REVIEW

Classification of CRISPR/Cas system and its application in tomato breeding

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Abstract
Remarkable diversity in the domain of genome loci architecture, structure of effector complex, array of protein composition, mechanisms of adaptation along with difference in pre-crRNA processing and interference have led to a vast scope of detailed classification in bacterial and archaeal CRISPR/Cas systems, their intrinsic weapon of adaptive immunity. Two classes: Class 1 and Class 2, several types and subtypes have been identified so far. While the evolution of the effector complexes of Class 2 is assigned solely to mobile genetic elements, the origin of Class 1 effector molecules is still in a haze. Majority of the types target DNA except type VI, which have been found to target RNA exclusively. Cas9, the single effector protein, has been the primary focus of CRISPR-mediated genome editing revolution and is an integral part of Class 2 (type II) system. The present review focuses on the different CRISPR types in depth and the application of CRISPR/Cas9 for epigenome modification, targeted base editing and improving traits such as abiotic and biotic stress tolerance, yield and nutritional aspects of tomato breeding.

Abbreviations

| Acronym | Definition |
|---------|------------|
| RNAi    | RNA interference |
| ZFN     | Zinc finger nuclease |
| TALEN   | Transcription activator-like effector nuclease |
| CRISPR  | Clustered regularly interspaced short palindromic repeats |
| GM      | Genetically modified |
| GMO     | Genetically modified organism |
| RNP     | Ribonucleoprotein |
| NHEJ    | Non-homologous end joining |
| PAM     | Protospacer adjacent motif |
| DBD     | DNA binding domains |
| DSB     | Double-stranded break |
| CRISPRi | CRISPR interference |
| crRNA   | CRISPR RNA |
| tracrRNA| Transactivating CRISPR RNA |
| gRNA    | Guide RNA |
| sgRNA   | Single guide RNA |
| MGE     | Mobile genetic element |
| RRM     | RNA recognition motif |
| RAMP    | Repeat-associated mysterious proteins |
| HEPN    | Higher eukaryotes and prokaryotes nucleotide-binding domain |
| PFS     | Protospacer Flanking Site |
| CP      | Coat protein |
| TYLCV   | Tomato yellow leaf curl virus |
| QTL     | Quantitative trait loci |

Introduction

Genetic diversity is a potential resource for a broad range of genetic research and trait improvement in plants. The gradual evolution in the plant breeding technologies and expansion of its possibilities is much required to cope up with the incessantly increasing needs of man (Xiong et al. 2015). With the aim of creating new varieties, breeders have developed novel methods to introduce heritable mutations into plant genomes. In the recent past, various mutagens like chemical compounds and irradiation were used to generate large pools of genetic variation in traditional breeding. Like all methods, these too have several drawbacks, such as the non-specific nature of the generated mutations, simultaneous
mutation of a large amount of nucleotides followed by deletion, duplication and rearrangement of lengthy genomic fragments (Hartwell et al. 2018), thus making the identification of the mutations a laborious process. Also the random mutagenesis methods usually prove to be less effective to improve traits in polyploid crops, because of their extreme genetic redundancy (Mao et al. 2019).

Although traditional breeding allowed the selection of unique crops with improved traits, enriched qualities and extended shelf life coupled with long breeding cycles yet lack of precision in hybridization, high ratio of heterozygosities along with low frequencies of the desirable mutation have led to the development of less/moderate resource-demanding technologies. Recombinant DNA technology has proved to be versatile since it can overcome the incompatibility issues between two species by the integration of foreign genes into the target plant genomes to get the desired traits (Genetically modified crops, GM crops). So far, GM crops constitute a significant proportion of the diet among the populations of America, Australia, China and other developing nations. However, several countries including India, oppose the use of GM crops for human consumption on the ground that it might pose risk to human health and environment. However, until now no long-term scientific study have been able to gather evidence that GM crops have profound adverse effect on human health and environment as compared to the existing plant breeding technologies (Council 2013). So this so-called debate seems to be far-fetched from the knowledge based upon solid scientific research and centered solely upon the moral traditions and political environment. Political hesitation regarding genetically modified organisms (GMOs) can easily be pointed out with the decision by seventeen European countries in 2015 to opt out of the possible commercialization of genetically modified food and fiber (Bonny 2003). The major concerns shown regarding GMOs is that they have been generated by introduction of a transgene into the host genome and the expression cassette (generally carrying viral promoter/terminator) often stayed there within the system (Li et al. 2019a). From that context, next-generation genome editing tools such as clustered regularly interspaced short palindromic repeats/Cas9 (CRISPR/Cas9) gain importance, as they are capable of introducing small insertion or deletion of nucleotides within the target gene itself, by using non-homologous end joining repair mechanism (NHEJ). Such genetic alteration does not involve the addition of any transgene and it is very much similar like the natural variation (Huang et al. 2016; Globus and Qimron 2018). Moreover, the expression cassette can be discarded from the progeny by genetic segregation or it is not at all required in case of ribonucleoprotein complex approach. Therefore, genome edited crops are very much different from GM crops in terms of their genetic properties and could be considered to have minimal/no risk to human life and its environment (Li et al. 2019a; Schulman et al. 2020). Spreading of more awareness and active cooperation will be required from both the scientific community and government agencies in order to gain public acceptance of the genome edited crops.

**CRISPR/CAS system**

In recent years, the RNA-programmable CRISPR/Cas9 technology of genome editing has caught the eyes of researchers and has traversed a long way in a very short period since it is an uncomplicated and effortless process. The field of biology (animal, plant and microbe) has undergone a massive transformation because of the immense potential of this powerful genome editing tool. CRISPR was first described by Japanese researchers in 1987 while working with iap genes in *Escherichia coli*, as a small stretch of highly homologous repeats interspaced with small spacers (Ishino et al. 1987). Years later, in 2005, three independent studies proved that these spacer sequences are acquired from external mobile genetic elements such as virus or plasmids (Bolotin et al. 2005; Poullet et al. 2005; Soria 2005). CRISPR was found out to be present in various other bacteria and archaea (Mojica et al. 2000) that serves primarily as bacterial adaptive defense mechanism (Makarova et al. 2006; Barrangou et al. 2007). The CRISPR loci consist of CRISPR repeats along with DNA-targeting spacers and a cas operon that contains all the Cas protein coding genes (Fig. 1a).

CRISPR/Cas-mediated adaptive immunity occurs in three different stages: i) new spacers from the invading organism are incorporated into the CRISPR array (Adaptation) (Barrangou et al. 2007), ii) precursor crRNAs (pre-crRNA) are transcribed from the CRISPR array which then undergoes maturation and becomes a set of CRISPR RNAs (crRNA) carrying a single targeted spacer flanked by repeat sequences. This maturation process is directed by trans-activating CRISPR RNA (tracrRNA) with the help of CRISPR-associated Csn1 protein and endogenous RNase III. These crRNAs along with Cas proteins get incorporated into ribonucleoprotein (RNP) complexes, which then start scanning for nucleic acid sequences that are complementary with the sequence coded by crRNA (Maturation) (Delcheva et al. 2011; Jinek et al. 2012; Zhang et al. 2012; Reeks et al. 2013). iii) Upon recognition, foreign sequences are cleaved by crRNA-guided Cas protein complex (Interference) (Semenova et al. 2011; Doudna and Charpentier 2014; Burmistrz et al. 2020) (Fig. 1b, c).

**How it is unique from other traditional gene editing tools**

Investigators have been using two complementary yet different approaches to elucidate the function of a particular gene. First is the forward genetics approach that depends on the observation of phenotypes, thereby trying
and linking it to its genetic basis; and another one is the reverse genetics approach, which creates insertion/deletion to induce mutation in the gene itself, thereby observing the change in the phenotype. RNA interference (RNAi) is the reverse genetics approach where investigator must have the prior knowledge of the gene sequence to be worked on. Irrespective of having been used extensively, it has been reported that RNAi often produce hypomorphic phenotype, which means there is a potential chance of getting a partial loss-of-function phenotype after targeting the gene of interest (Heigwer et al. 2018). To overcome this situation investigators need a more robust and unique reverse genetics approach and that is where zinc finger nuclease (ZFN) and transcription activator-like effector nuclease (TALEN) come in. As opposed to RNAi, these processes can achieve a complete loss-of-function phenotype by using customizable DNA-binding domains (DBDs) for recognition and targeting. These DBDs along with nucleases are capable of introducing double-stranded breaks (DSBs) into the targeted gene sequence causing frameshift mutation. The TALEN activity need not be maintained in the target cell, as these mutations are permanent in nature, whereas in RNAi mechanism, significant loss of siRNAs can dramatically reduce the loss-of-function phenotype.

On the other hand CRISPR/Cas9 system is the powerful genome editing tool that could be used both as a forward (Wang et al. 2014; Chen et al. 2015) and reverse genetics approach (Orack et al. 2015; Mianné et al. 2016). CRISPR interference (CRISPRi) has more profound knockdown effect (Gilbert et al. 2015) as opposed to RNAi. Probably the most striking difference between RNAi and Cas9 nuclease-mediated editing is that RNAi can induce transient knockdown of gene expression, whereas Cas9 nuclease can induce permanent damage in target DNA (Boettcher and McManus 2015). Compared to TALEN and CRISPR/Cas9, ZFNs are expensive to make, and considerable labor is required to construct these high-affinity DBDs. TALENs surpass this bottleneck since they need considerably fewer enhancements post-construction, but they do depend on the lengthy assembly process of the minute building blocks to generate artificial DNA-binding proteins. However, the lengthy process can be surpassed with the most recent CRISPR/Cas9 tool since it caters to a technically uncomplicated system of genome editing (Lozano-Juste and Cutler 2014). More than one gene editing technology has been used to target the same gene (Table 1) and CRISPR/Cas-mediated gene editing showed better results.
PAM-free CRISPR/Cas9 system

With the gradual evolution and advancement of the CRISPR/Cas technology, the modern researchers started manipulating this system according to the need of smooth working and eventually the possibility for relaxed protospacer adjacent motif (PAM) requirements emerged. The field has advanced by taking long strides targeting a purely PAM-free nuclease territory. PAM-free nucleases have been constructed through the strategies like natural ortholog mining and protein engineering (Colllias and Beisel 2021). Genetic engineering of Cas9 from the human pathogen *Streptococcus pyogenes* (SpyCas9) has relaxed its PAM profile considerably to one of two bases at a particular position. Numerous other Cas9 nucleases along with modified Cas12a nucleases are in the pipeline of being engineered having distinct properties like smaller size, high thermostability, etc.

The major advantage of a PAM-free nuclease is the ability to target any sequence (Fig. 2a). The selection of sites would be simplified and flexible with preferably high on-target and low off-target activity, thus producing disruptive insertions and deletions, which are highly predictable (Chakrabarti et al. 2019). The placement of the base-editing window can also be arranged over the specific nucleotide directly. These flexible features act to be truly beneficial during multiplex editing since only one nuclease comes into full action and targets all the sequences. There are serious derogatory results as well of the PAM-free system (Fig. 2c).

The guide RNAs (gRNAs) that are expressed from the DNA constructs would self-target the parent DNA immediately, ultimately leading to disastrous consequences. Another drawback of the PAM-free system is that a nuclease with no specific PAM will touch down on every sequence of that particular genome, thus taking a much longer time than required to finally find out its actual target and there would be increased risk of off-targeting (Fig. 2b). Apart from these potential drawbacks, this domain of PAM-free CRISPR/Cas system of genome editing is being taken seriously and constantly worked upon. The SpyCas9 group of nucleases have been engineered and made almost PAM-free, and the other remaining Cas nucleases like Cas9, Cas12a have much scope of relaxing their PAM recognition status and gradually move toward attaining a PAM-free status. To expedite the process of development for these above-mentioned nucleases, a combination of the procedures of ortholog mining and PAM engineering might prove to be highly effective and fruitful as already observed in the case of generating *Streptococcus canis* Cas9-Sc + + and another high-fidelity

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Fig. 2 Development toward a PAM-free CRISPR/Cas system. a A comparative account depicting target accessibility by Cas nucleases having relaxed or stringent PAM necessity. (N, any base; C, cytosine; T, thymine; A, adenine). b A comparative account between a repertoire of nucleases and PAM-free nuclease. The former recognize every possible sequence together. The repertoire is a collection of four nucleases that read one letter at position two. (N, any nucleotide; A, adenine; G, guanine; T, thymine; C, cytosine). c A Comparative qualitative account between nuclease repertoire, PAM-free nuclease, PAM-relaxed nuclease and a PAM-stringent nuclease. The comparison is on the based on the capacity of targeting performance under different conditions. Greater performance of the associated nuclease is represented by a more filled bar. Nuclease fidelity is studied on the capacity of the nuclease to bypass nontarget sequence having even a minimal match with the guide sequence. Multiplex ability is assessed on the capacity of the nuclease to be put to use in targeting any random set of sequences.

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Working on the not so well-characterized CRISPR/Cas types like type I and V could lead to major breakthrough by engineering type V–C nucleases, which have already been shown to recognize PAMs with a minimal of single base (Yan et al. 2019). An effective alternative to the entire issue of PAM-free nucleases in CRISPR/Cas system of genome editing is a nuclease repertoire (Fig. 5b), where the nuclease retains its property of recognition of a specific PAM sequence, that might be of a single base (e.g., NG) or a collection of bases (e.g., NAAA). This nuclease repertoire could be customized accommodating all the possible sequences, which would ultimately confer an overall PAM-free status to it. Since each nuclease retains its PAM recognition ability, it can bypass the shortfalls of the PAM-free system. Here the nucleases can be employed based on the required target on the basis of the flanking sequences. Hence, generation of a nuclease repertoire can be a potential solution by which any and every sequence can be targeted by CRISPR/Cas system.

### An efficient CRISPR toolkit for tomato breeding: Golden Gate

It has been proved now and again that CRISPR/Cas9 system of genome editing is the superpower in the domain of plant genome editing. The use of multi-single gRNAs (sgRNAs) to target single or multiple genes at a time has enabled in improving its gene-editing efficacy by leaps and bounds. An extremely handy and conveniently generated tool kit, which is used extensively in tomato breeding, is the Golden Gate Tool Kit. Englert et al. 2008, devised the Golden Gate cloning strategy. This strategy utilized the type IIS set of restriction enzymes that perform cuts far from their recognition sites. With neat and skillful design of cleavage sites, the two pieces cleaved by the above restriction enzymes can
be joined together resulting in an end product minus the original restriction site. Thus, the above strategy (restriction–ligation) is instrumental in producing in one-pot and one-step highly pure and correct recombinant plasmids in minimal time (Engler et al. 2008). In a nutshell, this strategy allows fast and flexible assembly of genetic constructs and combination of diverse functional modules, respectively, for various applications (Čermák et al. 2017).

Tomato has proved to be a perfect example of a model plant where CRISPR/Cas system has been an efficient tool in creating new varieties without compromising the plant genome with foreign genes. Hu et al. 2019 have devised a highly innovative and flexible modular system in the domain of plant genome engineering for functional genomics in tomato and other potential food and cash crops. This research group applied standardized BioBrick technology to make this system more flexible for upgradation and accommodating toward novel expression elements. BioBrick technology had been described as a synthetic biology technology that works on the basis of same-tail restriction enzyme where BioBrick modules are assembled without break, keeping the BioBrick sites intact. Their (Hu et al. 2019) research approach was to develop a Cas9 system comprising of two binary vectors pHNCas9 and pHNCas9HT (Fig. 3a, b). pHNCas9HT has the capacity to construct sgRNA expression cassettes bypassing the pivotal processes of PCR and direct Agrobacterium tumefaciens transformation. Golden Gate ligation strategy comes into the picture over here,
where pHNCas9 vector is used for the assembly of myriad sgRNA expression cassettes in one-pot. The type II restriction endonucleases used here are Esp3 I and Bsa I. They produce non-palindromic sticky end sites, which totally cuts off all possibilities of self-ligation and incompatible end ligation. So, this is a perfect method to link numerous DNA fragments in one-step and one-tube which can be simultaneously used for single site and multi-site editing. Apart from the two binary vectors the system provides with 5 separate Pol III-dependent promoters, sgRNA in pEASY cloning vector, 8 pairs of Bsa I-site primer along with a gene specific primer design aid. All the components together can make a perfect CRISPR tool kit (Fig. 3c–e).

In the experimental setup, a visual T-DNA marker was designed by utilizing a BioBrick. This T-DNA marker design used the overexpression of AtMYB75/PAP1 (having 35S promoter) as a template. The genome editing systems possessing visual markers makes screening of transformants smooth and hassle-free since a different plant color is the indicator of whether the T-DNA marker with CRISPR/Cas along with the visual BioBrick insert has passed on to the T0 progeny. According to Mendel’s law of segregation, heterozygous dominant transgenic lines and homozygous recessive lines (with no T-DNA and AtMYB75) can be easily screened based on the presence and absence, respectively, of purple color. To test the efficiency of this tool kit in tomato, Hu et al. 2019 targeted just 1 site in SIEIN2, SIERFE1 and SIARF2B gene, 2 sites in SIACS2 and SIACS4 genes and 3 sites in SIGRAS8 gene. This CRISPR tool kit can easily carry out single, multi-site editing along with multi-gene editing in tomato.

**Classification of CRISPR/Cas system**

In archaee and bacteria, the CRISPR/Cas systems render adaptive immunity against foreign nucleic acids. They have a plethora of variable features in various categories like protein composition, pre-crRNA processing and interference, genome locus architecture and mechanisms of adaptation, effector complex structure, etc. The CRISPR/Cas system has been broadly divided into two classes: Class 1 and Class 2, the former being more complex than the latter. Class 1 possesses multi-subunit effector complexes and Class 2 with single protein effector modules. Further experimentation and analysis of the Class 2 CRISPR/Cas system led to the discovery of two new types and multiple subtypes. Of the two newly discovered and characterized CRISPR type, the one that solely targeted RNA was the type VI systems. The class 2 systems in some cases display a unique feature where the effector protein is also found responsible in the processing of the pre-crRNA (Koonin et al. 2017).

The primary reason for the variability and fast evolution of the CRISPR/Cas systems is its constant battle with the viruses, which conditions the cas genes to evolve fast (Takeuchi et al. 2012), thus leading to a diverse array of gene repertoires and finally to the whole defense infrastructure (Makarova et al. 2011, 2015). More specifically we can conclude that the CRISPR/Cas system has diversified as a response to the competitive coevolution of the anti-CRISPR proteins (Bondy-Denomy et al. 2013, 2015; Pawluk et al. 2016a, b). Comparative sequence analysis, experimental data and structural studies strongly infer that despite being evolutionarily flexible, all CRISPR/Cas variants exhibit common architectural and functional principles and the principle building blocks too exhibit a common ancestry (Makarova et al. 2013). In 2015, a blueprint was proposed which combined the signature genes and the elements of the Cas loci. This assigned almost all the CRISPR/Cas loci identified till date, to certain specified subtypes. This updated classification system can be adopted to classify the new varieties of CRISPR/Cas variants discovered from new genomes (Koonin et al. 2017).

**Classes (class 1 and class 2), types and subtypes**

It is pretty complicated to classify CRISPR/Cas systems since there is a lack of universal Cas proteins that could have acted as phylogenetic markers. Eventually the classification is based upon multiple factors like signature cas genes, layout of the cas operons and phylogenies of the conserved Cas proteins (Koonin and Makarova 2019).

The two major classes of the CRISPR/Cas system, namely 1 and 2, have a strong basis of differentiation. The Class 1 group comprises of the multi-subunit crRNA effector complex, whereas the Class 2 group is a collection of the single crRNA effector complex. Diverging further the Class 1 CRISPR/Cas system has been divided into type I, III and IV that is classified into further subtypes. Similarly Class 2 has been divided into types II, V and VI, also subdivided into several subtypes. This classification is based on the mechanism of action of the CRISPR/Cas system, which has been broadly divided into three stages: adaptation, expression and maturation, interference (together form the effector module). The adaptation module performs the spacer acquisition, whereas the effector module performs the pre-crRNA processing, target recognition and cleavage. Both the ‘expression and maturation’ stage and the ‘interference’ stage in type I and type III systems are carried out by a multi-subunit protein complex also known as the Cascade (CRISPR-associated complex for antiviral defense) complex together with the Cas3 nuclease–helicase, the Csm or the Cmr complex for type I, III-A and III-B CRISPR/Cas systems respectively.
Fig. 4 Classification of Class-1 CRISPR/Cas system based on the organization of the CRISPR/Cas loci, domain architectures of the effector proteins and the target (predicted) nucleic acids along with their PAM/PFS sequences; SS—small subunit. The colored arrows represent the corresponding genes and the shaded area represent the Class 1 effector complex. The diversified and common type I, type III are included in the Class I systems found in archaea and less frequently in bacteria. The rare type IV possesses rudimentary CRISPR–Cas loci without the adaptation module. The type I and type III CRISPR–Cas effector complexes possess well-defined architecture, their backbone made of Cas 7 and Cas5, which are paralogous RAMPs (Repeat-Associated Mysterious Proteins, made up of RRM (RNA Recognition Motif) fold with ‘large’ and ‘small’ subunits additionally. Cas6, loosely bound with the effector complex, is an additional RAMP and acts as the repeat-specific RNase in the pre-crRNA processing.

| PAM/PFS                      | 5'-CCN-3'            |
|------------------------------|----------------------|
| 5'-TTC-3', 5'-ACT-3', 5'-TAA-3', 5'-TAG-3', 5'-CAC-3' | NA                  |
| 5'-NTTC-3'                  | NA                   |
| 5'-AAG-3'                   | NA                   |

Two consecutive GC base pairs

NA

NR

NR

NR

NR

NR

NR

NR

NR

NR

NA

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Fig. 5 Classification of Class-2 CRISPR/Cas system based on the organization of the CRISPR/Cas loci, domain architectures of the effector proteins and the target (predicted) nucleic acids along with their PAM/PFS sequences; TM-predicted transmembrane segment; HEPN—higher eukaryotes and prokaryotes nucleotide binding. The colored arrows represent the corresponding genes of Class 2 system. The Class 2 effector modules are made up of large, single multi-domain protein making it simpler and better organized than their Class I counterparts.
In type II, III, V and VI systems, both the expression and maturation stage are performed by a single large polypeptide like the Cas9 in case of type II and Cas10 for type III, whereas for type V systems the process is executed by Cpf1 or other related proteins (Amitai and Sorek 2016). Delving deep within the Class 1 CRISPR/Cas system, type I and type III is predominant in various permutations and combinations in phylogenetically diverse archaea and bacteria (compa- ratively less frequent), whereas type IV system is seen to be the rare type with a rudimentary CRISPR/Cas loci and does not possess the adaptation module. On the other hand, the distinct type II system of Class 2 is confined to bacteria only (Koonin et al. 2017; Munawar and Ahmad 2021). An interesting fact about CRISPR/Cas systems is that they have been found in viral genomes and plasmids, which suggest horizontal gene transfer (Makarova et al. 2006). Though the Cas proteins are vastly diverse among themselves, all can be grouped into four functional categories: nucleases/recombinases performing spacer acquisition, ribonucleases catalyzing the processing of crRNA guides, a plethora of proteins that bind with the RNA guides resulting in the formation of the crRNP complexes to carry out target surveillance and finally the nucleases that degrade the DNA or RNA targets (Van Der Oost et al. 2014).

Studies have suggested that type I and type III complexes possess some key similarities in their architecture, which points out to the chance of them having a common ancestry. They have a long backbone made of repeat-associated mysterious proteins (RAMPs). These proteins are made up of two parts: RNA recognition motif (RRM) fold plus large and small subunits. The large subunit located at the base of the backbone is Cas8 in case of most type I systems and Cas10 for type III system (Reeks et al. 2013). The backbone consists of multiple copies of Cas7 and a smaller Cas5 like protein. Cas5 couples with 5′ end of the crRNA to interact with the large subunit of both Cas8 (type I) and Cas 10 (type III). The gRNA is housed in the effector complexes that are made up of multiple Cas7 and one Cas5 subunit. It has been reported that the Cas5 subunit binds with the 5′-crRNA and far. The long pre-crRNA that results from the transcription of the CRISPR array is eventually processed by a cas6 type endoribonuclease, in type I and type III systems into separate crRNAs (Brouns et al. 2008; Carte et al. 2008). On the other side, for crRNA maturation, Cas9 aids the type II systems, along with host RNase III and a tracrRNA (Carte et al. 2014). During the interference stage of type I systems, the Cas proteins couple with the mature crRNAs forming a ribonucleoprotein complex, to degrade the foreign nucleic acid by Cas3 nuclease (Westra et al. 2012), whereas in type II and type III-B systems, this process is carried out by intrinsic nuclease activity of their crRNP complexes (Gasiunas et al. 2012; Jinek et al. 2012). It is interesting to note that the type I and II complexes target DNA while the type III-B complex targets RNA (Staals et al. 2014).

Fig. 6 Schematic representation of diverse range of applications of CRISPR/Cas9 technology in tomato breeding

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links up to the large subunit of Cas8 in type I and Cas10 for type III. Both type I and type III takes the help of a standalone Cas6 endonuclease in their crRNA processing pathways. The solo Cas6 endonuclease binds to the amorphous pre-crRNA cutting inside each repeat, thus generating intermediate crRNA possessing 5′-3′ repeat derived termini (Charpentier et al. 2015). A comparative study by Khan et al. 2019 (Khan 2019) gives us a clear idea that Cas5/Cas6 is a mandatory element in preprocessing of crRNA, Cas3 and Cascade for further cleavage and crRNA for interference, in case of type I CRISPR/Cas systems. Figure 4 gives a vivid detail of the distinct PAMs required by different subunits of the type I, III, IV systems to carry out target acquisition and recognition. A 5′-CNN-3′ PAM motif is required by type I-A systems (Gudbergsdottir et al. 2011) for interference. Six distinct PAMs are recognized by holoarchaea having type I-B systems. A Cas5-dependent crRNA maturation pathway identified and characterized in type I-C system recognizes

| Tool   | Gene       | Tomato cultivar | Effect                                                                 | References                          |
|--------|------------|-----------------|----------------------------------------------------------------------|-------------------------------------|
| CRISPR | IAA9       | Micro-Tom, Ailsa Craig | Regenerated mutants exhibited morphological changes in leaf shape and seedless fruit—a characteristic of parthenocarpic tomato | Ueta et al. (2017b)                  |
| CRISPR | PMR4       | Moneymaker      | Mutants showed increased resistance against powdery mildew          | Santillán Martínez et al. (2020)    |
| CRISPR | MAPK3      | Ailsa Craig     | Susceptibility to gray mold disease                                | Zhang et al. (2018)                 |
| CRISPR | DCL2b      | Ailsa Craig     | Susceptibility to Tomato Mosaic virus (ToMV)                        | Wang et al. (2018)                  |
| CRISPR | JAZ2       | Moneymaker      | Resistance to banana streak virus                                  | Ortigosa et al. (2019)              |
| CRISPR | Homolog of Arabidopsis SP, CLV3 | M82 | Modified inflorescence and plant architectures, Increase in locule number and fruit weight | Rodriguez-Leal et al. (2017)         |
| CRISPR | SP, WUS    | S. pimpinellifolium | Growth habit determination, enlarge fruit size                  | Li et al. (2018b)                   |
| CRISPR | RIN        | Ailsa craig     | Fruit ripening                                                      | Ito et al. (2015, 2017, 2020)       |
| CRISPR | ORRM4      | Micro-Tom       | Fruit ripening                                                      | Yang et al. (2017)                  |
| CRISPR | AGL6       | Line MP-1 (TYLCV tolerant) | Parthenocarpy                                                | Klap et al. (2017)                   |
| CRISPR | PSY1       | M82, Yellow flesh e3756, Bicolor e393, S. pimpinellifolium| Yellow colored fruits                                              | Hayut et al. (2017)                 |
| CRISPR | AGO7       | M82             | Change in leaf shape                                                | Brooks et al. (2014)                |
| CRISPR | BOP1, BOP2, BOP3, TFAM1, TFAM2 | M82 | Loss of floral organ abscission, defects in fruit shape, Altered leaf complexity | Xu et al. (2016)                    |
| CRISPR | DELLA      | Micro-Tom       | Reduction in serrated leaflets                                     | (Shimatani et al. 2017)             |
| CRISPR | PDS        | Micro-Tom       | Albino phenotype                                                    | Pan et al. (2016)                   |
| CRISPR | SHR        | Solanum spp.    | Short root                                                          | Ron et al. (2014)                   |

IAA9, Aux/IAA9 transcription factor; PMR4, Powdery Mildew Resistance 4; SIMAPK3, Solanum lycopersicum MAP kinase3; DCL2b, Dicer-like 2b; JAZ2, JASMONATE ZIM DOMAIN2; SP, SELF-PRUNING; CLV3, CLAVATA3; WUS, WUSCHEL; RIN, RIPENING INHIBITOR; ORRM4, organelle RNA recognition motif-containing protein4; AGL6, AGAMOUS-LIKE 6; PSY1, PHYTOENE SYNTHASE 1; AGO7, ARGONAUTE 7; BOP1/BOP2/BOP3, BLADE-ON-PETIOLE family; TFAM1/TFAM2, mitochondrial transcription factor A; DELLA, aspartic acid–glutamic acid–leucine–leucine–alanine; PDS, Phytoene desaturase; SHR, SHORT ROOT
an ‘NTTC’ consensus PAM sequence as found in *Bacillus halodurans* (Hyun et al. 2012; Sorek et al. 2013). Type I-E effector complex isolated from *Thermobifida fusca* and *E. coli* (Xiao et al. 2017) binds to a 5′-AAG-3′ PAM sequence. It has been found that type I-E effector complex consists of five Cas proteins, whereas type I-F (found in *Pseudomonas aeruginosa*) consists only four and type I-F targets foreign DNA with the aid of a PAM sequence having two G-C base pairs consecutively (Rollins et al. 2015).

Based on different adaptation, interference and recognition schemes, the type III system has been further divided into four significant subtypes. The type III-A subtype possesses the adaptation-related genes, whereas these are absent in type III-B, C and D systems. Hence, the later three systems incorporate new spacers with total reliance on other systems. Type III shares similarities with type I in pre-crRNA processing strategies and in the structural setup of the crRNPs complexes also called the Csm/Cmr complex (Rath et al. 2015). The Csm complex is present in both A and D subtypes while Cmr complex is found in both B and C subtypes. The Cas9 protein plays an integral part in pre-crRNA processing in type III just like type I systems.

The Class 2 CRISPR/Cas system possesses one large multi-domain protein as the effector complex; hence, it is inferred to be more neatly organized compared to the Class 1 system. The Class 2 system has been further subdivided into three subtypes: type II, type V and type VI (Fig. 5). The well-studied and characterized system in this Class is the type II system possessing the effector Cas9 endonuclease, utilized extensively in genome editing. The entire process of targeting DNA by recruiting Cas9 is orchestrated by crRNA. Yet the trajectory followed by tracrRNA, RNase III and various other factors to carry out the 5′ end processing of crRNA still remains an enigma (Munawar and Ahmad 2021). The most significant feature of type V CRISPR/Cas system is the presence of a single effector protein called Cas12, which has five reported subtypes A to E and a tentative subtype U (Yang et al. 2016). Type V-A CRISPR/Cas system possesses an active endonuclease called the Cas12a (previously known as Cpf1) which can carry out targeted cleavage without the aid of an extra tracrRNA (Zetsche et al. 2015). The ‘TTN’ PAM sequence is the signature sequence for target recognition in case of subtype V-A (Gleditzsch et al. 2019). AT-rich PAM sequence like ‘TTT/TTA/TTC’ is utilized by the type V-B systems, in the line of action to target dsDNA, with the assistance of both tracrRNAs and crRNAs. CasY (Cas12d) and CasX (Cas12e) are the respective characteristic proteins in subtype V-D and V-E and over here CasX needs tracrRNA during interference while CasY needs none. It has also been reported that CasY utilizes a signature ‘TA’ PAM sequence while CasX requires a 5′-TTCN-3′ PAM sequence for target recognition (Burrstein et al. 2017). Cas12 effector protein is found to be extremely advantageous in use because of its properties like comparatively smaller size, asymmetric cleavage sites and no requirement of tracrRNA. Hence, Cas12 is a much desired effector protein and experimentation is underway on ways and methods to broaden its horizon of targets by generating variants having different PAM specificities (Gao et al. 2017).

Higher eukaryotes and prokaryotes nucleotide-binding (HEPN) domains are a signature feature in the subtypes of type VI CRISPR/Cas system and it is predicted that they possess RNase activity only. Cas13a (C2c2) was the first protein to be characterized over here. The RNA-targeting activity was well demonstrated for *Leptotrichia shahii* Cas13a (LshCas13a). The presence of Protospacer Flanking Site (PFS) analogous to PAMs for RNA targets was an integral part for carrying out this interference activity (Gleditzsch et al. 2019). A significant difference in proteins among the different subtypes in the type VI system has been noticed. The gradual evolution of type VI-B proteins has been considered to take place from transmembrane proteins since corresponding transmembrane domains have been observed, which makes this subtype unique from type VI proteins (Shmakov et al. 2017; Smargon et al. 2017). Cas13b effector has been identified for *Bergeyella zoohelcum* (BzCas13b) and PFS identification also reported. In a nutshell, it has been inferred that type VI systems are comparatively less stringent than other types in the domain of substrate recognition as it targets RNA solely which has lesser harmful side effects on the cell.

The evolution and diversity observed in the CRISPR/Cas system, a potential arsenal of prokaryotic defense mechanisms against MGE, is the result of the novel effector proteins and novel molecular strategies. The diversity is pronouncedly vivid in the CRISPR/Cas systems of archaea and bacteria. These findings have helped and will help the scientific community for further in vivo and in vitro analyses to fully decipher the mechanisms and strategies on how the intriguing CRISPR/Cas variants function to protect the host cell from lethal MGE invasions.

### Application of CRISPR/Cas9 system

#### CRISPR/Cas9 system for resistant breeding and quality improvement in tomato

CRISPR/Cas9 has gained much attention in the last decade because of its ease of use and efficiency. It has been used in fleshy fruit model plant “tomato” to enhance several aspects
such as yield, nutritional value and tolerance against stress conditions (Fig. 6, Table 2).

**Resistance against abiotic stresses**

CRISPR/Cas9 proved to be a valuable tool for identifying previously unknown abiotic stress regulators in tomato plant. The plant hormone brassinosteroid has been known to be associated with various developmental and physiological processes such as cell division, cell elongation, reproduction and seed germination (Divi and Krishna 2009). Yin and colleagues showed that **BRASSINAZOLE-RESISTANT 1 (BZR1)** is a critical component of brassinosteroid signaling in tomato and it regulates **RESPIRATORY BURST OXIDASE HOMOLOG1 (RBOH1)** which in turn controls the apoplastic hydrogen peroxide (H$_2$O$_2$) production and heat stress tolerance (Yin et al. 2018). CRISPR/Cas9-mediated knockout of **BZR1** yields mutant lines with impaired **RBOH1** induction, reduced growth and heat tolerance. In another study, researchers were able to identify the potential role of tomato **NONEXPRESSOR OF PATHOGENESIS-RELATED GENE 1 (SlNPR1)** in drought stress responses which was previously thought to be involved only in plant’s defense response against pathogens (Li et al. 2019b). CRISPR-npr1 mutants showed an increased stomatal aperture, higher electrolytic leakage and a reduction in drought tolerance compared to wild-type plants. Wang and her team targeted **MITOGEN-ACTIVATED PROTEIN KINASES (SIMAPK3)** in tomato, which revealed its role in drought stress-related responses by transcriptional modulation of other stress-related genes and by protecting cell membranes from oxidative damage (Wang et al. 2017a). CRISPR/Cas9-mediated targeted mutagenesis of **LATERAL ORGAN BOUNDARIES DOMAIN (SlLBD40)** resulted in improved drought tolerance in tomato (Liu et al. 2020). Transcriptional activator **C-REPEAT BINDING FACTOR 1 (CBF1)** has long been associated with cold stress-related regulation of gene expressions (Kanaya et al. 1999; Gilmour et al. 2004), but the detailed mechanism is still not clear in tomato. CRISPR/Cas9-mediated knockout of tomato **SlCBF1** yields mutant lines with severe chilling injuries, higher electrolyte leakage and malondialdehyde levels compared to wild-type plants, contributing to a vivid insight into the molecular mechanism of **SlCBF1**-mediated tomato chilling sensitivity (Rui et al. 2018).

![Fig. 7 dCas9-mediated methylation/demethylation process.](image)

**a** dCas9-DNMT3A-mediated methylation of dsDNA resulting in decreased level of gene expression. **b** dCas9-TET1-mediated demethylation of dsDNA resulting in increased level of gene expression.
Resistance against biotic stresses

Pathogenic microbes such as bacteria, fungus or viruses cause severe damage in crop production. Disease-resistant smart crops can be the only sustainable way to cope with the massive demand in food supply worldwide. Enormous research has been done over the past 25 years to identify the key genes conferring disease resistance in crops. CRISPR/Cas9-mediated inactivation of tomato DOWNY MILDEW RESISTANCE 6 (SIDMR6-1) produced prematurely truncated protein conferring disease resistance against a wide variety of pathogens such as P. syringae, P. capsici and Xanthomonas spp (Zeilmaker et al. 2015; Paula de Toledo Thomazella et al. 2016). In another study, researchers created an improved variety of tomato that can resist powdery mildew fungal pathogen Oidium neolycopersici. They took a double guide RNA approach, targeting two regions within the MILDEW-RESISTANT LOCUS O (Mlo) in tomato for the required loss-of-function mutation (Nekrasov et al. 2017). One of the most devastating diseases in tomato is fusarium wilt disease caused by Fusarium oxysporum f. sp. lycopersici (Sacc.) causing huge losses in tomato production all across the globe. Complementation and knockout strategies using CRISPR/Cas9 revealed a novel tomato gene Solyco08g075770 as the primary reason behind tolerance to fusarium wilt (Prihatna et al. 2018). Another group of researchers demonstrated the potential of CRISPR/CAS9 system to target the coat protein (CP) sequence of tomato yellow leaf curl virus (TYLCV) genome in tomato and induce stable and efficient virus interference that remained active across multiple generations (Tashkandi et al. 2018). Methyl jasmonate plays a crucial role in the developmental processes and plant's defense response against various pathogens such as Botrytis cinerea. Transcription factor SIMY2C is the master regulator of methyl jasmonate signaling pathway (Kazan and Manners 2013). Knockout of SIMY2C significantly reduced the expression level of both disease defensive genes and genes related to jasmonic acid pathway, suggesting its prominent role in methyl jasmonate-induced disease resistance in tomato (Shu et al. 2020).

Improvement of yield and nutritional quality

Flowering in plants majorly depends upon the day length period, which varies from one season to another. This day length sensitivity limits the geographical range of crop cultivation and yield. CRISPR/Cas9-mediated mutagenesis in SELF-PRUNING 5G (SP5G) produced the loss of day-length-sensitive tomato lines with rapid flowering and enhanced yield, illustrating the power of this genome editing tool in crop improvement (Soyk et al. 2017). Long shelf life is inarguably the most elite characteristic to tomato breeders for storage and post-harvest produce distribution. Flowering in plants majorly depends upon the day length period, which varies from one season to another (Yu et al. 2017). Parthenocarpy is an industrially important trait in horticultural plants, and the key gene responsible for it is Aux/IAA transcription factor SI/AA9 (Wang et al. 2005). CRISPR/Cas9-mediated targeted mutagenesis in SI/AA9 produced parthenocarpic tomato lines and the trait is segregated into the next generation successfully (Ueta et al. 2017a). Wild crop varieties have been domesticated for decades to cope with the growing population and demand in the food supply, which leads to loss of genetic diversity and resistance against biotic and abiotic stresses. Researchers were able to domesticate a wild variety of tomato (Solanum pimpinellifolium) with improved size, number and nutritional value by targeting six important loci (SELF-PRUNING, OVATE, FACIATED, FRUIT WEIGHT, MULTIFLORA, LYCOPENE BETA CYCLASE) using CRISPR/Cas9 (Zsögön et al. 2018). GABA (γ-aminobutyric acid) homeostasis is crucial for plants’ developmental processes and is regulated via a GABA shunt (Takayama and Ezura 2015). Five key genes regulating this shunt were targeted using a multiplex pYLCRISPR/Cas9 system yielding mutant tomato lines with a significant increase in GABA accumulation in both leaves and fruits (Li et al. 2018a).

CRISPR/Cas9 system for targeting polyploid crops

CRISPR/Cas9 is the most widely utilized genome editing tool in plants, but its editing efficiency has varied widely and dramatically. The editing in polyploid crops is especially challenging because the paralogs and orthologs present there, having functional redundancy, require a cumbersome simultaneous knockout of all copies of genes, which are functionally similar. Two crucial factors that directly affect the mutagenic frequency of polyploid crops are: optimization of Cas9 codon, promoters and target sequence composition (Zhang et al. 2014; Ma et al. 2015; 2016; Yan et al. 2015; Mao et al. 2016). The most difficult task in this procedure is designing the sgRNAs, which is extremely challenging in case of polyploids than in diploids. Though a couple of sgRNA designing tools are prevalent in the market (CRISPR-P, CRISPR-P2.0), they cannot be used smoothly in polyploidy crops (Lei et al. 2014; Liu et al. 2017). The strategy best adopted for the simultaneous knockout of orthologs and paralogs would be to design sgRNAs picked up from a conserved domain, which targets the entire collection of gene copies. Manual designing of sgRNAs targeting one particular gene copy or all copies can be done after performing detailed sequence analysis. It happens sometimes that a large cross section of homologous genes do not possess a conserved site, those genes are divided into several groups and sgRNAs are designed on the basis of the conserved
site of each group (Zaman et al. 2019). Apart from targeted mutagenesis, another noteworthy application of CRISPR/Cas-mediated genome editing for improvement of polyploid crops is the targeted substitution of unwanted alleles with the desired ones (Schaart et al. 2021).

CRISPR-Cpf1, a new class of CRISPR system, is quite analogous to Cas9 and enables the editing of AT-rich regions like the 5' and 3' UTRs and promoter domains. The fundamental advantage of Cpf1 over Cas9 is that the Cpf1 crRNA is shorter than the spCas9 gRNA by 60 nucleotides, and at the same time, no tracrRNA is needed (Fonfara et al. 2016), hence facilitating multiplex gene editing (Wang et al. 2017b). Additional tracrRNA is not required by Cpf1 to form a mature crRNA. Unlike Cas9 that recognizes G-rich PAM sequences, Cpf1 recognizes T-rich PAM sequences. Finally, it was also observed that Cas9 endonuclease generates blunt ends, whereas Cpf1 endonuclease produces cohesive ends (Xu et al. 2019). Researchers have recently identified numerous expanded PAM of Cas9 and Cpf1 variants, which will eventually facilitate sgRNA designing for the genome editing of polyploid crops. Another noteworthy application of crop improvement is CRISPR/Cas-mediated precise base editing. An addition of a ‘base editing’ function to CRISPR/Cas9 can induce a C→T and G→A conversion. The base editing system of wheat can be superimposed to carry out site-specific modification in other polyploid crops.

Rodríguez-Leal et al. 2017 took a significant step ahead by combining the diverse behaviors of CRISPR/Cas9 to engineer quantitative trait loci variation (QTL) by performing mutations in the cis-regulatory regions. The huge collection of cis-regulatory alleles that was created revealed that a humongous number of quantitative variations could be achieved by remodeling the expression of individual genes. Thus, performing different permutations in promoters of various developmental regulators can modify diverse traits of crop plants. SlCLV3 promoter alleles were characterized, which consequently provided fundamental insights on numerous angles like the complex structure of the cis-regulatory regions, regulation of transcription and finally the control of quantitative traits. A study was performed, where a bidirectional strategy carried out (Li et al. 2018c) which involved promoting the lycopene content of tomato on the one hand and blocking the conversion of lycopene to β- and α-carotene. Five candidate genes regulating the carotenoid metabolic pathway were edited using the CRISPR/Cas9 system followed by Agrobacterium tumefaciens-mediated transformation. The breakthrough result obtained there was a 5.1-fold increase in the lycopene content of tomato. The homozygous trait was transmitted successfully to the succeeding generations. Breakthrough research by (Liang et al. 2017) on bread wheat makes it clear that it is more than possible to utilize CRISPR/Cas ribonucleoproteins (RNPs) for selection-free site-directed mutagenesis by embryo bombardment. To stop all possibilities of transgene integration and to minimize off-target mutations are the two essential points of optimization for making CRISPR/Cas9 system a highly precise method for crop breeding. This technique was a breakthrough since it involved the delivery of active Cas9-gRNA complexes embedded on gold particles into maize cells, along with the recovery of mutant plants without selection, makes the approach extremely user friendly for genome editing in major crops.

CRISPR/Cas9 for epigenome modification and base editing

Epigenetics refers to the heritable changes in the gene expression level, generating phenotypic variation without altering the nucleotides in the DNA sequence. Three mechanisms tightly control epigenetic regulation: DNA methylation, post-translational histone modification and the action of non-coding RNAs (short-interfering RNAs, siRNAs; micro-RNAs, miRNAs), which can lead to changes in chromatin structure without affecting the DNA sequence itself. This sort of gene regulation plays a crucial role in plant’s developmental stages and its response toward biotic and abiotic stresses (Fujimoto et al. 2012). DNA methylation in plants involves the addition of a methyl group on the 5′ carbon of cytosine base to form 5′methylcytosine and it can happen in both symmetric CG, CHG and asymmetric CHH context (H = A/C/T; A, adenine; G, guanine; C, cytosine; T, thymine) (Feng et al. 2010). DNA methylation within the coding sequence leads to altered gene expressions, whereas methylation in the promoter region results in gene silencing. A dynamic interplay between DNA methylation and demethylation is very crucial for silencing deleterious transposon insertions and regulating overall gene expressions during the developmental stages of plants (Van Oosten et al. 2014).

Since its inception, Cas9 has been very rapidly and widely accepted as a tool for genome editing. Initially dCas9—the nuclease inactivated Cas9 variant carrying point mutations in the HNH/RuvC-like catalytic domain was created to elucidate their roles in dsDNA cleavage (Jinek et al. 2012). But later on, it was adopted as a DNA-binding platform for a diverse range of functions such as epigenetic modifications (Hilton et al. 2015; Kearns et al. 2015) and gene-expression modulations (Cheng et al. 2013). The strategy to use CRISPR/Cas in epigenetic modification is to fuse the dCas9 with a transcriptional activator or repressor domain known to have epigenetic effects (epieffector). Specifically designed gRNA coupled with dCas9-epieffector complex can achieve the required methylation/demethylation at the DNA level for the desired change in phenotype or trait (Fig. 7a, b). The conventional CRISPR/Cas system has been successfully applied to a wide range of crops for improving yield, nutritional value, and stress and disease resistance. However, one of the
major concerns regarding this technology is the risk of off-targeting, resulting in unpredictable mutations. The conventional CRISPR/Cas system has been successfully applied to a wide range of crops for improving yield, nutritional value, and stress and disease resistance (Mlambo et al. 2018). Epigenetic modulation leads to genetic gain of function, which could speed up the process of domestication of wild species with improved yield, nutrition, fruit and seed numbers (Springer 2013). Induced methylation/demethylation could impact hybrid breeding and induce new gene expression patterns in next-generation offspring, thereby controlling their phenotypes (Stroud et al. 2013; Stelpflug et al. 2014). Papikian et al. 2019 successfully triggered early flowering phenotype in Arabidopsis thaliana using dCas9-SunTag system-mediated induction of FWA promoter. dCas9-mediated targeted induction of DNA methylation to alter gene expression level of those genes that negatively affects the desired traits and yield in crops could be a potential way to achieve food security in future.

Base editing refers to a novel CRISPR/Cas-mediated genome editing technique to mutate a single base without the need for double-stranded breaks (DSBs) or homology directed repair. With the aid of CRISPR/Cas base editors all four transition mutations (A to G, T to C, G to A and C to T) can be achieved. Conventional CRISPR/Cas-mediated indels by generating DSBs has greater risk of off-targeting, while fusing it with base editors to dCas9 or nCas9 (D10A nickase), desired point mutation can be achieved minimizing the risk of off-targeting (Komor et al. 2017). For instance, fusing cytidine deaminase which operates only on single-stranded DNA, to a nuclease inactivated dCas9 can achieve C to U conversion with great precision (Jiang et al. 2016; Komor et al. 2016). dCas9/nCas9-mediated adenine or cytosine base editing has been successfully applied to many plant species including tomato, rice, wheat, maize, brassica over last few years, proving it to be an alternative tool for crop improvement beside the conventional CRISPR/Cas9 (Shimatani et al. 2017; Zong et al. 2017; Hua et al. 2018; Kang et al. 2018).

Conclusion

As per ‘Food and Agricultural Organization’ (FAO), the world population is increasing in an exponential manner and it is estimated that it will reach approx. 9.1 billion by 2050 (United Nations World Population Prospects: FAO, 2019). The food production capacity needs to be increased by at least 70% to feed this huge population. Conventional breeding methods for agronomic crops such as tomato would not be able to keep up with the pace and improved variety of disease-resistant smart crops will be a necessity to address the food security of the world. In this context, next-generation genome editing technology like CRISPR/Cas gains importance, which has been used in tomato for enhancing the yield, nutritional values and tolerance to both biotic and abiotic stresses.

In recent years, researchers around the globe have been able to identify the presence of two different classes of CRISPR/Cas systems in bacteria and archaea. The diversity leading to the classification of CRISPR/Cas systems is the result of novel effector proteins, locus architecture and unique molecular mechanisms. The most mention worthy mechanisms being ‘the sole RNA targeting’ seen in the type VI systems establishing a direct link between CRISPR immunity and dormancy induction, the pre-crRNA processing by the type V-A, VI-A effector proteins and the activity of CRISPR-associated reverse transcriptase in type III systems carried out for RNA adaptation (Sukrit et al. 2016). The discovery of Casposons and type V-U loci opens doors for further investigation of the pathway of gradual evolution of the CRISPR/Cas system which finally leads to the formation of adaptive immunity from mobile genetic elements. The past two decades have witnessed an immense development in CRISPR/Cas research including PAM-free CRISPR/Cas system, dCas9-mediated epigenome modification and targeted base editing, yet several key questions remain to be answered like the missing link connecting interference and adaptation in primed spacer acquisition, the mystery behind the horizontal transfer of CRISPR and many more. A lot of breakthrough research is going on globally to elucidate numerous arcane and outstanding questions of this field. It appears that CRISPR/Cas system will not be barricaded by genomic complexity, GM controversy, government sanctions, etc. and is here to stay.

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Declarations

Conflict of interest The authors declare that they have no conflict of interest.

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