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Strigolactones and hormonal interaction in control of branching in *Zantedeschia* and other horticultural species

A thesis presented in partial fulfillment of the requirements for the degree of

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Abstract

Shoot branching that involves development of lateral buds into shoots is one of the important factors influencing crop productivity. Strigolactones have recently been found to be involved in the control of branching, but the actual bioactive compound/s that inhibits bud outgrowth is still unknown. A germination assay utilizing the seeds of a parasitic weed (*Orobanche minor*), detected strigolactones within the xylem exudates of different horticultural crop species; the strigolactone concentration negatively correlated with branching of cultivars or mutants. In *Zantedeschia* grown *in vivo*, the concentration of strigolactones was independent on the volume of guttation fluid (xylem exudates) suggesting the difference in concentration of strigolactones in high and low branched cultivars was due to the difference in potential of producing strigolactones between these cultivars and not due to differences in volume of guttation fluid. While identifying a bioactive compound using germination and branching assays in combination with liquid chromatography and mass spectrometry, compounds containing ‘N’ were detected in the low branched wild-type *Petunia*, but not in the highly branched mutant, suggesting the possibility of such compounds being SL-conjugates which may be associated with bud outgrowth inhibition.

In *Zantedeschia* (*in vitro*) and pea stems, strigolactone reduced the axillary shoot number stimulated by the cytokinin suggesting an antagonistic interaction between these two hormones on bud release. However, as cytokinin may stimulate subsequent growth of released buds by increasing the auxin transport out of the bud, strigolactone may have reduced subsequent growth by reducing auxin transport. Since GA₃ enhanced subsequent growth of buds in pea stems, but not the release, an antagonistic interaction between strigolactone and gibberellins on subsequent growth is possible. Interestingly, strigolactone successfully reduced adventitious bud formation in *Zantedeschia* grown *in vitro*, adding a new role for strigolactones in plant development.

Despite correlation between strigolactone and branching inhibition in different horticultural crops such as apple, kiwifruit, *Zantedeschia* and *Acer*, further studies relating to strigolactone and its interaction with other hormones on branching of these crops could be performed using *in vitro* techniques for a clear understanding of strigolactones’ role on branching inhibition. More importantly, quantification of strigolactones using the germination assay may have significant implications in horticultural crop breeding for obtaining desired shoot branching.
Extended abstract

Shoot branching, one of the important factors influencing crop productivity, involves development of lateral buds into shoots on an actively growing primary shoot. Recently, a new hormone, which may be a strigolactone, has been found that inhibits bud outgrowth, however, the precise chemical identity of the bioactive compound(s) is unknown. A bioassay based on the germination of a parasitic weed (*Orobanche minor*) was optimized to detect strigolactones. Although there has been controversy in the literature related to whether or not strigolactones are present in xylem exudates, in this thesis strigolactones were found in xylem exudates of a range of horticultural species. The strigolactone concentration correlated with branching of cultivars or mutants mainly at the stage of the growth cycle before the branches were visually evident.

As the germination assay detects all/most strigolactones, not necessarily specifically those associated with branching, a more specific bioassay based on branching was developed. This bioassay was combined with liquid chromatography and mass spectrometry in an attempt to identify a specific branching hormone, whether or not this was a strigolactone. In *Petunia*, four compounds containing ‘N’ were detected in the xylem sap of the wild-type, low branched, V26, but not in highly branched dad3 mutant, suggesting the possibility of such compounds being SL-conjugates which may be associated with branching inhibition. In *Zantedeschia* grown *in vivo*, since the concentration of strigolactones was independent of the volume of guttation fluid, it was suggested that difference in concentration of strigolactones in high and low branched cultivar was due to the difference in potential of producing strigolactones between these cultivars and not due to differences in volume of guttation fluid.

The synthetic strigolactone GR24 (0.1 or 1 mg L\(^{-1}\)) was able to reduce axillary shoot number stimulated by cytokinin in un-decapitated pea stems, and *Zantedeschia* grown *in vitro*, suggesting an antagonistic interaction between these two hormones on bud release, as opposed to subsequent growth. Likewise, strigolactone (1 mg L\(^{-1}\)) reduced decapitation-induced bud release, supporting the hypothesis that strigolactone may have interacted with endogenous cytokinin and/or sucrose. Strigolactone was also able to
reduce subsequent growth of the shoot, but the effect was stronger in buds of pea stems orientated horizontally, compared to those orientated vertically. Such disparity was possibly due to the fact that, in addition to vascular stream, the buds of horizontally orientated stems received strigolactones directly. Although exogenously applied cytokinin appeared to enhance subsequent growth of the released bud, the effect of cytokinin on subsequent growth may be via increasing the auxin transport out of the bud. Hence, rather than interacting with cytokinin, strigolactone may have reduced subsequent growth of the buds by reducing auxin transport. As GA$_3$ enhanced the subsequent growth of buds in pea stems, but not the release, an antagonistic interaction between strigolactone and gibberellins on subsequent growth is considered highly likely. Interestingly, strigolactone successfully reduced cytokinin-stimulated adventitious bud formation in *Zantedeschia* grown *in vitro*. Interaction studies of strigolactone with cytokinin, and probably ethylene, is recommended within highly branched cultivars of *Zantedeschia* spp. or other species, such as gentians and kiwifruit, to further explore the role of strigolactone in adventitious bud formation and development in order to obtain desirable shoots for commercial purposes.

Although strigolactone correlated with branching inhibition in different horticultural crops such as apple, kiwifruit, *Zantedeschia* and *Acer*, further studies for answering the direct role of strigolactone on bud outgrowth in these crops as well as its interaction with other hormones can be performed using *in vitro* techniques. More importantly, quantification of strigolactones using the germination assay may have significant implications in horticultural crop breeding for obtaining desired shoot branching. Since guttation fluid from *Zantedeschia* was found to give a true estimate of the concentration of strigolactones present in the xylem of the shoot system, future experiments may benefit through the use of guttation fluid for hormonal analysis and/or interaction studies *in vivo*. Consideration of the stages of shoot branching during such studies would be valuable for a clear understanding of the shoot branching mechanism and help modify the branching of commercially important crops.
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List of Abbreviations

ABA   Absisic acid
ANOVA Analysis of variance
BAP   Benzylaminopurine
BPC   Base peak chromatogram
BRC1  Branched1
CCD   Carotenoid cleavage dioxygenases
CK    Cytokinin
CRD   Completely randomised design
D27   Dwarf27
DAD   Decreased apical dominance
DMRT  Duncan's multiple range test
DW    Dry weight
FW    Fresh weight
GA    Gibberellin
GA₃   Gibberellic acid
GLM   General linear model
GR24  a synthetic strigolactone (3aR*,8bS*,E)-3-((R*)-4-methyl-5-oxo-2,5-
dihydrofuran-2-yl)oxy)methylene)-3,3a,4,8b-tetrahydro-2H-indeno[1,2-b]furan-2-one
HPLC  High performance liquid chromatography
HRMS  High resolution mass spectrometry
IAA   Indole-3-acetic acid
IPT   Isopentenyl transferase
LC/MS Liquid chromatography/ Mass spectrometry
LSD   Least significant difference
<table>
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<tr>
<td>M.9</td>
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<tr>
<td>MAX</td>
<td>More axillary shoot</td>
</tr>
<tr>
<td>MRM</td>
<td>Multiple reaction monitoring</td>
</tr>
<tr>
<td>NAA</td>
<td>Naphthalene 1-acetic acid</td>
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<tr>
<td>NMR</td>
<td>Nuclear magnetic resonance</td>
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<td>NPA</td>
<td>1-N-Naphthethylphthlamlic acid</td>
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