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GENETIC TRANSFORMATION AND HYBRIDIZATION

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Alcohol-inducible gene expression in transgenic *Populus*

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Abstract We tested the efficiency and optimized the con-7 ditions for controlled alcohol-inducible transgene expres-8 sion in *Populus* using gus as a reporter gene. Specificity 9 of induction, efficiency in different organs, effect of three 10 chemical inducers, and induction methods were tested us-11 ing up to 10 independent transgenic events generated in 12 two different Populus genotypes. The optimal inducer con-13 centration and the duration of induction period were deter-14 mined in dose-response and in time-course experiments. 15 Under *in vitro* conditions, β -glucuronidase (GUS) induc-16 tion was efficient both in the aerial parts and in the roots of 17 regenerated plantlets. Among the chemical inducers tested, 18 ethanol was the most effective activator with no apparent 19 phytotoxicity when concentrations were at or below 2%. 20 After 5 days of treatment, fluorometrically-determined the 21 GUS activity could be detected when inducing with ethanol 22 at concentrations as low as 0.5%. Prolonged induction by 23 ethanol vapors significantly increased the GUS activity in 24 leaves from both the tissue culture plants and greenhouse-25 grown plants. 26

27 Keywords Poplar · Aspen · Cottonwood · alc system ·
 28 Functional genomics

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Abbreviations Ac: Acetaldehyde · 2-B: 2-Butanone ·
BAP: 6-Benzylaminopurine · CaMV: Cauliflower mosaic
virus · CIM: Callus-induction medium · dpi: Days
post-induction · IBA: Indole-3-butyric acid ·
M: Agrobacterium-induction medium
LB: Luria-Bertani medium · MS: Murashige and Skoog
medium · RCF: Relative centrifugal force ·
SIM: Shoot-induction medium · TDZ: Thidiazuron ·
<i>uidA</i> : β -Glucuronidase, <i>gus</i> reporter gene ·
X-Gluc: 5-Bromo-4-chloro-3-indolyl-β-D-glucuronic
acid

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Introduction

Modifying endogene expression is an important tool for functional genomics of plants. However, the constitutive expression or silencing of a target gene often leads to pleiotropic effects, making it difficult to directly relate gene function with an observed phenotype. Developmental regulatory genes are particularly difficult to study because their constitutive over expression or silencing often interferes with the recovery of viable transgenic plants. To overcome these limitations, several chemically inducible gene-switch systems have been recently developed and tested in model plants (reviewed in Padidam 2003).

An alcohol-responsive system employing components of 52 the alc-inducible regulon of the fungus Aspergillus nidu-53 *lans* was effective in a wide range of model and crop plants 54 (Caddick et al. 1998; Roslan et al. 2001; Sweetman et al. 55 2002; Deveaux et al. 2003; Schaarschmidt et al. 2004; 56 Garoosi et al. 2005). The *alcR* gene encodes a protein that 57 regulates transcriptional activation of the alc-inducible reg-58 ulon, the product of which is required for the oxidation of 59 ethanol (Flipphi et al. 2001). The alcohol sensor protein, 60 AlcR, is a DNA-binding transcription factor (Mathieu et al. 61 2000). In the presence of inducers such as ethanol or alde-62 hydes, AlcR activates the transcription of alcohol dehydro-63 genase by binding to a specific motif in the *alcA* promoter 64 (Kulmburg et al. 1992). A transcriptional fusion of the alcA 65

promoter to a reporter gene activates the expression of the 66 reporter upon treatment with an appropriate inducing agent. 67 Using a 35S::alcR-alcA::uidA cassette, AlcR was re-68 ported to be an effective temporal and spatial regulator 69 of *uidA* (gus) gene expression in Arabidopsis (Roslan et al. 70 2001), tobacco (Sweetman et al. 2002; Schaarschmidt et al. 71 2004), potato (Sweetman et al. 2002; Junker et al. 2003), 72 oilseed rape (Sweetman et al. 2002), and tomato (Garoosi 73 et al. 2005). In plants, alcohol-inducible expression sys-74 tems responded to treatments by various chemical inducers 75 such as ethanol (Caddick et al. 1998), acetaldehyde (Ac) 76 (Schaarschmidt et al. 2004), and 2-butanone (2-B) (Junker 77 et al. 2003). 78

Apart from the preliminary observation made by Busov 79 et al. (2005), we know of no published reports of alc system being tested in Populus species (aspens and cot-81 tonwoods). Poplars are not only commercially important, 82 but they are also widely considered to be *the* model tree 83 for genomics and biotechnology (Brunner et al. 2004; 84 Strauss and Martin 2004). The availability of the com-85 plete Populus genome sequence (JGI, http://genome.jgipsf/poptr1/poptr1.home.html) adds to its utility as a model 87 system. The aim of this study was to determine whether 88 the *alc* system can function effectively in poplar, and to examine several of its properties. This includes the response 90 of the alc system to different chemical inducers, the dose-91 dependency and time-dependency of induction, different 92 methods of ethanol application, and whether the induction 93 could be observed in potted plants. We will show that the 94 alcohol-inducible system is functional and robust in trans-95 genic Populus. 96

97 Materials and methods

98 Plant genotypes, bacterial strains and vectors

⁹⁹ Hybrid aspen clones INRA 717-1B4 (female, *Populus trem-*¹⁰⁰ $ula \times P. alba$) and INRA 353-38 (male, *P. tremula* $\times P.$ ¹⁰¹ *tremuloides*) were used for all transformations.

Forty- to fifty-day-old, in vitro grown poplar plantlets 102 served as explant sources. Micro-cuttings of 717-1B4 103 and 353-38 were initially cultured on hormone-free, half-104 strength Murashige and Skoog medium (MS) (Murashige 105 and Skoog 1962). Shoot cultures were maintained on the 106 107 same medium and grown at 25°C under a 16-h photoperiod [fluorescent tubes (TL70, F25T8/TL735, Philips) at 108 a photon flux density of 45 $\mu E m^{-2} s^{-1}$]. The binary 109 vector pJH0143 was provided by Syngenta Biotechnol-110

ogy, Inc. (Research Triangle Park, NC). A diagrammatic representation of the T-DNA from pJH0143 is shown in Fig. 1. *Agrobacterium strain* C58/pMP90 (GV3101), a disarmed derivative of the nopaline strain (Koncz and Schell 1986), was transformed with pJH0143 using the freezethaw method of Holstein et al. (1978).

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Plant transformation

Agrobacterium cells harboring the binary vector were 118 grown for 24 h in Luria Butani (LB) medium (Weigel and 119 Glazebrook 2002) supplemented with 50 mg/l rifampicin, 120 50 mg/l kanamycin, and 50 mg/l gentamycin on an or-121 bital shaker at 28°C and 250 rpm. The cells were pel-122 leted by centrifugation at 3,500 rpm (1,992 RCF) for 30-123 40 min and then resuspended in sufficient Agrobacterium-124 induction medium (IM) (Han et al. 2000) to achieve an 125 OD_{600nm} of 0.5–0.6. Inter-nodal stem segments (3–4 mm 126 in length) and leaf discs (4 mm in diameter) were wounded 127 with multiple fine cuts and incubated in Agrobacterium 128 suspension with slow agitation for 1 h. The inoculated 129 explants were cocultivated in callus-induction medium 130 (CIM) [MS supplemented with 10 μ M naphthaleneacetic 131 acid (NAA) (Sigma, St. Louis, MO) and 5 µM N⁶-(2-132 isopentenyl)adenine (Sigma)] at 22°C in darkness for 2 133 days. Explants were then washed four times in sterile, 134 deionized water and once with wash solution (Han et al. 135 2000). For selection of transformed calli, explants were 136 transferred to CIM containing 50 mg/l kanamycin and 137 200 mg/l timentin for 21 days. Shoots were induced by cul-138 turing explants on SIM medium [MS containing 0.2 µM 139 TDZ (NOR-AM Chemical Co., Wilmington, DE), 100 mg/l 140 kanamycin, and 200 mg/l timentin (GlaxoSmithKline Inc., 141 Research Triangle Park, NC] for 2 to 3 months, subcultur-142 ing at 3- to 4-week intervals. For shoot elongation, explants 143 were transferred onto MS medium containing 0.1 µM 6-144 benzylaminopurine (BAP) (Sigma), 100 mg/l kanamycin, 145 and 200 mg/l timentin. The regenerated shoots were rooted 146 on half-strength MS medium supplemented $0.5 \,\mu$ M indole-147 3-butyric acid (IBA) (Sigma) and 25 mg/l kanamycin. Af-148 ter approximately 30 days, the rooted shoots were micro-149 propagated on the same medium. To ensure that the trans-150 formation events are independent, only a single clone per 151 individual explant was selected for further propagation af-152 ter confirmation of transgene presence by PCR. 153 154

All media were autoclaved at 120°C and for 20 min. Vitamins, growth regulators, and antibiotics were filtersterilized and added to medium after autoclaving. All media were adjusted to a pH of between 5.6 and 5.8 before





– gus reporter gene; t35S – CaMV 35S terminator; *pNOS* and tNOS – nopaline synthase promoter and terminator, respectively; *nptII* – neomycin phosphotransferase II. Genes are not drawn to scale

autoclaving except LB and IM, which were adjusted to pH7.0 and 5.0, respectively.

160 Genomic DNA isolation and PCR amplification

Genomic DNA was isolated from poplar leaves us-161 ing a Plant DNAeasy Kit (Qiagen, Valencia, CA) ac-162 cording to the manufacturer's instructions. Approxi-163 mately 25-50 ng of poplar DNA was used as template 164 for the polymerase chain reaction (PCR). The trans-165 gene presence was confirmed by using primers specific 166 for nptII (5'-ATCCATCATGGCTGATGCAATGCG-3' and 167 5'-CCATGATATTCGGCAAGCAGGCAT-3') and uidA 168 (5'-TGGGCATTCAGTCTG-3' and 5'-GTGATATCGTCC 169 ACCCA-3') to amplify 253- and 509-bp products, re-170 spectively. A 490-bp fragment encompassing the palcA: 171 *uidA* junction was amplified using the following: 172 primers: 5'-GTGCTCTCCTACCCCAGGAT-3' and 5'-173 TTCACACAAACGGTGATACG-3'. 174

To amplify the *nptII* junction and *alcA::uidA* junction specific fragments, the conditions used for 30 cycles were: 94°C for 1 min, 58°C for 1 min, and 72°C for 1 min. Conditions for amplification of the *uidA* gene for 30 cycles were: 94°C for 1 min, 61°C for 1 min, and 72°C for 1 min. The PCR products were separated on 1% agarose gels and stained with ethidium bromide.

182 Induction methods

For induction by a single pulse, we used a combination of 183 the root drench/vapor method. Four milliliters of the in-184 ducer solution (ethanol, acetaldehyde, or 2-butanone) were 185 applied directly to the solid medium in a sealed Magenta 186 box. Poplar tissues were harvested for the GUS activity 187 assays 5 days after induction. For continuous, long-term 188 induction, transgenic plants in the Magenta boxes were 189 treated with ethanol vapors for 30 days by dispensing 4 ml 190 of 2% ethanol into a small, open container on the bot-191 tom of the Magenta box; the ethanol was changed every 192 5 days. 193

In the greenhouse, 200 ml of aqueous solution of the 194 inducer was drenched into the soil (in a pot with an ap-195 proximate volume of 1.5 l). Pots and 3-month-old poplar 196 plants were then enclosed into sealed light-transparent plas-197 tic bags together with separate trays containing 200 ml of 198 the inducer solution for vapor induction. The inducer solu-199 tion in the tray was changed every 3 days. Soil in pots was 200 drenched with 200 ml of the inducer solution with a 3 days 201 202 interval during a 10-day induction period.

203 GUS activity assays

For histochemical GUS staining, tissues were incubated for 6–12 h in 2 mM of 5-bromo-4-chloro-3-indolyl- β -Dglucuronic acid (X-Gluc) solution at 37°C essentially as described (Weigel and Glazebrook 2002). After staining, β-Glucuronidase activity in vitro was determined in 215 triplicates using fluorometric assay and the GUS sub-216 strate 4-methylumbelliferyl β -D-glucuronide (4-MUG) 217 essentially as described at: http://www.markergene. 218 com/product_sheets/pis0877.pdf. GUS activity was calcu-219 lated as a mean of measurements from leaves (top, middle 220 and bottom) of three individual plants until stated other-221 wise. The release of 4-methyl umbelliferone (4-MU) was 222 measured using DynaQuant fluorometer (Amersham Bio-223 sciences Corp., Sunnyvale, CA) or Wallac Victor²1420 224 Multilabel counter (Perkin Elmer, Boston, MA) at 15-, 30-, 225 and 75-min time points. To determine the total protein con-226 centration and released 4-MU in the GUS quantification 227 assays, calibration curves were developed using Bradford 228 protein assay (Bradford 1976) and standard 4-MU dilu-229 tions, respectively. 230

Results

Populus transformation

Two genetically distinct Populus clones-717-1B4 and 233 353-38—were used for Agrobacterium transformation. The 234 T-DNA of pJH0143 vector (Syngenta Biotechnology, Inc., 235 Research Triangle Park, NC) carries alcohol-inducible 236 switch cassette 35S::alcR-alcA::uidA::t35S similar to de-237 scribed in Sweetman et al. (2002). The expression of AlcR 238 alcohol sensor is driven constitutively by cauliflower mo-239 saic virus (CaMV) 35S promoter (Franck et al. 1980). The 240 uidA reporter gene is transcriptionally fused to alcA pro-241 moter (Kulmburg et al. 1992) (Fig. 1). 242

A total of 33 independent kanamycin-resistant lines 243 (transformation events) yielded a PCR product of the ex-244 pected size for both *nptII* and *uidA* (Table 1). To confirm 245 the integrity of the T-DNA in each line we further used 246 PCR primers bracketing the *alcA::uidA* junction. We am-247 plified a product of the expected size (490 bp) from 15 248 transgenic events in clone 717 and 5 events in clone 353 249 (approximately 60% of all lines tested). 250

Alcohol inducibility

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A total of 20 selected events were subjected to testing 252 for ethanol induction of GUS activity by using the root 253 drench/vapor combination, as described in the section "Ma-254 terials and methods". Only some of PCR-positive events 255 showed β -glucuronidase (GUS) activity upon induction 256 with 2% ethanol. Induction was observed in 60% (9 out 257 of 15) of the lines produced in clone 717 and 20% (1 out 258 of 5) of those in 353-38. Ten out of 15 lines confirmed via 259

 Table 1
 Transformation frequencies and proportion of ethanolinducible independent poplar events transformed with pJH4103

Poplar clone	717-1B4	353-38
Total number of explants	1124	704
Explants rooted on kanamycin medium	107	25
PCR positive events		
509 bp (base pairs) <i>uidA</i> and 253 bp <i>nptII</i>	25	8
fragments		
490 bp <i>palcA::uidA</i> fragment	15	5
Number and frequency (%) of lines with	9 (60)	1 (20)
alcohol-inducible GUS activity		

PCR to contain the intact *alcA*::*uidA* junction also tested positive for the GUS activity (Table 1).

To ensure that there is no basal alcA promoter leakage,

the clones of all 10 selected lines were mock-induced with

water and histochemically stained for GUS. No consis-

tent GUS staining background (except rare occurrences of

weak, sporadic spots) was detected in all negative controls
(Fig. 2, data not shown) suggesting that the *alc* system is
tightly regulated in poplar. Three lines (the clones of events
191, 173 and 183) showing ethanol-inducible GUS staining
were selected for further histochemical and fluorometric
evaluation of GUS activity.266
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Distribution of GUS activity in ethanol-induced tissues

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Six-week-old poplar plantlets propagated from the primary transformants were treated with chemical inducers using two methods. In the vapor induction method, the inducer was poured into a small open container and then placed into the sealed Magenta boxes. For the root drench/vapor method, sterile inducer solution was poured directly onto the tissue culture medium.

Plants induced by 2% ethanol (v/w) using the root drench/vapor combination showed GUS activity in both leaf and root tissues (Fig. 2). Intense GUS staining was ob-



Fig. 2 Histochemical staining for GUS activity in tissues of transgenic *Populus* after induction by ethanol. **a** Control transgenic plantlet stained for GUS activity after mock treatment with water. **b** GUS staining of transgenic poplar induced by 2% ethanol. **c** Young leaf showing a gradient of GUS activity which is stronger in apical part and weaker at the leaf base. Direction of a gradient is indicated by *dashed arrow*. **d** GUS staining of emerging apical leaf (AL). GUS activity is evident only in apical portions of young and emerging

leaves. Petiole (P) and leaf base have limited GUS staining. **e** A close-up view of a mature leaf showing intense GUS staining in vascular bundles (V), hydathodes (H), and trichomes (T). **f** A close-up view of an adventitious root. GUS staining is evident in root vasculature and lateral roots (LR). All poplar plantlets (event 191, clone 717) were treated with 2% ethanol using a combination of vapor and root drench methods in enclosed Magenta boxes for 5 days. Scale bars: 1 cm (**a**–**b**), 0.2 mm (**f**), and 0.5 mm (**e**)

served in leaf vascular bundles, hydathodes, trichomes, and
in parenchyma cells. Young and emerging leaves showed a
gradient of GUS activity, which was strongest at the apex
and weakest at the leaf base. GUS staining was evident in
adventitious and lateral roots and to a lesser extent in root
hairs. The stems of a majority of induced plants showed
weaker GUS activity.

290 Optimal ethanol concentration and duration of291 induction period

Clones of three independent transgenic events with high 292 (line 191) and moderate (lines 183 and 173) GUS response 293 were used to determine dose dependency of induction. 294 Plants in the sealed Magenta boxes were treated by var-295 ious concentrations of ethanol using combination of the 296 root drench and vapor methods for 5 days. Fluorometri-297 cally determined GUS activity was detected in the leaves 298 of line 173 at ethanol concentrations as low as 0.5% (v/v) 299 (Fig. 3 and Supplementary Material S1). GUS activity in-300 creased with increasing ethanol concentration up to 2% in 301 a dose-dependent manner, but further increases in ethanol 302 303 concentration led to a reduction in GUS activity. Lines 191 and 183 demonstrated generally similar response except 304 ethanol-induced GUS activity in leaves of line 191 was 305 higher than in lines 173 and 183 (Fig. 3 and S1). Con-306 trol transgenic plants where ethanol was substituted by an 307 equal volume of sterile water consistently showed negli-308 gibly low levels of GUS activity for all the three lines 309 suggesting that alc system has a minimal leakage in poplar 310 transgenics. 311



Fig. 3 Dose dependency of ethanol induction. Poplar transgenics (event 173, 717-1B4) were induced with varying concentrations of ethanol for 5 days using combination of vapor and root drench methods in tissue culture conditions. Ctrl – control (transgenic plants mock-induced with water) (shown by filled bars). GUS activity of ethanol-treated plantlets is shown by *open bars. Numbers* indicate ethanol concentration. GUS activity was measured in triplicate in three leaves (harvested from lower, middle and top position of a single transgenic plantlet. *Error bars* indicate standard error of the mean



Fig. 4 Time course of continuous induction by ethanol vapor. Transgenic poplar plants (event 191, 717-1B4) were treated continuously for 30 days by ethanol vapor under *in vitro* conditions as described in the section "Materials and methods". GUS activity at each time point (5, 10, 15, 20, 25 and 30 days post-induction) represents a mean of measurements in the leaves harvested from three individual plantlets. *Error bars* indicate standard error of the mean. Ctrl – water-treated control

Continuous exposure to ethanol vapors resulted in a significant increase in induced GUS activity in the leaves of *alcA::uidA* plants, for up to 25 days, at which time GUS activity was approximately 10-fold higher than at day 5 (Fig. 4).

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Efficiency of various induction agents

The efficiency of *alcA* promoter activation was compared 318 for three chemical inducers: ethanol, Ac, and 2-B (Fig. 319 5). GUS activity was determined in leaf and root tissues 320 of transgenic poplars induced with 2% ethanol, 0.1% 321 Ac, or 2% 2-B. Ac and 2-B were chosen because both 322 compounds have been shown to activate the *alcA* response 323 system in transgenic potato (Junker et al. 2003). Ac was 324 also reported to be an effective tissue-specific activator of 325 the *alcA* promoter in tobacco (Schaarschmidt et al. 2004). 326 Acetaldehyde was used at a concentration that was 20-fold 327 lower than ethanol because concentrations higher than 328 0.1% consistently damaged in vitro-grown poplar plantlets 329 (data not shown). To achieve the simultaneous induction 330 of both roots and shoots in tissue culture, we used the 331 root drench-vapor combination described in the section 332 "Materials and methods". 333

Fluorometric determination of GUS activity at 5 days 334 postinduction (dpi) revealed that ethanol was the most ef-335 ficient inducer of the *alc* system in both leaves and roots, 336 when compared to Ac and 2-B (Fig. 5). GUS expression 337 was lower in roots than leaves but significantly higher than 338 in the control plants. Induction by 2-B had only a marginal 339 effect on GUS activity both in leaves and roots. Induction 340 by 0.1% Ac yielded GUS activity that was 3-fold lower 341



Fig. 5 Response of the *alc* system in *Populus* transgenics to different chemical inducers. GUS activity in the leaf and root tissues of poplar plantlets was induced using 2% 2-butanone (2-B), 0.1 % acetaldehyde (Ac), or 2% ethanol (EtOH). A combination of vapor and root drench methods was used for induction of propagated transgenic clones (event 191) during 5 days in tissue culture. GUS activity at each time point was measured in leaf tissues of three individual plantlets. Ctrl – control. *Error bars* denote standard error of the mean

in leaves, than the response seen for ethanol. GUS levels
induced by both ethanol and Ac in roots were significantly
higher than in water-treated control, however, the difference between ethanol and Ac induction was marginal. The
weaker induction of GUS activity by Ac vapors was consistent with weaker histochemical GUS staining (data not
shown).

349 Induction efficiency under greenhouse conditions

In a pilot study, we simultaneously tested the effect of a root 350 drench and vapors on *alc* induction in a greenhouse envi-351 ronment. For the latter treatment, plants in pots with soil 352 were watered with inducer solution enclosed with an open 353 tray containing the same inducer solution in a plastic bag 354 as described in the section "Materials and methods". Un-355 der these conditions, 2.5% ethanol was the most effective 356 inducer after 10 days of treatment. Again, Ac was a weaker 357 inducer at a concentration of 0.2% (Fig. 6). An Ac concen-358 tration of 2% was phytotoxic for both the root drench and 359 root drench/vapor methods (data not shown). Induction by 360 2% (v/v) 2-B had a marginal effect on the GUS expression. 361 No significant GUS activity was detected in the roots of 362 the greenhouse-induced transgenic trees when using root 363 drench method for all the three inductive agents. 364

When young trees (approximately 3-month-old) were treated continuously for 10 days using the root drench/vapor combination with aqueous solutions of 2.5% ethanol, 0.2% Ac, and 2% 2-B, there were no obviously detrimental effects on the plant growth and development. Apart from the short-term effects on shape due to confinement in tight plastic bags, both the control and the treated transgenic



Fig. 6 The *alc* system induction in transgenic poplar under greenhouse conditions. GUS activity in leaf and root tissues of poplar was induced with 2% 2-butanone (2-B), 0.2 % acetaldehyde (Ac), or 2.5% ethanol (EtOH). A combination of vapor and root drench methods was used for 10 days of continuous treatment as described in Materials and Methods. Three-month-old trees in pots with soil were treated by inducers in sealed bags as described in the section "Materials and methods". GUS activity was measured in the leaves and roots of the three cloned plants of transgenic event 191. Ctrl – control. *Error bars* denote standard error of the mean

trees appeared healthy when examined up to 40 days after the cessation of treatment (Fig. 7). 373

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Discussion

We investigated the function of an alcohol-responsive gene 375 induction system in transgenic Populus. Using multiple in-376 dependent transformation events in two different genetic 377 backgrounds we studied the efficiency, optimal concen-378 trations, and induction time. Under the conditions tested, 379 ethanol was the most effective inducer; both Ac and 2-380 B were weaker inducing agents. Even though the condi-381 tions of treatment (i.e., inducer concentration, application 382 method, and induction period) differed for poplar this con-383 clusion was generally consistent with the results seen for 384 potato (Junker et al. 2003). At concentrations higher than 385 0.1 and 0.2% (v/v), Ac was detrimental to plants both in 386 the tissue culture and in the greenhouse, causing obvious 387 tissue damage and/or necrosis. 388

Ac was reported to be an effective temporal local inducer 389 of GUS activity in tobacco roots at low concentrations 390 (0.05% or less) (Schaarschmidt et al. 2004), and activated 391 alc gene expression in potato tubers more rapidly and at 392 lower concentrations than did ethanol (Junker et al. 2003). 393 Therefore, Ac could be an inducer of choice in applications 394 requiring rapid or localized response such as the expression 395 or suppression of genes regulating the key aspects of 396 poplar development, particularly during regeneration or 397 in localized tissues after infiltration. Further, optimization 398 of Ac delivery to the root or leaf tissues may leads to the 399 Fig. 7 Recovery of poplar trees after chemical induction under greenhouse conditions. Picture shows the same transgenic trees used for determination of GUS activity in Fig. 6. Trees were grown for additional 40 days after termination of the 10-days induction period. No apparent changes in the plant size or morphology were evident in comparison with water-treated transgenic controls. 2-B - 2%2-butanone, Ac - 0.2%acetaldehyde, EtOH - 2.5% ethanol, and CTRL water-treated control



development of local *alc* induction system in poplar similar
to that described for tobacco (Schaarschmidt et al. 2004).
In contrast to Ac, ethanol, in concentrations up to 8%,
had no apparent effect on the *in vitro* or greenhouse-grown
plant. Detectable GUS activity was induced by ethanol in

tissue culture at concentrations as low as 0.5%. Optimal
concentrations for maximal ethanol induction ranged from
2 to 4%. The basal expression of GUS driven by the *alcA*promoter was minimal in the absence of the inducer, suggesting tight control of the *alc* system in poplar.

Induction appeared to be stronger in plants in tissue cul-410 ture than in those grown in the greenhouse, possibly be-411 cause of the limited exposure of cells in mature tissues to 412 the inducer. Inefficient induction of roots was likely the re-413 sult of low rates of inducer diffusion into the soil. A similar 414 problem was encountered for the developing potato tubers, 415 where the response time of the *alc* system was improved 416 by substituting ethanol with Ac (Junker et al. 2003). 417

The tissue and organ distribution of the reporter gene 418 activity in various *Populus* tissues suggest that an efficient 419 alcohol induction can be achieved in most of the aerial 420 organs. Unfortunately, little to no induction was observed 421 in stems. This may be a limitation of this system, as Pop-422 *ulus* represents an excellent model system for the study 423 of wood formation. Therefore, a modification of this sys-424 tem may be needed. Xylem-specific promoters have been 425 described (Wu et al. 2000) and tissue-specific alcohol in-426 ducible expression has already been demonstrated (Maizel 427 and Weigel 2004; Grandjean et al. 2004). 428

The suitability of different inducible systems for genomics and other technological applications depends on a number of factors. These include: responsiveness; expression levels in transgenics; basal, uninduced expression; toxicity; and environmental concerns for various inducing

chemicals (Padidam 2003). Histochemical staining showed 434 low background GUS activity in the absence of ethanol for 435 the majority of transformation events. Further, the fluoro-436 metric GUS quantification in the leaves of three selected 437 independent lines consistently confirmed low basal levels 438 of enzyme in the absence of inducer. The tight regulation 439 of the *alcA* promoter in poplar was consistent with the re-440 sults reported for a wide range of plant species, including 441 Arabidopsis (Roslan et al., 2001), tobacco (Salter et al., 442 1998), potato (Sweetman et al., 2002; Junker et al., 2003), 443 oilseed rape (Sweetman et al. 2002), and tomato (Garoosi 444 et al. 2005). Our results demonstrate that the alcohol-445 inducible system is highly efficient in *Populus*; simple, 446 with negligible basal expression; and low toxicity both for 447 the plant, the researcher, and the environment. The latter 448 is particularly important in *Populus*, where large physical 449 size requires testing in greenhouse and field conditions, 450 where confinement of hazardous inducing agents would be 451 problematic. 452

Chemically inducible gene switches, including this 453 alcohol-inducible system, could be especially valuable for 454 identifying and studying the functionality of candidate 455 genes. Overexpressing or silencing genes regulating key 456 developmental events could be lethal to the modified plants 457 or preclude regeneration altogether. Further modifications 458 of the *alcR-alcA* system by substituting 35S with the pro-459 moter from *LEAFY* allowed for inducible expression of the 460 GUS (Maizel and Weigel 2004) and green fluorescent pro-461 tein (Grandjean et al. 2004) reporter genes in specific cell 462 types during development of shoot apical meristems of Ara-463 *bidopsis*. The *alcR-alcA* system was also used successfully 464 to induce transient gene silencing via RNA interference 465 (Chen et al. 2003; Lo et al. 2005). The *alcR-alcA* appears to 466 be a sensitive and robust inducible system with promise for 467

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