Research Article

Identification of Xylem Occlusions Occurring in Cut Clematis (Clematis L., fam. Ranunculaceae Juss.) Stems during Their Vase Life

Agata Jedrzejuk, 1 Julia Rochala, 1 Jacek Zakrzewski, 2 and Julita Rabiza-Świder 1

1 Department of Ornamental Plants, Faculty of Horticulture and Landscape Architecture, Warsaw University of Life Sciences, Nowoursynowska 166, 02-787 Warsaw, Poland
2 Department of Forest Botany, Faculty of Forestry, Warsaw University of Life Sciences, Nowoursynowska 166, 02-787 Warsaw, Poland

Correspondence should be addressed to Agata Jedrzejuk, agata.jedrzejuk@wp.pl

Received 15 March 2012; Accepted 8 May 2012

Academic Editors: M. Edery and D. Granot

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During the vase life of cut stems obstruction of xylem vessels occurs due to microbial growth, formation of tyloses, deposition of materials in the lumen of xylem vessels and the presence of air emboli in the vascular system. Such obstructions may restrict water uptake and its transport towards upwards thus lowering their ornamental value and longevity of cut flowers. Clematis is a very attractive plant material which may be used as cut flower in floral compositions. Nothing is known about the histochemical or cytological nature of xylem blockages occurring in cut stems of this plant. This study shows that in clematis, tyloses are the main source of occlusions, although bacteria and some amorphic substances may also appear inside the vessels. A preservative composed of 200 mg dm$^{-3}$ 8-HQC (8-hydroxyquinolin citrate) and 2% sucrose arrested bacterial development and the growth of tyloses. This information can be helpful in the development of new treatments to improve keeping qualities of cut clematis stems.

1. Introduction

Clematis is used in Europe mostly as a climber plant, but, because of its beautiful flowers, this genus may also provide cut flowers for floral compositions. It is used as such in the United States; the European flower market still lacks suitable clematis cultivars and methods allowing to control thier postharvest quality. This creates a broad opportunity for the European growers and breeders of ornamental plants. Several Polish cultivars are proving themselves to be potential sources of a good cut ornamental material.

The postharvest life of clematis ranges between 2 and 14 days, and it depends on a cultivar. The standard preservative to effectively prolong the vase life of clematis is a solution of 200 mg dm$^{-3}$ 8-hydroxyquinolin citrate (8HQCI) with 2% sucrose [1], but more advanced studies are needed to develop preservatives and treatments suitable for clematis during all steps of the market chain. Proper water balance in cut stems is crucial for the flower postharvest longevity, and blockages occurring in vessels disturb it by limiting water uptake and transport to the flower. The main cause of reduced water uptake in cut stems is obstruction of xylem vessels by microbial growth, formation of tyloses, deposition of materials in the lumen of xylem vessels, and the presence of air emboli in the vascular system [2, 3].

The invasion of the dead lumens of tracheary elements by living parenchyma cells (formation of tyloses) is a well-known response to infection by pathogens and to wounding [4]. It is often accompanied or followed by the transformation of gums and tannins which add to the strength and durability of the composite polymers. The nature of such material was investigated cytologically, revealing the presence of pectic elements, callose, or lignin-like molecules [5–9]. Such material is produced by the plant in response to invasion by the bacteria [10–14] or in response to phytotoxins produced by bacteria [15].

This study was conducted to provide cytochemical and immunohistochemical information on vessel occlusions and
the involvement of tyloses, gels, or gums in their formation in cut clematis stems kept in different vase solutions.

2. Material and Methods

2.1. Plant Material. The study was done on flowering stems of clematis (Clematis L.) kindly provided by Mr. Szczepan Marczynski and Wladyslaw Piotrowski from the plant nursery in Duchnice near Warsaw. The choice of the cultivar was based on observations of the vase life length made by Skutnik and Rabiza-Świder [1, 16] and previous anatomical observations of stems of five different cultivars, of which two were short-lasting (“Andromeda” and “Viola”), one medium lasting (“Isago”), and two long-lasting (“Solidarność” and “Silver moon”). Additionally, anatomical studies of stem blockage formation were done in all five cultivars, while the histochemical, immunohistochemical and cytological identification of the nature of blockages was done in only one, cv. “Solidarność.”

Flowering stems were harvested at the same stage of development, immediately transferred to laboratory and trimmed to 20 cm. Shoots were placed in distilled water or the standard preservative composed of the bactericide 8HQC + sucrose (SUC) which was tested as the most effective preservative [1, 16]. There were eight shoots in each treatment, individually tagged and treated as individual replications. The experiments were conducted at 18–20°C and a 12 h photoperiod, provided by luminescence light with a quantum irradiance of 25 mol m\(^{-2}\) s\(^{-1}\). The relative air humidity was maintained at 60%.

2.2. Anatomy, Histochemistry, and Immunolocalization. Stem ends ca 5 mm long were sampled on three dates: just after harvest (control, day 0), after 7 days (wilting of flowers kept in distilled water, term I), and after 12 days, when wilting and loss of a decorative value occurred in flowers placed into the preservative (term II). On terms I and II, the stem fragments were collected from both treatments (distilled water, term I), and after 12 days, when wilting of flowers kept in distilled water, term II). On terms I and II, the stem fragments were collected from both treatments (distilled water, preservative).

The specimens were fixed in the PFA fixative: 4% paraformaldehyde (Sigma), 0.4% DMSO (Sigma), 0.05M phosphate-buffered saline (PBS) (pH 7.0), DEPC-treated water (Sigma) for 12 h under 0.6 atm. Fixed samples were washed twice for 30 min. each in the phosphate-buffered saline (PBS), dehydrated in the graded ethanol series (30%, 50%, 70%, 80%, 95%, 100%), each series for 1 h in RT (room temperature), and twice in Histoclear (Histochoice Clearing Agent, Sigma) for 30 min each. Paraplast pellets (Sigma) were added to the last series of Histoclear in the paraffin oven, twice a day for 5–7 days, in temperature 56–58°C, until the Histoclear evaporated completely. In the last step, specimens were embedded in clear Paraplast (Sigma). Semithin sections (10 μm) were sectioned on a rotary microtome (Reichert Jung). All preparations were made on the RNase, DNase-free objective slides (Thermo Scientific MenzelGläser, Superfrost Plus), and dried at 42°C for 2–4 days.

For general anatomical identification of xylem occlusions in clematis stems, permanent slides were stained using the safranin—fast green method. For the histochemical identification of xylem occlusions, slides were stained as listed in Table 1.

For some cell wall components, sections were incubated with monoclonal primary antibodies Jim 5, Jim 7 (detection of homogalacturonans), Jim 11, Jim 12, Jim 20 (detection of extensins) synthesized by Dr Knox, Centre of Plant Sciences, University of Leeds, Leeds, UK (details at http://www.plantprobes.net/). Primary antibodies diluted 1:20 in PBS were applied for 2 h at 37°C. Secondary, antiRat IgG antibody labeled with the alkaline phosphatase (SIGMA) was applied for 2 h in 37°C, and slides were incubated with the nitroblue tetrazolium chloride and 5-bromo-4-chloro-3′-indolyphosphate p-toluidine salt (NBT/BCIP) (Sigma) diluted in 100 mM Tris, 100 mM NaCl, 50 mM MgCl\(_2\), for 2 h in the dark. All observations were made using olympus BX41 bright field microscope.

2.3. Electron Microscopy (EM). For conventional EM observations, stem fragments were fixed for 6 h in 2.5% glutaraldehyde (Sigma) buffered with 0.1 M cacodylate buffer, pH 7.2, rinsed in the same buffer and postfixed for 2 h in 1% osmium tetroxide (Merck). Samples were dehydrated in a graded series of alcohol followed by dehydration acetone and embedded in Epon (Fluka). After thin sectioning, samples were stained with 3% uranyl acetate and Reynold’s lead citrate and examined under a JEOL JEM100C transmission electron microscope.

2.4. Statistical Analyses. The xylem vessel data were tested using analysis of variance (Anova 1) with the Statgraphics 4.1. program. Means were compared using the Duncan’s multiple range test at \(P = 0.95\).

3. Results and Discussion

3.1. Anatomical Organization of Clematis Stems. Stems of different cultivars of clematis contain between six to twelve primary vascular bundles with the diameter in the metaxylem between 17.9 and 110.5 μm (see Table 2). Stems of cv. “Solidarność” contain six primary vascular bundles with well visible cambium between the phloem and xylem zones (Figure 1). Cambium consists of 3-4 meristematic cells in
Table 2: Diameter of xylem lumen in 5 different cultivars of clematis stems.

| Cultivar       | Minimum | Maximum | Mean  | Standard error |
|----------------|---------|---------|-------|----------------|
| Andromeda      | 17.9    | 53.9    | 34.8  | ±2.32          |
| Viola          | 20.4    | 53.4    | 37.4  | ±7.02          |
| Isago          | 25.0    | 66.7    | 44.7  | ±2.69          |
| Solidarnośc    | 27.0    | 69.9    | 46.4  | ±2.26          |
| Silver moon    | 28.9    | 110.5   | 59.0  | ±1.48          |

![Figure 1](image)

Figure 1: Transverse section of Clematis "Solidarnośc" stem, p: pith, pr: pith rays, ph: phloem, px: protoxylem, *: vessel of metaxylem, l: xylem, e: endodermis, c: cortex with lamellar collenchymas, pc: pericycle.

3.2. Cytological Identification of Xylem Occlusions. In freshly harvested control stems, the xylem vessels were free of occlusions. The thickness of cell walls ranged from 1.0 to 1.2 µm (Figure 2(a)). After seven days in distilled water (date 1), xylem vessels were blocked primarily by tyloses and to a lesser extent by bacteria. Tyloses contained mostly amyloplasts, and their nuclei showed well-advanced fragmentation (Figure 2(b)). On this collection date, no amorphic or jelly substances were observed in the vessel lumen, but, in 5–7% of the specimen examined, some bacteria were present. On the second collection date, 12 days of vase life in water, tyloses filled nearly the entire volume of xylem vessels. Tyloses contained all components of parenchymatous cell matrix, for example, plastids (Figure 2(c)), mitochondria, plastids, degenerated lipid bodies (data not shown), and plenty of amorphic substances probably originating from degenerating cytoplasm (data not shown). Bacteria were responsible for 27–30% of blocked vessels. In most cases, the vessels blocked by bacteria were free of tyloses.

According to van Meeteren et al. [17], the thickness of cell walls ranged from 1.0 to 1.2 µm (Figure 2(a)). After seven days in distilled water (date 1), xylem vessels were blocked primarily by tyloses and to a lesser extent by bacteria. Tyloses contained mostly amyloplasts, and the nuclei showed well-advanced fragmentation (Figure 2(b)). On this collection date, no amorphic or jelly substances were observed in the vessel lumen, but, in 5–7% of the specimen examined, some bacteria were present. On the second collection date, 12 days of vase life in water, tyloses filled nearly the entire volume of xylem vessels. Tyloses contained all components of parenchymatous cell matrix, for example, plastids (Figure 2(c)), mitochondria, plastids, degenerated lipid bodies (data not shown), and plenty of amorphic substances probably originating from degenerating cytoplasm (data not shown). Bacteria were responsible for 27–30% of blocked vessels. In most cases, the vessels blocked by bacteria were free of tyloses.

According to van Meeteren et al. [17], the vessel diameter may affect the duration of the postharvest life. This is based on the fact that wide vessels are more efficient in water transport [2, 18], and the xylem occlusions do not block the entire lumen of the vessel. Our preliminary research (Table 2) showed that diameter of primary xylem vessels seems to be associated with the length of postharvest life. Observations on all five cultivars showed a similar architecture of stem anatomy, except for the number of primary xylem vessels and their diameters. According to Skutnik and Rabiza-Świder [1, 17] the cut stems of cv. “Solidarnośc” are long-lasting and their wilting in distilled water occurs after 10 days.

In both treatments, young tyloses (those from collection date 1) usually were globular in shape and elongated during the tylose development. According to Clerivet et al. [15], globular-shaped tyloses are outgrowths of the vessel-associated parenchyma cells, which balloon through pit cavities into adjacent vessel elements. They are generally considered as a primary defense mechanism during vascular attacks and hamper the pathogen transportation within xylem vessels [18–21]. Our observations show that the main source of xylem blockage in cut clematis stems is tyloses and that their development is delayed when stems are kept in a standard preservative containing 8HQC + SUC. Even...
**Figure 2:** Ultrastructural identification of xylem occlusions in stems of *Clematis “Solidarno´s´c”* (a) empty vessel, control, (b) young tylose, stems kept in distilled water, date I, (c) mature tyloses, stems kept in distilled water, date II, (d) young tylose, stems kept in 8HQC with 2% sucrose, date I, (e) and (f) mature tylose, stems kept in 8HQC with 2% sucrose, date II, (g) bacteria in blocked vessels, stems kept in 8HQC with 2% sucrose, date II. VL: vessel lumen, ←: cell wall, *: amyloplast, →: degrading nuclei, ⇒: tylose cell membrane, P: plastid, ic: intact cytoplasm, v: vacuole, ↑: mitochondria, ↑↑: lipid bodies, b: bacteria, Control: term 0, Term I: after 7 days of postharvest life (wilting of flowers kept in distilled water), Term II: and after 12 days, when wilting and loss of a decorative value occurred in flowers placed into the preservative.
Identification of Xylem Occlusions. 

3.3. Anatomical, Histochemical, and Immunohistochemical Identification of Xylem Occlusions. In the freshly harvested stems (collection date 0), no traces of mechanical vessel blockage or gel occlusions were observed in any of the five cultivars studied (Figures 3(a) and 3(b)). After seven days of vase life (collection date I), in stems kept in water, completely blocked vessels presented ca 5.06%–7.88% and half-blocked vessels ca 8.64–11.36% of the total vessel number in all observed cultivars (Figures 3(c) and 3(d), Table 3). Blocked vessels were observed only in the metaxylem. After 12 days of the vase life in distilled water (collection date II), completely blocked vessels represented about 10–18.94% and the half-blocked vessels about 9.5–16.6% of the total vessel number (Figures 3(e) and 3(f), Table 4). Completely and half-blocked vessels were present both in the proto- and in metaxylem (Figures 3(g) and 3(h), Table 4).

In stems kept in the standard preservative on the collection date I, completely blocked vessels were blocked in 5.7% in the short-lasting cv. “Andromeda” and in 3% in cv. “Isago.” The remaining three cultivars had less than 1% vessels blocked (Table 3); half blocked vessels were present in 6.2–33.8%, and they were seen in the proto and metaxylem (Table 3). On the collection date II, completely blocked and half-blocked vessels represented 3%–29% and 22.3–37.5% of the total number of vessels observed, respectively, and they were present both in the proto- and in metaxylem (Figures 3(g) and 3(h), Table 4).

There was no correlation between the vessel diameter and proportion of blocked or half-blocked vessels. In stems stored in distilled water (collection date I), the short-lasting cv. “Andromeda” had the highest number of completely blocked vessels (7.8%), but cv. “Viola,” another short-lasting cultivar had the lowest number of completely blocked vessels (5.1%). For the collection date II, the highest number of completely blocked vessels was in the mid-lasting cv. “Isago,” around 19%, and the lowest number of completely blocked vessels was in the short-lasting cv. “Andromeda” and the long-lasting cultivar “Solidarnośc” (12.7%). Stems stored in 200 mg dm$^{-3}$ 8HQC with 2% sucrose in sampling date II showed a clear effect of the preservative: the short-lasting cv. “Andromeda” had 28.3% of completely blocked vessels while the long-lasting cv. “Silver Moon” had 2.9% of such completely blocked vessels. Skutnik and Rabiza-Świder [1, 17] rate cv. “Andromeda” and “Viola” as short-lasting, and both had the lowest diameters of the metaxylem vessels, 17–54 µm. Long-lasting cultivars had larger diameters of xylem vessels, 27–110 µm. Their better postharvest longevity may be associated with a better water hydraulic conductivity through wider lumen of the vessels. We have observed that on sampling date II, the number of completely blocked vessels was higher in stems kept in 8HQC + SUC than in stems kept in distilled water in only one, short-lasting cv. “Andromeda” (Table 4). In cv. “Viola” and “Isago,” the number of completely blocked vessels in the stems kept in 8HQC + SUC was lower when compared with stems kept in distilled water and it was significantly lower in the long-lasting cv. “Solidarnośc” and “Silver Moon” (Table 4). 8HQC

| Cultivar      | Completely blocked vessels | Half-blocked vessels |
|---------------|----------------------------|----------------------|
|               | Distilled water | 8HQC + SUC | Distilled water | 8HQC + SUC |
| Andromeda     | 7.88g          | 5.7e        | 8.64b          | 33.84g     |
| Viola         | 5.14d          | 0.74b       | 8.72b          | 18.85d     |
| Isago         | 7.06f          | 3.0c        | 11.0c          | 21.4f      |
| Solidarnośc   | 5.66e          | 0.1a        | 9.34b          | 20.24e     |
| Silver moon   | 7.14f          | 0.1a        | 11.36c         | 6.2a       |

Numbers followed by the same letter do not differ significantly at $P = 0.95$, according to Duncan’s multiple range test, $P \leq 0.05$. $P$: the least significant difference.

| Cultivar      | Completely blocked vessels | Half-blocked vessels |
|---------------|----------------------------|----------------------|
|               | Distilled water | 8HQC + SUC | Distilled water | 8HQC + SUC |
| Andromeda     | 10.2c           | 28.28h      | 15.88b         | 37.54e     |
| Viola         | 14.18f          | 11.7d       | 11.26a         | 35.48e     |
| Isago         | 18.94g          | 11.4d       | 16.6b          | 22.36c     |
| Solidarnośc   | 12.7e           | 3.74b       | 9.52a          | 28.64d     |
| Silver moon   | 13.74f          | 2.9a        | 15.24b         | 22.6c      |

Numbers followed by the same letter do not differ significantly at $P = 0.95$, according to Duncan’s multiple range test, $P \leq 0.05$. $P$: the least significant difference.
Figure 3: Identification of xylem occlusions in short-lasting (Andromeda) and long-lasting (Solidarność) cultivars of Clematis by staining with fast green and safranin. (a) and (b) Control stems with xylem vessels free of any occlusions in Andromeda (a) and Solidarność (b). (c) and (d) Stems kept in distilled water, term I with xylem vessels half and completely blocked by the occlusions in Andromeda (c) and Solidarność (d). (e) and (f) Stems kept in distilled water, term II with xylem vessels half and completely blocked by the occlusions in Andromeda (e) and Solidarność (f). (g) and (h) Stems kept in 8HQC + 2% sucrose, term II with xylem vessels half and completely blocked by the occlusions in Andromeda (g) and Solidarność (h). c: cortex with lamellar collenchyma, p: pith, pr: pith rays, ph: phloem, arrow: xylem vessel, ∗: blocked xylem lumen. The choice of the cultivar was based on observations of the vase life length made by Skutnik and Rabiza-Świder [1, 16] and previous anatomical observations of stems of 5 different cultivars. Two of them were characterized as short-lasting ("Andromeda”, “Viola”), one medium (”Isago”), and two long-lasting (”Solidarność” and “Silver moon”) cultivars. Control: term 0, term I: after 7 days of postharvest life (wilting of flowers kept in distilled water), term II: after 12 days, when wilting and loss of a decorative value occurred in flowers placed into the preservative.
+ SUC significantly reduced the numbers of completely blocked vessels for the sampling date in all studied cultivars, when flowers were still decorative and at the onset of wilting (Table 3). The number of half-blocked vessels in the stems treated by 8HQC + SUC was higher than in stems stored in distilled water in all observed cultivars in sampling dates I and II (Tables 3 and 4). This suggests that 8HQC + SUC delays the development of tyloses compared to distilled water but does not stop it completely.

3.4. Histological and Immunohistological Identification of Xylem Occlusions in cv. "Solidarność"

3.4.1. Polysaccharides. In controls (sampling date 0) free of any xylem pollutants, polysaccharides were detected in cortex, proto- and metaxylem cell walls with parenchymatic cells of the primary rays. The red color of the PAS reaction was also visible in pith rays, but the color intensity suggested lower accumulation of polysaccharides in pith cells. No color reaction was observed in the phloem and cambium. The epidermis stained brown which is the natural color of the tissue and not the test reaction (Figure 4(a)).

In distilled water after 7 days of vase life (collection date I), the PAS reaction produced clear, red color of blocked vessels. The color intensity ranged from light in the half-blocked vessels to intense in the completely blocked vessels (Figure 4(b)). On the second collection date (after 12 days), strongly red stained vessels were present also in protoxylem (data not shown).

On both collection dates (after 7 and 12 days), in stems kept in the standard preservative, only light red color was visible in the blocked and half-blocked vessels. In this treatment, only metaxylem showed weak color coming from the reaction (data not shown).

Homogalacturonan epitopes were recognized by JIM 5 and JIM 7 antibodies [22]. Histological immunolocalization of homogalacturonans showed strong, blue coloration in the phloem, cambium, pith, and particular cells of primary rays when specimens were incubated with the Jim 5 antibody. No evidence of homogalacturonans was present in xylem occlusions observed in stems placed either in distilled water (Figure 4(c)) or in 8HQC + SUC on any collection date. The Jim 7 antibody gave a much weaker signal than Jim 5 antibody (data not shown).
Strong red color of the xylem occlusions after the PAS reaction in the stems kept in distilled water confirmed a high concentration of polysaccharides in the tyloses. Histoimmunochemistry did not localize any homogalacturonans in tyloses nor in the tubular material blocking xylem vessels. According to the literature [8, 15, 23], pectins concentrate in gels which may occlude xylem vessels, but their presence in tyloses is rather rare. In our study, no pectins were detected on the histochemical level, neither in tyloses nor in amorphous, extracellular material occluding xylem vessels. However, in some plants, tylose differentiation may correlate with the accumulation of pectins in parenchymatic cells of pith [15, 24]. In the stems of *Clematis*, accumulation of pectin epitopes in pith rays was quite evident, but they were not observed in tyloses.

3.4.2. Proteins. The presence of proteins in xylem occlusions was checked using the Bradford reagent. In control stems, an intensive blue color was visible in phloem, cambium, and the parenchymatic cells surrounding the primary xylem. No evidence of proteins was observed in pith rays, xylem vessels, and collenchyma (Figure 5(a)). On both collection dates during the vase life, in stems kept in distilled water, proteins were visible in all blocked vessels (the half- and completely blocked) both in the proto- and metaxytlem (Figure 5(b)).

In stems kept in the standard preservative blocked vessels colored light blue on the sampling date I and strongly blue on the sampling date II. The coloration intensity was equally strong in the completely and half-blocked vessels. Proteins are very often present in the gels or gums occluded into the lumen of the vessel. Tyloses mostly include polyphenols with
Figure 6: Histochemical identification of lignins in xylem occlusions in stems of Clematis “Solidarnośc” by using Azur B and phloroglucinol-HCl methods. (a) and (b) Identification of lignins in xylem of control stems; (c) and (d) Identification of lignins in xylem of stems kept in distilled water; p: pith, pr: pith rays, ph: phloem, arrow: xylem vessel, c: cortex with lamellar collenchyma, *: blocked xylem lumen.

their serious antiseptic properties. The tyloses observed here also included proteins probably transported as an extracellular material from the parenchymatic cells surrounding the primary xylem.

Extensins were identified with the extensin-specific antibodies (Jim 11, Jim 12, Jim 20). None of the antibodies bound to the xylem occlusions. When Jim11 was used, the dark blue color indicating the presence of extensin was not detected in the phloem and endodermis (Figures 5(c) and 5(d)). Jim 12 and Jim 20 did not detect any extensins at the histological level. The synthesis of extracellular structural proteins after an injury or a pathogen attack, as well as their subsequent incorporation into cell walls via oxidative cross-links, has frequently been reported [25–28]. During their immobilization in the cell wall, these proteins can be linked to other extracellular compounds [29, 30] or to other proteins [31–33]. In this study, even though proteins were detected in blocked xylem vessels, extensins were not localized either in tyloses or in the amorphous material occluding xylem vessels.

3.4.3. Lignins. Azur B showed intensive blue, and the HCl phloroglucinol showed intensive red coloration in control xylem cell walls (Figures 6(a) and 6(b)) indicating the presence of lignins. On both sampling dates, the occlusions present in stems kept in distilled water gave strong, blue coloration in metaxylem when stained with Azur B (Figure 6(c)). In phloroglucinol staining, xylem occlusions preserved their natural, orange color in contrast to strongly stained vessels (Figure 6(d)). This phenomenon appeared in xylem vessels from stems kept both in distilled water and in the preservative solution. Azur B staining of xylem blockage in stems placed in the preservative produced blue coloration of the occlusions, but the intensity was lower than in stems kept in distilled water. These confirms cytological observations that tyloses are the main cause of xylem blockage in clematis stems. According to Soukup and Votruba [34] and preceding authors [15, 35], tyloses contain plenty of polyphenols which are main components of lignins and their role is mainly antibacterial. In clematis here, staining with Azur B was more specific for lignins relative to the HCL phloroglucinol test. The test on lignin’s presence in blocked vessels clearly shows that 8HQC + SUC arrests the development of tyloses in cut stems of clematis, as shown by weaker color of stained occlusions.

3.4.4. Suberins. Sections prepared from freshly harvested stems (sampling date 0) and from those kept in the solution
of 8HQC + SUC, did not show any evidence of suberins (Figures 7(a) and 7(b)). Suberins were only present in xylem occlusions from stems kept in distilled water. On sampling dates I and II, suberins were present only in blocked vessels (Figure 7(c)).

4. Conclusions

The main reasons of xylem blockage in cut flowers are air embolism, tyloses, plant and soil microparticles present in the water, bacteria developing in old, dirty water and spreading into vessels, and gums and gels formed in response to cutting [41, 42]. In this study, we have focused mainly on the identification of xylem occlusions appearing in the *Clematis* stems during their postharvest life, depending on the keeping solution. According to Skutnik and Rabiza-Świder [1, 17], solution composed of 8HQC + SUC efficiently prolongs *Clematis* vase life. 8HQC acts as a bactericide, and it may be responsible for the reduced formation of tyloses and other artifacts. According to Van Doorn et al. [43], in some woody ornamentals such as common lilac for cut flower, tylose formation is suppressed by 8HQC (8-hydroxyquinoline citrate) and AVG (aminoethoxyvinylglycine). These results were confirmed by Jedrzejuk and Zakrzewski [44] on stems of common lilac stored in distilled water, 200 mg dm$^{-3}$ 8HQC, and Chrysal Professional. This study shows that tyloses, when they appear in *Clematis* stem kept in 8HQC, they never occlude the entire lumen of a vessel as they do in stems kept in water. Tyloses produced in stems treated by 8HQC + 2% sucrose did not show the presence of degraded

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**Figure 7**: Histochemical identification of suberins in xylem occlusions in stems of *Clematis* “Solidarność” by using Sudan IV. (a) Identification of suberins in xylem of control stems; (b) identification of suberins in xylem of stems kept in 8HQC + 2% sucrose; (c) identification of suberins in stems kept in distilled water. P: pith, pr: pith rays, ph: phloem, arrow: xylem vessel, c: cortex with lamellar collenchyma, *: blocked xylem lumen, ↗: epidermis.
lipids or autophagosomal vacuoles, which is indicative of less degeneration of stem structures in comparison to stems kept in water. Histological tests revealed that the occlusions are mainly composed of proteins, lipids, polysaccharides, phenolics, and lignin-like material. These compounds were reported to appear both in woody and in herbaceous plants [24]. In Clematis, all these components were detected in occluded vessels of the stems kept both in water and in the preservative solution. However, when stems were placed in distilled water, the percentage of blocked vessels was higher than in the stems kept in the preservative. This is demonstrated by relative differences in staining intensities in quantitative color reactions employed in this study. The 8HQC + SUC combination also significantly arrested bacterial proliferation: on the second sampling date, the percentage of vessels containing bacteria was only one half that of stems kept in distilled water (27–30%).

This study confirms that the preservative composed of 200 mg dm$^{-3}$ 8HQC, and 2% sucrose arrests the development of bacteria in the vessels of cut Clematis stems and to some extent also reduces the growth of tyloses. We also identify the origin of the xylem occlusions in Clematis and compare their development in water and in the preservative solution. Such information can be useful to develop new treatments aiming to improve keeping qualities of cut Clematis stems.

**List of Abbreviations**

8HQC: 8-Hydroxyquinoline citrate
8HQC + SUC: 200 mg dm$^{-3}$ + 2% sucrose
DEPC: Diethylypyrocarbonate
PEA: Paraformaldehyde
DMSO: Dimethylsulfoxide
NBT/BCIP: Nitroblue tetrazolium and 5-bromo-4-chloro-3’-indolyphosphate.

**Acknowledgment**

This research was supported by Grant of Ministry of Science and Higher Education no. 0893/B/P01/2009/36.

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