ohba et al. 2009). These cytotoxic proteins called Parasporins are mainly classified into six groups according to the homology of amino acid sequence by the Committee of Parasporin Classification and Nomenclature (PS1 through PS6). Human cancer cells from diverse origin illustrate cytotoxic activity with the six parasporins with Cry numbers Cry31A, Cry41A, Cry45A, Cry46A, Cry63A and Cry64A (Ohba et al. 2009). In addition to the δ-endotoxins, various isolates of *Bt* produce proteins such as vegetative insecticidal proteins (Vip) and secreted insecticidal proteins (Sip) that are secreted into the medium during the vegetative growth phase and have been found to have insecticidal properties against pests (Estruch et al. 1996). *Bt* also produces other potential insecticidal toxins like non-proteinaceous β-exotoxins, which are active against insect pests (Li et al. 2014).

**Bacillus thuringiensis toxin nomenclature**

In 1993, a committee was established in order to revise the nomenclature of δ-endotoxins of *Bt*, formerly devised in 1989 by (Höfte and Whiteley 1989). The modern classification of these δ-endotoxins was completely based on the identity of the amino acid sequence (Xu et al. 2014).
This systematic nomenclature helps the researchers to avoid the bioassay of every novel protein that is identified and ranked with close proximity. The original nomenclature has biological specificity. In the previous system, toxins and their corresponding genes were denoted in Roman numerals namely CryI for lepidopterans, CryII for coleopterans and dipterans, CryIII for coleopterans; and CryIV toxins specific to dipterans were illustrated in the previous classification. In the present classification Roman numerals have also been exchanged for Arabic numerals in the primary rank (e.g. CryIIIa became Cry3A) (Kaur and Allam 2006) (Fig. 1).

According to the pairwise amino acid identity, every novel toxin is given a four-rank name. In this new system of nomenclature, no implications were made according to the structure, host range or even mode of action of the newly identified toxins. Arabic numerals are given the first and fourth rank, upper- and lower-case alphabets were used for second and third ranks. In this manner, proteins sharing less than 45% pairwise identity are assigned a different primary rank. A completely new toxin might therefore, be assigned the name Cry76Aa1. To be more convenient enough it is proposed that the quaternary rank is an option and it gives the variation among proteins that are less than 5 and 95% alike in the protein sequence and is only used for elucidation. To make a note of quaternary ranks are allotted to all independently sequenced toxin genes, therefore even though in reality that several toxins encompass varied quaternary ranks—they could in fact be identical. This nomenclature in applied to δ-endotoxin proteins (encoded by the cry genes) and cytotoxic proteins (encoded by the cyt genes) and a class of insecticidal proteins secreted during the vegetative phase includes vegetative insecticidal proteins (vip) and secreted insecticidal protein (sip) (Crickmore et al. 2018). There are other potential insecticidal toxins discussed in brief below (Fig. 2).

Cry toxins (cry genes)
Currently more than 800 cry genes were classified into 75 classes of δ-endotoxins based on the amino acid sequence similarities. These classes consist of several subclasses i.e., cry1A, cry1B ... cry1Y) which are subdivided into subfamilies or variants (cry1Aa, cry1Ab, cry1Ac, etc.). The genes included in the class had 45% similarity to each other. A restricted spectrum of activity can be observed by specific cry gene products limited to the larval stages of a small number of species. Updated list of δ-endotoxin genes can be found at http://www.lifesci.sussex.ac.uk/home/Neil_Crickmore/Bt/intro.html (Crickmore et al. 2020).

The identity of these Cry proteins has nowhere in relation to its spectrum of activity. For instance, the genes 84% identical to each other like cry1Aa and cry1Ac proteins among which only cry1Aa is toxic to silkworm (Bombyx mori Linnaeus) (Atsumi et al. 2005). On the other hand, cry3Aa and cry7Aa genes 33% identical to each other show toxicity against the Colorado potato beetle (Leptinotarsa decemlineata Say) (Lichtfouse et al. 2009). The typical illustration of a bacterium with this protein pattern is Bt subsp. israelensis, used all over the world as commercial preparations against dipteran larvae (Cantón et al. 2011). However, other than this set of
proteins, in some cases, other complementary proteins such as Cry1 and Cry2 have been detected that enhance the toxic effect against Diptera (Ben-Dov et al. 1997). Some of Cry toxins i.e., Cry5, Cry6, Cry12, Cry13, Cry14, Cry21 protein classes are inactive against insects but show toxicity against invertebrates such as nematodes (Palma et al. 2014). More recently, binary toxins from Bt designated as Cry34Ab1/Cry35Ab1, active against various coleopteran insect pests of the Chrysomelidae family have also been characterized. In spite of their little homology with the other members of the Cry toxins, binary toxins were assigned a Cry designation. The Cry34A and Cry35A are 14-kDa and 44-kDa proteins, respectively, that function as binary toxins showing activity on the western corn rootworm (Diabrotica virgifera virgifera LeConte) (Ellis et al. 2002). Lepidopterans usually consume toxins through ingestion. Bt toxins classified and studied with insecticidal activity in this group are Cry1, Cry2, Cry9, and Cry15. However, the toxins viz., Cry1, Cry2, and Cry9 groups were reported with insecticidal activity to velvet bean caterpillar (Anticarsia gemmatalis Hübner) (Bel et al. 2019).

Cry proteins are not only insecticidal but some of the Cry proteins of Bt strains B622 and B626 exhibited their toxicity on human pathogenic protozoan Trichomonas vaginalis and also produces lectin-like effect on rabbit erythrocytes (Palma et al. 2014). However, it did not show any insecticidal effect against the lepidopteran insect diamondback moth (Plutella xylostella Linnaeus) and the dipteran London underground mosquito (Culex pipiens L.). The cry genes expressed in genetically modified (GM) crops like cry1Aa and cry1Ca showed synergism against tobacco budworm (Heliothis virescens Fabricius), sugarcane stalk borer (Diatraea saccharalis Fabricius) and fall armyworm (Spodoptera frugiperda J. E. Smith) (Lemes et al. 2014). Bt Cry toxins have shown their toxicity against Asian tiger mosquito (Aedes aegypti Skuse) (Crickmore et al. 1998) and Bt subsp. israelensis to blood worm (Chironomus tepperi Skuse) (Hughes et al. 2005). According to (Yamagiwa et al. 2001), larvae tested on northern house mosquito (Culex pipiens Linnaeus) hasn’t shown any toxicity against individual Cry protoxin, but the toxins turned active when larvae were exposed to two toxins together. This shows that the toxicology of the Cry proteins depends upon the formation of active toxin complex. The activity of Bt Cry toxins may differ widely depending upon Cry protein intake by herbivores and may be related to time, location, and amount of expression of the toxin in the plant (Devo et al. 2012). A new B. thuringiensis subsp. Galleriae producing Cry8Da has been commercialized recently which is active against coleopterans. The efficacy of this Bt product has been tested against the Alfalfa weevil (Hypera postica Gyllenhal) shown up to 40% reduction in number (Shrestha et al. 2018). The Bt toxins were termed to be versatile as they show undesirable activities such as haemolytic activity described for Cry15A toxin (Estruch et al. 1996). The Cry toxins also exhibit antibacterial activity which can hamper cloning and/or adequate expression (e.g., the three-domain toxins Cry13A and Cry14A) (Wei et al. 2003). Bt subsp., displayed antibacterial activity upon proteolytic activation against species of the anaerobic Gram-positive genus Clostridium and to an archaeal species (Yudina et al. 2007). Nevertheless, several Cry toxin genes have been incorporated into transgenic crops, providing an effective way to control insect pests in agriculture and lowering the worldwide use of field-applied chemical pesticides. The incorporation of Bt toxins saved up to US$30 per ha through 70% cutback in the usage of insecticides and also tremendous increase in the yield up to 80–87% (Qaim 2009). Other Cry and Cyt-like proteins have also been reported to be bactericidal (Yudina et al. 2007).

**Structural features of crystal proteins**

The microscopic observation of the Bt under the phase contrast displays one or more crystalline inclusion (parasporal crystal) bodies during the sporulation of its growth cycle (Bechtel and Bulla 1976). These crystalline inclusions, for example, are synonymously called insecticidal crystal proteins (ICPs), Cry toxins or δ-endotoxins. These parasporal crystals consist of proteins, which exhibit highly toxic insecticidal activity. On the other hand, actively growing cells lack the crystalline inclusions, so that they are not toxic. The δ-endotoxins fall into two categories; Cyt and Cry. These two types of δ-endotoxins do not share significant sequence homology, although, both seem to work through pore formation that leads to cell lysis and irreversible damage of the insect midgut (Chang et al. 1993). The three dimensional structures of the four δ-endotoxins (Cry1, Cry2, Cry3 and Cry2A) were resolved by X-ray crystallography (Grochulski et al. 1995). The Cry1, Cry2, and Cry3 are remarkably similar, each of them consisting of three domains. The N-terminal Domain I consists of seven α-helices with a central core helix surrounded by six amphipathic helices. Domain II consists of three β-sheets with three-fold symmetry and the conformation is called ‘Greek Key’. The C-terminal, domain III, consists of two anti-parallels of β-sheets in a ‘jelly-roll’ formation. Each domain of the Cry toxin has a role in the mode of action of the toxin i.e., Domain I is involved in membrane insertion and pore formation, Domains II and III are involved in receptor reorganization and binding (De Maagd et al. 2001).
**Action mechanism of insecticidal cry toxins**

*Bt* produces crystal proteins during their sporulation which are toxic to the host and also show specificity. Consequently, each type of Cry protein is capable of toxins to one or more particular insect species. In contrast to the chemical pesticides crystal proteins do not affect many beneficial insects, plants and animals including humans due to its specific toxicity. The specificity of these insecticidal crystal proteins (ICPs) derives from their mode of action (Gill et al. 1992). The crystals produced during the sporulation of *Bt* consist of the ICPs which exist as protoxins. Following the ingestion of parasporal crystals by the susceptible insect pest, these crystals are dissolved in alkaline conditions (pH 10–12) of the midgut of susceptible insect pest, producing 130 to 135 kDa protein chains called protoxin. The gut proteases truncate the protoxins into actual toxic fragments of 60–65 kDa (Höfte and Whiteley 1989). Thus, the activated toxin binds to the specific receptors of the larval mid-gut epithelium. Upon binding of the activated toxins, pores are created in the cell membrane. The formation of pore triggers off the osmotic shock and disturbs the ion channels. Consequently, the cell membrane breaks down, paralysis arises and as a result, the insect stops feeding and dies from starvation. *Bt* subsp. *israelensis* is highly toxic to different Aedes, Culex and Anopheles mosquito species that acts as vectors of several human infectious diseases (including chikungunya), produces crystal inclusions composed of Cry4Aa, Cry4Ba, Cry10Aa, Cry11Aa, Cyt1Aa and Cyt2Ba toxins (Fernández-Luna et al. 2010). As mentioned previously, the mosquitoicidal active Cry proteins Cry11Aa, Cry4A and Cry4B share similar structures with the lepidopteran active toxin Cry1Aa suggesting a similar mode of action of these Cry proteins in mosquitoes (Dai and Gill 1993).

The Cry toxins produced by *Bt* have two different models. The most studied is the classical model also known as sequential binding model and the recent model is Signaling pathway or alternative mode of Cry toxins. In both the models, after the ingestion of Cry toxins, they get dissolved in the insect midgut at alkaline pH forming an activated Cry toxin. These activated Cry toxins are monomers formed by the proteolysis at N- & C- terminal ends by trypsin and chymotrypsin like proteases of the insect midgut (Bravo et al. 2007). The major role in receptor binding is played by the C terminal end of the Cry toxin. Till date different insect midgut receptors including ABC transporters, glycophosphatidylinositol (GPI) anchored alkaline phosphatases (ALP), cadherin like proteins (CADR), GPI anchored aminopeptidases (APN) were identified as essential for activation of Cry toxins (Yudina et al. 2007). The cleaved Cry toxin oligomerizes to form pre-pore complex, which binds to the GPI anchored receptors and leads to pore formation through membrane insertion which in turn damages the midgut epithelium leading to the death of pest insect larvae. The acceleration of intracellular apoptotic pathway after the attachment of Cry toxin to the Cadherin receptors directing to the cell destruction damages midgut epithelium leads to larval death (Heckel 2020).

**Cyt toxins**

Apart from Cry proteins among δ-endotoxins another significant insecticidal protein are Cyt (cytotoxic) proteins, coded by cyt genes in *Bt*. The predominant specificity to dipterans and its cytolytic (hemolytic) activity makes the Cyt proteins different from that of Cry proteins. Cyt1Aa and Cyt2Ba showed three-dimensional structures with a single domain and of three-layer alpha–beta proteins (Palma et al. 2014). The Cyt1Ca protein encoded by the pBtoxis plasmid of *Bt* subsp. *israelensis* different in having a further domain with homology to the carbohydrate binding domain of ricin, attached to the C-terminal end of the Cyt domain but no larvicidal or hemolytic activity has been observed with this toxin. Cytotoxic proteins identified till date could only be kept in three different families Cyt1, Cyt2 and Cyt3; with primary rank acknowledged in the *Bt* Toxin Nomenclature Committee which are lethal mostly against some mosquitoes and black flies. A wide range of toxic activity is observed in insects of Diptera, Lepidoptera and Coleoptera with the cyt genes present the diverse strains of *Bt* subsp., e.g., subsp. *morrisoni* (Guercichioff et al. 2001). The Cyt2C protein showed a significant toxicity against nematodes (Rhabditida) and cancer cells. The Cyt proteins also play a vital role in insect resistance management by synergizing the insecticidal activity of other Cry or Vip3 toxins in some insect species. For instance, cotton leaf beetle (*Chrysomela scripta* Fabricius) pest is controlled by Cyt1Aa toxin and this toxin also inhibits the resistance of Cry3Aa (Federici and Bauer 1998). In the same way, Cyt1Aa is competent to hold back resistance to Cry4 and Cry11Aa toxins in larvae of laboratory selected southern house mosquito (*Culex quinquefasciatus* Say) populations (Soberón et al. 2013). The Cyt proteins bind to Cry proteins to show synergism for instance binding of Cry11Aa to Cyt1Aa aids in cooperation of oligomerization of Cry11Aa toxin and pore formation. Moreover, toxins Cyt1Ab and Cyt2Ba from *Bt* subsp. *medellin* and subsp. *israelensis* enhanced the insecticidal activity of *Lysinibacillus sphaericus* (*Bacillus sphaericus*) against *A.aegypti* and resistant *C.quinquefasciatus* larvae (Soberón et al. 2013). Cyt1Aa demonstrated to have a synergistic activity, when combined with Mtx1 toxin from *L. sphaericus*, against *C. quinquefasciatus*. Two different modes of action have been proposed for the Cyt
group of proteins: one suggests a pore-formation model whereas the other supports a less specific detergent action mechanism (Palma et al. 2014). For toxins like Cyt1Aa, with a typical cytolysin fold and a specific hemolytic pattern that differs from ionic and non-ionic detergents, a pore-forming mechanism was further suggested.

**Secreted toxins**

**Vip**

Some strains of *Bt* produce proteins during their vegetative growth phase into the growth medium, found to have insecticidal properties against a number of insects, extending the overall host range of this bacterium. These insecticidal proteins include vegetative insecticidal proteins (Vip) and secreted insecticidal protein (Sip) and do not share any sequence or structural homology with the Cry toxic proteins. These secreted proteins had a specific conserved signal peptide sequence that are commonly cleaved before or after the secretion process is completed. At present, the *Bt* Toxin Nomenclature Committee has classified Vip proteins into four different families namely: Vip1, Vip2, Vip3 and Vip4 based on the amino acid sequence similarity. Among these Vip toxins, Vip1 and Vip2 act as binary toxins exhibiting toxicity against insects of Hemiptera and Coleoptera orders. While Vip3 proteins have no sequence and structural similarity with either Vip1 or Vip2 and are toxic to lepidopteran insects. Intriguingly, studies showed that the insecticidal mode of action of Vip3 toxins resembles with that of the Cry proteins. The target host insect for the Vip4 family insecticidal proteins remains to be identified (Chakroun et al. 2016).

**Sip**

The secreted insecticidal protein (Sip) constitutes one member Sip1Aa1 demonstrated toxicity against coleopteran larvae till 2017 (Fernández-Chapa et al. 2019). A recent study in China reported Sip1Ab gene from a native *Bt* strain QZL38 and its insecticidal activity against *Colaphellus bowringi* Baly (Coleopteran) (Sha et al. 2018). Sip proteins were initially isolated from supernatants of the *Bt* strain EG2158 and was as Sip1Aa1. The extent of sip1Aa1gene is 1104 bp and it encodes 367 amino acid protein and ~41 kDa. Sip1Aa1 exhibits typical predicted Gram-positive consensus secretion signal 30 amino acids long. However, the protein was found N-terminally processed, with its first 43 amino acids eliminated by active proteases present in the culture medium. It demonstrated little but considerable match to the 36-kDa Mtx3 mosquitocidal toxin (a member of the ETX_MTX2 family of toxins) from *L. sphaericus*. This homology toxins strongly recommends that Sip1Aa1 toxicity possibly be caused by pore formation, but its mode of action remains unknown (Palma et al. 2014). Sip1Aa1 is lethal for Colorado potato beetle (*Leptinotarsa decemlineata*) (Coleoptera: Chrysomelidae) and inhibits growth of spotted cucumber beetle (*Diabrotica undecimpunctate* Howardi) (Coleoptera:Chrysomelidae) and Western corn rootworm (*Diabrotica virgifera virgifera*) (Palma et al. 2014). The only protein reported till 2017 is Sip1Aa1 with 90% homology and the recent novel protein Sip1Ab identified in 2018 showed insecticidal activity towards Coleopterans (Sha et al. 2018).

**Insect resistance to Bt**

The pests often become resistant to the *Bt* toxins if they are exposed to selection pressure by the toxin during several consecutive generations of the pest. The hindrance of resistance is essential to formulate a sustainable pesticide. There could be two possible ways to overcome the resistance mechanism. The primary strategy is to switch toxins hoping that the resistance besides first toxin is lost during the usage of the second toxin. The other strategy is to use several toxins altogether assuming that the development of resistance is quite impossible to the toxin combinations. These strategies only be productive against one of the toxins does not lead to resistance to the other toxins Lee et al. (1996) examined insecticidal activity of different toxins Cry1Aa, Cry1Ab, and Cry1Ac against lepidopteran pests like *Lympantria dispar* (gypsy moth) and *Bombyx mori* (silkworm) by force-feeding bioassays (Lee et al. 1996). The investigations demonstrated the synergism between the mixture of Cry1Aa and Cry1Ac in gypsy moth (*L. dispar*) pests whereas antagonistic effects were observed in Cry1Aa and Cry1Ab toxins. In the case of silkworm (*B. mori*) no synergistic effect was observed. The Cry toxins which were insensitive larvae to African cotton leaf worm (*S. littoralis*) were associated with the endochitinase ChiAll in order to increase the insecticidal effect even at low concentrations a synergistic toxic effect was observed. The interaction between the crystal proteins (Cry) and the vegetative insecticidal proteins (Vip) showed synergism when applied together. The toxins Cry9Aa and Vip3Aa exhibited high affinity in binding assay and showed elevated insecticidal activity against, the Asiatic rice borer (*Chilo suppressalis* Walker) (Wang et al. 2018).

Torres-Quintero et al. (2018) constructed hybrid-Cyt1Aa mutants expressing the loop3 of crystal protein Cry1Ab-domain II in varied demarcated regions of the Cyt1Aa toxin. The three hybrid variants of Cyt1Aa, Cyt3-Loop6, Cyt3-Loop7 and Cylc3-Loop9 exhibited considerable binding to amino peptidase-N1, Alkaline phosphatase and Cadherin receptors in comparison to the control Cyt1Aa toxin. A significant toxicity was observed in two different lepidopteran larvae, tobacco
are recognizable agricultural products with well documented toxicity expressed in trichomes, the plant showed enhanced resistance range. As acknowledged earlier Vip3A proteins were inhibited by the transgenic plants with an increased inhibiting effects of five different Vip3A proteins on selected lepidopteran pests (Hernández-Martínez et al. 2013). The insect- resistant corn Bt transgenic plants have set a case in point as a potential biotechnological commercial success story for the people. Moreover transgenic Bt cotton, maize and rice which are resistant to different lepidopteran pests are the present keys which are recognizable agricultural products with well documented insect virulence coupled with an extreme degree of antibiosis. Both lepidopteran and coleopteran pests were inhibited by the transgenic plants with an increased resistance range. As acknowledged earlier Vip3A proteins are excellent contenders for gene pyramiding in transgenic crops to combat development of resistance against the currently deployed genes. A synthetic plant-preferred codon-optimized novel vip3Aa44 was cloned into pBI-NAR plant transformation vector and tobacco explants were transformed with leaf disc co-cultivation method to evaluate toxicity of this gene against cotton bollworm (Helicoverpa armigera) and cotton leafworm (S. litura) were highly potential (Kalia and Kaur 2019).

**Plant secondary metabolites**

Plant secondary metabolites are a diverse and large number of specialized compounds which are not primarily useful in the growth and development of plants but are essential for the plant to survive. There are about 200,000 compounds which are mainly classified into three groups including flavonoids and allied phenolic and polyphenolic compounds, terpenoids and nitrogen-containing alkaloids and sulphur-containing compounds. These metabolites are toxic and deterrent to insect pests and few compounds function also indirect protection by attracting parasitic wasps or other natural enemies of insects feeding on plants. For instance, the two genes zFPP and ShZIS coding enzymes that synthesized the sesquiterpene 7-epizingiberene were transferred from tomato (Lycopersicon esculentum) wild variety to cultivated variety expressed in trichomes, the plant showed enhanced protection against multiple insects (Douglas 2018).

**Arcelins**

Arcelins are insecticidal proteins, obtained from wild accessions of the common bean (Phaseolus vulgaris Linnaeus), with resistance against bruchid beetles. Arcelin protein purified from hyacinth bean (Lablab purpureus) had the ability to manage storage pest in cereals transformed with L. purpureus defense related gene (Janarthanan et al. 2008). The chemical composition of arcelin has many similarities with lectin including agglutinating activity. Till date, different allelic variants (designated Arc-1–7) of arcelin proteins have been described, with molecular weight in the range of 27–42 kDa. Of great interest are the insecticidal properties of arcelin variants toward bruchid pests and, in particular, their inhibitory effect on the larval development of the Mexican bean weevil (Zabrotes subfasciatus Boheman) (Karuppiah et al. 2018).

**RNA interference**

Designing insecticides that are different from the toxic proteins in terms of mode of action and specificity can be acquired through RNA interference (RNAi). The incorporated double-stranded RNA (dsRNA) particular to an essential gene of an insect pest into the cell turns out to be small interfering RNA (siRNA) molecules by dicer enzymes. Thus, the complementary mRNA is degraded by siRNA which guides Argonaute protein of the RNA-induced silencing complex (RISC) and in few occasions it also interferes with target mRNA. The crop protection in corn against the western corn rootworm and Colorado potato beetle in potato L. decemlineata was ensued through the orally delivered RNAi. In the United States, research is being carried out on processing of insecticidal RNAi against the western corn rootworm (Niu et al. 2017). Explicitly, SmartStax PRO (Monsanto) incorporates a dsRNA sequence in antagonistic to the Snf7 gene of the western corn rootworm, stacked with cry3Bb1 and a herbicide resistance gene (Moar et al. 2017). The Snf7 protein is a class E vacuolar sorting protein, and it’s down regulation resulted in perturbation to protein deubiquitination and autophagy in the insect midgut and fat body. More significantly, expression of dsRNA directed against suitable insect target genes in transgenic plants showed a protection against pests, opening the way for a new generation of insect-resistant crops (Niu et al. 2018).

**CRISPR-CAS9 system**

CRISPR-Cas9 (Clustered Regularly Interspaced Short Palindromic Repeats) systems have rapidly transitioned from intriguing prokaryotic defense systems to powerful and versatile bio molecular tools. The efficiency of this technology can be easily demonstrated in plants. Recent applications in bacteria have centered on multiplexed
genome editing, programmable gene regulation, and sequence-specific antimicrobials, while future applications can build on advances in eukaryotes, the rich natural diversity of CRISPR-Cas9 systems, and the untapped potential of CRISPR-based DNA acquisition (Jinek et al. 2012). The model plant for CRISPR based systems is thale cress or Arabidopsis (*Arabidopsis thaliana*) (Bechtold and Pelletier 1998). In fact, this is an exception because in vitro transformation is the general methodology to produce stable transformed plants, which is a laborious process that requires suitable facilities and moreover, regeneration procedure in a varied species ranging from months to a year making it a time taking process (Busov et al. 2005). The genes that confer the resistance to antibiotics or herbicides (Miki and Mchugh 2004) are the key elements in transgene integration which also raises the concerns on biosecurity (Darbani et al. 2007). To remove these marker genes many alternative strategies were developed that are complex. Consequently, rapid exclusion of the transgene left a challenge in plant biotechnology after genome editing, particularly for prolonged life cycle crops and multiploidy, to avoid transgene position effects, minimize the probability of off-target mutation appearance, and to deliver consumers with plants free of the recombinant gene editing machinery (Yau and Stewart 2013).

CRISPR-Cas9 applications include a hassle in counter selection based on resistance marker genes as the plants without the transgene can’t survive the selection and evaluating the transgene for more than two generations is an obligation. *Arabidopsis thaliana* was successfully tested for the presence of transgene prior to the germination with the expression of fluorescent proteins as selective markers (Stuitje et al. 2003). Moreover, it has also been used in combination with CRISPR in Arabidopsis (Gao et al. 2016). Despite its clear advantage, investigating on species such as tomato or rice is vital, and also need special requirements of in vitro transformation protocols (Zlobin et al. 2020).

**As a proof of concept**

Abbas et al. (2018b) has constructed a gene encoding IAA methyl transferase (IAMT) as a gene editing target in Arabidopsis (*Arabidopsis thaliana*), rice (*Oryza sativa* Linnaeus) and tomato (*Solanum lycopersicum* Linnaeus), given that loss of function results only in difficulty for hypocotyl reorientation after gravistimulation (Abbas et al. 2018a) and increased pollen tube growth rate (Abbas et al. 2018b), neither of which are traits that can bias our identification of mutations by direct observation unless specific tests are performed. The two plant species Arabidopsis (*A. thaliana*) and rice (*O. sativa*) were identified by a gene at5g55250 and os04g56950, respectively that encodes IAMT activity (Qin and Partridge 2005). On the other hand, in tomato two orthologues of IAMT1 Solyc07g64990 and Solyc12g14500 were identified by phylogenetic analysis. Therefore, tests were conducted on diverse editing strategies in each of the three selected species: targeting only one gene with one sgRNA (single guide RNA) in rice, simultaneously targeting two genes with two sgRNAs (in tomato), and targeting different regions of a single gene (in Arabidopsis) to evaluate the efficiency of the vectors when looking for multiple mutations and larger deletions. Preferably, the primary transformed plants include one copy of the transgene that would be segregated in the next generation in parallel to any CRISPR induced mutations in germ line, thus it could use DsRED, basic red fluorescent protein visualization as marker of transgene existence to select non fluorescent seeds and subsequently look for mutations. Virus interference in plants can be impacted by molecular tools, the CRISPR/Cas system of genome editing (Chaparro-Garcia et al. 2015). Ali et al. (2015) reported the CRISPR/Cas9 approach for protection to plants against Gemini viruses. The plant viruses Gemini viruses species including BCTV (Beet curly top virus), TYLCV (Tomato yellow leaf curl virus), and MeMV (Merremia mosaic virus) displayed enhanced resistance against the given viruses through this system (Chaparro-Garcia et al. 2015).

**Commercial success**

Since 1995, several new traits and combinations of traits have been developed and marketed in cotton cultivars. In 1996, Bollgard cotton hit the market, which had a transgene from bacterium *Bt* producing Cry1Ac which is an endotoxin (Begemann 1996). Bollgard cultivars control major lepidopteran pests such as the tobacco budworm (*H. virescens*), the pink bollworm (*Pectinophora gossypiella* Saunders), and suppress populations of bollworm (*Helicoverpa zea* Boddie). In 1998, cultivars possessing both the Cry I Ac *Bt* endotoxin and the Roundup Ready technology, often referred to as stacked gene cultivars, were introduced (Kerby and Voth 1998). Until 2004, no new types of transgenic technologies were introduced for cotton. In 2004, second generation of transgenic cultivars commenced. Cultivars were introduced that made improvements in both *Bt* and glyphosate resistance technologies. Bollgard II, a genetic technology whereby two *Bt* endotoxins were expressed by the cultivar, had an enhanced spectrum of control of lepidopteran insect pests compared to the single-gene *Bt* cultivars. Bollgard II technology was commercially released in 2004 in cultivars that also expressed the RR technology (Jost et al. 2008). These stacked traits along with the refuge strategy slowed down the evolution
of Bt resistance in bollworm for more time than predicted. In 2005, another two gene Bt technology were released by Dow Agrosciences. In 2006, an improved version of the Roundup Ready technology known as Roundup Ready Flex (RF), was made commercially available (Murdock and Mullins 2006). The genetic event conferring glyphosate resistance in the RR technology only fully protects cotton fruiting forms if glyphosate was applied to cotton foliage before the 4 leaf stage (Pline et al. 2002). Roundup Ready Flex technology permits application of glyphosate over the top of cotton throughout the period of fruit set. In 2006, Bollgard II technology in conjunction with Liberty-Link technology was introduced. Breeding procedures involved in the development of multi-stacked traits, some of the advantages and disadvantages associated with multi-stacked traits were detailed in Also, a list of multi-stacked traits, private industries who own them and the gene involved in each stack were detailed in Que et al. (2010).

In India, cotton (Gossypium hirsutum Linnaeus) and soybean (Glycine max Linnaeus) are the approved genetically modified crops. In 2014 GEAC (Genetic Engineering Appraisal Committee) approved 11 crops for the field trials which includes maize, rice, wheat, groundnut sorghum and cotton. A Moratorium was laid on Bt Brinjal in 2010 by the Indian Government which crippled the research on transgenic crops. The data generated by India was taken by the Bangladesh and 25,000 farmers cultivated Bt Brinjal and made it a success. USA, Brazil, Argentina, Canada, and India altogether have occupied 91% of the global biotech crop area (Brookes and Barfoot 2017). According to ISAAA (The International Service for the Acquisition of Agri-biotech Applications), the USA has 203 GM crops approved with 21 variants, cultivating food crops like maize, soybean, canola, sugar beet, papaya, squash, potato, livestock feed like alfalfa and a commercial crop cotton in nearly 70.1 million hectares in USA, followed by Brazil, Argentina, Canada and India. The information from ISAAA proclaims that around 2.7 billion hectares of biotech tech crops planted since 1996. Malawi, Ethiopia and Nigeria have planted Bt cotton for the first time in 2019.

Conclusions
Integrated pest management combined with other control methods like chemical, physical and planting both Bt and non-Bt together is more effective. Moreover, new molecular techniques have to be applied in order to overcome insect pest resistance by merging RNAI to Bt may considerably delay resistance, especially in bollworm. Molecular tools like CRISPR-based gene drivers can be employed to extend the target genetic elements through large numbers of inhabitants in combination with Bt-transgenic crops may lead to effective pest resistance management with ecofriendly methods.

| S. No | Event name | Genes incorporated | Source | Function |
|-------|------------|--------------------|--------|----------|
| 1     | BNLA-601   | cry1Ac             | Bacillus thuringiensis subsp. kurstaki strain HD73 | Confers resistance to lepidopteran insects by selectively damaging their midgut lining |
| 2     | JK 1 TRADE NAME | cry1Ac & nptII*   | Bacillus thuringiensis subsp. kurstaki strain HD73 | Confers resistance to lepidopteran insects & allows transformed plants to metabolize neomycin and kanamycin antibiotics during selection |
| 3     | GFM Cry1A  | cry1Ab-Ac delta endotoxin (fusion protein) nptII* uidA* | synthetic fusion gene derived from Bacillus thuringiensis | Confers resistance to lepidopteran insects & produces blue stain on treated transformed tissue, which allows visual selection |
| 4     | MLS 9124  | cry1C delta endotoxin | synthetic fusion gene derived from Bacillus thuringiensis | Confers resistance to lepidopteran insects, specifically Spodoptera |
| 5     | Bollgard II™ Cotton | cry1Ac & cry2Ab nptII* uidA, aad* | Bacillus thuringiensis subsp. kumamotoensis & Bacillus thuringiensis subsp. kurstaki strain HD73 | Confers resistance to lepidopteran insects by selectively damaging their midgut lining |
| 6     | Bollgard™ Cotton, Ingard™ | cry1Ac & cry2Ab nptII* aad* | Bacillus thuringiensis subsp. kurstaki strain HD73 | Confers resistance to lepidopteran insects by selectively damaging their midgut lining |
### Abbreviations

**Bt**: *Bacillus thuringiensis*; **SIT**: Sterile Insect Technique; **ICPs**: Insecticidal Crystal Proteins; **Cry**: Crystal toxins; **VIP**: Vegetative Insecticidal Proteins; **ALP**: Anchored Alkaline Phosphatases; **CADR**: Cadherin Like Proteins; **APN**: aminopeptidase N; **GPI**: Glycophosphatidylinositol; **Cry**: Crystal proteins; **Cyt**: Cytotoxic proteins; **VIP**: Vegetative Insecticidal Proteins; **CrisPR/Cas9**.

### The table gives information regarding genetically modified crops approved in India with *Bt* genes incorporated. Out of 11 approved GM crops 6 are cotton crops *Gossypium hirsutum* L and rest 5 crops are Soybean *Glycine max* L but Intacta™ Roundup Ready™ 2 Pro is the only *Bt* gene incorporated crop. All genes confer resistance to lepidopteran pest, MLS 9124 confers resistance to lepidopterans. *'* are the marker/reporter genes incorporated in the event crop. (Source: https://www.isaaa.org/gmapprovaldatabase/default.aspx).

| S. No | Event name | Genes incorporated | Source | Function |
|-------|------------|--------------------|--------|----------|
| 7     | Intacta™ Roundup Ready™ 2 Pro | cry1Ac & cp4 epsps* | Bacillus thuringiensis subsp. *kurstaki* strain HD73 | Confers resistance to lepidopteran insects by selectively damaging their midgut lining, conferring increased tolerance to glyphosate herbicide |

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#### Competing interests

The authors declare that they have no competing interests.

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