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Evidence of genotype dependency within *Agrobacterium tumefaciens* in relation to the integration of vector backbone sequence in transgenic *Phytophthora infestans*-tolerant potato

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In this study the effect of *Agrobacterium tumefaciens* genotype of two strains AGL1 and LBA4404 was investigated in regard to the propensity for backbone integration during the transformation of potato for blight tolerance conferred by the resistant to blight (*RB*) gene carried by the vector pCLD04541. A PCR based walking approach was employed to identify left and right backbone sequences as well as for selected genes carried on the plasmid backbone. It was found that adjacent to the left border insertion site, the integration of backbone sequence was greater for AGL1 than for LBA4404; however, the opposite was observed with regards to the right border T-DNA junction. Considering both T-DNA borders LBA4404 was found to have a two fold greater integration potential for backbone than the AGL1. The possibility of only backbone integration in T-DNA negative plants was also investigated with the average rate of integration between the two strains calculated at 4.2% with LBA4404 recording a three fold greater occurrence of backbone integration than AGL1. In summary, evidence of *Agrobacterium* genotype dependency showed that LBA4404 has greater potential to integrate non-T-DNA vector sequence than AGL1 and this should be taken into account when utilising the listed *A. tumefaciens* genotypes in generating transgenic potato. Additionally, the application of a PCR and primer walking system proved to be reliable and allows for fine detailed studies of backbone sequence integration of transgenic plant.

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[Key words: Agrobacterium; Transformation; Potato; Transgenic; RB; Backbone integration]

Agrobacterium tumefaciens-mediated transformation (ATMT) is the main process employed for plant transformation. Established initially on dicot species (1-4), it has been extended to monocots with an ever increasing efficiency (5-10). The ATMT of plant species occurs through the delivery and integration of the T-DNA containing a gene(s) of interest, in to the target cell/tissue. T-DNA transfer is unspecific with regard to its insertion site within the host genome; however studies have shown a preference for T-DNA integration to occur in highly transcribed regions (11-13). The right and left border repeat sequences of the T-DNA are required for its integration, which requires the T-DNA to be cut precisely at the right border. Integration can also result in the nicking of the left border T-DNA sequence which can occur in an imprecise fashion leading to the delivery of non-T-DNA sequences from the vector backbone (14–16). Integration of a transgene into a plant genome will have intended effects, which are those that are targeted to occur from the introduction of the gene(s) (17). On the contrary, the erroneous integration of vector backbone sequence might have "unpredictable and unintended effects" (17). Specifically, the integration of backbone sequence might participate in the silencing of endogenous/non-endogenous gene(s) (18, 19). Additionally, it has been suggested that it could facilitate the horizontal transfer of genes between a transgenic plant and a microorganism based on the availability of DNA microhomology (20, 21).

Evidence of backbone integration post-ATMT has been accumulated from several plant species: tobacco (14), rice (22, 23) Arabidopsis (24) strawberry (25) cherry rootstock (26) and barley (27). In regards to potato, the fourth most important crop worldwide (28) and one of the first crops to be transformed (29, 30), two studies (31, 32) have been published one of which, investigated the application of in situ hybridization of extended DNA fibres to evaluate the integration patterns of T-DNA and backbone sequence (32). In general, only small populations of transgenic plants have been analysed and to date, no study has examined the effect of bacterial genotype on the propensity for backbone integration. The aim of this study was therefore to investigate the influence of bacterial genotype on the propensity to deliver non-T-DNA (backbone) sequence in a large population of putative transgenic potato plants (n=327). This was achieved using two A. tumefaciens strains (AGL1 and LBA4404) both of which were equipped with the vector pCLDO4541 to confer resistance to Phytophthora infestans through expression of the RB gene (33, 34);

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furthermore described as the resistant to blight (*ResBl*) gene to avoid confusion with the T-DNA right border (RB). Furthermore, we investigated the presence or absence of backbone sequences in T-DNA negative plants known as escapes.

MATERIALS AND METHODS

The plant material used in this investigation was Plant and vector materials Solanum tuberosum L., cv. Désireé and consisted of internodal explants taken from sterile in vitro plantlets of 4 to 6 weeks old which, according to their length, were cut into segments of circa 5 mm. For this study, an average of 250 stem fragments per treatment was employed for each ATMT. The vector used was pCDLO4541 (35), (Accession number: AF184978) which was kindly provided by Dr. Jiming Jang, University of Wisconsin, Madison, USA. This vector (27.6 kb), contains the ResBl gene (8.6 kb), which confers increased resistance to P. infestans (33, 34). Transgenic plant selection was assessed through resistance to kanamycin (50 µg/ml), as conferred by the *nptII* gene. Bacterial selection was conferred by the Tet gene located on the pCLDO4541 backbone. The plasmid was introduced into A. tumefaciens strains AGL1 (36) and LBA4404 (37) by the freeze-thaw method (38). Regeneration, transformation and selection of Désireé explants were carried out as in Petti et al. (unpublished data). Briefly, on following bacterial exposure $(OD_{600} = 1.0)$ of the explants, they were transferred to MS media and co-cultivated for 2/3 day. A. tumefaciens overgrowth was washed off (MS medium plus cefotaxime-500 µg/ml) and blot-dried samples were transferred to callus inducing media consisting of MS basal salts (4.414 µg/ml, Apollo Scientific, Stockport, Cheshire, UK), ZR (0.8 µg/ml; Apollo Scientific, UK), thiamine-HCl (10 µg/ml; Sigma, St. Louis, MO, USA), pyridime-HCl (1 µg/ml; Sigma), nicotinic acid (1 µg/ml; Sigma), NAA (0.3 µg/ml; Sigma), BAP (2.25 µg/ml; Sigma) and 2, 4-D (0.05 µg/ml; Sigma), sucrose (3%; Sigma) and agar technical 3 (0.6%, Oxoid, Basingstroke, Hampshire, UK), cefotaxime (500 µg/ml; Apollo Scientific) and the plant selective antibiotic, kanamycin (50 µg/ml; Apollo Scientific). Fragments were sub-cultured every 2 weeks on to fresh media. Each treated fragment was evaluated for the formation of independent calli (callus formed in noncontiguous areas) indicating putative independent transformation events; these were transferred to shoot inducing media (MS basal salts 4.414 µg/ml), ZR (0.8 µg/ml; Apollo Scientific), thiamine-HCl (10 µg/ml), pyridime-HCl (1 µg/ml), nicotinic acid (1 µg/ml), gibberellic acid (GA3, 0.8 µg/ml; Sigma) sucrose (3%) and agar technical 3 (0.6%; Oxoid) till shoot formation. From each independent event, a single shoot was transferred to root inducing media (RIM), as for SIM without GA3 supplemented with indoleacetic acid (IAA, 0.1 µg/ml; Sigma) at full kanamycin selection (100 µg/ml) and incubated at 22 °C for 16/8 h day length. For DNA extraction, the growing shoots were excised near the base and put into a 2.2 ml eppendorf and a glass bead (3 mm diameter; Qiagen, Crawley, West Sussex, UK) added. The samples were pre-frozen at -80° C for 1-2 h and freezedried for 24-48 h. The plant material was disrupted in a mixer mill (Retsch GmbH, MM400, Haan, Germany) and the DNA extracted via a modified CTAB method to accommodate the 2.2 ml sample size.

Primer design

Verification of transgenic lines The molecular diagnosis of the putative *ResBl* lines was completed through single and multiplex PCR. Single PCR evaluation was completed using *ResBl* gene-specific primers (*ResBl-F*, *ResBl-R*: Table 1), which were designed according to the principles of Mismatch Amplification Mutation Assay (MAMA) PCR and described elsewhere (39). Multiplex PCR screening targeted: *ResBl-nptII* and 35S-*nptII* which amplified 812 bp and 535 bp, and 515/535 bp and 1070 bp fragments respectively (Fig. 2). Primer sequences required for the detection of the *nptII ResBl* genes and 35S promoter are given in Table 1. The PCR conditions for the *ResBl* primers and for the multiplex PCRs were: first cycle 96 °C for 6 min, 94 °C for 15 s, 57 °C for 30 s, 68 °C 1 min, 35 additional cycles and 5 min final extension at 72 °C. The PCR was carried out with 20 ng of DNA, 200 µM of dNTP's, 10 pM of primers and 0.20 units/µl of Taq polymerase (NEB, Wilbury Way, Hitchin, Herts, UK) made up to a 20 µl reaction with 1× buffer and PCR-grade water.

Synchronous and asynchronous backbone analysis Synchronous backbone analysis (SBA hereafter) aims to determine the presence/absence of pCLDO4541 backbone sequences, be it adjacent to the right and/or left T-DNA border, in the ResBl positive plants. The asynchronous analysis investigated the possibility that T-DNA negative lines contained pCDLO4541 backbone integrated within their genome. For both analyses, a PCR based primer walking approach was utilised. A single forward primer was designed within the T-DNA and upstream of the left border (Accession number: AY234326) while the reverse primers (12 sets) were designed downstream of the left border and out on the plasmid backbone with an incremental size from 150 bp to 621 bp (Fig. 1). The primer sequences and the expected products are given in Table 1. Furthermore, two more downstream primers (IN1 and IN2) were designed at 730 bp (IN1) and at 1840 bp from the left border (IN2, Table 1). The primers (lb 1-12, IN1 and IN2) were designed to have a similar melting temperature and a generic program was adequate to amplify each of the above sets using the vector DNA as a control template (illustrated in Fig. 3). PCR conditions were: cycle 1, 96 °C, for 5 min; 96 °C for 50 s, 54 °C for 35 s, 72 °C for 50 s, for 30 cycles and a final extension of 5 min at 72 °C. PCR reaction carried out as previously described. To analyse the frequency of integration of the backbone sequences on the right border, only two primers were designed, rb1 and rb2 (Fig. 1 and Table 1). The forward primer was internal to the T-DNA sequences and located at 271 bp upstream of the position of the right border (AY456904) while two external

TABLE 1. Sequences of primers adopted for the synchronous and asynchronous

backbone analysis						
Primer	Sequence	Size product (bp)				
lb1-12-F	5'-ATAATAACGCTGCGGACATCT	150				
lb1-R	5'-GGCGGCAGGATATATTCAA					
lb2-R	5'-GCGTAGAAACCAACATGCAA	195				
lb3-R	5'-GCCTAACCGGCTCAGTTCTG	204				
lb4-R	5'-GCACATGGCTCAGTTCTCAA	246				
lb5-R	5'-CCGTCGCTCACCGTGTT	274				
lb6-R	5'-AGTCCCATGTGGATCACTCC	300				
lb7-R	5'-CGGATCCGACGGATGATGTT	324				
lb8-R	5'-GTCGGCTTGATCTCACCGA	362				
lb9-R	5'-ATCGTGTTGCGCGCCTG	387				
lb10-R	5'-CGTCGGCTCGATTGTACCTG	424				
lb11-R	5'-TAGCTTGCTTGGTCGTTCCG	454				
lb12-R	5'-CGAAATCCTTGGGCCTCCAC	621				
rb-F	5'-TTGCCTTTTCTTATCGACCA	379				
rb-R1	5'-TCGGCAGTTCATCAGGGCTA					
rb-R2	5'-AATATGCGTCCCTTTGGAGA	539				
IN1-F	5'-AGCAGTACCAGCAGGAATGG	330				
IN1-R	5'-TTGGTTGAACATAGCGGTGA					
IN2-F	5'-CAGGGCATCAAGAAGATGGT	485				
IN2-R	5'-GTCCAGGTCCAGCTCGATAG					
OriV-F	5'-GCAGCCCTGGTTAAAAACAA	449				
OriV-R	5'-GATACCTCGCGGAAAACTTG					
Tra-F	5'-GCTGATCCTGCATACCATCA	359				
Tra-R	5'-ATAACGAGGCCCACACCAC					
Tet-F	5'-ATGCTGGCGGAGAATCATAC	452				
Tet-R	5'-TCAACGTTCCTGACAACGAG					
TrfA-F	5'-ATCGACACGCGAGGAACTAT	409				
TrfA-R	5'-CCTGGCAAAGCTCGTAGAAC					
VirD2-F	5'-GCAGAGCGACCAATCACATA	360				
VirD2-R	5'-GGCTTCAGCGACATAGGAAG					
At-F	5'-TGCTACCCCTTCATCAAACC	375				
At-R	5'-GTTGTTTGTCCGCCTTGTTT					
ResBl-F	5'-CATCTTGAGAGAGTGAAGAATGATCT	812				
ResBl-R	5'-CTAGTGCGCAACACAATTGAA					
NPTII-F	5'-GATGTAGGAGGGCGTGGATA	535				
NPTII-R	5'-GATGTTGGCGACCTCGTATT					
35S-F	5'-AAACCTCCTCGGATTCCATT	515				
35S-R	5'-GCTCAACACATGAGGGAAAC					

(reverse primers) to the T-DNA and on the backbone sequence were designed at 106 bp and 268 bp downstream the right border (Fig. 1). PCR conditions and reactions were as for the left border analysis and were previously described.

For the asynchronous backbone analysis (AsBA hereafter), the DNA of T-DNA negative lines was checked for the presence of 4 backbone genes. These genes were *TrfA* (AY204478), *Tet* (AY204478), *Tra* and *OriV* (AY204478) and their primer sequences and the expected amplicon sizes are reported in Table 1. The PCR conditions for these sets of primers were 95 °C for 10 min, 94 °C for 1 min 60 °C for 30 s, -1 °C per cycle, 72 °C for 1 min, nine cycles with an additional 24 cycles at 94 °C for 30 s, 55 °C for 30 s, 72 °C for 30 s and a final extension of 10 min at 72 °C. The PCR reaction was carried out as for the SBA test previously described. Two distinct sets of primers for *A. tumefaciens* were employed; the first designed specifically for the detection of *A. tumefaciens* chromosomal sequences (http://www.agrobacterium.org, accession number: AE007870) while the second set was based on sequences for the *A. tumefaciens virD2* gene (AF242881).

Southern analysis was also completed on the 3 AsBA positive lines using a PCR generated amplicon of the *Tet* gene from the vector pCLD04541 as a probe. This was not possible on 2 LBA4404 lines due to their accidental destruction in the glasshouse. The PCR amplicon was gel purified (QIAquick, Qiagen), following the manufacturers indications and utilised in a random priming reaction using a direct alkaline phosphatase reaction (Alkphos, Amersham, UK) following the manufacturers instructions. Signal was generated by the application on the membranes of the enhanced chemifluorescence (ECF) substrate and a molecular scanner (Thyphoon T9410, Amersham, Little Chalfont, Buckinghamshire, UK) was employed to record the fluorescent signal.

RESULTS

Genotypes of *A. tumefaciens* influenced transformation **rate** An initial 481 potato stem explants were exposed to *A. tumefaciens*; 254 treated with AGL1 and 227 with LBA4404. Both strains harboured the vector pCLD04541 containing the *ResBl* gene, to enhance tolerance to *P. infestans* (33, 34). A total of 455 independent putative transgenic calli formed on the treated fragments (Table 2) from which 327 lines were regenerated with 209 PCR-positive for the



FIG. 1. Schematic representation detailing location of diagnostic primers (lb1–lb12; lb1– 12-F; IN1, IN2-F, IN1, IN2-R; rb-F, rb1 and rb2) for pCDLO4541.

ResBl transgene (as illustrated in Fig. 2), equating to a transformation rate of 43.4% (209/481) with respect to the overall fragments exposed. However, with respect to the *A. tumefaciens* strains employed the transformation rate was 47.6% for AGL1 and 38.7% for LBA4404. The T-DNA negative lines, which were PCR-negative to all the T-DNA genes: *ResBl*, NPTII and 35S promoter, accounted for a total of 118 plants with 79 derived from the AGL1 treatment and 39 from LBA4404.

Synchronous backbone analysis (SBA): evidence for A. tumefaciens genotype dependency A chromosome walking approach was employed for the SBA (Fig. 3) to determine the extent of backbonesequence integration at either the left or right border of the T-DNA insertion site. Prior to proceeding with the SBA evaluation, the 209 T-DNA-ResBl positive lines were examined for the presence of A. tumefaciens chromosome specific sequence and for the Ti plasmidcarried virD2 gene sequences. In 10% of the transgenic lines (n=20, data not shown) we found evidence of such sequences and those lines were eliminated from the investigation. Of the remaining 189 lines (109 AGL1-derived and 80 LBA4404-derived), which were initially analysed at the left border only of the insertion site, we found that with respect to the Désireé transgenic lines 6.4% (7/109) of the AGL1-derived ResBl lines contained backbone sequence whereas only 3.8% (3/80; line 6-2, 7-2, 8-2) tested positive within the LBA4404-derived population (Table 3). The majority of these AGL1 and LBA4404 derived lines tested positive for all the 12 left border primers tested with an integrated non-T-DNA



FIG. 2. Representative PCR screening of *ResBl* lines. (A) Multiplex PCR to identify *ResBl* and the *nptll* gene from 5 transgenic lines. (B) The same lines tested using a multiplex PCR for the 35S and for the *nptll* gene. 6: blank lane, 7: untreated plant DNA, 8: pCDL04541 DNA.

sequence of 621 bp. Three of the 8 AGL1 lines tested positive for a lower number of primer sets. More specifically 2 lines (2-1 and 3-1) for 11 primer sets with 454 bp backbone sequence integrated and 1 line (5-1) for 10 primer sets corresponding to 424 bp of backbone sequence integrated. An analogous pattern was also reported for the LBA4404derived population with 2 out of 8 (line 4-2 and 5-2) being positive to 11 sets of primers and 454 bp of backbone sequence integrated. Furthermore, we found evidence, for each of the AGL1 and LBA4404derived populations, of backbone sequence amplified up to 1800 bp downstream (with IN1 and IN2 primer pairs) from the left border. This represented 6.4% (7/109) and 10% (8/80) of the AGL1 and the LBA4404 treated populations respectively. Considering that the IN2 was designed 1800 bp downstream of the left border and the resulting amplicon was an additional 485 bp, it implies that for these specific lines at least 2.2 kb of backbone sequence was integrated into the host chromatin.

The presence of non-T-DNA sequences, solely on the right border, was detected only in 1 case (line 9-1) for the AGL1-derived lines (0.90%; 1/109) but in 4 LBA4404 lines (5%, line 9-2, 10-2, 11-2 and 12-2, Table 3). Moreover, an AGL1-derived line which was left-border-positive was also found to be right-border-positive whereas for the LBA4404-derived *ResBl* lines 5 tested positive for the left and right border sequences (Table 3).

The overall frequency of integration for backbone sequences (either left or right border) was, with respect to the AGL1 recovered *ResBl* transgenic lines, calculated at 8.2% (9/109) whereas with regards to the LBA4404 lines, it equated to 15% (12/80).

Asynchronous backbone analysis (AsBA): backbone transgenesis of potato plants A total of 118 Désireé lines were T-DNA negative (PCR-negative to *ResBl, nptII* and 35S promoter) following treatment with either *A. tumefaciens* genotypes (Table 2). When tested none of this population possessed either *A. tumefaciens* chromosomal DNA

TABLE 2. Details of the transformation experiments on Désireé fragments by A. tumefaciens strain AGL1 and LBA4404 carrying the vector pCLD04541

Strain	Fragments	Fragments Independent calli/fragment					Lines ^b	ResBl +,	ResBl –,		
		0	1	2	3	4	≥5	Total ^a	nptII+ ^c (nptII+ ^c (%)	nptII- ^d (%)
AGL1	254	107	46	49	42	9	1	311	200	121 (60.5)	79 (39.5)
LB4404	227	136	54	25	8	4	0	144	127	88 (69.3)	39 (30.7)
Total	481	243	100	74	50	13	1	455	327	209 (63.9)	118 (36.1)

^a Total number of calli resulting from the overall number of exposed fragments. This was computed by multiplying the number of fragments that developed calli per the corresponding number of independent calli formed on each fragment.

^b Indicates the number of independent shoots recovered from regenerating calli.

^c Indicates presence of *ResBl* and *nptll* transgene as verified by PCR.

^d Equates to absence of *ResBl* and *nptll* transgene.



FIG. 3. Illustration of PCR based assay to verify Ib1-Ib12 and IN1, IN2 primers as completed in duplicate on two independent pCLD04541 plasmids, a: 100 bp ladder.

sequence or the Ti plasmid encoded *virD2* gene (data not shown). Of interest the AsBA indicated that 5 T-DNA negative lines (two-AGL1 and three-LBA4404-derived) were characterised by the presence of backbone sequence belonging to 4 major genes (Fig. 4) carried by the plasmid pCLDO4541, which also included the tetracycline antibiotic resistance gene. Integration rates for *Tet*, *Tra*, *OriV* and *TrfA* were calculated at 2.5% for the AGL1 lines and 7.7% for the LBA4404 lines, implying that an estimated 27 kb circa was integrated into the DNA of each individual.

PCR results were supported by Southern analysis (Fig. 5) with 2 AGL1 and one LBA4404-derived line containing a single backbone integration event. As expected, no hybridization signal was detected for the control non-*Agrobacterium* exposed Désireé line.

DISCUSSION

Since the first report of ATMT in potato (29, 30) numerous protocols have been published (40–44) aiming to improve the rate of transfor-

TABLE 3. Degree of vector backbone sequence integrated into transgenic potato

 following treatment via AGL1-based or LBA4404-based A. tumefaciens-mediated

 transformation

Genotype	T-DNA border	<i>ResBl</i> -potato line ^a	Backbone sequence length (bp)
AGL1	Left	1-1	621
AGL1	Left	2-1	454
AGL1	Left	3-1	454
AGL1	Left	4-1	621
AGL1	Left	5-1	424
AGL1	Left	6-1	621
AGL1	Left	7-1	621
AGL1	Left	8-1	621
AGL1	Right	1-1	539
AGL1	Right	9-1	539
LBA 4404	Left	1-2	621
LBA 4404	Left	2-2	621
LBA 4404	Left	3-2	621
LBA 4404	Left	4-2	454
LBA 4404	Left	5-2	454
LBA 4404	Left	6-2	621
LBA 4404	Left	7-2	621
LBA 4404	Left	8-2	621
LBA 4404	Right	1-2	539
LBA 4404	Right	2-2	539
LBA 4404	Right	3-2	539
LBA 4404	Right	4-2	539
LBA 4404	Right	5-2	539
LBA 4404	Right	9-2	539
LBA 4404	Right	10-2	539
LBA 4404	Right	11-2	539
LBA 4404	Right	12-2	539

^a Same code number indicates the same plant line.

mation, to reduce the genotype dependency and to eliminate the risk for somaclonal variation. Only relatively recently particle bombardment (45, 46) mediated transformation has been employed (47, 48). However, the use of the entire plasmid, in general favoured to the use of cassettes during particle bombardment, is often associated with complex concatamer integration patterns and the incorporation of backbone sequences, which complicate downstream risk assessments. Previous reports have noted how A. tumefaciens will integrate not only T-DNA sequences but also vector backbone sequences and this has been shown to occur adjacent to both borders of the T-DNA, with predominance for the left border (49, 50) even though the opposite was also reported (14). An initial assessment of potential backbone integration was done on 3 transgenic Karnico lines (32) and showed a complex pattern of repeat T-DNA and backbone sequence integration often separated by potato DNA. In this study we utilised a chromosome walking PCR-based approach to investigate not only the frequency of backbone integration on a large population of transgenic potato plants but also the extent of this erroneous sequence integration in relation to the bacterial strains utilised. Furthermore, we investigated the possibility that asynchronous sequence integration could occur resulting in A. tumefaciens delivering the plasmid backbone instead of the T-DNA in to the plant genome.

The initial transformation experiment employed 484 Désireé stem explants and two *A. tumefaciens* bacterial strains: AGL1 and LBA4404. As expected the number of independent calli regenerated following *A. tumefaciens* exposure was greater for the hypervirulent strain AGL1 (36) than for the LBA4404 (37) genotype. However, this manifested as a substantially higher number of independent transformed calli per treated fragment rather than an increase in the total number of fragments transformed (p>0.05). An average transformation rate



FIG. 4. Illustration of asynchronous backbone integration in the T-DNA negative lines 1, 2, 3, 7 and 8; 4: water, 5: untransformed plant and 6 vector pCLDO4541.



FIG. 5. Southern analysis indicating the stable integration of pCLDO4541 backbone sequence in T-DNA negative transgenic potato. Lane 1, water control; lane 2, untransformed potato; lanes 3, 4, AGL1 derived lines; lane 5, LBA4404 derived line.

was determined at 43.3% based on the PCR results. The application of a low selection pressure (kan, 50 μ g/ml) reduced the toxicity of the selection increasing the recovery of transgenic plants but it also resulted in a substantial increment in the rate of false positive, or escapes (35%, 118/327).

The genotype of the transforming bacterial strain clearly had an impact on potential backbone sequence integration. Remarkably, despite the higher virulence of the AGL1 strain, the overall rate of backbone integration accounted for just 8.2% of the transgenic ResBl potato population which was nearly half of the rate associated with the LBA4404-derived population (15%). This trend was not only observed for the delivery and integration of backbone sequence at the right and left border of the T-DNA but was also recorded for the case of asynchronous delivery and integration of the full plasmid backbone sequences; 2.5% of AGL1 population against 7.7% of LBA4404-derived population. It seems evident that the genomic composition of the bacterial strain affects the way the vector borders are read and therefore what is in fact, being delivered. The importance of the left border as the T-strand terminator has already been reported in a study carried out on rice transformation (51) which showed that multiple repeats of the left border reduced the rate of backbone transformation. This is further supported through evidence that the left and the right borders can both act as initiator and terminator of T-strand production, where the most effective combination was the right-left and the less effective was the double pairing of the left.

In this study we showed that integration of the complete vector backbone occurred in 4.2% (5/118) of the T-DNA negative lines, based on PCR analysis. This result is in accordance to what Gilbertons has shown with regards to the possibility of the LB and RB borders being able to act as initiator/terminator of T-DNA strand production (Gilbertons L., presented at the Symposium on Plant Transformation Technologies, Vienna, Austria, February 4–7, 2007). Hence, suggesting that errors in the recognition of the RB/LB borders can occur and the synthesis of the T-DNA takes place in the opposite (via backbone) direction. The LBA4404 strain was most prone to generate full backbone transformants. Indeed, Ramanathan et al. (15) generated tobacco calli resistant to kanamycin from a vector created with the *nptII* gene located outside the left border. In our system, no selection was applied

to determine backbone transformed calli/plants. In fact, the intention was not to select for backbone transformed plants, rather to verify if non-transgenic plants or more accurately T-DNA negative plants could be harbouring vector DNA. As much as 27.6 kb of pCDL04541 sequence was in fact delivered into the genome of 5 individual lines (4.2% of untransformed population) which were morphologically identical to the wild type (data not shown). Preliminary analysis of these plants using interspersed-single sequence repeats (I-SSR), (52, 53) noted that the amplified DNA patterns were equivalent amongst transgenic lines and wild type (data not shown), indicating that no major genetic alterations had occurred following transformation with the backbone.

The possibility of utilising backbone transformed plants might have potential if the backbone is plant based; cisgenic (54, 55) or intragenic vector (56, 57) and the vector sequences at the left and right borders are regulatory elements for the introduced genes or matrix attachment regions (MARs). These sequences could greatly increase the efficiency by which transgenes are introduced and expressed, which could result in great benefit for those species, such as cereals, which are still difficult to transform for major traits.

The aim of this project was to investigate the potential genotype dependency of Agrobacterium in delivering non-T-DNA sequence into transgenic potato populations transformed for tolerance to P. infestans and to verify whether T-DNA negative plants could have been backbone transformed. Critically, this analysis forms part of the risk assessment required for these transgenic potato populations prior to their introduction into field trials to assess their tolerance to P. infestans (EC 2001/18). Significantly we have shown that LBA4404 was more prone to deliver backbone sequence either at the left or the right border compared to the more virulent strain, AGL1. Furthermore we proved the propensity for LBA4404 to facilitate full backbone integration (up to 27.6 kb) more efficiently than AGL1. We also showed that a PCR based approach with chromosome walking is an effective system to identify backbone transformed plants. This information should be considered by the current cohort of scientist working on the development of disease resistant potato using the LBA4404 A. tumefaciens genotype.

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