Coevolution of Two Sulfonylurea-Resistant Common Chickweed (Stellaria media) Biotypes with Different Mutations in the Acetolactate Synthase Gene

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Abstract

Common chickweed [Stellaria media (L.) Vill.] is an annual weed found in grain fields and pastures. This creeping weed produces a mat-like canopy that competes with crop seedlings for light and nutrients. This species is almost exclusively autogamous. Acetolactate synthase (ALS)-inhibiting herbicides are frequently used to control this broadleaf weed, and two mutations in the S. media ALS gene (Pro-197-Gln and Trp-574-Leu) have been reported to confer herbicide resistance. Seeds from several plants that survived an application of thifensulfuron-methyl/tribenuron-methyl (16 g ai ha−1) were collected from the same field in Quebec, Canada. Seedlings were grown, tested for the presence of mutations in the ALS gene, and treated with different rates of foramsulfuron, flumetsulam, and imazethapyr, each belonging to different families of ALS inhibitors. Two herbicide resistance–confering mutations were identified in this population. One biotype had the previously reported Pro-197-Gln, while a new mutation, Pro-197-Ser, was identified in different plants from the same population. The new mutation (Pro-197-Ser) confers a lower level of resistance to sulfonylureas than Pro-197-Gln, and both biotypes are susceptible to imidazolinone and triazolopyrimidines.

Introduction

Common chickweed [Stellaria media (L.) Vill.] is present in all but the most extreme habitats. This plant is native to Europe and is now nearly cosmopolitan, being found on every continent. Its range extends from Arctic and sub-Antarctic regions to warmer temperatures of the Equator, although it generally prefers cool temperatures (Turkington et al. 1980). Stellaria media is found from coast to coast in Canada and is more common in British Columbia and in eastern Canada than in the prairies (Turkington et al. 1980). The plant will flower and set seed throughout the year. The seeds produced will germinate at any time during the year when conditions are suitable; however, most germinate in early spring or late autumn. For that reason, plants of all ages can be found in the field. Turkington et al. (1980) reported that an S. media plants can produce from 500 to 2,500 seeds, whereas Lutman (2002) described reports of up to 15,000 seeds plant−1.

Stellaria media is cleistogamous, essentially homogamous, self-pollinated and self-fertilized (Mulligan and Findlay 1970; Mulligan and Kevan 1973; Salisbury 1974). Anther dehiscence takes place in the bud before the flower opens, thus ensuring self-fertilization under the most favorable circumstances with complete protection from the weather (Bennett 1869). This inbreeding mechanism limits the amount of genetic exchange passed on through reproduction. The capacity of the plant to adapt to a wide range of growing conditions is attributed to an intrinsic variability that is attested by its polymorphic characteristics (Salisbury 1974). A chromosome count of 20 (2n = 40) has been described, but counts of 28, 36, 40, 42, and 44 have also been reported (Turkington et al. 1980). Scholte (1978) reported that S. media is a allotetraploid (2n = 44) with a subsequent loss of chromosomes (Sobey 1981; Verkleij et al. 1980).

While there are some beneficial uses of S. media in maintaining soil structure and as animal feed, it is mostly considered a weed of grain fields and cultivated areas. The growth of young crop seedlings will be stifled by the mat-like growth of S. media canopy and competition for light and nutrients (Turkington et al. 1980). For example, although increasing the density of barley (Hordeum vulgare L.) planting in pots reduces losses due to S. media competition, the costs caused by the weed amount to nearly four-fifths of the grain yields in closely planted fields (Mann and Barnes 1950). When barley is more sparsely seeded, its growth may be reduced to less than 10% of its normal growth in the absence of the weed (Mann and Barnes 1950).
Stellaria media has the potential to release water-soluble compounds into its rhizosphere that may cause allelopathic interference to the seedling growth of wheat (Triticum aestivum L.) (Inderjit and Dakshini 1998). Cover crops (white mustard [Sinapis alba L.], radish [Raphanus sativus L.], common vetch [Vicia sativa L.], and mixtures) have been shown to reduce the plant density of S. media (Sturm et al. 2016).

Acetolactate synthase (ALS) inhibitors (WSSA Group 2 herbicides) are used to control S. media. Resistance to ALS inhibitors in S. media has been described and attributed to two mutations in the ALS gene in codons 197 and 574 (codon numbers according to the Arabidopsis thaliana ALS gene sequence [Genbank Accession AY124092]) (Kudsk et al. 1995; Marshall et al. 2010). The Pro-197-Gln mutation confers resistance to sulfonylureas, whereas the Trp-574-Leu mutation confers resistance to both sulfonylureas and triazolopyrimidines (Heap 2018). The mechanism of resistance is a conformational change in protein structure following amino acid substitutions at codons 197 and 574, which impairs herbicide binding while allowing for protein function to be maintained (McCourt et al. 2006). Other cases of Group 2 herbicide–resistant S. media have been reported in Canada, China, Denmark, Ireland, New Zealand, Norway, South Africa, and the United States with no mention of the mechanism conferring resistance (Heap 2018; Kudsk et al. 1995; O’Donovan et al. 1994). Resistance in biotypes from Belgium, Latvia, Germany, and Finland is said to be due to an altered target site, but no further information is available (Heap 2018; Uusitalo et al. 2013). It has been reported that a biotype from France is resistant to Group 2 herbicides due to a Trp-574-Leu mutation in the ALS gene (Heap 2018). Resistance to mecoprop (WSSA Group 4) was first reported by Lutman and Lovegrove in 1985, but the mechanism of resistance remains elusive (Lutman and Heath 2013). It has been reported that a biotype resistant to atrazine (WSSA Group 5), but again there is no known information about the mechanism of resistance (Heap 2018).

In 2016, an S. media biotype resistant to a mixture of thifensulfuron-methyl/tribenuron-methyl was reported in a wheat field in Saint-Camille-de-Lellis, QC, Canada. The objectives of the following research were: (1) to confirm resistance and determine the resistance index to Group 2 (WSSA) herbicides, (2) to detect resistance-conferring mutations, and (3) to provide a quick genetic test to rapidly diagnose the presence of herbicide-resistant S. media.

Materials and Methods

Plant Material

Six small leaves of S. media plants suspected to be resistant to Refine® M (E.I. du Pont Canada, Mississauga, ON, Canada), a combination of Groups 2 and 4 herbicides, were collected and sent to the Agriculture and AgriFood Canada (AAFC) Research and Development Center in Saint-Jean-sur-Richelieu for investigation. These samples were collected from different locations within a wheat field in Saint-Camille-de-Lellis, QC, Canada. The ALS gene sequencing revealed the presence of two mutations: Pro-197-Gln and Pro-197-Ser. Stellaria media seeds were there-after collected from individual plants still growing in the same field. For negative controls, seeds from plants outside the field were also collected. Susceptibility for these seeds was not confirmed. Stellaria media seeds from the AAFC collection were also used as negative controls.

DNA Extraction, ALS Gene Amplification, and Sequencing

Genomic DNA of 27 S. media plants was extracted using Qiagen DNeasy Plant Mini Kit (Qiagen, Mississauga, ON, Canada) according to the manufacturer’s instructions. The different regions of the ALS gene where previous mutations have been reported were amplified using four different primer pairs, ALS-F01/R01, ALS-F02/R02, ALS-F03/R03, and ALS-F04/R04, designed based on ALS gene sequences of S. media and water starwort [Myosoton aquaticum (L.) Moench] (GenBank accession numbers: HE998774 and KF588981, respectively). A polymerase chain reaction (PCR) amplification was performed using the Titanium Taq PCR kit from Clontech Laboratories (Takara Bio USA, Mountain View, CA, USA). The primer pair sequences and the amplification conditions of each primer pair are indicated in Table 2. The PCR-amplified products were visualized on a 1% agarose gel, and the amplicons were sequenced by the dideoxynucleotide chain–termination method (Sanger et al. 1977) at the Genome Quebec Innovation Centre, Montreal, QC, Canada. The sequence alignment and mutation visualization were performed using MEGA (Molecular Evolutionary Genetics Analysis v. 5.05 ) and the Staden Package (Bonfield et al. 1995).

Detection of Pro-197-Gln Using Nuclease Assay

The Pro-197-Gln mutation in S. media–resistant biotypes has been previously reported to confer resistance to mesulfuron (Marshall et al. 2010). A first nuclease assay (TaqMan®) was therefore designed and used to discriminate between S. media wild type and the resistant biotype Pro-197-Gln (Holland et al. 1991). Primers and probes (Thermo Fisher Scientific, Cambridge, MA, USA) were designed based on the S. media ALS gene sequence generated through sequencing. The two primers (ALS-F: 5′-543GGTCCCGATCTGTTGCAATTACG583-5′; ALS-R: 5′-604GGAAACGATCAGTCCCCGATCACG582-5′) were designed to amplify a 62-bp fragment containing the single-nucleotide polymorphism (SNP). Two TaqMan® probes were designed and labeled with fluorescent dyes to detect either the wild type (FAM-labeled: 567ACAAG TCCCGGACG581) or the resistant biotype Pro-197-Gln (VIC-labeled: 567CAAGTCCGAGACG581). Once the S. media ALS gene sequence is aligned with its homologous gene in a closely related species (M. aquaticum), the position of the oligonucleotides S. media ALS gene sequence refers to its positions relative to the start codon in the M. aquaticum ALS gene sequence. The SNP genotyping PCR reaction (10 µl) contained 5 µl of 2X ABI TaqMan® Genotyping Master Mix, 0.25 µl of 40X ABI Custom Assays (a mix of primers...
The sequencing results of the ALS gene revealed the presence of a new mutation, Pro-197-Ser (proline to serine), in some S. media resistant biotypes. This new ALS mutation (Pro-197-Ser) can be distinguished from the other resistant biotypes (Pro-197-Gln) using either the cleaved amplified polymorphic sequence (CAPS) method or a nuclease assay (Konieczny and Ausubel 1993).

The single nucleotide mutation that occurred in the resistant biotype Pro-197-Ser (CCCGAG to CTGGAG) created a XhoI restriction enzyme site. This enzyme recognizes the nucleotide sequence CTCGAG and cuts between the cytosine and thymine codons. To validate this CAPS method, the mutation-containing region of the ALS gene in the three biotypes (wild type, Pro-197-Gln, and Pro-197-Ser) was amplified using the primer pair ALS-F01/R01 under the conditions indicated in Table 2. The amplicons were digested by mixing 5 µl of PCR product with 10 µl of sterilized deionized water, 2 µl of enzyme buffer (10X), 10 µl of enzyme buffer (10X), and 3 µl of XhoI (New England BioLabs, ON, Canada). After incubation at 37 °C for 2 h, the product of the digestion was visualized on a 2 % agarose gel (Figure 1).

Detection of Pro-197-Ser Mutation Using CAPS Method or Nuclease Assay

The sequencing results of the ALS gene revealed the presence of a new mutation, Pro-197-Ser (proline to serine), in some S. media resistant biotypes. This new ALS mutation (Pro-197-Ser) can be distinguished from the other resistant biotypes (Pro-197-Gln) using either the cleaved amplified polymorphic sequence (CAPS) method or a nuclease assay (Konieczny and Ausubel 1993).

The single nucleotide mutation that occurred in the resistant biotype Pro-197-Ser (CCCGAG to CTGGAG) created a XhoI restriction enzyme site. This enzyme recognizes the nucleotide sequence CTCGAG and cuts between the cytosine and thymine codons. To validate this CAPS method, the mutation-containing region of the ALS gene in the three biotypes (wild type, Pro-197-Gln, and Pro-197-Ser) was amplified using the primer pair ALS-F01/R01 under the conditions indicated in Table 2. The amplicons were digested by mixing 5 µl of PCR product with 10 µl of sterilized deionized water, 2 µl of enzyme buffer (10X), and 3 µl of XhoI (New England BioLabs, ON, Canada). After incubation at 37 °C for 2 h, the product of the digestion was visualized on a 2 % agarose gel (Figure 1).

For the nuclease assay, the two primers (ALS-F: 5′-TCTGCAGAGGAACCTGGCG-3′; ALS-R: 5′-CGAATTTGGGGATTCTC-3′) were designed to amplify a 116-bp fragment containing the nucleotide mutations. Two custom

Table 1. Correspondence between sequencing results and genetic tests (nuclease assays and cleaved amplified polymorphic sequence [CAPS] method).

<table>
<thead>
<tr>
<th>Stellaria media samples</th>
<th>Biotypes</th>
<th>Nuclease assay for Pro-197-Gln</th>
<th>Nuclease assay for Pro-197-Ser</th>
<th>Sequences (CAPS)</th>
</tr>
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<tbody>
<tr>
<td></td>
<td></td>
<td>Wild-type FAM</td>
<td>Mutant VIC</td>
<td>Wild-type FAM</td>
</tr>
<tr>
<td>SM001</td>
<td>Wild type</td>
<td>+</td>
<td>−</td>
<td>+</td>
</tr>
<tr>
<td>SM002</td>
<td>Wild type</td>
<td>+</td>
<td>−</td>
<td>+</td>
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<td>+</td>
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<tr>
<td>PL12-7</td>
<td>Resistant</td>
<td>+</td>
<td>+</td>
<td>+</td>
</tr>
</tbody>
</table>

aSM001 to SM004: susceptible controls from Agriculture and AgriFood Canada collection; SM005 to 008: suspected susceptible biotype from the Saint-Camille-de-Lellis field; PL10-1 to PL10-12 and PL12-1 to PL12-7: plants from seeds collected from suspected resistant biotypes PL10 and PL12, respectively.

bSequences were determined using the dideoxynucleotide methodology described by Sanger et al. (1977); CAPS fragments are italicized.

cFluorescent dies used to label the TaqMan probes.
Table 2. Acetolactate synthase (ALS)–specific primer pair sequences and amplification conditions used in this study.

<table>
<thead>
<tr>
<th>Fragment length (bp)</th>
<th>Primer pairs (5'→3')a</th>
<th>Amplification conditions</th>
</tr>
</thead>
</table>
| 408                  | ALS-F01: 385CGCTATCGGTACGTCCTC <sup>506</sup>  
ALS-R01: 174CAGTCCGAAATCGCA <sup>704</sup>  
ALS-R02: 113CCTGGAGATGTCAGGAAACAA <sup>793</sup>  
ALS-F02: 507TAGTATGCGACTCCGTAAG <sup>1128</sup>  
ALS-R03: 110CCCAACGCGCCACACCAAA <sup>1503</sup>  
ALS-R04: 120GATGGGAGGTAGTTGGAAG <sup>1325</sup>  
ALS-RO4: 1177ATGGCTGAATCATCGGAAGG <sup>1760</sup> | 95 C: 5 min;  
35 cycles for 95 C: 25 s; 54 C: 25 s; 68 C: 40 s;  
68 C: 5 min;  
95 C: 3 min;  
35 cycles for 95 C: 30 s; 58 C: 30 s; 68 C: 1 min;  
68 C: 8 min;  
95 C: 5 min;  
35 cycles for 95 C: 30 s; 53 C: 30 s; 68 C: 35 s;  
68 C: 5 min;  
95 C: 5 min;  
35 cycles for 95 C: 25 s; 53 C: 25 s; 68 C: 35 s;  
68 C: 5 min |
| 594                  | ALS-F03: 893GATGGGAGGTAGTTGGAAG <sup>1319</sup>  
ALS-R03: 150CAGATCGGAGAGGCGTTATTAT <sup>1483</sup>  
ALS-F02: 507TAGTATGCGACTCCGTAAG <sup>1128</sup>  
ALS-R03: 150CAGATCGGAGAGGCGTTATTAT <sup>1483</sup>  
ALS-R04: 120GATGGGAGGTAGTTGGAAG <sup>1325</sup>  
ALS-RO4: 1177ATGGCTGAATCATCGGAAGG <sup>1760</sup> | 95 C: 5 min;  
35 cycles for 95 C: 30 s; 53 C: 30 s; 68 C: 35 s;  
68 C: 5 min;  
95 C: 5 min;  
35 cycles for 95 C: 25 s; 53 C: 25 s; 68 C: 35 s;  
68 C: 5 min |
| 607                  | ALS-R02: 110CCCAACGCGCCACACCAAA <sup>1503</sup>  
ALS-F02: 507TAGTATGCGACTCCGTAAG <sup>1128</sup>  
ALS-R03: 150CAGATCGGAGAGGCGTTATTAT <sup>1483</sup>  
ALS-F03: 893GATGGGAGGTAGTTGGAAG <sup>1319</sup>  
ALS-R03: 150CAGATCGGAGAGGCGTTATTAT <sup>1483</sup>  
ALS-R04: 120GATGGGAGGTAGTTGGAAG <sup>1325</sup>  
ALS-RO4: 1177ATGGCTGAATCATCGGAAGG <sup>1760</sup> | 95 C: 5 min;  
35 cycles for 95 C: 30 s; 53 C: 30 s; 68 C: 35 s;  
68 C: 5 min;  
95 C: 5 min;  
35 cycles for 95 C: 25 s; 53 C: 25 s; 68 C: 35 s;  
68 C: 5 min |
| 476                  | ALS-F01: 385CGCTATCGGTACGTCCTC <sup>506</sup>  
ALS-R01: 174CAGTCCGAAATCGCA <sup>704</sup>  
ALS-R02: 113CCTGGAGATGTCAGGAAACAA <sup>793</sup>  
ALS-R02: 110CCCAACGCGCCACACCAAA <sup>793</sup>  
ALS-F02: 507TAGTATGCGACTCCGTAAG <sup>1128</sup>  
ALS-R03: 150CAGATCGGAGAGGCGTTATTAT <sup>1483</sup>  
ALS-R04: 120GATGGGAGGTAGTTGGAAG <sup>1325</sup>  
ALS-RO4: 1177ATGGCTGAATCATCGGAAGG <sup>1760</sup> | 95 C: 5 min;  
35 cycles for 95 C: 30 s; 53 C: 30 s; 68 C: 35 s;  
68 C: 5 min;  
95 C: 5 min;  
35 cycles for 95 C: 25 s; 53 C: 25 s; 68 C: 35 s;  
68 C: 5 min |

aThe superscript numbers indicate the nucleotide position respective to the start codon of the ALS gene in Myosotis aquaticum (GenBank accession number KFS89891).

nuclease assay probes were designed (Integrated DNA technologies, Coralville, IA, USA) to detect either the wild type (FAM-labeled: 567CAAGTCCGGA <sup>577</sup>) or the resistant biotype Pro-197-Ser (HEX-labeled: 561CAAGTCTGGA <sup>569</sup>). The SNP genotyping PCR reaction was performed using QuantiNova Multiplex PCR Kit (Qiagen). The 10 µl of reaction contained 2.5 µl of 4X Multiplex PCR Master Mix, 0.4 µl of each primer (ALS-F and ALS-R) at 10 µM, 0.4 µl of each probe at 8 µM, 4.9 µl of sterilized water, and 1 µl of genomic DNA. The amplification was performed under the following conditions: 2 min at 95 C, followed by 40 cycles of 5 s at 95 C and 30 s at 60 C in an AriaMx Real-Time PCR System (Agilent Technologies).

Resistance Evaluation and Dose Response

Following the identification of a new mutation by sequencing, dose–response experiments were performed in a greenhouse at AAFC’s Saint-Jean-sur-Richelieu Research and Development Center. For each of the three populations (susceptible, resistant Pro-197-Gln [P10-12 population], and resistant Pro-197-Ser [P12-7 population]), two groups of nine individuals were treated with each dose of the herbicides in a randomized complete block design. This entire experiment was repeated three times for a total of 54 plants per treatment. Plants from susceptible and suspected resistant biotypes were treated using a DeVries Manufacturing (Hollandale, MN, USA) moving-nozzle cabinet sprayer equipped with a 8001E-VS even-banding nozzle calibrated to deliver 180 L ha <sup>-1</sup> of spray solution at 207 kPa. Doses of 0, 0.5, 1, 2, and 4 times the recommended dose for each of the three herbicides tested were applied. The recommended doses for foamsulfuron, flumetsulam, and imazethapyr were 35.1, 87.5, and 100.8 g ai ha <sup>-1</sup>, respectively. Shoot dry biomass was collected and weighed 28 d after treatment. Log-logistic dose–response analysis was performed as described by Seefeldt et al. (1995), where the response in dry biomass (Y), expressed as a percentage of the untreated control, is related to the herbicide dose (x) and used to determine the value of the slope (b) and GR50 value that would best fit the distribution of the values according to Equation 1 (C, lower asymptote; D, upper asymptote; GR50, dose generating a 50% reduction in dry biomass).

\[
Y = C + \left\{ \frac{(D-C)}{\left[1 + (x/GR50)^{b}\right]} \right\}
\]

[1]

The resistance factor was calculated by dividing the GR50 value of the resistant biotype by the GR50 value of the susceptible biotype. Dose–response calculations were performed in SigmaPlot (Systat Software, San Jose, CA, USA).

Results and Discussion

To investigate the possible resistance mechanism allowing the collected S. media biotype to survive field application of thiensulfuron-methyl/tribenuron-methyl, seeds from different plants suspected to be resistant and from a susceptible biotype were sown and genomic DNA extracted from plantlets. Four fragments totaling 1,361 bp of the ALS gene were amplified using four primer pairs (Table 2) and sequenced (Figure 1). As expected, a CCG codon (Figure 1A), coding for proline, was found in the wild type at position 197. For seeds collected from the PL10 population, several individuals (PL10-12 shown in Figure 1B) were polymorphic (A/C) for the second nucleotide of the codon. This polymorphism creates two different codons, CCG and CAG coding for proline and glutamine, respectively. The mutation giving rise to a glutamine-197 is known to confer resistance to sulfonylureas as reported by Marshall et al. (2010). Seedlings from PL12 were also polymorphic at codon 197, where the first base was heterozygous (C/T), again giving rise to the proline codon CCG but also a serine codon TCG.

Figure 1. Acetolactate synthase gene sequencing chromatograms. Codon 197 is under the blue bar; (A) wild type is CCG = proline; (B) PL10-12 has two different codons: CCG = proline and CAG = glutamine; and (C) PL12-7 has two different codons: CCG = proline and TCG = serine.

Myosoton aquaticum (GenBank accession number KFS89891)
this allele (unpublished data). A serine at position 197 has been reported to confer resistance in a plethora of species: kochia [Bassia scoparia (L.) A. J. Scott], low falsepimpernel [Linaria dubia (L.) Pennell], Linaria micrantha D. Don, common falsepimpernel [Linaria procumbens (Krock.) Philcox], prostrate pigweed (Amaranthus blitoides S. Watson), wild radish (Raphanus raphanistrum L.), corn poppy (Papaver rhoes L.), ripgrit brome (Bromus diandrus Roth), crown daisy [Glebionis coronaria (L.) Cass. ex Spach], monochoria [Monochoria vaginalis (Burm.f.) C. Presl ex Kuntz], wild mustard ([Sinapis arvensis L.), rock burrush [Schoenoplectus juncoides (Roxb.) Pall], hare barley [Hordeum marinum ssp. leporinus (Link) Arcang.], rigid ryegrass (Lolium rigidum Gaud.), shepherd’s-purse [Capsella bursapastoris (L.) Medik.], flxweed [Descaria sophia (L.) Webb ex Prantl], mayweed chamomile (Anthemis cotula L.), horseweed (Erigeron canadensis L.), silky windgrass [Apera spica-venti (L.) Beauv.], M. aquaticum, threeleaf arrowhead (Sagittaria trifolia L.), common groundsel (Senecio vulgaris L.), annual sedge (Cyperus compressus L.), smallflower umbrella sedge (Cyperus difformis L.), and Palmer amaranth (Amaranthus palmeri S. Watson); but this is the first report for S. media (Heap 2018).

A dose–response experiment was conducted to determine the level of resistance to three families of Group 2 herbicides conferred by this new Pro-197-Ser mutation, in comparison to a susceptible biotype and the Pro-197-Gln population of S. media (Figure 2). After it was determined that S. media was on the label rate of flumetsulam and imazethapyr (0.2 vs. 0.6 and 0.2 vs. 1.1, respectively, Table 3). Population PL10-12 had a lower resistance factor value when compared with the biotype PL12-7 when treated with both flumetsulam and imazethapyr (0.2 vs. 0.6 and 0.2 vs. 1.1, respectively, Table 3). This result indicates that PL10-12 is more susceptible to the herbicide treatment than the biotype PL12-7 and the susceptible control (because values are <1). It is unknown whether this difference in susceptibility is due to the mutation per se or other genetic factors influencing the outcome of this experiment. This difference could also be attributed to the data used for the regression. A treatment with 0.25 times the label rate might have changed the log-logistic regression in a way that would have reduced this apparent difference in susceptibility. In either case, it is clear that the label rate of flumetsulam and imazethapyr is more than sufficient to control all three biotypes. In both flumetsulam and imazethapyr, both flumetsulam and imazethapyr were able to control all biotypes effectively; the calculated resistance factors for biotypes Pro-197-Ser and Pro-197-Gln were close to or below 1 and the susceptible control (because values are <1). It is unknown whether this difference in susceptibility is due to the mutation per se or other genetic factors influencing the outcome of this experiment. This difference could also be attributed to the data used for the regression. A treatment with 0.25 times the label rate might have changed the log-logistic regression in a way that would have reduced this apparent difference in susceptibility. In either case, it is clear that the label rate of flumetsulam and imazethapyr is more than sufficient to control all three biotypes. When treated with foramsulfuron, the PL10-12 and PL12-7 biotypes show resistance factors of 231.3 and 24.2, respectively. The results obtained for the Pro-197-Gln population (PL10-12) agrees with those reported by Marshall et al. (2010), although a different herbicide of the same chemical family was used in that study. The authors reported that this mutation conferred a resistance factor greater than 48 for metsulfuron, another sulfonylurea. It is also interesting to note that the Pro-197-Ser biotype is almost 10 times less resistant to foramsulfuron than the Pro-197-Gln biotype. Again, it is unclear whether other genetic factors change the resistance factor positively for the Pro-197-Gln biotype where our results suggest that the identity of the changed codon impacts the level of resistance. Such a finding was reported for 197 was made for P. rhoes, A. cotula, and S. vulgaris treated with either triazolopyrimidines or imidazolinone (Heap 2018). In addition, we report here that none of the codon 197 mutations tested so far (Pro-197-Ser and Pro-197-Gln) confer resistance to one imidazolinone herbicide, imazethapyr. Marshall et al. (2010) also reported that a Trp-574-Leu amino acid change confers broad-spectrum resistance to five different classes of Group 2 herbicides ([A] foramsulfuron, [B] flumetsulam, and [C] imazethapyr) at five different concentrations (0 × , 0.5 × , 1 × , 2 × , and 4 × times the label rate). Dry biomass was collected at 28 d after treatment.

![Figure 2. Dose–response curves for susceptible (dark points), Pro-197-Ser (light points), and Pro-197-Gln (dark triangles) Stellaria media biotypes treated with three different Group 2 herbicides ([A] foramsulfuron, [B] flumetsulam, and [C] imazethapyr) at five different concentrations (0 × , 0.5 × , 1 × , 2 × , and 4 × times the label rate). Dry biomass was collected at 28 d after treatment.](https://www.cambridge.org/core)
resistance to Group 2 herbicides. This example clearly illustrates the importance of knowledge about the resistance-conferring mutation. In our study the use of Group 2 herbicides not belonging to the sulfonylurea family will provide control of the resistant *S. media* biotype, and this information should be directly applicable in the field.

Resistance confirmation is the first step to undertake in order to select management practices to control weeds, and genetic tests provide a quick method to identify resistant biotypes for which the mechanism of resistance is known. For this purpose, we have developed assays to identify both Pro-197-Ser and Pro-197-Gln mutations. Table 1 summarizes results obtained with the nuclease assay for Pro-197-Ser and the two nuclease assays have produced results (Table 1). The results of sequencing, CAPS, and nuclease tests, based on the nuclease assay and on CAPS, can be used to determine at which frequency this convergent evolution takes place. Westwood and Weller (1997) have identified field bindweed (*Convolvulus arvensis*) biotypes from the same field that vary in their level of tolerance to glyphosate, and those authors propose that the mechanisms can combine to achieve higher tolerance factors. Similarly, there are two different mechanisms of resistance to glyphosate for *L. rigidum* (altered translocation and a target-site mutation), and both can be seen in the same field (Preston et al. 2017). Because *L. rigidum* is an outcrosser, it is likely that individuals show either or both resistance mechanisms. Alarcon-Reverte et al. (2015) also suggest that multiple mechanisms can be present in a glyphosate-resistant jungerlic *[Echinochloa colona]* (L.) population. These examples, along with the *S. media* biotypes described in this paper, argue that plants will try to adapt in every way they can to herbicide selection, and all mechanisms can be tested in parallel. Selection of the fittest biotypes continues for as long as there is herbicide pressure.

The two mutations occurred in the same codon, which is surprising, because there are at least eight bases in the *ALS* gene known to confer resistance once mutated (Beckie and Tardif 2012). There are 86 mutations in codon 197 out of a total of 171 mutations reported on WeedScience.org, which suggests that this locus is highly variable (Heap 2018). We can only speculate on the reasons why this region is so inclined to mutate; maybe the methylation level at this site is lower than for the rest of the gene. A relationship between methylation state and mutation rate has been observed for humans (Xia et al. 2012). A relationship between methylation state and mutation rate has been observed for humans (Xia et al. 2012). A relationship between methylation state and mutation rate has been observed for humans (Xia et al. 2012).

Diploid Mendelian segregation indicates that at least five samples were expected to be homozygous for the mutation out of the 23 resistant plants tested, but we found none in our populations (Table 1). This result is expected, because *S. media* is autogamous, the plants tested are autotetraploids, and there are two copies of the *ALS* gene, one in each parent genome. It is impossible, with our results, to determine whether the two resistance-causing mutations are found in the same copy of the gene.

It is interesting to see two separate selection events occurring in the same field under the same selective pressure. Because not all resistance cases have been studied in depth, it is impossible to determine at which frequency this convergent evolution takes place. Westwood and Weller (1997) have identified field bindweed (*Convolvulus arvensis*) biotypes from the same field that vary in their level of tolerance to glyphosate, and those authors propose that the mechanisms can combine to achieve higher tolerance factors. Similarly, there are two different mechanisms of resistance to glyphosate for *L. rigidum* (altered translocation and a target-site mutation), and both can be seen in the same field (Preston et al. 2017). Because *L. rigidum* is an outcrosser, it is likely that individuals show either or both resistance mechanisms. Alarcon-Reverte et al. (2015) also suggest that multiple mechanisms can be present in a glyphosate-resistant jungerlic *[Echinochloa colona]* (L.) population. These examples, along with the *S. media* biotypes described in this paper, argue that plants will try to adapt in every way they can to herbicide selection, and all mechanisms can be tested in parallel. Selection of the fittest biotypes continues for as long as there is herbicide pressure.

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We report here a new mutation that confers resistance to Group 2 herbicides to *S. media*, Pro-197-Ser. This mutation was identified in a field biotype growing alongside another Group 2–resistant biotype harboring a Pro-197-Gln mutation. Genetic tests, based on the nuclease assay and on CAPS, can be used to rapidly provide information about the presence of resistance alleles in a field population. Dose–response experiments show that the new mutation confers intermediate resistance to sulfonylureas but the biotype is susceptible to imidazolinone and triazolopyrimidines.

**Table 3.** Dose required to reduce dry biomass by 50% (GR50) for the susceptible biotype and the two resistant biotypes PL10-12 and PL12-7 treated with foramsulfuron, flumetsulam, and imazethapyr.

<table>
<thead>
<tr>
<th></th>
<th>Foramsulfuron</th>
<th>Resistance factor</th>
<th>Flumetsulam</th>
<th>Resistance factor</th>
<th>Imazethapyr</th>
<th>Resistance factor</th>
</tr>
</thead>
<tbody>
<tr>
<td>Wild type</td>
<td>2.6</td>
<td>14.7</td>
<td>12.8</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>PL10-12</td>
<td>231.3</td>
<td>97.4</td>
<td>31.1</td>
<td>0.2</td>
<td>2.7</td>
<td>0.2</td>
</tr>
<tr>
<td>PL12-7</td>
<td>24.2</td>
<td>9.2</td>
<td>9.3</td>
<td>0.6</td>
<td>14.4</td>
<td>1.1</td>
</tr>
</tbody>
</table>

**Figure 3.** Agarose gel electrophoresis of uncleaved and cleaved amplicons by XhoI restriction enzyme of *ALS* gene in *Stellaria media* biotypes. Lanes: 1, 100-bp DNA ladder; 2, SM001; 3, SM006; 4, PL10-3; 5, PL10-4; 6, PL10-5; 7, PL10-6; 8, PL10-7; 9, PL10-8; 10, PL10-9; 11, PL10-10; 12, PL10-11; 13, PL10-12; 14, PL12-1; 15, PL12-2; 16, PL12-3; 17, PL12-4; 18, PL12-5; 19, PL12-6; and 20, PL12-7. The cleaved amplicons are harboring the Pro-197-Ser mutation. The uncleaved amplicons might be wild type or Pro-197-Gln mutants.

**Supplementary material.** To view supplementary material for this article, please visit https://doi.org/10.1017/wsc.2018.26
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