A composite transcriptional signature differentiates responses towards closely related herbicides in *Arabidopsis thaliana* and *Brassica napus*

Malay Das · Jay R. Reichman · Georg Haberer · Gerhard Welzl · Felipe F. Aceituno · Michael T. Mader · Lidia S. Watrud · Thomas G. Pfleeger · Rodrigo A. Gutiérrez · Anton R. Schäffner · David M. Olszyk

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**Abstract** In this study, genome-wide expression profiling based on Affymetrix ATH1 arrays was used to identify discriminating responses of *Arabidopsis thaliana* to five herbicides, which contain active ingredients targeting two different branches of amino acid biosynthesis. One herbicide contained glyphosate, which targets 5-enolpyruvylshikimate-3-phosphate synthase (EPSPS), while the other four herbicides contain different acetolactate synthase (ALS) inhibiting compounds. In contrast to the herbicide containing glyphosate, which affected only a few transcripts, many effects of the ALS inhibiting herbicides were revealed based on transcriptional changes related to ribosome biogenesis and translation, secondary metabolism, cell wall modification and growth. The expression pattern of a set of 101 genes provided a specific, composite signature that was distinct from other major stress responses and differentiated among herbicides targeting the same enzyme (ALS) or containing the same chemical class of active ingredient (sulfonylurea). A set of homologous genes could be identified in *Brassica napus* that exhibited a similar expression pattern and correctly distinguished exposure to the five herbicides. Our results show the ability of a limited number of genes to classify and differentiate responses to closely related herbicides in *A. thaliana* and *B. napus* and the transferability of a complex transcriptional signature across species.

**Keywords** Differentiation of herbicides · Gene expression signature · Transfer from model to crop plant · Acetolactate synthase · 5-enolpyruvylshikimate-3-phosphate synthase

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M. Das · J. R. Reichman · L. S. Watrud · T. G. Pfleeger · D. M. Olszyk
National Health and Environmental Effects Research Laboratory, Western Ecology Division, U.S. Environmental Protection Agency, Office of Research and Development, Corvallis, OR 97333, USA

G. Haberer
Institute of Bioinformatics and Systems Biology, Helmholtz Zentrum München, German Research Center for Environmental Health, 85764 Neuherberg, Germany

G. Welzl
Institute of Developmental Genetics, Helmholtz Zentrum München, German Research Center for Environmental Health, 85764 Neuherberg, Germany

F. F. Aceituno · R. A. Gutiérrez
Departamento de Genética Molecular y Microbiología, Facultad de Ciencias Biológicas, Pontificia Universidad Católica de Chile, Santiago, Chile

M. T. Mader
Institute of Stem Cell Research, Helmholtz Zentrum München, German Research Center for Environmental Health, 85764 Neuherberg, Germany

A. R. Schäffner
Institute of Biochemical Plant Pathology, Helmholtz Zentrum München, German Research Center for Environmental Health, 85764 Neuherberg, Germany

e-mail: schaeffner@helmholtz-muenchen.de

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Introduction

Herbicides contain structurally diverse chemicals designed to inhibit targets in susceptible weeds, and, thus, help reduce crop yield losses or assist the selective growth of genetically modified plants. In general, important plant-specific processes, such as photosynthesis and the biosynthesis of cell walls, lipids and amino acids are the primary targets of these chemicals (Anderson 1996). Inhibitors of acetolactate synthase (ALS) have become one of the most important herbicide groups because of their large spectrum weed control activity, high crop selectivity, low application rates and low mammalian toxicity (Zhou et al. 2007). ALS is the first enzyme in the biosynthesis of the three branched chain amino acids—leucine, isoleucine, and valine. Several structurally divergent herbicide chemical families including sulfonylurea, imidazolinone, triazolopyrimidine, pyrimidinyl-thiobenzoate, and sulfonylamino-carbonyltriazolinone compounds possess ALS as their common primary target. Glyphosate is another amino acid biosynthesis inhibiting herbicide widely used on both grass and broadleaf weeds. It targets 5-enolpyruvyshikimate-3-phosphate synthase (EPSPS) involved in the biosynthesis of the aromatic amino acids phenylalanine, tyrosine, and tryptophan (Duke and Powles 2008). Glyphosate is widely used as a herbicide also in conjunction with genetically modified Roundup-Ready™ crops (http://monsanto.com).

A number of studies involving transcriptional analyses have been undertaken to understand the primary mode of action, detoxification mechanisms or diagnostic detection of different herbicides. Reverse transcription-PCR analysis in Arabidopsis thaliana identified three glutathione-S-transferases (GST) (AtGSTF2, AtGSTU1, AtGSTU24) and two isoforms of 12-oxophytodienoate reductase (OPR1, OPR2), which are putatively involved in the metabolism of chloroacetanilide herbicides and explosive compounds (Mezzari et al. 2005). Kelley et al. (2004, 2006) analyzed GH3, an auxin-regulated gene, as a potential candidate to assess and diagnose differential soybean injury caused by various plant growth regulator herbicides. Plant responses to various chemicals used as herbicides, herbicide-additives, or inhibitors have also been studied in Arabidopsis thaliana at the transcriptome level by using array hybridization technologies (affymetrix.arabidopsis.info/narrays/experiment browse.pl; Glombitza et al. 2004; Madhou et al. 2006; Manfield et al. 2004; Raghavan et al. 2005, 2006; Manabe et al. 2007). Glombitza et al. (2004) used a focused DNA array comprised of only 267 target sequences related to secondary metabolism to determine the differential effects of bromoxynil, primisulfuron and prosulfuron herbicides and other biotic and abiotic stresses on A. thaliana. Similarly, a microarray containing 600 barley cDNAs was used to study the impacts of 2,4-dichlorophenoxyacetic acid (2,4-D), cinidion-ethyl and tribenuron-methyl on wheat transcriptomes under controlled environment and field settings (Pasquer et al. 2006).

Affymetrix ATH1 expression analysis of Arabidopsis suspension cultured cells habituated to cellulose synthesis inhibitor isoxaben, implicated novel genes in cell wall assembly (Manfield et al. 2004). Treatment with 1 mM 2,4-D for 1 h regulated Arabidopsis genes involved in auxin response (IAA1, IAA5, IAA13, IAA19), ethylene signaling (ERS, CTR1, AtERF4, AtERF8), as well as ABA biosynthesis (NCED3), signaling and response (Raghavan et al. 2005). A follow-up study using a range of 2,4-D concentrations from auxinic (1 μM) to herbicidal (1 mM) activity revealed that in contrast to the response to high 2,4-D concentrations, both ABA biosynthesis and signaling was repressed at auxinic concentrations (Raghavan et al. 2006). By studying the global transcriptome expression of imidazolinone-sensitive (wild-type) and imidazolinone-resistant (csr1-2D) mutant Arabidopsis plants, it has been observed that imazapyr did not significantly alter gene expression in csr1-2D mutants and hence inferred that imidazolinone herbicides act solely by targeting CSR1, a catalytic subunit of ALS (Manabe et al. 2007). In contrast, their time course analysis in wild-type plants showed a dynamic change in gene expression pattern with induction of mostly detoxification genes at early time points, while genes related to amino acid biosynthesis, secondary metabolites and rRNA were induced at a later stage.

In contrast to these analyses, which either allowed single compound—single plant species interactions and/or considered only the active ingredients as the test herbicides, we compared Arabidopsis transcriptional responses to five related herbicides and attempted to transfer results obtained in the model plant to a related crop species. To allow better herbicidal uptake by the plants (cells) comparable to realistic field situations, field applicable commercial formulations were used that contains inert and non-toxic coformulants in addition to the active compounds. It is required by e.g., US EPA regulations that tests for non-target plant phytotoxicity are to be conducted with the typical end-use product and not with the technical grade of the active ingredient only (Electronic Code of Federal regulations, ECFR 2009). Four ALS-inhibiting herbicides and a glyphosate containing herbicide were chosen, targeting two different branches of plant amino acid biosynthesis (Table 1). Among the selected ALS-inhibiting herbicides two had active ingredients from the same chemical family, which contained a sulfonylurea backbone (sulfometuron and primisulfuron). The other two ALS-inhibiting herbicides contained active ingredients from other chemical families with completely unrelated structures: the imidazolinones and triazolopyrimidines. The major objectives of our study were: (a) to study the impact
of these agronomically used herbicides on Arabidopsis transcriptome, (b) to identify a robust transcriptional signature in Arabidopsis that allows differentiating and classifying the response to closely related herbicides targeting the same enzyme (ALS), yet having different active compounds, or having even the same chemical backbones (sulfonylurea) and (c) to test if such a complex transcriptional signature identified in Arabidopsis could be transferred to a related, major agronomic species, Brassica napus. The latter objective might lay the groundwork for future research to apply gene expression signature in environmental risk characterization, such as non-target movement of herbicides. Moreover, it might also provide a general case of translational research from the model towards a related crop plant.

**Results and discussion**

Gene expression analyses in response to herbicides containing ALS- and EPSPS-inhibitors in A. thaliana

To identify effects of closely related herbicides at the gene expression level, Arabidopsis plants were treated with one of four ALS-inhibiting herbicides or a herbicide containing the EPSPS-targeting active ingredient glyphosate (Roundup Original™) at non-lethal concentrations producing an approximately 50% reduction in shoot dry weight (EC$_{50}$, Table 1). Two of the ALS-targeting herbicides Oust™ and Beacon™, have active ingredients sulfometuron and primisulfuron respectively, with similar chemical structures derived from a sulfonylurea backbone. The other ALS-targeting herbicides contain completely unrelated active ingredients; Arsenal™ with the imidazolinone chemical imazapyr, and FirstRate™ with the triazolopyrimidine chemical cloransulam. Altered transcript levels in response to the herbicides were analyzed using Affymetrix ATH1 GeneChips. The array expression analyses were independently verified by quantitative real-time reverse transcription polymerase chain reaction. Although the magnitude of changes varied in a few cases between the two techniques, the majority (54/62) of cases either confirmed the directionality of change or verified the lack of a significant alteration (Table S1).

**Table 1  Characteristics of the five herbicides used in Arabidopsis thaliana (Col-0) and Brassica napus (Westar) gene expression studies**

| Trade name       | Manufacturer                                  | Mode of action | Chemical family | Active ingredient (a.i.) |
|------------------|-----------------------------------------------|----------------|----------------|-------------------------|
| Arsenal™         | BASF Corporation, Research Triangle Park, NC, USA | ALS inhibitor | Imidazolinone   | Imazapyr               |
| Beacon™          | Syngenta Crop. Protection Inc., Greensboro, NC, USA | ALS inhibitor | Sulfonylurea    | Primisulfuron-methyl    |
| FirstRate™       | Dow AgroSciences LLC, Indianapolis, IN, USA   | ALS inhibitor | Triazolopyrimidine | Cloransulam-methyl |
| Oust™            | E.I. du Pont de Nemours and Company, Wilmington, DE, USA | ALS inhibitor | Sulfonylurea    | Sulfometuron methyl    |
| Roundup original™ | Monsanto Company, St. Louis, MO, USA       | EPSPS inhibitor | –              | Glyphosate             |

Common and distinct gene expression patterns induced by ALS- and EPSPS-inhibiting herbicides in A. thaliana

Only sixteen genes were significantly affected by Roundup Original™ application (Table S2), while 478 genes were significantly and coordinately regulated by all four ALS-inhibiting herbicides at the EC$_{50}$ concentration (Table S3). In our study eight Roundup Original™ induced genes were
linked to secondary metabolism and defense of which seven were also induced by ALS-inhibitors. All of them were potentially related to herbicide metabolism and detoxification, e.g., CYP71A13, UGT74E2, ABC transporter gene ATPDR12 and two MATE transporter genes (Table S2).

Among the 478 genes responsive to ALS-inhibiting herbicides, 82 were down-regulated and the remaining 396 were up-regulated. According to TAIR classifications, genes globally classified as cellular biosynthetic processes were enriched among differentially expressed genes (Table 2). The significant enrichment of included subcategories was more instructive. An effect on protein biosynthesis was noticed such as genes encoding ribosome associated proteins or protein synthesis initiation (eIF4A, eIF4E and eIF5), which were mostly up-regulated. Amino acid metabolic process was also found enriched, but at a lower stringency level ($P$-value 0.015). Only one gene (threonine ammonia lyase, AT3G10050) related to the BCCA group was induced along with six other genes related to amino acid biosynthesis. Other enriched categories included indole derivative metabolic process, camalexin metabolic process, and response to temperature, e.g., members of cytochrome P450 monoxygenases (CYP), GST and UDP-glycosyltransferases (UGT), which are also involved in plant secondary metabolism (Wagner et al. 2002; Messner et al. 2003; Brazier-Hicks et al. 2007). Some of the most highly induced genes were members of these families with unknown substrates such as CYP89A2, CYP81D8, CYP91A1, GSTU9, GSTU24, GSTZ1, UGT74E2, UGT76B1, UGT87A2 (Table S3). They could be related to the metabolism of ALS-inhibiting herbicides irrespective of their chemical structure.

Changes in the expression of genes involved in central energy pathways were also observed. A number of genes encoding components of mitochondrial transporters indicated enhanced organellar membrane exchange and metabolic activity. The up-regulation of two mitochondrial alternative oxidases ATAOX1a (AT3G22370) and ATAOX1b (AT3G22360), along with two mitochondrial NADH dehydrogenases, NDB2 (AT4G05020) and NDB4 (AT2G20800) point to a specific, concerted action

### Table 2 Functional classification according to TAIR ontology

| Functional category                  | Observed out of 477* | Observed frequency (%) | Expected out of 21,947 | Expected frequency (%) | $P$ value |
|-------------------------------------|----------------------|------------------------|------------------------|------------------------|-----------|
| Cellular biosynthetic process       | 57                   | 11.90                  | 1,304                  | 5.90                   | 8.87E-05  |
| Organelle organization and biogenesis | 34                  | 7.10                   | 652                   | 3.00                   | 5.80E-04  |
| Translation                         | 29                   | 6.10                   | 556                   | 2.50                   | 3.18E-03  |
| Ribonucleoprotein complex biogenesis and assembly | 23              | 4.80                   | 201                   | 0.90                   | 2.04E-08  |
| Ribosome biogenesis and assembly    | 20                   | 4.20                   | 122                   | 0.60                   | 4.48E-10  |
| RNA processing                      | 16                   | 3.40                   | 205                   | 0.90                   | 2.52E-03  |
| tRNA processing/metabolic process   | 9                    | 1.90                   | 25                    | 0.10                   | 3.31E-07  |
| Indole and derivative metabolic process | 7                  | 1.50                   | 30                    | 0.10                   | 6.3E-04   |
| Camalexin metabolic process         | 3                    | 0.60                   | 4                     | 0.00                   | 8.83E-03  |
| Response to temperature stimulus    | 19                   | 4.00                   | 291                   | 1.30                   | 5.02E-03  |
| Syncytium formation                 | 4                    | 0.80                   | 10                    | 0.00                   | 9.17E-03  |

478 genes regulated similarly by four ALS-inhibiting herbicides were tested for over-representation against Arabidopsis thaliana ATH1 (23,334 genes) background population by using BioMaps tool at VirtualPlant (http://virtualplant.bio.nyu.edu/cgi-bin/vpweb2). Categories enriched at $P < 0.01$ are listed here. The method used to calculate $P$ value of over-representation was hypergeometric distribution

* One gene was not assigned and hence total number of observed genes was reduced to 477
Fig. 1 Composite signature of transcriptional responses of 101 marker genes to herbicide treatments and selected stresses. a Clustering of the responses to five herbicide treatments (Table S4). Group 1 marker differentiates all ALS-inhibitors from EPSPS-inhibitor and group 2 marker differentiates among four ALS inhibitors and towards EPSPS-inhibitor. The TIGR/TAIR 6 genome annotation code was used as the gene identifier. The two top correlated stress response patterns with each herbicide (Table S8) were assorted to the gene matrix of panel A showing a signature clearly distinct from all herbicides in all seven cases. Red color indicates up-regulation of transcripts in treatment group compared to control; blue indicates down-regulation and black indicates no change. Abbreviations: ARS [Arsenal™, imazapyr active ingredient (a.i.)], BEA (Beacon™, primisulfuron a.i.), FIR (First-Rate™, cloransulam a.i.), OUS (Oust™, sulfometuron a.i.), ROU (Roundup Original™, glyphosate a.i.), Nor norflurazon, UV-B ultraviolet B, P infest: Phytophthora infestans; P syrin p: Pseudomonas syringae pv. phaseolicola; P syrin t: Pseudomonas syringae pv. tomato avr Rpm1; Met jas: methyl jasmonate
to prevent oxidative damage (Table S3). A similar co-expression pattern of ATAOX1a and NDB2 had previously been observed in response to imidazolinone (Manabe et al. 2007). The treatment with sulfonylurea and imidazolinone herbicides triggered induction of AOX enzyme activity in sycamore cells, while no such change was observed with glyphosate treatment (Aubert et al. 1997).

Molecular signature for differentiating herbicide responses

Besides providing insight regarding their molecular effects, herbicide-specific changes in gene expression provided an opportunity to analyze whether the combined expression signature of any genes could establish a collective molecular marker for differentiating responses among closely related herbicides. Metabolite profiling has been useful to diagnose herbicide treatments on plants (Aranibar et al. 2001; Sauter et al. 1988). However, metabolites may differ depending on the actual enzymatic activities in different plants or species and measuring mRNAs is much more standardized than quantifying metabolite levels. Therefore, identifying mRNA markers to develop a signature that is sensitive enough to classify even related herbicides that have either similar chemical backbone or a common primary target would be very useful to diagnose herbicide treatments on plants. To identify such a signature, we used stringent statistical thresholds to obtain a set of gene transcripts that could characterize each herbicide response (Fig. 1a, Table S4, “Materials and methods”). First, a change larger than twofold in at least one experiment with opposite or no response in at least one other treatment was required. Besides differentiating the herbicide responses, this measure also antagonized the selection of general stress responsive genes. Second, genes had to be well expressed in control experiments in order to select reliable
alterations in transcription, although some potent markers may be eliminated thereby. A total of 101 candidate marker genes were selected and classified into two major groups. Group 1 contained 46 candidate genes that differentiated the ALS-inhibiting group from the EPSPS group, of which 34 were up-regulated and 12 were down-regulated. Group 2 contained 55 candidates that differentiated the responses among the four ALS-inhibiting herbicides, as well as towards glyphosate. Thirteen different combinations of gene expression patterns were observed for group 2 markers (Fig. 1a, Table S4).

Cluster analysis revealed a close similarity between Beacon™ (a.i. primisulfuron) and FirstRate™ (a.i. cloransulam) mediated transcriptional alteration. Arsenal™ (a.i. imazapry) also was associated with this pair of herbicides, while Oust™ (a.i. sulfometuron) was not. Thus, surprisingly, the effects of two SU herbicides (Beacon™ and Oust™) on gene transcription could be differentiated. This analysis suggested that the composite signature from these 101 genes could differentiate the plant’s responses towards herbicides that even possess structurally related active compounds.

Testing the validity of the identified molecular signature

To test the potential of these diagnostic genes as a marker set of herbicide exposure in Arabidopsis, we first calculated correlations of individual herbicide experiments. Correlations between individual samples of different treatment groups were, except for FirstRate™ versus Beacon™, considerably lower than within group correlations (Table S5). Roundup Original™ did not show a correlation to any of the other treatment groups. To test the reproducibility of independently conducted experiments as well as a prediction of how well any single experiment would identify the correct herbicide signature we employed a classification strategy. Diagonal discriminant analysis has been tested useful in gene expression data (Dudoit et al. 2002). Therefore, this method was applied to our data set. The performance of the classification to the herbicide treatments was assessed by a leave-one-out cross-validation. The cross-validation rate was 100%, i.e., each 101-gene-signature of the biologically independent replicates was correctly classified to the corresponding herbicide group (Fig. 2a). Hence, the global similarity of FirstRate™ and Beacon™ as measured by their high correlations (Table S5) was resolved by using the supervised learning algorithm. These results confirmed the
high reproducibility of the independent replicates as well as the distinctive properties and robustness of the signature provided by the 101 marker set.

Distinguishing herbicide responses from other chemical and stress treatments

To compare the expression pattern of the identified molecular markers for herbicides with other major stress responses, expression data were collected from quality-controlled Affymetrix ATH1 microarray experiments for a wide range of abiotic, biotic and plant growth regulators stresses. In particular, the herbicide marker expression pattern was compared to a broad range of chemical compounds that inhibit diverse physiological processes, such as photosystem I (methyl viologen), photosystem II (PN08), carotenoid biosynthesis (norfurazon), gibberellin biosynthesis (prohexadione, uniconazol, paclobutrazol, propiconazol), auxin transport (TIBA, NPA) and brassinosteroid biosynthesis (Brz220) (Table S6, “Materials and methods”). All selected experiments had been performed using leaves or seedlings of the Col-0 genetic background similar to this study. In addition, 72 out of 101 marker genes displayed a constitutive expression across four developmental stages (seedling until rosette) that cover all these experiments (Table S7).

Determination of Pearson correlation coefficients by comparing the expression patterns altered by herbicides and diverse stress treatments clearly demonstrated that expression data from none of the other stress experiments were similar to the studied herbicide treatments. Herbicide-to-herbicide correlations were significantly better in all cases (Tables S5, S8). Even an outlier herbicide replication for herbicide correlations were significantly better in all cases similar to the studied herbicide treatments. Herbicide-to-sion data from none of the other stress experiments were diverse stress treatments clearly demonstrated that expres-

Correct classification of B. napus herbicide exposure even based on a reduced set of genes

In total, 23 B. napus genes were identified that exhibited a similar response pattern to the herbicides like that of their homologous Arabidopsis genes. Since only three groups (groups 2c, 2h, 2i) out of fifteen different expression patterns in the original 101 marker set were not represented, it was examined whether such a reduced set could still allow a correct classification of the herbicide responses. Indeed, a diagonal discriminant analysis applying only 23 Arabid-opis genes again separated the five herbicides and a 100% cross-validation rate was found by a leave-one-out cross-validation assessment (Fig. 2b). Next, the B. napus genes
Table 3 Comparative expression analyses of *A. thaliana* marker genes and *B. napus* best BLASTN hit ESTs

### a. Single BLAST hits

| AGI code | Best B. napus hit | E-values |
|----------|------------------|----------|
| AT1G5420 | EL592039         | 0.00E+00 |
| AT1G31970| TA12650_3708     | 0.00E+00 |
| AT5G4210 | CD89979          | 0.00E+00 |
| AT5G69890| TA17549_3708     | 0.00E+00 |
| AT2G9310 | TA15948_3708     | 0.00E+00 |
| AT3G16250| CN28678          | 0.00E+00 |
| AT5G6930 | TA13498_3708     | 0.00E+00 |
| AT2G3750 | TA18409_3708     | 0.00E+00 |
| AT5G28920| ES909818         | 2.00E-170|
| AT5G3610 | EV056117         | 0.00E+00 |
| AT1G24440| DY017660         | 0.00E+00 |
| AT5G51820| EL5920           | 0.00E+00 |
| AT3G14600| TA11777_3708     | 0.10E-177|
| AT1G28520| CD85294          | 0.10E-177|
| AT5G40440| DY007055         | 0.00E+00 |
| AT5G13630| TA12380_3708     | 0.00E+00 |
| AT5G18520| BN250771         | 1.00E-164|
| AT5G17170| EV096573         | 0.10E-164|
| AT2G9660 | EV102535         | 0.00E+00 |
| AT4G66970| EV023792         | 0.00E+00 |

### b. Multiple BLAST hits

| AGI code | Best B. napus hit | E-values |
|----------|------------------|----------|
| AT3G14600| TA17777_3708     | 0.00E+00 |
| AT4G14110| TA13545_3708     | 0.10E-177|
| AT5G52820| CD82595          | 1.00E-177|
| AT1G02410| CN360387         | 0.00E+00 |
| AT1G35880| CA16913_3708     | 0.00E+00 |
| AT1G77120| CD83163          | 1.00E-177|
| AT2G28970| TA16197_3708     | 0.00E+00 |
| AT4G22920| CD838267         | 0.00E+00 |

Changes in gene expression of herbicide treatment versus control are displayed as log2-values. Orange (>0.8) and red (>1) indicate induction, light blue (<-0.8) and blue (<1) highlight suppression; non-significant changes (−0.5 < ratio < 0.5) are labeled grey

Non-overlapping independent ESTs may represent 5'- and 3'-part of the orthologous gene. *Abbreviations* ARS (Arsenal™, imazapyr active ingredient (a.i.)), BEA (Beacon™, primisulfuron a.i.), FIR (FirstRate™, cloransulam a.i.), OUS (Oust™, sulfometuron, a.i.), ROU (Roundup Original™, glyphosate a.i.)

herbicide-dependent expression patterns were classified with these *Arabidopsis* data and each *Brassica* pattern was correctly associated with the corresponding herbicide pattern. Thus, a robust gene expression signature for the differentiation of herbicide responses could be developed in *A. thaliana* and *B. napus*.
Conclusions

In this study changes in gene expression were identified, that provided enhanced molecular insight regarding the effects of five herbicides targeting two different branches of amino acid biosynthesis. A set of 101 gene transcripts could be extracted from these analyses that provided a composite diagnostic signature to differentiate among Arabidopsis responses to related herbicides, which was clearly distinct from those provoked by other stress treatments and chemical compounds. The possible specificity of the diagnostic signature was corroborated by the finding that none of the herbicide-to-stress correlations reached the quality of herbicide-to-herbicide correlations. Interestingly, even herbicides with active ingredients having the same enzymatic target (ALS) or similar chemical structures (sulfonylurea) could be distinguished. However, using the commercial herbicide formulations precludes an unambiguous assignment of these differences due to the active chemicals only, since we cannot rule out that non-active ingredients used in the herbicide formula might have an effect on plant gene expression. On the other hand, in contrast to studies using pure active ingredients this work provided the opportunity to establish gene expression changes and marker signatures, which are associated with the formulated herbicides as they are used in agriculture and as they facilitate better access to the treated plant (cells). Furthermore, the successful transfer of a diagnostic composite transcript signature from the model plant into the crop species Brassica napus could be demonstrated—despite the triplicate nature of the Brassica genome and the lack of a complete genome or transcriptome sequence. In the particular case studied here a robust gene expression pattern has been established that allows the specific differentiation and classification of closely related herbicides. Its ability to classify the correct herbicidal exposures highlights future implications of such transcript-based signatures in environmental studies such as non-target movement of herbicides, or in a broader perspective in analogous analyses of the impact of chemicals on organisms.

Materials and methods

Herbicide treatment and isolation of RNA

Prior to carrying out gene expression studies, range-finding experiments were conducted to identify concentrations of the five herbicides that would produce similar growth responses in 14-day-old seedlings of Arabidopsis thaliana (accession Col-0) and Brassica napus cv. Westar (Table 1). Four of the herbicides were ALS-inhibitors. Two of them Oust™ and Beacon™, have active ingredients (a.i.), sulfofeturon and primisulfuron, respectively, with similar chemical structures derived from a sulfonyleurea backbone. The other ALS-targeting herbicides contain completely unrelated active ingredients; Arsenal™ with imazapyr, and FirstRate™ with cloransulam. The fifth herbicide Roundup Original™ contained an EPSPS-targeting active ingredient (glyphosate). A surfactant (Preference™, Agriliance, LLC, St. Paul, MN, USA) was added to each of the herbicide solutions as recommended on the label to enhance herbicide efficacy. An additional set of plants was sprayed only with the surfactant and used as a control. Plants were maintained at 22–23°C with a 16 h photo-period (175–210 μmol m⁻² s⁻¹).

For expression analyses plants were grown and treated with the concentration of each herbicide producing a 50% reduction in shoot dry weight (Table 1) or the surfactant solution as a control. Hence, the application of a comparable functional dose (EC₅₀) would allow direct comparison of herbicide regulated gene expression patterns in these two species. Three independent biological replicates (individual pots) were used per herbicide or control treatment. Leaves without visible symptoms were collected at 24 h from at least ten plants post treatment. Total RNA was isolated from approximately 200 mg of pooled leaf tissue using TRizol reagent (Invitrogen, USA), DNase treated and further purified with RNAeasy mini spin columns (Qiagen, USA).

RNA labeling, hybridization and scanning

RNA quality analyses (Agilent Bioanalyzer 2100, Agilent Technologies, USA) and array hybridizations were conducted at the Center for Genome Research and Biocomputing Core Laboratories of Oregon State University, USA. A. thaliana whole genome oligonucleotide microarray (ATH1, Affymetrix) were used for hybridization of each replicon and scanned with an Affymetrix GeneChip™ Scanner 3000.

Microarray data analysis

Array images were first visually screened following each scan to correct for signal artifacts, scratches or debris. Then the arrays were processed with MAS 5.0 algorithm to check for data quality based on the four important parameters: scale factor (<3), average background level (<100), percentage of genes present and ratios derived from 3′- and 5′-feature of housekeeping genes. CEL files were imported to ChipInspector (Genomatix, Germany) to check if dynamic range of any array deviates from the average. Data was normalized on the single probe level (Cohen et al. 2008). Each raw value of an array was adjusted to make the mean intensity the same for each individual array in an analysis. Probe-sets were defined
following Haberer et al. (2006) based on TIGR/TAIR 6 genome annotation release. Statistical algorithm for permuted t-test with false discovery rate (FDR) calculation was similar to SAM (Tusher et al. 2001).

Multi-class analysis was performed in ChipInspector (Genomatix, Germany) based on exhaustive matching combination within an experimental class. Statistical thresholds applied were: FDR-0, median number of falsely called features—0, chosen delta value for regulation—0.14 and minimum probe coverage—3. Transcripts having ≥ twofold (1.0 in log2) changes in abundance with P-values ≤ 0.05 (Tables S2, S3) were functionally categorized by MapMan 2.0.0 (Usadel et al. 2005) and TAIR (www.arabidopsis.org) gene classifications. The BioMaps tool at www.virtualplant.org was used to identify statistically over-represented groups based on TAIR classifications. Significance is based on P-values corrected for multiple testing by hypergeometric distribution (Table 2).

Stringent statistical cut-off criteria were applied to check whether transcript profiling could be used to identify markers for differentiating responses of *A. thaliana* to the applied herbicides. A change of ≥ twofold (1.0 in log2) in at least one treatment group and opposite or no deregulation in at least one other treatment group at *P* ≤ 0.05 was set as an initial cut-off level (Table S4). In addition, an average expression threshold value of 7.0 (MAS 5.0, normalized, log scale, target value 200) was introduced to exclude transcripts of low abundance and low reliability. Hierarchical cluster analysis (HCL) of the markers was conducted by average linkage agglomerative clustering using the Euclidean distance (Fig. 1a) according to Rainer et al. (2006).

Candidate markers genes were analyzed for other stresses responses across the ATH1 Affymetrix based microarray experiments available from the NASCArrays database as described previously (Aceituno et al. 2008). Expression data from 17 abiotic, 14 biotic, 8 plant growth stresses and 14 chemical compound treatments at the 2-leaved rosette to early flowering stage were selected (Table S6). The basal expression patterns of the markers were checked at www.genevestigator.ethz.ch (Table S7).

The similarity of the transcriptional response to herbicides and selected stresses was analyzed by Pearson correlation coefficients (Tables S5, S8). For the statistical analysis and classification of samples into treatment groups based on microarray data several methods have been suggested (e.g., diagonal discriminant analysis (DDA), support vector machines (SVM), and k-nearest neighbour). Misclassification rates for many discriminant methods have been compared using real or simulated data. Dudoit et al. (2002) found that DDA has smallest error rates. For typical data matrices as encountered with a small number of microarrays with numerous gene expression values the crucial methodological aspect is a stable estimation of the covariance matrix. In this situation DDA is a robust method, because only the diagonal of the covariance matrix (variances) is estimated.

The method diagonal discriminant analysis was trained with the gene marker sets given in Table S10 or in Table 3. Leave-one-out cross-validation (LOOCV) was performed to check the efficiency of the diagnostic markers in predicting the particular herbicide to which *Arabidopsis* plants had been exposed. It is a common choice for the small sample sizes usually encountered in microarray studies. Each sample is left out, one at a time. For each training set constructed in that way, i.e., 14 experiments in this project, a DDA was calculated and the discriminant function was used to predict the class of the left-out sample. This prediction was then counted as being correct or incorrect. By repeating this procedure for each sample the total number of classification errors was determined (Simon et al. 2003).

Quantitative real time RT-PCR (RT-qPCR) analyses

Fourteen *Arabidopsis* genes were selected for validation of microarray data by RT-qPCR using the same RNA samples as for the array experiments (Table S11). A QuantiTect SYBR Green RT-PCR kit (Qiagen, USA) was used for one-step real time RT-PCR amplification in an ABI prism 7700 (SDS 1.9.1, Applied Biosystems, USA). The herbicide-regulated expression of candidate *B. napus* homologs was also analyzed by RT-qPCR assay. Primers were designed based on EST sequences (Table S12). The real-time expression data were subjected to analysis by geNORM (Vandesompele et al. 2002), which identified ACTIN2 (*AT3G18780* and UBQ10 (*AT4G05320*) as the two most stably expressed genes in *A. thaliana*, while ACTIN7 (*AF111812*) and Actin (AY570244) for *B. napus*. Hence, ACTIN2 and ACTIN7 were used as a single reference gene for data normalization for *Arabidopsis* and *Brassica*, respectively. As a reciprocal control, we checked for the relative ACTIN2 expression also among the ATH1 analyses; indeed the gene proved to be highly stable with relative expression levels of 0.952–0.974 in case of all five herbicide treatments. Relative fold change in gene expression was estimated following the ΔΔ Ct method with the carriersprayed sample as calibrator (Livak and Schmittgen 2001).

Each value represents the mean of three biological replicates with two technical replicates each (Tables 3, S1).

Identification of homologous genes in *Brassica napus*

Candidate homologs between the *Arabidopsis* coding sequences (TAIR6 annotation; Poole 2007) and the TIGR
transcript assemblies of *Brassica napus* comprising 35,460 singletons and tentative consensus sequences were identified (Childs et al. 2007). In the absence of syntenic- or map-based genome data, candidate homologs between *Arabidopsis* and *Brassica* were defined as reciprocal best BLASTN hit (RBH; Hirsh and Fraser 2001; Haberer et al. 2006). BLASTN provides a higher selectivity by e.g., recognizing also synonymous changes in case of species from one family. In case of two or more reciprocal top scoring matches showing equal scores (*e*-values) between *Brassica* ESTs and an *Arabidopsis* gene model, we considered all top scoring matches for further analysis and designated such relations as ‘multiple’ (bi-directional best BLAST) hits.

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Author contributions

MD, DO, TF, JR, LW, and AS conceptualized the research; MD performed expression analyses in Arabidopsis and B. napus; JR assisted in expression analyses; DO and TP were responsible for herbicide treatments and related analyses; MD, TS, RG, FA, GH, MM, GW performed statistical and comparative data analyses, MD wrote the manuscript with contributions from AS, RG, and DO.