Water stress and triclopyr on clopyralid efficacy in honey mesquite

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Abstract

Water stress may affect herbicide efficacy in herbaceous and woody plants. Chamber studies were conducted to evaluate the influence of water stress (−1.3 to −2.8 MPa) and triclopyr on the absorption and translocation of clopyralid in greenhouse-grown honey mesquite (Prosopis glandulosa Torr.). Xylem water potential was determined in honey mesquite at time of herbicide application. Absorption and translocation of clopyralid was determined at low (−1.3 MPa), medium (−2.2 MPa), and high (−2.8 MPa) water stress at 4 h after application for 1.5-mo-old plants, while only translocation was determined at either a low (−1.4 or −1.6 MPa) or a high (−2.4 MPa) water stress treatment at 24 hours after herbicide application for 3-mo-old plants. Water stress did not affect (P < 0.05) absorption or translocation of clopyralid alone in either study. With 1.5-mo-old plants, the addition of triclopyr to clopyralid increased (P < 0.05) clopyralid absorption in leaves at low (63 µg) and medium (54 µg) water stress compared to high water stress (33 µg) but did not affect (P > 0.05) translocation at 4 hours after application. On 3-mo-old plants, triclopyr decreased (P < 0.01) clopyralid translocation 24 hours after treatment at high water stress. The reasons for reduced uptake and 24 hours post-treatment translocation of clopyralid when applied with triclopyr at high water stress are unclear, but have implications for field applications.

Key Words: Prosopis glandulosa, herbicide, growth chamber, xylem water potential, gas chromatographic analysis

The monoethanolamine salt of clopyralid (3,6-dichloro-2-pyridinecarboxylic acid) is a highly effective herbicide for honey mesquite (Prosopis glandulosa Torr.) control when used alone or in mixtures with the butoxyethyl ester of triclopyr ([3,5,6-trichloro-2-pyridyl] oxalacetic acid) (Bovey et al., 1988, Bovey and Whisenant 1991, Jacoby et al. 1981). Davis et al. (1968) studied the effect of water stress on two other auxin-type herbicides, picloram (4-amino-3,5,6-trichloro-2-pyridinecarboxylic acid) and 2,4,5-T ([2,4,5-trichlorophenoxy]acetic acid). They found that water stress reduced foliar uptake of picloram but not 2,4,5-T in honey mesquite. However, water stress sufficient to slow growth markedly reduced the translocation of both herbicides. Clopyralid, however, may be less sensitive to water stress since it has been effective on honey mesquite during periods of drought. Bovey and Meyer (1986) detected high clopyralid concentrations in stem phloem and xylem when applications were made during drought or fall months. Kloppenburg and Hall (1990a) found that the absorption and translocation of the acid and ester formulations of clopyralid were not reduced in water stressed wild buckwheat (Polygonum convolvulus L.) but absorption of the monoethanolamine and potassium salt formulations were reduced.

The objectives of this study were to determine the effect of plant water stress on the absorption and translocation of the monoethanolamine salt of clopyralid alone and in mixtures with the butoxyethyl ester of triclopyr in greenhouse-grown honey mesquite.

Resumen

El estrés hídrico puede afectar la eficacia de los herbicidas en las plantas herbáceas y leñosas. Se condujeron estudios en cámaras para evaluar la influencia del estrés hídrico (−1.3 a −2.8 MPa) y el tricloropirimidina en la absorción y translocación de clopiralid en mezquite (Prosopis glandulosa Torr.) desarrollado en invernadero. El potencial del agua del xilema se determinó en el mezquite al momento de la aplicación del herbicida. La absorción y translocación del clopiralid se determinó 4 h después de la aplicación en plantas de 1.5 meses de edad sujetas a un estrés hídrico bajo (−1.3 MPa), medio (−2.2 MPa) y alto (−2.8 MPa), mientras que en plantas de 3 meses de edad, sometidas a estrés hídrico bajo (−1.4 o −1.6 MPa) o alto (−2.4 MPa), solo se determinó la translocación 24 h después de la aplicación. En ningún estudio se afectó el estrés hídrico (P < 0.05) la absorción o translocación del clopiralid aplica solo. En plantas de 1.5 meses de edad, la adición de tricloropirimidina incrementó (P < 0.05) la absorción de clopiralid en los niveles de estrés hídrico bajo (63 µg) y medio (54 µg) comparado con el estrés hídrico alto (33 µg), pero no afectó (P > 0.05) la translocación a las 4 h después de la aplicación. En plantas de 3 meses de edad, en el tratamiento de estrés hídrico, el tricloropirimidina disminuyó (P < 0.01) la translocación del clopiralid 24 h después de la aplicación. Las razones para la reducida absorción y la translocación del clopiralid 24 h después de tratadas las plantas, cuando se aplica con tricloropirimidina, no son claras, pero tienen implicaciones para las aplicaciones en campo.
Materials and Methods

Growth environment

Honey mesquite were grown from seed in the greenhouse under natural light in pots (12.7-cm diam x 12.7-cm deep) containing a mixture of Bleiberville clay (fine montmorillonitic Udic Pellusterts), sand, and peat moss (1:1:1, v/v/v) from March to June 1990. Daytime temperature was 35°C and night temperature was 25°C. Two plants were grown per pot, and each had a single woody stem. Plants treated at 1.5-mo old averaged 20-cm tall with approximately 8 to 10 leaves per plant. Plants treated at 3-mo old averaged 36-cm tall with approximately 17 leaves per plant. Pots were watered daily to saturation.

Plants were transferred to a growth chamber 2 weeks prior to initiating water stress. Chamber temperature was set at a day/night regime of 35/30°C with a 16-hour day length and relative humidity of 75%. Light irradiance of the Na/Hg lamps ranged from 600 to 700 μmol m-2 sec-1 at the plant apex. Water content of the soil at field capacity was determined by weighing the pots after watering to saturation and allowing the pots to drain.

Stress measurements

Xylem water potential in the honey mesquite was determined by the Scholander apparatus (Scholander et al. 1965). A separate batch of plants were grown along with the test plants to establish water stress categories. Water was withheld from selected sets of plants at staggered times coincidently with selected sets of the test plants. Various plants were measured for xylem potential from these selected sets of extra plants until a clear separation of water stress categories emerged. At the time of treatment, 1 of the 2 plants in each pot was cut with a razor blade 1 cm from the soil level. The cut stem was inserted through the lid of the pressure bomb with the cut end exposed to detect negative hydrostatic pressure in the xylem of honey mesquite. Stem diameter varied from 1.5 to 2.0 mm in 1.5-mo old plants to 3 to 4 mm in 3-mo old plants. The pressure at which liquid first wet the cut surface was recorded. Based on these measurements, 3 stress categories were created that represented low (~1.3 MPa), moderate (~2.2 MPa), and high (~2.8 MPa) water stress. Each pressure stress category represented the mean xylem potential ± 0.2 MPa. The average xylem potentials for each stress category ranged from −1.3 to −2.8 MPa.

Absorption and Translocation 4 hours After Herbicide Application

The paired, uncut honey mesquite plants (1.5-mo old) in every pot were treated with 60 μg plant⁻¹ of the monoethanolamine salt of clopyralid in 20 μl aqueous solution or 60 μg plant⁻¹ of clopyralid plus 20 μg plant⁻¹ of the butoxyethoxyester of triclopyr. Commercial formulations were used. The final treating solutions contained 0.025% (v/v) surfactant¹. Surfactant concentration of 0.025% had little biological activity but was necessary for application of the aqueous solution on the leaf cuticle. Herbicides were applied to the two youngest mature leaves at the apex in 10 μl aqueous solution (30 μg clopyralid) leaf⁻¹ of a plant that consisted of approximately 8 to 10 leaves. Honey mesquite seedlings were treated at each stress category based on xylem potential readings as described previously.

Plants were harvested 4 hours after treatment. Samples included washed treated leaves, treated leaf washes, and stem which included the entire plant minus treated leaves and roots. The leaf wash was accomplished by washing the detached leaves 2 times in 50 ml basic water (1 ml concentrated NH₄OH 1 liter⁻¹ distilled H₂O) for 30 sec.

To extract clopyralid from plant samples, samples were ground in a Waring blender with acidified acetone [(0.5 ml HCl liter⁻¹ acetone and water mixture (7:1 v/v)]. Samples were then filtered through a Büchner funnel and 5 ml NH₄OH were added to the filtered solution.

The acetone was evaporated at room temperature and the H₂O fraction was adjusted to pH 12 with concentrated NH₄OH and extracted once with ether. The ether fraction was discarded, and the H₂O fraction was adjusted to pH 2 with HCl and saturated with approximately 12 g of NaCl to assist in the extraction. This fraction was then extracted 3 times with 50 ml of ether. Ether fractions were passed through an anhydrous Na₂SO₄ column to remove moisture, then combined and completely dried. A 50-ml portion of the leaf wash was adjusted to pH 2 with HCl, saturated with NaCl (12 g) and extracted in the same manner as the H₂O fraction. A butyl ester derivative of the clopyralid was formed by adding 1 ml 1-butanol and 6 drops H₂SO₄ to the residue and heating for 30 min in boiling water (Cotterill 1978). The mixture was cooled and 20 ml H₂O and 5 ml hexane were added to the test tube. The mixture was shaken for 1 min, and the hexane layer was dispensed.

The hexane fraction was analyzed by a gas chromatograph (GC) using a 2-m long glass column packed with 3% OV 210 on 80 to 100 mesh Supelcoport. The GC was equipped with an electron capture detector and GC settings were: column-160°C, injector-290°C, and detector-300°C. Herbicide concentration was determined by comparing samples to a standard of known concentration. Treatment solutions were also analyzed by preparing a derivative of a 10-μl aliquot. Percent recovery for the entire procedure was also determined by fortifying untreated plant samples with known amounts of clopyralid (3 μg and 30 μg). Clopyralid recovery was approximately 80% and was easily detected to 0.05 μg g⁻¹. Triclopyr was not determined in these experiments because it had been previously determined that it did not interfere in extraction and analysis of clopyralid as determined by analysis of fortified plant material with clopyralid alone and clopyralid with triclopyr.

The experiment utilized a completely randomized design. Four plants were used per replication with 4 replications. The entire experiment was repeated. Data were subjected to analysis of variance (SAS 1989). Means for the herbicide amount (μg) and concentration (μg g⁻¹) for the 2 herbicide treatments were compared using Fisher’s protected LSD at the 5% level (Steel and Torrie 1980).

Translocation 24 hours After Herbicide Application

Three-mo-old honey mesquite plants were sprayed with 0.28 kg ha⁻¹ of the monoethanolamine salt of clopyralid or clopyralid at 0.28 kg ha⁻¹ plus 0.14 kg ha⁻¹ of the butoxyethoxyester of triclopyr. Herbicides were applied in water with 0.025% surfactant (v/v) of the spray solution at the equivalent volume of 93.5 liter⁻¹ ha⁻¹ with a laboratory spray chamber (Bouse and Bovey 1967). Plant growth environment and water stress measurements were similar to those in previous experiments except that plants were treat-


²Supelco Inc., Supelco Park, Bellefonte, Penn. 16823.
ed at only 2 stress levels; low (~1.4 or 1.6 MPa) and high (~2.4 MPa). The upper canopy was sprayed while the lower 10 cm of canopy and stem were protected by fitting split styrofoam cups and cotton over the lower plant and soil surface to prevent herbicide contact. The cup and cotton were removed after spraying. Plants were harvested 24 hours after treatment since previous studies showed that maximum clopyralid translocation occurred by this time (Bovey et al. 1987, 1989, et al. 1990). Plants were analyzed for clopyralid content in the lower canopy only. Three replications with 3 plants per replication were used in a completely randomized design. The entire experiment was repeated. Data were subjected to analysis of variance and means were compared using Fisher’s protected LSD at the 5% level (Steel and Torrie 1980).

Results and Discussion

Absorption and Translocation 4 hours After Herbicide Application

Data were pooled for presentation since no interaction between date and treatment were found. Also, there was no herbicide treatment x moisture stress interaction for leaf wash, treated leaves, or stem. When clopyralid was applied alone, clopyralid amounts were higher in the leaf wash under high (-2.8 MPa) than at low (-1.3 MPa) moisture stress levels 4 hours after treatment (Table 1). There was also a higher fresh weight concentration of clopyralid at the high versus low stress level (data not shown). However, there were no differences in clopyralid content absorbed by the treated leaf or translocated to the lower stem.

Clopyralid remaining in the plant tissue represented 63 to 77% of the original amount applied 4 hours after treatment after adjusting for the percent recovery of the method, however, the roots were not analyzed (Table 1). Data were consistent with other studies that indicated rapid translocation of clopyralid to the lower stem and roots of plants within 4 hours or more after treatment (Bovey et al. 1987, 1988, 1989, 1990; Bovey and Maryueh 1980, Devine and Vanden Born 1985, Kloppenberg and Hall 1990a, O’Sullivan and Kossatz 1984). Although movement is rapid, some clopyralid may be lost by metabolism (Hall and Vanden Born 1988). Devine and Vanden Born (1985) found that after 144 hours, 29% of the 14°C-clopyralid was recovered in roots and developing root buds of Canada thistle [Cirsium arvense (L.) Scop.] plants. T. O’Sullivan and Kossatz (1984) using 14°C-clopyralid in Canada thistle only recovered about 85% of the radioactivity after harvesting the entire plant 2 days after treatment. Based on this earlier work, the discrepancy in the amount of clopyralid remaining was probably due to translocation to the root tissue within the first 4 hours.

When clopyralid was combined with triclopyr, more clopyralid was detected in the treated leaves at low and medium stress compared to high stress (Table 1). However, no differences occurred in clopyralid recovered among water stress levels in the leaf wash or the amount of clopyralid translocated to the stem 4 hours after application of the clopyralid:triclopyr mixture (Table 1).

Data in this report agree with field research which indicated more clopyralid was detected in leaves when triclopyr was combined with clopyralid (0.28 + 0.28 kg ha\(^{-1}\)) than when clopyralid was applied alone at 0.28 kg ha\(^{-1}\) 4 hours after treatment (Bovey et al. 1988). After 24 hours, more clopyralid was detected in the upper stem phloem when combined with triclopyr than when applied alone (Bovey et al. 1988).

Translocation 24 hours After Herbicide Application

Data were pooled for presentation since no date by treatment interaction occurred. However, a herbicide treatment x moisture stress interaction was present (Table 2). There was no difference in clopyralid translocation to the lower stem when applied alone in 3-mo-old honey mesquite 24 hours after treatment at low and high stress levels (Table 2). However, there was less clopyralid detected in both amount and concentration in high stress plant stems when applied with triclopyr (Table 2). The reason for reduced translocation of clopyralid when combined with triclopyr at high stress is not clear. Kloppenberg and Hall (1990b) found that acid forms of clopyralid and triclopyr and the butoxymethyl ester of triclopyr readily partitioned through chemically isolated cuticular membranes into agar but ester formulations of clopyralid were retained by the cuticular membrane. However, they did not look at the influence of one herbicide on the behavior of another herbicide (Kloppenburg and Hall 1990b). Bovey et al. (1983) found that triclopyr was readily

<table>
<thead>
<tr>
<th>Herbicide</th>
<th>Water potential (MPa)</th>
<th>Leaf wash Amount ((\mu)g)</th>
<th>Leaf wash Conc. ((\mu)g (g^{-1}))</th>
<th>Treated leaves(^1) Amount ((\mu)g)</th>
<th>Treated leaves(^1) Conc. ((\mu)g (g^{-1}))</th>
<th>Stem(^2) Amount ((\mu)g)</th>
<th>Stem(^2) Conc. ((\mu)g (g^{-1}))</th>
</tr>
</thead>
<tbody>
<tr>
<td>Clopyralid</td>
<td>1.3</td>
<td>154 b(^1)</td>
<td>416 b</td>
<td>27 a</td>
<td>73 ab</td>
<td>2.6 b</td>
<td>1.0 a</td>
</tr>
<tr>
<td></td>
<td>2.2</td>
<td>120 ab</td>
<td>368 ab</td>
<td>28 a</td>
<td>78 ab</td>
<td>2.5 ab</td>
<td>1.0 a</td>
</tr>
<tr>
<td></td>
<td>2.8</td>
<td>222 c</td>
<td>659 c</td>
<td>23 a</td>
<td>69 a</td>
<td>3.1 b</td>
<td>1.4 b</td>
</tr>
<tr>
<td>Clopyralid + triclopyr</td>
<td>1.3</td>
<td>89 a</td>
<td>229 a</td>
<td>63 b</td>
<td>165 c</td>
<td>2.4 b</td>
<td>0.9 a</td>
</tr>
<tr>
<td></td>
<td>2.2</td>
<td>99 ab</td>
<td>231 a</td>
<td>54 b</td>
<td>117 b</td>
<td>2.3 ab</td>
<td>0.8 a</td>
</tr>
<tr>
<td></td>
<td>2.8</td>
<td>129 ab</td>
<td>381 ab</td>
<td>33 a</td>
<td>101 ab</td>
<td>1.5 a</td>
<td>0.8 a</td>
</tr>
</tbody>
</table>

\(^1\)Monoethanolamine salt of clopyralid and the butoxymethyl ester of triclopyr applied at 60 \(\mu\)g and 20 \(\mu\)g, respectively, to the 2 youngest mature leaves of each plant. Four plants/replication were treated and 4 replications were included in the study.

\(^2\)Plant minus treated leaves and roots.

\(^3\)Values in columns followed by the same letter are not significantly different at the 5% level using Fisher’s protected LSD. The values represent mean total amounts and concentrations from 4 plants per replication. 

Table 2. Clopyralid detected in the lower stem 24 hours after spray application of clopyralid on 3-mo-old, greenhouse-grown honey mesquite.
absorbed from either ester or amine formulations and that triclopyr translocation was rapid in greenhouse-grown honey mesquite. As indicated earlier, certain rates of clopyralid plus triclopyr are synergistic in controlling honey mesquite (Bovey and Whisenant 1991), but the benefit of adding triclopyr to clopyralid may not always be attained when plants are under high water stress.

Conclusion

From these data, we concluded that clopyralid absorption and translocation in honey mesquite plants is not altered by water stress extremes when evaluated 4 or 24 hours after treatment. Data from the field support these conclusions since high concentrations of clopyralid were detected in upper and basal stem phloem and xylem both 3 and 30 days after treatment during periods of water stress that ranged between −3.1 and −1.9 MPa during midday (Meyer and Bovey 1986).

In the field, even during periods of high water stress at midday, predawn water stress in honey mesquite may be < −1.0 MPa (Meyer and Bovey 1986, Haas and Dodd 1972) and may permit significant clopyralid translocation. In these studies, water stress was constant, and clopyralid uptake and transport was not reduced under high water stress. It is evident that clopyralid is highly stable and mobile within honey mesquite plants. Absorption through the symplast and translocation through the phloem are essential for activity. The uptake of clopyralid has been classified as nonfacilitated diffusion by Devine et al. (1987). An ion-trap mechanism was shown to be responsible for retaining the undissociated herbicide in the cytoplasm. Since clopyralid was reversibly bound, it can easily be moved into the phloem for transport.

The reasons are unclear for reduced uptake of clopyralid when applied with triclopyr at high water stress. This mechanism should be investigated since clopyralid:triclopyr mixtures are sometimes synergistic in controlling honey mesquite plants and similar mechanisms may exist for other weeds and herbicides.

Literature Cited


