SYMBIOGENETICS

New naturally transgenic plants: 2020 update

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Abstract

Agrobacterium-mediated gene transfer leads to crown gall or hairy roots disease, due to expression of transferred T-DNA genes. Spontaneous plant regeneration from the transformed tissues can produce natural transformants carrying cellular T-DNA (cT-DNA) sequences of agrobacterial origin. In 2019, based on genomic sequencing data, cT-DNA horizontally transferred from Agrobacterium were found in two dozen species of angiosperms. This made it possible to evaluate the spread of this phenomenon, as well as make some generalizations regarding the diversity of horizontally transferred genes. The presented research is a continuation of work in this field. It resulted in the description of new naturally occurring transgenic species Aeschynomene evenia C. Wright, Eperua falcata Aubl., Eucalyptus cloeziana F. Muell., Boswellia sacra Flueck., Kewa caespitosa (Friedrich) Christenh., Pharnaceum exiguum Adamson, Silene noctiflora L., Nyssa sinensis Oliv., Vaccinium corymbosum L., Populus alba L. × Populus glandulosa Moench. The previously identified patterns regarding the frequency of the occurrence of natural transformants and the general properties of the cT-DNAs were confirmed in this study.

Keywords: cT-DNA, horizontal gene transfer, naturally-transgenic plants

Introduction

Agrobacterium-mediated transformation is the most common method for obtaining genetically modified plants. It is based on the ability of these soil bacteria to transfer a fragment of their plasmid (T-DNA, transferred DNA) and integrate it into the chromosome of the host plant. In nature, such a transfer leads to the development of two types of diseases: crown gall and hairy root diseases. These neoplasms are transgenic tissues on a non-transgenic plant. Scientists have managed to replace T-DNA genes with the sequences they need, transfer them using agrobacterial vectors into plant cells, and regenerate whole plants from such transgenic cells (Nester, 2014). It turned out that similar processes occur in nature, since plants were found to contain sequences homologous to the T-DNA of Agrobacterium in their genomes (Chen and Otten, 2017; Matveeva, 2018). This T-DNA was named cellular T-DNA (cT-DNA). The first such plants were found within the genus Nicotiana (White et al., 1983), and more than 20 years later in the genomes of Linaria and Ipomoea (Matveeva et al., 2012; Kyndt et al., 2015). Until 2019, the list of naturally transgenic plants was limited to these three genera. Digressing slightly from the main topic, we want to note that we are aware that the phylogeny of the genus Agrobacterium has been revised since the first discovery of T-DNA in wild plants (Young et al., 2001, 2003; Farrand et al., 2003); however, in the text of the manuscript we will use the collective term Agrobacterium as a tribute to tradition, and also because of the impossibility of accurately identifying the type of bacteria that participated in the transformation of the plant millions of years ago. The small fragments of T-DNA present in plant genomes are not sufficient for this. At the same time, further in the text of the manuscript, when indicating the closest of the modern strains, we will provide their modern name.

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The development of genomic sequencing and bioinformatics methods have opened up new opportunities for the search for new natural GMOs. Such a search was crowned with success in 2019 (Matveeva and Otten, 2019): another two dozen species, the ancestors of which underwent Agrobacterium-mediated transformation during their evolution, were described within the genera Eutrema, Arachis, Nissolia, Quillaja, Euphorbia, Parasponia, Trema, Humulus, Psidium, Eugenia, Juglans, Azadirachta, Silene, Dianthus, Vaccinium, Camellia and Cuscuta. Analysis of transcriptome data revealed an additional list of natural transformants. However, the degree of confidence in natural transgenicity based on transcriptomic data is lower than that based on results of genome sequencing and assembly. This is due to the lack of information about the localization site of the sequences, which leads to the possibility that the sequences result from Agrobacterium DNA contamination. The most interesting results of transcriptome assembly were several T-DNA-like sequences of the representatives of the genus Diospyros, containing a combination of opine and *plast*-genes. Matveeva and Otten's (2019) study was done exclusively using bioinformatic analysis of published sequences of plant genomes. A few months later, an article was published in which molecular methods confirmed the presence of T-DNA in plants of the genus Cuscuta, previously identified by bioinformatics means (Zhang et al., 2020). Numerous new examples of natural transformants show that at least 7% of the dicotyledonous species are naturally transformed plants, and provide valuable material for studying the role of horizontal gene transfer in plant evolution (Matveeva and Otten, 2019). These results also serve as an important argument in support of GMOs.

A year has passed since the publication of Matveeva and Otten (2019). During this time, new plant genomes were sequenced and deposited in the NCBI database (O'Leary et al. 2016). The aim of this work was to update the list of naturally transgenic plants taking into account new NGS data, and generalize all the results obtained.

Material and methods

The search for T-DNA-like sequences was done based on National Center for Biotechnology Information (NCBI) Whole-Genome Shotgun (WGS) contigs of all plant genomes sequenced since April 2019 to date, using the TBLASTN algorithm with default settings. In the second step, Vir protein sequences were used to search for possible *Agrobacterium* contaminations in those genomes. In the third step, contigs that potentially encoded T-DNA-like protein sequences with identity levels 30% or higher were analyzed further. They were used as queries in BLASTX with default settings to detect the closest protein homologs and to identify proteins encoded by plant genes surrounding the cT-DNA. All query sequences are detailed in our previous paper (Matveeva and Otten, 2019). The Vector NTI AdvanceTM software was used to build the cT-DNA maps.

Phylogenetic analysis of *rolB/C* homologs was done in MEGA 7.0 (Kumar et al., 2016) by using the Maximum Likelihood method based on the JTT matrix-based model (Jones et al., 1992) (In addition, the Dayhoff matrix based model (Schwarz and Dayhoff, 1979), Poisson correction model (Zuckerkandl and Pauling, 1965) and Equal Input model (Tajima and Nei, 1984) were used for more reliable conclusions). The bootstrap consensus tree inferred from 500 replicates was taken to represent the evolutionary history of the taxa analyzed (Felsenstein, 1985). Branches corresponding to partitions reproduced in less than 50% bootstrap replicates were collapsed. Initial tree(s) for the heuristic search were obtained automatically by applying Neighbor-Join and BioNJ algorithms to a matrix of pairwise distances estimated using a JTT model, and then selecting the topology with superior log likelihood value. The analysis involved 19 amino acid sequences. All positions with less than 95% site coverage were eliminated. That is, fewer than 5% alignment gaps, missing data, and ambiguous bases were allowed at any position. There were a total of 140 positions in the final dataset.

The supplementary materials present a similar analysis performed by UPGMA method (Sneath and Sokal, 1973) and neighbor-joining method (Saitou Nei, 1987).

Results and discussion

Since April 2019 (Matveeva and Otten, 2019), the genomes of another 206 angiosperm species have been sequenced. New examples of natural GMOs were identified in 10 species (about 5%) from 10 genera, 9 families and 7 orders, according to the previously described methodology (Matveeva and Otten, 2019). They are listed in Table 1. Schemes of extended cT-DNAs are shown in Figure 1.

For representatives of two genera, the cT-DNA structure was specified. At the same time, their transgenic nature was described earlier.

Until recently, two variants of cT-DNA have been characterized in plants of the genus *Ipomoea* (Kyndt et al., 2015; Quispe-Huamanquispe et al., 2019). In our study, based on the genome sequences of *I. trifida* (Kunth) G. Don and *I. batatas* (L.) Lam., a new cT-DNA variant was discovered. It contains *mas2*'-like and *mas1'*like sequences. The fragment that we found in *I. trifida* was named *It*-TDNA3. A similar (86%) fragment was also found in *I. batatas*. At the same time, the boundary sequences of plant origin are 97% similar, showing that they result from the same transformation event. The database also contains short contigs containing *mas2'* homologues. However, it is not possible to attribute them to any extended sequence. Further research is required



Fig. 1. Structure of cT-DNA plant species. (Wide green arrows show sequences similar to Agrobacterium T-DNA genes, blue arrows show inverted repeats, green thin arrows show direct repeats. Red arrows show short repeating sequences).

to clarify the nature of these sequences. Therefore, they are not currently listed in the results table.

We predicted a cT-DNA in *Diospyros lotus* L. (dateplum) based on the analysis of the TSA database (Matveeva and Otten, 2019). Analysis of the results of genome assembly made it possible to describe seven variants of cT-DNA in this species, representing footprints of several independent transformation events in the evolution of this species (Fig. 1). *Dl*-TDNA1 and 2 are located close to the boundaries of the assembled sequences. They share 99% similarity and may be part of the same cT-DNA. If so, then this is the youngest cT-DNA in the genome of this species, which can be dated by the repeat structure. It is followed by *Dl*-TDNA5, 7 and 6. *Dl*-TDNA6 is the oldest one. Other traces of multiple acts of agrobacterial transformation in the evolution of ancestral forms of

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	Similarity	level between 2 arms of the cT-DNA	92				n/a	n/a	95								
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	ldentity	% of identity	49	60	49	86	62	54	70	69	61	7147	77	61	66	44	56
-		position	98346 -99995	102808 -102180**	106748 -104516	112142 -112940	25927086 — 25926118	1339-149	19003305 — 19004414	19005239 -19004732	19029872 -19028766	19030390 -19031235 19046200 -19046439	19047360 -19046844	19064603 — 19063528	19065068 — 19066309	19091525 — 19090591	19097721 -19096760
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	Gene homolog		orf13-like	<i>rolB</i> -like	<i>orf13-</i> like	acs-like	mis-like	nos-like	mas1'-like	ags-like	<i>mas2'-</i> like	mas1'-like	ags-like	<i>mas2'-</i> like	mas1'-like	<i>mas2</i> '-like	<i>mas2</i> '-like
Ilysis of WGS database		Accession #	SMNX01000141.1				RYYW0100009.1	CWNJ01257379.1	JABKBO010000005.1								
cT-DNAs detected by ana		species, curuvar, line, isolate	<i>Populus alba × Populus glandulosa</i> isolate Bei- jing Shi				Aeschynomene evenia isolate CIAT22838	Eperua falcata	<i>Eucalyptus cloeziana</i> iso- late ANBG68772								
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level to proteins from NCBI	Organism and protein ID	WP_034521016.1 Agrobacte- rium sp.	WP_174054196.1 R. rhizogenes	WP_174054196.1 R. rhizogenes	AAA22094.1 R. rhizogenes	WP_149743959.1 <i>Rhizobium</i> sp.	AIM40179.1 R. rhizogenes	AIM40178.1 Nicotiana tomen- tosiformis	BAB85949.1 Nicotiana glauca	WP_176453671.1 R. rhizogenes	WP_176453671.1 R. rhizogenes	BAB85949.1 Nicotiana glauca	BAB85949.1 Nicotiana glauca	BAB85949.1 Nicotiana glauca	BAB85949.1 Nicotiana glauca	WP_156551602.1 <i>Allorhizobium vitis</i> (Ophel and Kerr 1990)	WP_156551602.1 A. vitis	WP_156551602.1 A. vitis	WP_174084799.1 A. vitis	WP_071208191.1 A. vitis				
Identity	% of identity	69	47	45	40	40	31	29	65	62	54	51	66	62	59	60	64	64	63	54	68	63	62	72
	position	4554 — 5602	15557 -13686	17274 — 19107	19840 -20193	21778 — 22257	23456 — 23151	23561 -23151	509 — 1510	1503 — 2552	1924 — 2417	4964 — 5356	26894 — 27826	7318 — 8232	14903 — 15822	1170610 — 1176104	1182125 — 1183057	1210326 — 1209418	37722 — 36790	5037 — 5411	25613 -24951	28697 -27765	35431 -34499	6031 — 5510
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:	Species, cultivar, line, isolate	<i>Boswellia sacra</i> isolate BS-S2							Kewa caespitosa				Pharnaceum exiguum			<i>Silene noctiflora</i> isolate OPL-1.1								
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Continuation of the Table 1

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orde	meì	isolate	Accession #	Gene homolog	tact*	position	% of identity	Organism and protein ID	2 arms of the cT-DNA	1
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səli		Diospyros lotus cv. Kun-	D/-T-DNA1 BEWH01006414.1	<i>cus</i> -like		41404 — 42306	73	WP_156551602.1 A. vitis	n/a	
Ēricā		Sensin		orf14-like	ı	44357 -43779	56	WP_174054201.1 R. rhizogenes		
			DI-T-DNA2 BEWH01000237.1	<i>cus</i> -like	+	334877 — 335779	73	WP_156551602.1 A. vitis	n/a	
				orf14-like		337830 -337252	56	WP_174054201.1 R. rhizogenes		
				acs-like	+	347771 — 348979	81	GAJ95539.1 R. rhizogenes		
			D/-T-DNA3	acs-like	+	8509 — 7220	80	GAJ95539.1 R. rhizogenes	n/a	
			BEWHUI000419.1	<i>orf3</i> -like		10262 — 11610	70	KEA04445.1 R. rhizogenes		
				IS5 transposase	ı	12185 -12934	83	WP_184141638.1 <i>Shinella</i> <i>fusca</i> Vaz-Moreira et al. 2010		
				<i>iaaM</i> -like	1	13558 -15830	45	WP_034521028.1 Agrobacte- rium sp.		
			<i>D</i> /-T-DNA4 BEWH01000029.1	orf8-like	+	2523369 — 2522746	68	WP_116979321.1 Agrobacterium salinitolerans Yan et al. 2017	n/a	
				<i>orf3</i> -like		2525305 -2523957	72	KEA04445.1 R. rhizogenes		
				orf2-like		2525577 -2526392		WP_174054193.1 R. rhizogenes		
				acs-like	1	2526938 — 2528244	73	KEA0447.1 Agrobacterium sp.		1
			D/-T-DNA5 BEWH01004217.1	sus-like	+	21622 — 22629	77	WP_174080856.1 R. rhizogenes	97	
				IS630		25887 -26667	83	WP_165826447.1 Rhizobium wuzhouense Yuan et al. 2018		
				C -like	1	28060 -26786	76	WP_174054195.1 R. rhizogenes		
				C -like		30052 — 31252	76	WP_174054195.1 R. rhizogenes		
				IS630	,	32218 -31443	83	WP_165826447.1 R. wu- zhouense		
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Identity	% of identity		58	60	68	70	47	73	60	55	38	50	48	41
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Similarity	2 arms of the cT-DNA		n/a							n/a		n/a
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Identity	% of identity		71	72	77	75				77	68	63
	position		13842355 -13841152	13842876 -13842715	13845955 -13846602	13847831 — 13848160	ed by Kyndt et al. (2015)		ed by Kyndt et al. (2015)	54319 -53674	56951 — 58017	27628510 — 27627275
<u>!</u>	In- tact*						describe		describe			
	Gene homolog	Contains <i>Ib-</i> TDNA2	<i>mas2</i> '-like	mas2-like'	mas1'-like	mas1'-like	Contains <i>Ib</i> -TDNA1,		Contains <i>Ib-</i> TDNA2,	mas1'-like	<i>mas2</i> '-like	<i>mas2</i> '-like
	Accession #	SMMV01000602.1	<i>lt</i> -TDNA3 SMMV01000003.1				NXFB01008336.1	FLTB01041015.1	NXFB0100007.1	<i>lb</i> -TDNA3 NXFB01000244.1		NXFB01000002.1
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** gene location on the negative strand



Fig. 2. Molecular phylogenetic analysis of *rolB/C* homologs from *Rhizobium*, *Ensifer*, *Laccaria*, *Ipomoea*, *Vaccinium* and *Nyssa* species by Maximum Likelihood method based on the JTT matrix-based model. (Dayhoff matrix based model, Poisson correction model and Equal Input model resulted to the same topology of the tree). The cluster containing new *rolB/C*-like gene is outlined in red.

modern species have been previously described within the genera *Nicotiana* and *Parasponia* (Chen et al., 2014; Matveeva and Otten, 2019)

All new species of naturally transgenic plants belong to the same orders where natural GMOs were previously described. Vaccinium corymbosum L. and Silene noctiflora L. belong to genera in which natural GMOs were previously found. They contain sequences similar to those described earlier, which can be further used for phylogenetic studies based on the T-DNA structure. Our study also confirms the prevalence of opine genes in natural transformants. As before, we observe extended cT-DNAs organized as repeats. Inverted repeats may be generated during the process of T-DNA transfer and integration into plant chromosomes. Direct repeats may possibly be explained by DNA rearrangements associated with transposons found around the repeated cT-DNA regions. An interesting feature of eucalyptus T-DNA is that relatively short fragments of agrobacterial origin with similar opine genes are interspersed with extended DNA fragments of plant origin. A large number of repeats of the same opine genes, that are found in Silene species, Kewa caespitosa (Friedrich) Christenh. and Pharnaceum exiguum Adamson is another feature that requires further study; it may result from the insertion of multiple copies during the initial transformation

event, or from amplification of integrated copies at a later stage.

The data on the fine structure of cT-DNA in representatives of different taxa obtained earlier and in the present work can be further used to search for patterns of host specificity of modern agrobacterial strains. This issue can be investigated both from a phylogenetic and from an ecological point of view, since the idea of coevolution of symbionts is gaining in importance (Matveeva et al., 2018). We can already illustrate this thesis with the case of an unusual *plast* gene, which we described for the first time in the genomic sequence of Vaccinium macrocarpon Aiton. This fragment attracted our interest because it was closer to fungal *plast*-genes than agrobacterial ones. In the present work, a similar sequence was found in Nyssa sinensis Oliv. Figure 2 shows that Nyssa, Vaccinium and Laccaria sequences cluster together with rolB/C-like gene of Ensifer sp. from the Rhizobiaceae family. Phylogenetic trees constructed by other methods (Supp. Fig. 1) have a similar topology, which confirms the reliability of this cluster. The genera Nyssa and Vaccinium are not related, but these plants share similar habitats, characterized by increased moisture (https:// www.hortweek.com; Song and Hancock, 2011). Perhaps the search for an Agrobacterium strain similar to those that transformed these species will lead to the discovery of bacterial determinants that are important for the survival of such strains in wet habitats.

Conclusion

Thus, in this study, new natural GMOs were described in 10 species (*Aeschynomene evenia, Eperua falcate, Eucalyptus cloeziana, Boswellia sacra, Kewa caespitosa, Pharnaceum exiguum, Silene noctiflora, Nyssa sinensis, Vaccinium corymbosum, Populus alba* × *Populus glandulosa*) belonging to 10 genera, 9 families and 7 orders. The new type of cT-DNA was described in *Ipomoea trifida* and *Ipomoea batatas*, and the structure of cT-DNAs of *Diospyros lotus* cv. Kunsenshi was clarified. The previously identified patterns regarding the frequency of the occurrence of naturally transgenic plants and the general properties of the cT-DNAs were confirmed. The data obtained can be used further for genetic engineering, plant phylogeny and evolutionary research.

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SUPPLEMENTS

Supplementary



Comparison of the topology of phylogenetic trees of *rolB/C* homologs constructed by

- A Maximum Likelihood method based on the JTT matrix-based model (as in fig. 1)
- B Neighbor-joining method based on the JTT matrix-based model
- C UPGMA method based on the Poisson correction model
- D UPGMA method based on the JTT matrix-based model