

# We are IntechOpen, the world's leading publisher of Open Access books Built by scientists, for scientists

6,900

Open access books available

185,000

International authors and editors

200M

Downloads

Our authors are among the

154

Countries delivered to

TOP 1%

most cited scientists

12.2%

Contributors from top 500 universities



WEB OF SCIENCE™

Selection of our books indexed in the Book Citation Index  
in Web of Science™ Core Collection (BKCI)

Interested in publishing with us?  
Contact [book.department@intechopen.com](mailto:book.department@intechopen.com)

Numbers displayed above are based on latest data collected.  
For more information visit [www.intechopen.com](http://www.intechopen.com)



# Molecular Identification of Genetically Modified Crops for Biosafety and Legitimacy of Transgenes

*Shahid Nazir, Muhammad Zaffar Iqbal and Sajid-ur-Rahman*

## Abstract

Crops undergo artificially DNA modifications for improvements are considered as genetically modified (GM) crops. These modifications could be in indigenous DNA or by introduction of foreign DNA as transgenes. There are 29 different crops and fruit trees in 42 countries, which have been successfully modified for various traits like herbicide tolerance, insect/pest resistance, disease resistance and quality improvement. GM crops are grown worldwide and its area is significantly increasing every year. Many countries have very strict rules and regulations for GM crops and are also a trade barrier in some situations. Hence, identification and testing of crops for GM contents is important for identity and legitimacy of transgene to simplify the international trade. Normally, molecular identification is performed at three different levels, i.e., DNA, RNA and protein, and each level has its own importance in testing about the nature and type of GM crops. In this chapter, current scenario of GM crops and different molecular testing tools are described in brief.

**Keywords:** biotechnology, genetic engineering, transgenic plants, molecular testing, polymerase chain reaction, enzymes linked Immuno-sorbent assay

## 1. Introduction

Biotechnology is a set of scientific tools in which living organisms are used for the welfare of mankind. This technique is efficiently used to modify and improve plants, animals and other microorganisms to increase their value. Biotechnology has a very wide range of applications and almost every field of daily science get benefit from this technology. Application of biotechnology in the field of agriculture has been practiced for a long time as people have wanted to improve agriculturally important crops by selection and breeding. In 1970s with the advancements in molecular biology, researchers were able to modify DNA which is a chemical building block and specify the features of living organisms at molecular level. This modification in genetic material or DNA is called as recombinant DNA technology or genetic engineering [1]. With the involvement of genetic engineering in agriculture, one can transfer useful hereditary/genetic information from distant sources into targeted crop which was not possible through traditional breeding methods. This genetic information is coded in the form of DNA or genes. Genes from any living organisms (human, animal, plant and microorganism) could be easily manipulated

and transferred into other organisms to enhance their value. Organisms artificially modified at genome level using genetic engineering tools are termed as genetically modified organisms (GMOs). Microorganisms, i.e., bacteria and viruses have been genetically modified for the production of different kinds of medicines, pharmaceuticals and food ingredients [2]. Genetic engineering also has a great role in the field of agriculture by developing the transgenic crops for various traits. For example, a useful gene from bacteria, fungi and animals etc. could be isolated, cloned and integrated into desired crop to develop resistance against diseases and pests, drought and salinity tolerance or to improve the quality related traits etc. and are known GM crops [3]. After transformation, the transgenes replicate with indigenous plant genes and produce specific protein [4]. Biotechnology supports in practical exploitation of genetic material for the betterment of mankind. By using latest trends in genetic engineering one can create the new face of existing cultivars with improved and desirable characteristics. In addition to the improvement of agronomic traits, scientists are also looking in the production and expression of commercially valuable protein in plants like spider silk protein and polymers used in surgery [5]. A huge number of human vaccines, antigens and other pharmaceutical products are very efficiently expressing in transgenic plants. GMO offer many benefits to humans, but at the same time people also worry about the possible threats of using GMOs. These risks include the possible introduction of allergens in GM foods and transfer of selection marker genes which are normally antibiotic resistant genes to gut flora [6–8].

With the introduction of foreign genes, there are also some biosafety issues linked with GM crops. Such crops are often unintentionally or intentionally used for food and feed production. In some conditions, GM crops spread globally by trading, transportation and storage either intentionally or unintentionally and contaminate GM free items. Many countries have very strict rules and regulations for the development, cultivation, commercialization and labeling of GM crops and is also a trade barrier in some situations [9]. For example, USA has an optional labeling of GM in food items, whereas European Union has very strict rules for approval, cultivation and use of GM crops, including a compulsory labeling system [10]. They require very comprehensive information about such crops like type of targeted crop and transgene, safety for humans, environment, animals and effects on other related non-modified crops [11–13]. The increase in GM crop production has been coupled with an intricate and asynchronous international regulatory approval system, requiring identification and testing of food and agricultural products for the presence of GM content to simplify international trade. Molecular identification of GM crops confirms the identity and type of modified product at each stage and assures compliance with import for GM food and feed [14]. The testing of GM crops could be performed in open field or under controlled laboratory conditions that depends upon type of samples and sensitivity of test performed. Normally, molecular identification and testing of GM crops is performed at three different stages, i.e., DNA, RNA and protein. Each testing level has its own importance in testing the nature and type of GM crops. Generalized GM development methodology, global status, testing methods, possible biosafety issues and other benefits etc. are discussed in brief.

## **2. Global scenario of GM crops**

The rapid acceptance of GM crops shows the significant benefits realized by large and small growers in both developed and under-developed countries growing GM crops commercially. Around 99% of global GM crops area is occupied by four major crops, i.e., soybean, maize, cotton and canola. USA is leading in the area under GM crops with 75.0 million hectares followed by Brazil and Argentina with 50.2 and 23.6 million hectares, respectively. In 2017, 24 countries planted 189.8

million hectares with an increase of 3.0% than 2016 [15]. Despite the possible health risks, cultivation area of GM crops is regularly increasing and introduction of new GM crops is continued. There are 29 different crops and fruit trees in 42, which countries have been successfully modified for various traits. A brief detail of GM crops, targeted/GM traits, number of GM events with responsible transgenes has been given in **Table 1**. Among GM trait distribution, herbicide

| Sr. # | GM crops                                   | GM events | GM traits   | Transgenes  |
|-------|--|-----------|---|---|
| 1     | Alfalfa                                    | 05        | Herbicide tolerant, Modified Product Quality  | <i>cp4 epsps (aroA:CP4)</i> , <i>ccom1 (inverted repeat)</i>  |
| 2     | Apple                                      | 03        | Modified Product Quality  | <i>PGAS PPO suppression gene</i>  |
| 3     | Argentine Canola ( <i>Brassica napus</i> ) | 41        | Modified Product Quality, herbicide tolerant, Pollination control system  | <i>te</i> , <i>gat4621</i> , <i>pat</i> , <i>cp4 epsps (aroA:CP4)</i> , <i>goxo247</i> , <i>bar</i> , <i>barnase</i> , <i>barstar</i> , <i>phyA</i> , <i>bxn</i> , <i>Lackl-delta12D</i> , <i>Picpa-omega-3D</i> , <i>Micpu-delta-6D</i> , <i>Pyrco-delta-6E</i> , <i>Pavsa-delta-5D</i> , <i>Pyrco-delta-5E</i> , <i>Pavsa-delta-4D</i> ,  |
| 4     | Bean                                       | 01        | Disease resistance  | <i>ac1 (sense and antisense)</i>  |
| 5     | Carnation                                  | 19        | Herbicide tolerance , Modified flower color, Delayed ripening/senescence  | <i>dfr</i> , <i>hfl (f3'5'h)</i> , <i>surB</i> , <i>bp40 (f3'5'h)</i> , <i>dfr-diaca</i> , <i>cytb5</i> , <i>acc (truncated)</i>  |
| 6     | Flax                                       | 01        | Herbicide tolerance   | <i>als</i>  |
| 7     | Maize                                      | 231       | Male sterility , Fertility restoration , Modified alpha amylase , Herbicide tolerance, Insect resistance, Phytase production, Modified amino acid, Increased Ear Biomass, Drought stress tolerance, | <i>ms45</i> , <i>zm-aa1</i> , <i>amy797E</i> , <i>cry1Ab</i> , <i>pat</i> , <i>mepsps</i> , <i>mcry3A</i> , <i>ecry3.1Ab</i> , <i>cry1Fa2</i> , <i>cry1F</i> , <i>cry34Ab1</i> , <i>cry35Ab1</i> , <i>vip3Aa20</i> , <i>cp4 epsps (aroA:CP4)</i> , <i>cry3Bb1</i> , <i>aad-1</i> , <i>dam</i> , <i>zm-hra</i> , <i>gat4621</i> , <i>vip3A(a)</i> , <i>cry2Ab2</i> , <i>cry1A.105</i> , <i>phyA2</i> , <i>bar</i> , <i>cry9C</i> , <i>pinII</i> , <i>cry1Ac</i> , <i>2mepsps</i> , <i>cordapA</i> , <i>goxo247</i> , <i>athb17</i> , <i>dvsnf7</i> , <i>dmo</i> , <i>cspB</i> , <i>barnase</i> , <i>mocry1F</i> <i>epsps grg23ace5</i> |
| 8     | Cotton                                     | 60        | Herbicide tolerance, insect resistance  | <i>S4-HrA</i> , <i>pat</i> , <i>cry1Ac</i> , <i>cry1F</i> , <i>vip3A(a)</i> , <i>aad-12</i> , <i>cp4 epsps (aroA:CP4)</i> , <i>bar</i> , <i>bxn</i> , <i>cry2Ab2</i> , <i>cry2Ae</i> , <i>2mepsps</i> , <i>cry1Ab-Ac</i> <i>cry1C</i> , <i>dmo</i> , <i>CpTI</i> ,  |
| 9     | Potato                                     | 48        | insect resistance , Modified starch/carbohydrate, Reduced Acrylamide Potential , Reduced Black Spot, Viral disease resistance, Fungal Disease Resistance, Herbicide Tolerance                       | <i>cry3A</i> , <i>gbss (antisense fragment)</i> , <i>asn1</i> , <i>ppo5</i> , <i>pvv_cp</i> , <i>plrv_orf1</i> , <i>plrv_orf2</i> , <i>cp4 epsps (aroA:CP4)</i> , <i>Rpi-vnt1</i>   |
| 10    | Linseed                                    | 01        | herbicide tolerance   | <i>als</i>  |
| 11    | Eucalyptus                                 | 01        | Volumetric Wood Increase  | <i>cel1</i>   |
| 12    | Eggplant                                   | 01        | insect resistance   | <i>Cry1Ac</i>   |
| 13    | Creeping Bentgrass                         | 01        | Herbicide tolerance   | <i>cp4 epsps (aroA:CP4)</i>   |
| 14    | Chicory                                    | 03        | Herbicide Tolerance, Male sterility   | <i>bar</i> , <i>barnase</i>   |
| 15    | Soybean                                    | 40        | Modified Product Quality, Herbicide Tolerance, Insect resistance,   | <i>gm-fad2-1 (silencing locus)</i> , <i>pat</i> , <i>csr1-2</i> , <i>2mepsps</i> , <i>aad-12</i> , <i>cp4 epsps (aroA:CP4)</i> , <i>cry1Ac</i> , <i>cry1F</i> , <i>gm-fad2-1 (partial sequence)</i> , <i>gat4601</i> , <i>gm-hra</i> , <i>hppdPF W336</i> , <i>Hahb-4</i> , <i>fatb1-A (sense and antisense segments)</i> , <i>fad2-1A (sense and antisense)</i> , <i>dmo</i> , <i>bbx32</i> , <i>cry1A.105</i> , <i>cry2Ab2</i> , <i>Pj.D6D</i> , <i>Nc.Fad3</i> , <i>avhppd-03</i> , <i>bar</i>   |
| 16    | Poplar                                     | 02        | Insect Resistance   | <i>cry1Ac</i> , <i>API</i>  |
| 17    | Plum                                       | 01        | Viral disease resistance  | <i>ppv_cp</i>   |
| 18    | Petunia                                    | 01        | Modified Product Quality  | <i>chsA</i>   |
| 19    | Papaya                                     | 02        | Viral disease resistance  | <i>prsv_rep</i> , <i>prsv_cp</i>  |
| 20    | Melon                                      | 02        | Delayed ripening/senescence   | <i>sam-k</i>  |
| 21    | Rose                                       | 02        | Modified Product Quality  | <i>5AT</i> , <i>bp40(f3'5'h)</i>  |
| 22    | Rice                                       | 08        | Modified Product Quality, Insect Resistance, Herbicide Tolerance  | <i>ctrl</i> , <i>psy1</i> , <i>7crp</i> , <i>cry1Ab</i> , <i>cry1Ac</i> , <i>bar</i> , <i>cry1Ab (truncated)</i>  |
| 23    | Squash                                     | 02        | Viral disease resistance  | <i>cmv_cp</i> , <i>zymv_cp</i> , <i>wmv_cp</i>  |
| 24    | Sugar beet                                 | 03        | Herbicide tolerance,  | <i>cp4 epsps (aroA:CP4)</i> , <i>goxo247</i> , <i>pat</i>   |
| 25    | Sugarcane                                  | 04        | Insect resistance, Drought stress tolerance   | <i>cry1Ab</i> , <i>EcBetA</i> , <i>RmBetA</i>   |
| 26    | Sweet pepper                               | 01        | Viral disease resistance  | <i>cmv_cp</i>   |
| 27    | Tobacco                                    | 01        | Nicotine reduction  | <i>NtOPT1 (antisense)</i>   |
| 28    | Tomato                                     | 11        | Delayed ripening/senescence , Insect resistance, Delayed fruit softening, Modified Product Quality, Viral disease resistance  | <i>acc (truncated)</i> , <i>sam-k</i> , <i>cry1Ac</i> , <i>accd</i> , <i>pg (sense or antisense)</i> , <i>anti-efe</i> , <i>cmv_cp</i>  |
| 29    | Wheat                                      | 01        | Herbicide tolerant  | <i>cp4 epsps (aroA:CP4)</i>   |

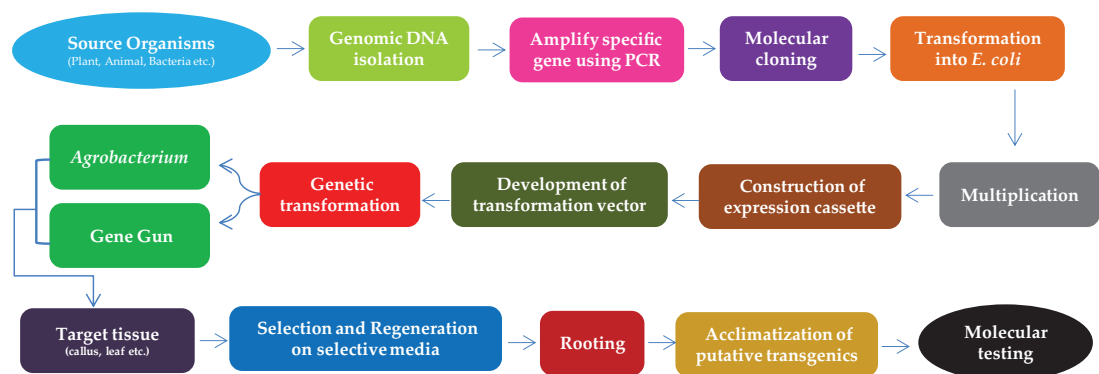
**Table 1.**  
Summary of GM crops with modified traits and introduced transgenes.



tolerance (HT) enjoys the top position with 47% of the GM crops area. Stacked traits and insect resistance (IR) occupy 41 and 12% of the cultivated area of GM crops in 2017, respectively. The cultivation area under stacked traits, i.e., HT/IR is increasing very fast and various stacked gene products were got approved for food/feed and general commercialization. Soybean, maize and cotton are major crops developed with stacked traits [16]. Countries approving GM crops for food, feed and general cultivation are also increasing every year. In year 2017, 18 countries issued 176 approvals regarding GM crops cultivation, commercialization and use as food/feed [15].

### 3. GM crop's development methodology

Plants, in which one or more foreign genes are introduced artificially instead of plant getting them under natural conditions of cross-breeding or normal recombination, are known as GM plants. The introduced gene, known as transgene, could be from identical species or from different species within the same kingdom or other kingdom [17]. The process of introducing the transgene is called as genetic transformation that has become an important tool for crop improvement. Different steps are involved in the genetic transformation work like selection and identification of gene of interest (transgene), isolation from source organisms, cloning into suitable plasmid vector. Followed by development of expression vector containing all regulatory elements, i.e., promoters and terminators for regulation of transgene expression in targeted plants [18]. In addition, another gene cassette of selection is also the part of expression vector which serves as the primary selection of putative transgenic cells on artificial plant media. Normally two types of selection markers are used, antibiotic and visual selection markers, which depend upon the type of work. Final expression cassette is multiplied in suitable bacterial media and verified using various molecular biology techniques before transformation [19]. Integration of final expression cassette into plant can normally be achieved by two methods: (i) direct DNA delivery system, i.e., using biolistic gene gun by coating DNA on gold or tungsten particles and shooting on plant tissue with a specific pressure of helium gas (ii) introduction of gene by using biological vectors like disarmed Ti-plasmid of *Agrobacterium tumefaciens*. Both methods have successfully been used for the introduction of transgenes in plants [20]. Following genetic transformation, the transformed tissues are initially screened for transgene integration using selective plant tissue culture media. The regenerated plantlets on selective media supposed to have the transgenes and called as putative transgenics. Because there are three possibilities that the developed plantlets may be (i)



**Figure 1.**  
General methodology for the development of genetically modified (GM) plants.

true transgenics (ii) escapees (iii) mutants. Hence, various molecular biological techniques like PCR, blotting, *Enzyme-linked ImmunoSorbent Assay* (ELISA) etc. are used to confirm transgene integration and true transgenics. The overall methodology for gene isolation, cloning, transformation and selection of putative transgenics has been shown in **Figure 1**.

#### **4. Molecular test methods for the identification of GM crops**

Introduction of GM crops and their products in markets required to be monitored and need to know the presence and type of GM elements. Labeling rules and trade requirements vary from country to country which necessitates for the development of reliable methods for the detection, identification and quantification of GM crop varieties and their products. GM crops can be tested by identifying either transgenes at DNA level, at transcriptional level by mRNA of transgene or using resulting trans-protein. There are many other methods like chromatography and mass spectrometry etc. which have their own importance in GMO testing. An overview of test methods used for detection and identification for GM crops has been given in **Figure 2**. Every test method has its own significance and value towards the final conclusion of GM crops. A brief summary of these methods has been shown in **Table 2**.

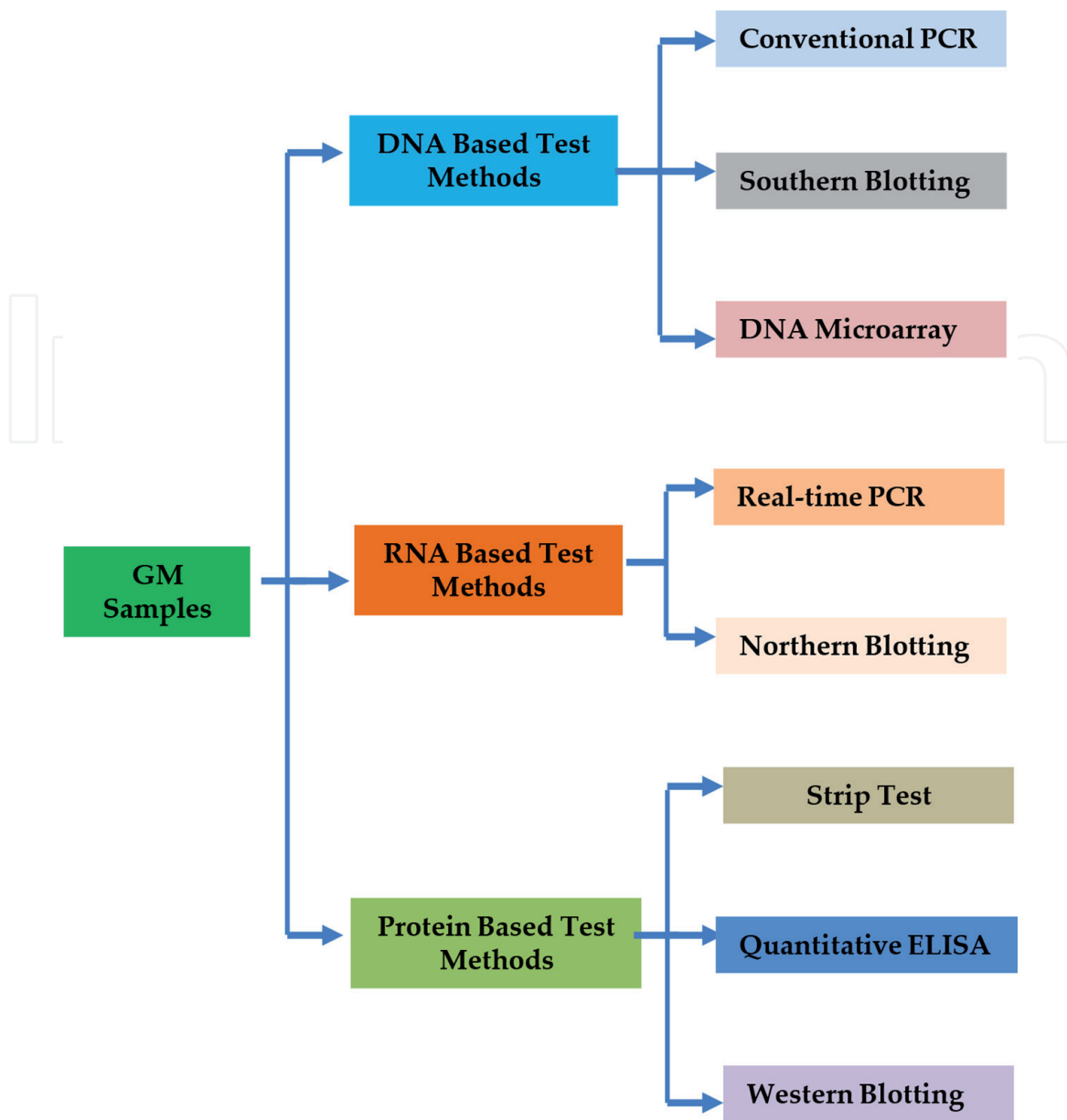
##### **4.1 DNA based test methods**

There are three main types of DNA based GMO testing methods.

###### *4.1.1 Qualitative PCR*

Polymerase Chain Reaction (PCR) is a primary method for screening of GM crops at DNA level. Qualitative analysis comprises of specific detection of target DNA sequence in test samples. Qualitative results clearly validate the presence or absence of GM elements under study, comparative to suitable controls and within the detection limits of analytical technique used, and test portion analyzed [21, 22]. This method has found very broad and wide applications in GMO detection as commonly accepted tool for regulatory purposes. In this method target gene/GM element multiplied to millions or billions by using gene specific primers. PCR process is basically comprised on three main steps, i.e., denaturation, annealing and extension in one cycle. In first step the double stranded DNA is separated into two single strands, primers then identify their homologous sequence and are annealed to each strand in second step. Third and final step involves making two identical copies of original DNA strand by adding exact nucleotides with the help of DNA polymerase at an appropriate temperature. These cycles repeated normally 40–50 times which results in an exponential amplification of target DNA/gene. Amplification of target gene occur in-vitro through a reaction catalyzed by a DNA polymerase in the presence of oligonucleotide primers and deoxyribonucleoside triphosphates in a defined reaction buffer [23, 24]. This amplified DNA can be visualized by using gel electrophoresis techniques. The results of this method will be either positive or negative for specific GM elements.

There are four testing methods which includes (i) Target-taxon specific (ii) Screening (iii) Construct-specific and (iv) Event-specific, these methods are generally used for the detection and identification of GM crops using PCR. Selection of specific and suitable primers is the most critical step in GMO detection which depends upon the testing method used. Brief detail of qualitative PCR based testing methods is given below:



**Figure 2.**  
*Diagrammatic presentation of molecular test methods of GM crops.*

**4.1.1.1 Target-taxon specific method**

PCR with various barcoding methods normally used for plant identification from mixed food samples particularly prepared from different plants. DNA barcode is broadly used technique for the detection and identification plants, animals or fungi taxa by sequencing an optimized short DNA fragment. PCR and barcoding approaches identify specific taxa very intelligently within samples of different origins [25–27]. This approach also plays very important role in the detection of mislabeled species and accidental or intentional species exchanges in food samples [28, 29]. The success of this method for identification and detection of species depends on the selected loci, because DNA barcode constitute a small portion of genome coupled with other PCR limiting factors, no single locus has been selected as universal DNA barcode region for all plant identification. For example lectin gene Le1 for soybean [30], chloroplast trnL intron for the identification of multicopy

| Test methods  | Test Name                  | Testing sample                                    | Sensitivity /limit of detection | Conditions required     | Advantages/ disadvantages  |
|---------------|----------------------------|---|---------------------------------|-------------------------|--|
| DNA based     | Conventional PCR           | DNA from Leaf/seed/processed food/feed items etc. | Highly sensitive/ 0.01%         | Laboratory              | Easy, broad range, used worldwide for GMO analysis   |
|               | Southern blotting          |   | Highly sensitive                |                         | Require more time, sophisticated equipment/expertise, expensive, not used frequently   |
|               | DNA Microarray             |   | Highly sensitive                |                         |  |
| RNA based     | Quantitative Real-time PCR | RNA converted in to cDNA before used in PCR       | Highly sensitive                | Laboratory              | Broad range, allow actual quantification, require sophisticated equipment/expertise, expensive                                       |
|               | Northern blotting          | RNA   | Highly sensitive                |                         | Require more time, sophisticated equipment/expertise, expensive, not used frequently   |
| Protein based | Strip test                 | Protein   | Low sensitive/ 0.1-1.0%         | May be perform in field | Easy, quick, cheap, not suitable for processed food  |
|               | Qualitative ELISA          |   | Highly sensitive/ 0.01-0.1%     | Laboratory              | Not suitable for processed food, require expertise, expensive, results may vary from person-to-person and lab -to-lab of same sample |
|               | Western blotting           |   | Highly sensitive                |                         | Require more time, sophisticated equipment/expertise, expensive, not used frequently   |

**Table 2.**  
Brief summary of GMO test methods.

DNA sequences in plants [31], polygalacturonase gene (PG gene) codes for a PG-enzyme that is linked with ripening in GM Zeneca tomato etc. [32].

4.1.1.2 Screening method

This is a most generalized method and widely used for the screening of GM crops from non-GM materials. This is not crop specific and can detect the GM elements even in raw and processed matrices like food and feed products developed from GM crops. In this method promoter, terminator and selection marker genes are the target elements in PCR. These are the bacterial gene sequences used to regulate the transgenes and selection of transgenic cells on artificial plant media [33, 34]. These genetic elements include cauliflower mosaic virus CaMV 35S promoter, *Agrobacterium tumifaciens* nopaline synthase NOS terminator and neomycin phosphotransferase NPTII etc. present in most commercialized GM crops in market. Hence, one can easily detect and identify the presence of GM crop by using specific primers of these genetic elements in PCR [35, 36].



#### 4.1.1.3 Construct-specific method

In this method specific primer pairs normally got designed from the transformed gene construct. These construct could be transformed more than one crop for genetic improvement. The construct-specific detection method involves targeting the junction between two elements, and it is not able to distinguish two different events transformed with the same plasmid [37]. These methods either DNA or protein based. For practical reasons, several DNA sequences are shared by many GM crops and protein based methods detect the product of inserted DNA like Bt toxin in GM crops. Since different GM crops may produce the identical protein, this test method can detect a sample for several GMOs in one step. For examples GTS-40-3-2 GM construct for the development of roundup ready soybean, Zeneca F282 GM tomato, Bt11, Bt176 and T25 for GM maize etc.

#### 4.1.1.4 Event-specific method

The junction sequences in the transgene integration points in the plant genome can be used to identify and detect the specific transformation event. The transgene integration site usually unique and specific for each transformation event due to lack of homologous recombination. Hence, different GM crops could be produced with similar gene construct and this event-specific detection method will be the only approach to differentiate between GM crops having similar transgenic cassette. This method can distinguish legitimate transgenic events from related unauthorized genotypes/varieties having identical transgene construct, thus this approach frequently used to assess the legality of GM crops [24]. Examples are Mon-531 event for Bollgard cotton, Mon-1445 event for Roundup Ready cotton, Mon-89,034 event for YieldGard VT Pro maize etc.

#### 4.1.2 Southern blotting

Another DNA based GM crops identification techniques is southern blotting which was described by Southern in 1975 [38]. This test method is frequently used for the identification of specific DNA fragments transformed into the genome of transgenic plants or its products. This method could also be used in gene discovery and mapping, evolution and developmental studies, diagnostics and forensics etc. This test method involved five steps (i) DNA isolation and enzyme restriction (ii) electrophoresis for DNA separation (iii) shifting and fixing of separated DNA on suitable membrane (iv) hybridization with labeled probe and (v) detection by chemiluminescence or radioactive methods. This is very reliable method that provides the molecular evidence of the transgene integration and also estimates the copy number of introduced gene into the GMO genome. In comparison with PCR, this method associated with some limitations like it requires large amount of DNA, expensive, requires more time, proper infrastructure and trained manpower etc.

#### 4.1.3 DNA microarray

A microarray is a laboratory method used to identify the expression of more than one gene in a single test. It is DNA based and new in comparison to previous protocols. This test method has been included in GMO screening as a method for simultaneous detection of more than 250,000 targets in single assay/chip [39–41]. This method consists of pre-amplification step of the desired targets, followed by hybridization on a chip having specific probes, and then detection step [42, 43]. So far, it is used for qualitative information of GMO, sometime semi-quantitative.

Use of microarray technology for the GMO detection is restrained as it requires very special and costly equipment for scanning microarrays, chances of cross contamination and laborious in comparison with other techniques.

## **4.2 RNA based test methods**

Transgenic DNA must be translated into protein to be an effective and have effects in an organism. This translation process occurs when DNA is transcribed into messenger RNA (mRNA), and is considered as the intermediate step transferring information contained in DNA to protein. The presence of mRNA is directly associated with gene expression. Different molecular biology techniques used to monitor and study the gene expression in GMOs include real-time PCR, northern etc. These methods could be used to identify the transgene expression in various plant tissues and at different developmental phases in GMOs.

### *4.2.1 Quantitative real-time PCR*

Gene expression is normally verified in RT-PCR using isolated mRNA from GMOs. This test method is based on reverse transcription of mRNA and synthesis of complementary DNA (cDNA) which is then used as template in PCR amplification of target gene. The amplified fragment is electrophoresed and visualized using agarose gel under UV. Intensity of amplified band in agarose gel gives some indications of target mRNA in tested sample [44]. Quantitative RT-PCR is an up-to-date method, principally based on RT-PCR and is generally known as qRT-PCR. It is more robust, specific and sensitive, provides good quantitative results. The process of amplification is presented in real-time by capturing a fluorescent signal in a more sophisticated way. In real-time assay of transgene in GMOs, the amplification and detection occur simultaneously [45].

### *4.2.2 Northern blotting*

Similar to RT-PCR, northern blotting also requires mRNA as tested material from GMOs. This is a standard method for the analysis of size and level of target RNA in a complex GMO sample. Likewise southern blotting, it also composed of five steps, only difference is that the starting material is mRNA instead of DNA and the labeled probe is complementary DNA (cDNA), which hybridizes the RNA. It gives comparative amount of gene expression at the RNA level. This is comparatively simple to perform, cheap and not overwhelmed by artifacts [46]. Recent advancements of hybridization membranes and buffers have resulted in increased sensitivity, closing the gap to the more laborious nuclease protection experiments. It is considered that this test method can study gene expression for a limited number of genes per analysis. This can be very useful to monitor the up- or down regulation of transgene for specific problem, but is not useful in monitoring the up- or down regulated genes which are unknown.

## **4.3 Protein based test methods**

Immunoassay protocols for the detection of GMOs by antibodies are the impressive for the detection of various types of proteins either qualitatively or quantitatively [47]. Two types of antibodies, i.e., monoclonal and polyclonal could be used depending on the need and specificity of detection method. Normally, Enzyme-linked Immunosorbent Assay (ELISA) and western blot methods have been used for the protein analysis in GMOs.

#### 4.3.1 Qualitative strip test

Most common antibody based test for GMO screening is strip test method also known as lateral flow or dipstick test. It is qualitative in nature and gives the information about the presence or absence of specific proteins in tested samples. In this method, thin strip made-up of nitrocellulose membrane used which protected by a sample pad on one end and a wicking pad on other end. Test samples normally homogenized in suitable buffer solutions and membrane on strip wicks up the solution and it will move upward via capillary movement and protein will bind to its specific antibody. The results shown in the form of visible lines on the strip depicting that the specific protein is present in test sample. There are normally two lines appears on the strip, one for tested protein and second of control line showing the authenticity of all test procedure and strip used. The appearance of only control line on the strip, shows that sample is negative for transgenic protein, but the test was performed accurately [48]. This is very quick method to test GMOs which normally take 5–15 minutes to gives results [49]. In addition, it is cheap, easy to perform and not require specific equipment and special trained manpower. It can be performed in open field as well. Currently, strips are available to detect multiple proteins in single assay [50].

#### 4.3.2 Quantitative ELISA test

Another more sensitive antibody-based protein identification method is Enzyme-linked ImmunoSorbent Assay (ELISA) also called a plate test or quantitative ELISA. It gives information about the quantity of protein in tested samples. In this assay protein specific antibody coated multi-well plate is used to identify and quantify the specific protein. Specific protein present will bind to antibody, following washing, another antibody specific for protein of interest and tagged with an enzyme is added to well [51]. The enzyme linked identification antibody will bind with specific protein and unbound antibody removed by washing. The color of the solution will change from blue to yellow by the addition of substrate for enzyme. Intensity of yellow color is directly proportional to amount of protein present in well. This GMO test method is more sensitive in comparison with strip test and can detect target protein even in very low concentrations. However, it requires more time, trained manpower and good laboratory facilities in contrast to strip test.

#### 4.3.3 Western blotting

This is very specific method and provides the qualitative results of the target protein in GM crop sample. This method is very useful to analyze the insoluble proteins [47, 50]. Like other blotting techniques samples are solubilized with detergents and reducing agents and separated by electrophoresis and shifted to membrane. Binding immunoglobulin sites on membrane are blocked by dried nonfat milk and specific sites are probed with antibodies. Detection carried out using different staining agents silver nitrate of Coomassie, alkaline phosphatase etc. [18]. Its detection limit varies with test ample like 0.25% for seeds and 1.0% for toasted meal [52]. In comparison with other protein based assays, it is difficult method, and is capable of studying only a few samples at a time. Therefore, it is not frequently used in GMO testing activities but it is more used in research purpose to verify initial results generated by other testing method.

## 5. Certified/standard reference materials for GMO testing

Validity and authenticity of GMO testing results is doubtful until the use of positive and negative controls at each testing step. Use of certified reference material (CRM) or standard reference material (SRM) during testing produce not only validate the testing results but at the same time, assess the performance of test method, equipment, personnel and other environmental conditions in which testing being performed [52]. CRM must contain the certificate of analysis, should be prepared by following ISO-Guide 34, have information about which GM events or elements present and what is its concentration, storage requirements, preparation and expiry date etc. While SRM have all the similar information but lacks the certificate of analysis and was not prepared by a certified company. Both CRM and SRM could be used to validate the testing results but CRM is more reliable and globally acceptable. Each GMO needs specific CRM which is used in testing and conclusion about the presence of specific GM event/element in testing samples. Normally seeds of GM and Non-GM crops are mixed at specific percentage and homogenized to make powder before analysis [51]. The availability of CRM is presently restricted due to some concerns of IPR and expenses [53]. The Institute of Reference Materials and Measurements at the Joint Research Center (JRC) in Geel, Belgium, FAPAS Fera Science Ltd., Sand Hutton UK, American Oil Chemists Society (AOCS), Urbana, Illinois, USA etc. are authorized companies to produce and sale of CRMs for GMO testing.

## 6. Pros and cons of GM crops

The most common improvement by the introduction of GM crops is the increase in yield and quality. There are many yield limiting factors like insect/pest, diseases, drought, heat, salinity, rapidly changing climatic conditions etc. Conventional approaches like irrigations, sprays and use of fertilizers etc. done a great job but the problem was increasing day-by-day. By the introduction of recombinant DNA technology in agricultural sector, scientists successfully develop the new face of existing cultivars with improved and desirable traits. The GM technologies increase the opportunities for plant breeders to develop crops that are protected from climatic stresses and attacks of insects and diseases [54, 55]. The crops have been successfully modified for herbicide tolerant, insect/pest resistance, disease resistance, abiotic stress tolerance, micronutrient enrichment etc. Furthermore, this technology helping us to improve the nutritional quality, longer shelf life, foods that are more appealing to eat and easier to transport. Development of various biopharmaceuticals and expression of human therapeutic proteins in plants also a great contribution of GM technology to improve the human life [56].

On the other hand there are also some biosafety issues linked with the use of GM crops. Biosafety means the need to protect human and animal health from possible adverse effects of GM technology. There are some reports about the potential threats linked with the use of GMOs like risks of allergenicity, development of herbicide tolerant weeds and resistant insects, harms to non-target organisms, selection marker gene could induce antibiotic resistant and reduce the effectiveness of antibiotics to cure disease etc. [7, 57–58]. Turning on of certain genes due to the use of strong promoters and might be harmful in humans, effects on the nutritional profiling, transgene may flow from non-target crops/weeds etc. Biosafety is an essential to modern biotechnology and the adoption of biotech products requires to be balanced with acceptable biosafety safeguards. Participation of different



stakeholders and dissemination of information and knowledge in public about GM products is much important to safe use of this technology.

## 7. National scenario of GM crops, biosafety, labeling and trade aspects

Agriculture sector of Pakistan plays a dominant role in the economy with 18.9% contribution in Gross Domestic Product (GDP) and engages 42.3% labour force. It is also a chief source of foreign exchange earnings and provide raw material for progress of other sectors [59]. Pakistan stands at seventh position among 26 countries growing GM crops, and insect resistant GM cotton of Mon-531 event is the only crop grown in the country with adoption reaching 2.9 million hectares of total 3.0 million hectares cotton crop area. Mon-531 is the only approved commercialized GM event in the country having insecticidal *Cry1Ac* gene of *Bacillus thurengiensis* to control the lepidopteron insects. In 2015, US\$398 million economic gain was estimated with the adoption of GM cotton [60]. Moreover, field trials of GM maize hybrids have successfully been conducted for single and stacked insect resistant (IR) and herbicide tolerances (HT) traits. For single HT trait, Monsanto event NK-603 was tested, while for stacked traits, i.e., IR/HT, Mon-89,034 x NK-603, TC-1507 x Mon-810 x NK-603 and TC-1507 x NK-603 were studied. These GM traits were officially approved for commercial cultivation by National Biosafety Committee in 2016. Field performance trials were completed as the part of regulatory requirements and varietal registration by Federal Seed Certification and Registration Committee of National Food Security and Research ministry [15]. In near future, GM maize having IR and HT traits will be grown by farmers, and it will be the second approved GM crop in the country.

Pakistan is signatory to World Trade Organization (WTO) and has sanctioned Convention on Biological Diversity (CBD) in 1994 and Cartagena Protocols in 2009. Different legislations under the Agreement of Trade Related Aspects of Intellectual Property Rights have been disseminated in the country. In addition, Pakistan Biosafety rules were designed in 2005, which are responsible for safe use of GM technology, manufacturing, import and storage of GMOs. Following these, National Biosafety Guidelines were developed in which the procedures to undertake all linked activities to GMO work were highlighted. These guidelines were framed in accordance with the recommendations of Food and Agriculture Organization (FAO), World Health Organization (WHO), United Nations Industrial Development Organization (UNIDO) and United Nations Environment Programme (UNEP). As per these guidelines the biosafety aspects of GMO work are monitored at three different levels, i.e., Institutional Biosafety Committee (IBC), Technical Advisory Committee (TAC) and National Biosafety Committee (NBC). The applications related to GMO work are submitted to IBC, and after thorough evaluation, the case is submitted to TAC for assessment and recommendations, while NBC is the final body to take further action regarding its approval or rejection. NBC is responsible to look after the laboratory research, field studies, commercial release, imports, exports and sale/purchase of GMOs and their products [61].

Pakistan exports rice, cotton, fruits (oranges and mangoes), vegetables and fish to its neighboring states, Middle East and Central Asian countries. IR cotton of Mon-531 event is the only one GM crop officially approved for general cultivation in the country. Very comprehensive testing procedures are adopted to test and verify the status of approved events in the candidate cotton varieties. Around 49 universities and 07 research institutes are actively involved in the teaching and research related to the development and testing of GM crops in the country [62]. In Pakistan, Agricultural Biotechnology Research Institute (ABRI) at Ayub Agricultural Research Institute (AARI), Faisalabad and SGS Laboratories Karachi have GMO testing labs, which have been accredited by Pakistan National Accreditation Council

(PNAC) for ISO-17025. These labs are efficiently working on the testing and identification of GM crops and are equipped with state-of-the-art facilities needed for the detection, identification and quantification of GMOs. All crop seeds being imported from other countries are first tested for the presence and type of GM elements from these laboratories and then allowed for cultivation in the country. All the import and export activities are strictly monitored with reference to GMOs.

## 8. Conclusion

Testing of GM crops is important issue for the legitimacy, biosafety and regulatory purposes. The area under GM crops is increasing very rapidly and many new genes are being introduced in major crops. For the safety of humans, environment, animals and other related micro-flora, a comprehensive molecular testing of newly developed GMO is very important before commercial release. Regulatory processes for GM crops approval need comprehensive risk analysis for each case separately. The detection and identification of GMOs is also of great value in identifying the purity of sample, labeling food and trade reasons. Therefore, combined use of more than one testing methods would be advantageous for complete analysis, authenticity and biosafety assessment of GM samples.

## Author details

Shahid Nazir\*, Muhammad Zaffar Iqbal and Sajid-ur-Rahman  
Agricultural Biotechnology Research Institute, Ayub Agricultural Research Institute (AARI), Faisalabad, Pakistan

\*Address all correspondence to: [shahidnazir97@yahoo.com](mailto:shahidnazir97@yahoo.com)

## IntechOpen

© 2019 The Author(s). Licensee IntechOpen. This chapter is distributed under the terms of the Creative Commons Attribution License (<http://creativecommons.org/licenses/by/3.0>), which permits unrestricted use, distribution, and reproduction in any medium, provided the original work is properly cited. 

## References

- [1] Harlander SK. The evolution of modern agriculture and its future with biotechnology. *Journal of the American College of Nutrition*. 2002;**21**:161S-165S. DOI: 10.1080/07315724.2002.10719260
- [2] Atherton KT. Safety assessment of genetically modified crops. *Toxicology*. 2002;**181**:421-426. DOI: 10.1016/S0300-483X(02)00485-7
- [3] Glass S, Jessica F. Genetic modification Technology for nutrition and improving diets: An ethical perspective. *Current Opinions in Biotechnology*. 2017;**44**:46-51. DOI: 10.1016/j.copbio.2016.11.005
- [4] Beyaz R, Selcen ED, Aycan M, Kayan M, Yıldız M. A novel method for high-frequency transgenic shoot regeneration via *Agrobacterium tumefaciens* in flax (*Linum usitatissimum* L.). *Journal of Plant Biotechnology*. 2016;**43**:240-247. DOI: 10.5010/JPB.2016.43.2.240
- [5] Ma JKC, Drake PMW, Christou P. Genetic modification: The production of recombinant pharmaceutical proteins in plants. *Nature Reviews Genetics*. 2003;**4**:794-805. DOI: 10.1038/nrg1177
- [6] Yaqoob A, Ahmad AS, Tahir RS, Rao AQ, Khan MAZ, Sana T, et al. Risk assessment of Bt crops on the non-target plant-associated insects and soil organisms. *Journal of the Science of Food and Agriculture*. 2016;**96**:2613-2619. DOI: 10.1002/jsfa.7661
- [7] Mesnage R, Clair E, Gress S, Then C, Székács A, Séralini GE. Cytotoxicity on human cells of Cry1ab and Cry1ac Bt insecticidal toxins alone or with a glyphosate-based herbicide. *Journal of Applied Toxicology*. 2013;**33**:695-699. DOI: 10.1002/jat.2712
- [8] Bawa AS, Anilakumar KR. Genetically modified foods: Safety, risks and public concerns—A review. *Journal of Food Science and Technology*. 2013;**50**:1035-1046. DOI: 10.1007/s13197-012-0899-1
- [9] Smyth SJ. Genetically modified crops, regulatory delays, and international trade. *Food and Energy Security*. 2017;**6**:78-86. DOI: 10.1002/fes3.100
- [10] Smart RD, Blum M, Wesseler J. Trends in approval times for genetically engineered crops in the United States and the European Union. *Journal of Agricultural Economics*. 2016;**68**:182-198. DOI: 10.1111/1477-9552.12171
- [11] Lu H, Katherine AM, John CB. Messages promoting genetic modification of crops in the context of climate change: Evidence for psychological reactance. *Appetite*. 2017;**108**:104-116. DOI: 10.1016/j.appet.2016.09.026
- [12] Baig S, Khan MZU, Nadeem Z, Ahmed S, Mushtaq AS. GM crops and their environmental risks: A mini overview. *Fuuast Journal of Biology*. 2017;**7**:193-198
- [13] Klümper W, Matin QA. Meta-analysis of the impacts of genetically modified crops. *PLoS One*. 2014;**9**:e111629. DOI: 10.1371/journal.pone.0111629
- [14] Miraglia M, Berdal K, Brera C, Corbisier P, Holst-jensen A, Kok E, et al. Detection and traceability of genetically modified organisms in the food production chain. *Food and Chemical Toxicology*. 2004;**42**:1157-1180. DOI: 10.1016/j.fct.2004.02.018
- [15] ISAAA. Global Status of Commercialized Biotech/GM Crops in 2017: Biotech Crop Adoption Surges as Economic Benefits Accumulate in 22

Years. Ithaca, NY: ISAAA; 2017. ISAAA Brief No. 53

[16] Gressel J, Aaron JG, Micheal ODK. How well will stacked transgenic Pest/herbicide resistances delay pests from evolving resistance? *Pest Management Science*. 2017;**73**:22-34. DOI: 10.1002/ps.4425

[17] Tripathi L. Techniques for detecting genetically modified crops and products. *African Journal of Biotechnology*. 2005;**4**:1472-1479. DOI: 10.4314/ajfand.v4i13.71830

[18] Sambrook J, Russell DW. *Molecular Cloning: A Laboratory Manual*. New York: Cold Spring Harbor Laboratory Press, Cold Spring Harbor; 2001

[19] Brown TA. *Gene Cloning and DNA Analysis: An Introduction*. 7th ed. Chichester, West Sussex ; Hoboken, NJ: John Wiley and Sons Ltd; 2016

[20] Rao AQ, Allah B, Sarfraz K, Kamran S, Ahmad AS, Husnain T, et al. The myth of plant transformation. *Biotechnology Advances*. 2009;**27**:753-763. DOI: 10.1016/j.biotechadv.2009.04.028

[21] Vijayakumar KR, Martin A, Gowda LR, Prakash V. Detection of genetically modified soya and maize: Impact of heat processing. *Food Chemistry*. 2009;**117**:514-521. DOI: 10.1016/j.foodchem.2009.04.028

[22] Saiki RK, Scharf S, Faloona F, Mullis KB, Horn GT, Erlich HA, et al. Enzymatic amplification of  $\beta$ -globin genomic sequences and restriction site analysis for diagnosis of sickle cell anemia. *Science*. 1985;**230**:1350-1354. DOI: 10.1126/science.2999980

[23] Mullis KB, Faloona FA. Specific synthesis of DNA in vitro via a polymerase-catalyzed reaction. *Methods in Enzymology*. 1987;**155**:335-350. DOI: 10.1016/0076-6879(87)55023-6

[24] Holst-Jensen A, Rønning SB, Løvseth A, Berdal KG. PCR technology for screening and quantification of genetically modified organisms (GMOs). *Analytical and Bioanalytical Chemistry*. 2003;**375**:985-993. DOI: 10.1007/s00216-003-1767-7

[25] Coghlan ML, Haile J, Houston J, Murray DC, White NE, Moolhuijzen P, et al. Deep sequencing of plant and animal DNA contained within traditional Chinese medicines reveals legality issues and health safety concerns. *PLoS Genetics*. 2012;**8**:e1002657. DOI: 10.1371/journal.pgen.1002657

[26] Newmaster SG, Meghan G, Dhivya S, Sathishkumar R, Subramanyam R. DNA barcoding detects contamination and substitution in north American herbal products. *BMC Medicine*. 2013;**11**:222. DOI: 10.1186/1741-7015-11-222

[27] Tillmar AO, Barbara DA, Jenny W, Gunilla H. A universal method for species identification of mammals utilizing next generation sequencing for the analysis of DNA mixtures. *PLoS One*. 2013;**8**:e83761. DOI: 10.1371/journal.pone.0083761

[28] Galimberti A, Fabrizio AM, Alessia L, Ilaria B, Silvia F, Maurizio C, et al. DNA barcoding as a new tool for food traceability. *Food Research International*. 2013;**50**:55-63. DOI: 10.1016/j.foodres.2012.09.036

[29] Ferri G, Corradini B, Ferrari F, Santunione AL, Palazzoli F, Alu M. Forensic botany II, DNA barcode for land plants: Which markers after the international agreement? *Forensic Science International: Genetics*. 2015;**15**:131-136. DOI: 10.1016/j.fsigen.2014.10.005

[30] Meyer R, Chardonnens F, Hübner P, Lüthy J. Polymerase chain reaction (PCR) in the quality and safety assurance of food: Detection of soya in



processed meat products. *Zeitschrift für Lebensmittel-Untersuchung und Forschung*. 1996;**203**:339-344. DOI: 10.1007/BF01231072

[31] Taberlet P, Gielly L, Pautou G, Bouvet J. Universal primers for amplification of three noncoding regions of chloroplast DNA. *Plant Molecular Biology*. 1991;**17**:1105-1109. DOI: 10.1007/BF00037152

[32] Gierson E, Tucker GA, Keen J, Ray J, Bird CR, Schuch W. Sequencing and identification of a cDNA clone for tomato polygalacturonase. *Nucleic Acid Research*. 1986;**14**:8595-8603. DOI: 10.1093/nar/14.21.8595

[33] Pietsch K, Waiblinger HU, Brodmann P, Wurz A. Screening methods for the identification of “genetically modified” food of plant origin. *Deutsche Lebensmittel Rundschau*. 1997;**93**:35-38

[34] Hemmer W. Foods derived from genetically modified organisms and detection methods. In: *BATS Report by Centre for Biosafety and Sustainability*. Basel, Switzerland; 1997. pp. 40-44

[35] Wolf C, Scherzinger M, Wurz A, Pauli U, Hübner P, Lüthy J. Detection of cauliflower mosaic virus by the polymerase chain reaction: Testing of food components for false-positive 35S-promoter screening results. *European Food Research and Technology*. 2000;**210**:367-372. DOI: 10.1007/s002170050565

[36] Deisingh AK, Badrie N. Detection approaches for genetically modified organisms in foods. *Food Research International*. 2005;**38**:639-649. DOI: 10.1016/j.foodres.2005.01.003

[37] Holst-Jensen A. Testing for genetically modified organisms (GMOs): Past, present and future perspectives. *Biotechnology Advances*.

2009;**27**:1071-1082. DOI: 10.1016/j.biotechadv.2009.05.025

[38] Southern EM. Detection of specific sequences among DNA fragments separated by gel electrophoresis. *Journal of Molecular Biology*. 1975;**98**:503-517. DOI: 10.1016/S0022-2836(75)80083-0

[39] Leimanis S, Hernández M, Fernández S, Boyer F, Burns M, Bruderer S, et al. A microarray-based detection system for genetically modified (GM) food ingredients. *Plant Molecular Biology*. 2006;**61**:123-139. DOI: 10.1007/s11103-005-6173-4

[40] Kim JH, Kim SY, Lee H, Kim YR, Kim HY. An event-specific DNA microarray to identify genetically modified organisms in processed foods. *Journal of Agriculture and Food Chemistry*. 2010;**58**:6018-6026. DOI: 10.1021/jf100351x

[41] Elenis DS, Kalogianni DP, Glynnou K, Ioannou PC, Christopoulos TK. Advances in molecular techniques for the detection and quantification of genetically modified organisms. *Analytical and Bioanalytical Chemistry*. 2008;**392**:347-354. DOI: 10.1007/s00216-008-1868-4

[42] Querci M, Bulcke MV, Jana Ž, Guy VE, Hermann B. New approaches in GMO detection. *Analytical and Bioanalytical Chemistry*. 2010;**396**:1991-2002. DOI: 10.1007/s00216-009-3237-3

[43] Von Gotz F. See what you eat--Broad GMO screening with microarrays. *Analytical and Bioanalytical Chemistry*. 2010;**396**:1961-1970. DOI: 10.1007/s00216-009-3204-z

[44] Heid CA, Stevens J, Livak KJ, Williams PM. Real time quantitative PCR. *Genome Research*. 1996;**6**:986-994. DOI: 10.1101/gr.6.10.986

[45] Higuchi R, Fockler C, Dollinger G, Watson R. Kinetic PCR analysis:

- Real time monitoring of DNA amplification reactions. *Biotechnology*. 1993;**11**:1026-1030. DOI: 10.1038/nbt0993-1026
- [46] Ferreira JL, Geraldo MAC, Aluizio B, Wellington SG, Tesfahun AS. Biosafety and detection of genetically modified organisms. In: Çiftçi YO, editor. *Transgenic Plants—Advances and Limitations*. Intechopen; 2012. pp. 427-448. DOI: 10.13140/RG.2.1.3470.2245
- [47] Brett GM, Chambers SJ, Huang L, Morgan MRA. Design and development of immunoassays for detection of proteins. *Food Control*. 1999;**10**:401-406. DOI: 10.1016/S0956-7135(99)00082-1
- [48] Conceição FR, Moreira AN, Binsfeld PC. Detecção de organismos geneticamente modificados em alimentos e ingredientes alimentares. *Ciência Rural*. 2004;**36**:315-324. DOI: 10.1590/S0103-84782006000100053
- [49] Urbanek-Karlowska B, Fonberg-Broczek M, Sawilska-Rautenstrauch D, Badowski P, Jedra M. Usefulness of an immunoassay test trait for detection of genetically modified roundup ready soybean in food products. *Rocz Panstwowy Zaklad Higieny*. 2001;**52**:313-320
- [50] Lipton CR, Dautlick JX, Grothaus GD, Hunst PL, Magin KM, Mihaliak CA, et al. Guidelines for the validation and use of immunoassays for determining of introduced proteins in biotechnology enhanced crops and derived food ingredients. *Food and Agriculture Immunology*. 2000;**12**:153-164. DOI: 10.1080/095401000404094
- [51] Kuiper HA. Summary report of the ILSI Europe workshop on detection methods for novel foods derived from genetically modified organisms. *Food Control*. 1999;**10**:339-349. DOI: 10.1016/S0956-7135(99)00072-9
- [52] Ahmed FE. Detection of genetically modified organisms in foods. *Trends in Biotechnology*. 2002;**20**:215-223. DOI: 10.1016/S0167-7799(01)01920-5
- [53] Serageldin I. Biotechnology and food security in the 21st century. *Science*. 1999;**285**:387-389. DOI: 10.1126/science.285.5426.387
- [54] Areal FJ, Riesgo L, Rodriguez-Cerezo E. Economic and agronomic impact of commercialized GM crops: A meta analysis. *The Journal of Agricultural Science*. 2013;**151**:7-33. DOI: 10.1017/S0021859612000111
- [55] Finger R, El Benni N, Kaphengst T, Evans C, Herbert S, Lehmann B, et al. A meta analysis on farm-level costs and benefits of GM crops. *Sustainability*. 2011;**3**:743-762. DOI: 10.3929/ethz-b-000042242
- [56] Brookes G, Barfoot P. Global income and production impacts of using GM crop technology. 1996-2014. *GM Crops and Food*. 2016;**7**:38-77. DOI: 10.1080/21645698.2016.1176817
- [57] Ives AR, Cate P, Andrew H, Sharon D, David AA, Ralph H, et al. Spatio-temporal variation in landscape composition may speed resistance evolution of pests to Bt crops. *PLoS One*. 2017;**12**:e0169167. DOI: 10.1371/journal.pone.0169167
- [58] Finamore A, Marianna R, Serena B, Giovanni M, Roberto A, Aida T, et al. Intestinal and peripheral immune response to Mon810 maize ingestion in weaning and old mice. *Journal of Agricultural and Food Chemistry*. 2008;**56**:11533-11539. DOI: 10.1021/jf802059w
- [59] Economic Survey of Pakistan 2017-18. Chapter-2 Agriculture, Economic Adviser's Wing, Finance Division, Government of Pakistan. pp. 13-32

[60] ISAAA. Global Status of Commercialized Biotech/GM Crops: 2016. ISAAA Brief No. 52. Ithaca, New York: ISAAA; 2016

[61] Gabol WA, Ahmed A, Bux H, Ahmed K, Mahar A, Laghari S. Genetically modified organisms (GMOs) in Pakistan. *African Journal of Biotechnology*. 2012;**11**:2807-2813. DOI: 10.5897/AJBX11.001

[62] Khan MS, Joyia FA. Biotechnology and GM crops. In: Khan IA, Khan MS, editors. *Developing Sustainable Agriculture in Pakistan*. Taylor and Francis, USA: CRC PressTaylor and Francis; 2018. pp. 355-368