Sucrose synthase expression pattern in the rhythmically growing shoot of common oak (*Quercus robur* L.)

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Abstract – In the search for a trophic control of rhythmic growth of young oak seedlings, sucrose synthase accumulation, localization and activity were studied in the apex, underlying internodes and leaves, during the second flush of shoot growth. The use of an anti-SuSy antibody raised against the *Vicia faba* protein allowed the detection of SuSy, which showed a 90 kDa subunit. This antibody, used to immunolocalize the SuSy protein in oak revealed a positive signal in the reserve parenchyma tissues of the apex and in the leaf vascular tissues and underlying internodes. The study of SuSy activity along with the growth pattern of the different organs, suggested that SuSy may be involved in the control of rhythmic growth through the mobilization of sucrose in storage tissues and through loading/unloading processes. Such activities would help to bring enough nutrients to support the high morphogenic and growth processes during the rhythmic growth of the shoot.

common oak / rhythmic growth / bud / sugar metabolism / sink strength

1. INTRODUCTION

Common oak (*Quercus robur* L.) shows an endogenous growth rhythm of the shoot axis under natural conditions [15]. This growth rhythm, which can easily be reproduced under controlled conditions, is characterized by the production of a flush every 25 days, after an active growth period of 14 days characterized by elongation of the stem axis and production of different types of leaves [5]. Several hypotheses have been suggested to explain the determinism of rhythmic growth: trophic, hormonal or hydric control [5, 8, 9]. Among these hypotheses, control through trophic competition has most often been investigated. Barnola et al. [4] studied cDMO accumulation (a lipophilic acid whose accumulation indicates an increase in membrane permeability) and showed that the shoot apex, the stem and the leaves displayed successively the largest sink strength during the flush. These authors therefore suggested, that the successive access to more nutrients for each organ of the shoot explained its temporarily increased growth activity and induced the rhythm. To increase our understanding of trophic control of rhythmic growth, we are currently studying carbohydrate metabolism in this system. Here, we report on sucrose synthase (UDP-d-glucose:d-fructose2-α-glucosyl-transferase, SuSy, EC 2.4.1.13), which catalyses the reversible conversion of sucrose and UDP into UDP-glucose and fructose. SuSy is ubiquitous in higher plants and assigns a variety of important roles. It is the predominant sucrose cleavage enzyme in cereal endosperm and in storage organs such as tubers, fruits, seed or immature leaves [24, 28]. Sucrose synthase in these carbohydrate sink organs provides substrates for respiration [29],

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starch [7] and cell wall synthesis [3]. A positive correlation between sucrose synthase activity, sink strength and sucrose import has been demonstrated in many sink organs, suggesting a role for this enzyme in carbohydrate partitioning between sink and source organs [24, 28, 30]. In addition, SuSy appears to play a key role in supplying energy for loading and unloading in phloem by providing substrate for respiration [16].

In this paper, sucrose synthase is studied through the complementary aspects of protein accumulation, tissue localisation as well as activity within the apex, the underlying internodes, the young leaves and the mature leaves during the second flush of growth. Changes in sucrose synthase distribution and activity are discussed in the light of its possible role in the control of rhythmic growth.

2. MATERIALS AND METHODS

2.1. Plant material

Common oak (Quercus robur L.) seedlings were grown from acorns. After a two weeks’ germination in pots filled with moist vermiculite at 16 °C, young seedlings were transferred to hydroponic culture in INRA-Morizet nutrient solution [18] in a greenhouse at 25 °C (± 2 °C) with a 16 h photoperiod. The irradiance from 3 fluorescent tubes averaged 190 μmol m⁻² s⁻¹. Under these culture conditions, common oak produced a new flush of growth every 25 days.

2.2. Sampling procedure

Three key stages of oak development, described by Alatou et al. [2], were chosen for all the studies: the 2nd day with bud swelling, the 7th day, with linear stem elongation, small and epinastic leaves and the 14th day, with maximal leaf expansion and cessation of stem elongation. For each stage, three organs were collected from the second flush: (i) apex (shoot apical meristem plus leaf primordia), (ii) green expanding leaves and (iii) underlying internodes (portion of stem between the first developing green leaf and the apex). Mature leaves of the first flush were added. For enzyme extractions, each sample was made up from at least fifteen plants and the results are means ± the standard deviation of two different cultures × three replicates. All samples were harvested within the same 1-h period in the morning in order to reduce variations due to daily fluctuations of plant metabolism. Tissues were immediately frozen in liquid nitrogen and stored at −80 °C until assay.

For measurements of sucrose and glucose + fructose concentrations, statistical analysis was performed using the Instat software and the Student’s t test (p ≤ 0.05).

2.3. Protein gel blot analysis

Liquid nitrogen-frozen plant tissues were ground, and the proteins from 20 mg of powder were extracted on ice with 10 volumes (w/v) of extraction buffer (50 mM Tris-HCl, pH 8.5, 1% [w/v] polyvinylpyrrolidone (PVPP), 20 mM β-mercaptoethanol, 1% [w/v] SDS, 5 mM EDTA, 5 mM EGTA, 1 mM DTT, and 1 mM phenylmethylsulfonyl fluoride). After centrifugation for 30 min at 16 000 g at 4 °C, the supernatant was collected and resuspended in loading buffer (500 mM Tris, pH 6.8, 10% SDS, 10% glycerol, 1% DTT and 0.01% [v/v] bromophenol blue) to a final concentration of 1 μg/μL. Protein samples (20 μg per lane) were separated by 10% SDS-PAGE and blotted onto PVDF membranes. The immunological detection of SuSy was performed using a polyclonal antibody raised against the denatured SuSy protein of Vicia faba (a gift from Dr Ross H.A. [22]) diluted at 1/1000 prior to incubation with anti-rabbit IgG (Sigma). The protein-antibody complex was detected using chemiluminescence ECL, western-blotting system (Bio-rad).

2.4. Enzyme extraction and assays

For each sample, an extract was obtained after grinding frozen tissues with a glass rod in an Eppendorf tube containing fine sand (diameter < 10 μm). Extraction buffer (50 mM Hepes-NaOH (pH 7.5), 1% [w/v] polyvinylpyrrolidone, 5 mM MgCl₂, 1 mM NaN₃EDTA, 2.6 mM dithiothreitol, 0.02% Triton X100, 0.6% bovine serum albumin) was added twice to a final volume of 200 μL for apices and underlying tissue samples or of 500 μL for leaf samples. The extract was centrifuged for 3 min in a microcentrifuge at 12 000 g and 4 °C. After centrifugation, the supernatant was desalted on G25 Sephadex columns equilibrated with 50 mM Hepes-NaOH (pH 7.5), 1 mM MgCl₂, 1 mM NaN₃EDTA, 2.6 mM dithiothreitol and 0.1% bovine serum albumin. SuSy activity was assayed on 50 μL of desalted extract in a reaction mixture composed of 50 mM Hepes-NaOH (pH 7), 10 mM MgCl₂, 1 mM EDTA, 2.6 mM DTT, 100 mM sucrose, 1 mM ATP, 0.4 mM NAD, 4.2 U hexokinase, 3.5 U phosphoglucosomerase; 2 U glucose-6-P-deshydrogenase. The reaction was initiated by adding 10 mM UDP to the mixture and incubated at 30 °C and the kinetic formation of NADH was measured at 340 nm.

2.5. Sucrose and reducing sugars (glucose + fructose) contents

The supernatant previously obtained was boiled for 3 min and centrifuged at 12 000 g for 5 min. Sucrose and glucose + fructose contents were then determined by enzymatic assays, as described previously [19].

2.6. Immunohistochemistry of SuSy

Samples were fixed for 4 h in F4A (3.7% [v/v] formalin, 5% [v/v] acetic acid, and 50% [v/v] ethanol, in water) at 4 °C. The fixed tissues were then dehydrated in an alcohol series (50 to 100%), embedded in paraffin and sectioned at 10 μm thickness. Sections were incubated overnight at room temperature in a blocking solution (1% BSA [w/v], 0.3% Triton X-100 [v/v] in TBS). In order to reduce the aspecific linkage, slides were incubated for 20 min with an unconjugated anti-rabbit antiseraum raised in goat diluted 1/50 in TBS (Sigma) before incubation for 2 h at room temperature with the primary antibody (anti-rabbit polyclonal antibody given by Ross et al. [22]) or with the pre-immune serum diluted 1/500 in blocking solution. The slides were then washed six times in TBS and incubated for 20 min in unconjugated anti-rabbit antiseraum in goat diluted 1/50 in TBS before incubation for 1 h at room temperature with the secondary antibody (Goat anti-rabbit IgG alkaline phosphatase conjugate, Promega, France) diluted 1/7500 in TBS. After five washes in TBS, sections were incubated with NBT/BCIP. Once color had developed on the sections (2–3 h exposure), sections were rinsed in stop buffer (100 mM Tris, pH 8, 1 mM EDTA), dehydrated, mounted in Entellan (Merck) and photographed.

3. RESULTS

3.1. Protein accumulation (Figs. 1A–1D)

The use of the polyclonal antibodies raised against Vicia faba SuSy, on total protein extracts from apices (Fig. 1A), underlying tissues (Fig. 1B), young leaves (Fig. 1C) and
mature leaves (Fig. 1D) of oak, showed that this antibody recognized a single band with a molecular mass of 90 kDa. This size of the oak protein subunit was slightly smaller than that of *Vicia faba*, estimated to be 93 kDa (Fig. 1). Western blots revealed that the SuSy subunit was present in all organs of common oak, and at all stages tested. Densitometry analysis of protein bands revealed very weak difference in subunit accumulation between dates in apices, underlying nodes and young leaves of the second flush. Only a decrease in mature leaves of the first flush between days 2 and 7 were observed (Fig. 1D).

### 3.2. Protein SuSy localisation (Fig. 2)

In order to investigate the distribution of the SuSy protein during the second flush of growth, an immunohistological analysis was performed in apices, underlying internodes and leaves. Identical localizations were evidenced whatever the stage. In the apices, SuSy protein accumulated in the medular and cortical parenchyma whereas it was absent of the differentiating vascular tissue or of the shoot apical meristem (Figs. 2A and 2B). Unlike in the apices, analysis of underlying internodes sections showed that SuSy protein accumulated in the vascular tissues (Figs. 2C and 2D) and more precisely in the phloem companion cells and in the xylem parenchyma (Figs. 2E and 2F). In mature leaves of the first flush (data not shown), as well as in young leaves of the second flush, a vascular localization of SuSy protein was also observed (Figs. 2G and 2H).

<table>
<thead>
<tr>
<th>Plant part</th>
<th>Enzyme activity (nmol NADH h⁻¹ g⁻¹ FW)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Apex of the second flush</td>
<td>410 ± 140</td>
</tr>
<tr>
<td>Underlying nodes of the second flush</td>
<td>UD</td>
</tr>
<tr>
<td>Young leaves of the second flush</td>
<td>260 ± 80</td>
</tr>
<tr>
<td>Mature leaves of the first flush</td>
<td>UD</td>
</tr>
</tbody>
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<table>
<thead>
<tr>
<th>Plant part</th>
<th>7th day</th>
<th>14th day</th>
</tr>
</thead>
<tbody>
<tr>
<td>Apex of the second flush</td>
<td>UD</td>
<td>UD</td>
</tr>
<tr>
<td>Underlying nodes of the second flush</td>
<td>UD</td>
<td>UD</td>
</tr>
<tr>
<td>Young leaves of the second flush</td>
<td>260 ± 80</td>
<td>260 ± 80</td>
</tr>
<tr>
<td>Mature leaves of the first flush</td>
<td>UD</td>
<td>600 ± 200</td>
</tr>
</tbody>
</table>

### 3.3. Sucrose synthase activity (Tab. I)

SuSy activity was not detected in all organs (Tab. I). Indeed, in apices, SuSy was active only on the 2nd day although its presence was detected by protein gel blot and immunolocalisation after this stage. Similarly in the underlying internodes, presence of SuSy protein but not its activity could be detected. In contrast, SuSy activity was detected in young leaves of the second flush, and its value was stable during the 7th and the 14th days. In the mature leaves of the first flush, enzymatic activity of SuSy was only detected on the 14th day of the second flush.
Figure 2. Localisation of SuSy protein in cross sections of apex (A-B), underlying internodes (C-F) and young leaves (G-H) on the 14th day of the second flush in common oak (*Quercus robur* L.). A, C and G: cross sections incubated with pre-immune serum. B, D, E (detail from D), F (detail from E), and H cross sections incubated with the anti-SuSy antibody of *Vicia faba*. Cc: companion cell, cp: cortical parenchyma, fp: foliar primordium, m: meristem, mp: medular parenchyma, ph: phloem, pr: procambium, v: vasculature, xy: xylem, xyp: xylem parenchyma. Bars = 100 µm (A, B), 300 µm (C, D), 150 µm (E), 30 µm (F), 350 µm (G, H).
3.4. Soluble sugars content (Tabs. II and III)

The sucrose concentration of the apex, underlying nodes, young leaves of the second flush and mature leaves of the first flush measured during the 2nd, 7th and 14th day of the second flush, is shown in Table II. While no significant variations in sucrose concentration were measured in each organ during these developmental stages, Table II shows that for all stages, the sucrose concentration of mature leaves of the first flush is 3× higher than in other organs. The lowest sucrose concentration is measured in underlying nodes of the second flush, with a mean of 4.5 mg g⁻¹ FW.

Variations in reducing sugar (glucose + fructose) concentrations are shown in Table III. As for sucrose concentration, no significant variations were found in each organ during these developmental stages, Table II shows that for all stages, the sucrose concentration of mature leaves of the first flush is 3× higher than in other organs. The lowest sucrose concentration is measured in underlying nodes of the second flush, with a mean of 4.5 mg g⁻¹ FW.

Tables II and III also show that at each stage, the apex of the second flush contains more sugar (sucrose and reducing sugar) than the underlying nodes of the same flush. Comparison between the apex and young leaves of the second flush show that young leaves displayed slightly larger sucrose and reducing sugar concentrations than those of the apex. In mature leaves of the first flush, reducing sugar concentration is significantly lower than in young leaves of the second flush (Tab. III) while for sucrose concentration, an inverse pattern is measured (Tab. II).

4. DISCUSSION

4.1. Detection of SuSy in a heterologous system

Plant sucrose synthases are generally reported to be homo- or heterotetrameric proteins with native molecular masses ranging from 280 to 540 kDa and subunit molecular masses in the range of 83–100 kDa. A single 90 kDa band was detected in oak protein samples using the anti-SuSy antibody from *Vicia faba*. Since in the same western blot, the antibody also recognized the 93 kDa subunit of *Vicia faba*, described in the literature [12], it is very likely that the 90 kDa band corresponds to an oak SuSy subunit. Recently, the release of the complete genome sequence of *Arabidopsis* has shown that the SuSy subunit is encoded by six distinct genes [6]. Each of these genes is differentially expressed depending on the organ type and the environmental conditions, suggesting a transcriptional regulation of SuSy [6]. At the protein level, SuSy is also controlled by numerous posttranslational modifications (for review [11]). In cell cultures of sycamore, for example, western blot analyses of SuSy did not show noticeable differences in protein levels while changes in the SuSy activity were measured. Phosphorylation of SuSy seems to inactivate it in this species [21].
mechanism could be suggested in oak and therefore explain the fact that SuSy was detected in all organs and at each date, but not its activity.

4.2. Differential activity and localization of SuSy during the second flush

The data reported here indicate that sucrose synthase activity varies during the second flush of growth and that a different cellular localization of SuSy protein is observed among organs.

4.2.1. Apex

In oak apices, SuSy activity was only detected during the 2nd day of the flush. At this date, SuSy activity was higher than in other shoot organs. Since SuSy is considered a good indicator of the sink strength of an organ [24], the shoot apex on the 2nd day of the flush appears therefore to be the strongest sink of the shoot. This fact is supported by the non-vascular localization of SuSy protein observed here. Indeed, in maize, this localization is linked with sink tissue activity [10]. Furthermore, measurements of sugar concentrations (sucrose and glucose + fructose) show that in comparison with the underlying nodes, the apex has a higher concentration of these components. Our results are in agreement with those of Barnola et al. who showed, by using the incorporation of DMO as a indicator of organ sink strength in oak, that on the 2nd day, the apex is a stronger sink than the other organs [4].

We also localized the SuSy protein in storage tissues of the bud, i.e., medular and cortical parenchyma, but not in the shoot apical meristem (SAM). Correlation between the localization of this protein and its high activity at a stage when the organogenic activity of the SAM is highest, suggests that SuSy activity could contribute to sucrose mobilization in storage tissues in order to supply hexoses and energy to support morphogenetic and growth activity in the SAM. Similar conclusions on the contribution of SuSy to the maintenance of growth processes, through the hydrolysis of sucrose from storage tissues, were shown previously [28].

Our results also show that SuSy did not accumulate in the shoot apical meristem sensu stricto. In other species, the localization of SuSy has seldom been observed in the meristem region. At the transcript level, two opposite localization patterns are described. In maize and potato, no accumulation of SuSy transcripts is observed in the apical meristem [10, 26] while in tomato Pien et al. [20] showed localization of SuSy transcripts in the SAM and more precisely in a localized region of the meristem flank which includes the earliest stages of primordium bulge formation.

4.2.2. Leaves and underlying internodes

In contrast to the apices, in young and mature leaves as well as in the underlying internodes, SuSy accumulated preferentially in the vascular tissues of the phloem and in particular in phloem companion cells of the underlying internodes. Such localization is in agreement with the results of Kleines et al. [13], who showed a high level of SuSy transcripts in phloem cells in the stem of Craterostigma. Similarly, in mature leaves of eggplant and in young fruit of lemon [25] a high level of SuSy activity was also measured in vascular bundles. SuSy was also localized specifically in the companion cells in maize leaves [17] and in castor bean [27]. This localization in common oak leads also to suggest an involvement of SuSy in controlling sugar fluxes in the vascular system. Indeed, it has been suggested that the hydrolysis of sucrose by SuSy in phloem cells could either constitute a source of energy for loading/unloading of the sucrose [27] or for the provision of UDP-glucose for callose synthesis in the sieve plates [14]. In common oak, the leaves of the first flush are source organs for young leaves of the second flush [1]. Our results show that SuSy activity in mature leaves is high on the 14th day and localized in phloem cells. This suggests that the hydrolytic activity of SuSy in mature leaves could provide energy for sucrose loading in the phloem, thus contributing to the export of sucrose towards the young expanding leaves of the second flush. This hypothesis is reinforced by results obtained in young leaves. Indeed, in this organ, SuSy activity was detectable from the 7th day to the 14th day of the flush and localized in the vascular tissues. At this stage, young leaves seemed to be strong sink and SuSy could participate in the unloading of the sucrose provided by the mature leaves in order to support the rapid growth of young leaves.

5. CONCLUSION

By a multiple approach (enzymatic activity, western blot and immunolocalization), this study provides a detailed analysis of sucrose synthase expression in a woody species, common oak. Results obtained in this rhythmically growing species showed that Susy activity was highest in apices on the 2nd day, followed by young leaves on the 7th day and finally mature leaves on the 14th day. The apices and young leaves organs also contained high soluble sugar concentrations. These facts support the hypothesis of a trophic competition between apices and young leaves. Differential tissue localization of SuSy was also shown in these organs. In the light of these results, this localization was linked with two major roles for SuSy: mobilization of sucrose in storage tissues (apex) and involvement in loading/unloading processes (leaves and underlying nodes). In order to determine if trophic competition is the primary signal that triggers rhythmic growth, further studies should be made on carbohydrate metabolism in the early stages of a flush (for example, each day between the 2nd and 7th days). Moreover, the role of other enzymes involved in the hydrolysis of sucrose in plants should be studied. Indeed, enzymes such as invertases also have an important role in the regulation of source-sink relationships [23] and should also be considered. These questions are currently being addressed using this system and will be reported elsewhere.

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REFERENCES


