Flow-induced reconfiguration of aquatic plants, including the impact of leaf sheltering

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Abstract

Many aquatic plants are flexible and bend in response to current. This reconfiguration can reduce the drag on the plant, both by reducing the frontal area and by creating a more streamlined shape. Previous studies have considered how the buoyancy and rigidity of a plant impact the drag reduction. This study additionally considered how reconfiguration impacts the sheltering between leaves on a plant and how this, in turn, impacts the drag on the plant. The posture and drag of single-stemmed, leaved plants were studied through a combination of laboratory experiments and theoretical modeling using both plastic Rotala bonsai and live Bacopa caroliniana. The laboratory experiments measured drag and posture on individual plants over a range of channel velocity. The theoretical model calculated plant posture and drag based on a force balance that included buoyancy, the restoring force due to stem stiffness, and leaf drag modified to account for sheltering between leaves. Leaf sheltering was characterized by a sheltering coefficient, $C_s$, which is a function of the plant posture, leaf angle, leaf spacing, and leaf width. $C_s$ decreased from 1 to a minimum value, $C_{sb}$, associated with a fully deflected, horizontal stem posture. Once validated, the model was used to explore a range of leaf configurations, following examples found in real plants. The modeling and experiments revealed conditions for which drag increased with reconfiguration, and also that the drag reached a finite, limiting value for horizontal stem posture. Neither trend has been described in previous reconfiguration models.

Aquatic plants are common features in many environments, including rivers, flood plains, coastal regions, and shallow lakes. They provide many ecosystem services, such as producing and storing carbon (Greiner et al. 2013; Tang et al. 2018), enhancing bed and embankment stability (Barbier et al. 2011; Arkema et al. 2017), and providing shelter (Costanza et al. 1997) and food (Waycott et al. 2005) for fish and aquatic invertebrates. Aquatic plants can be adversely impacted by hydrodynamic drag forces which can damage (Duan et al. 2006) or uproot (Schutten et al. 2005) them. Flexible plants can diminish their exposure to drag by bending in response to flow, which is called reconfiguration (e.g., Vogel 1989; Sand-Jensen 2003). The reconfiguration of a plant depends on its stiffness and material density. Natural plants exhibit a wide range of behavior, from very high current.

The change in drag due to reconfiguration has been quantified by a dimensionless parameter called the Vogel number, $\nu$, defined within the context of the power law dependence of the drag force, $F$, with the velocity, $F = \nu^2 U^{2 + \epsilon}$ (e.g., Vogel 1984; Gosselin et al. 2010; de Langre et al. 2012). A negative Vogel number indicates that the drag dependence on velocity is weaker than quadratic, i.e., drag is reduced relative to a rigid body. Harder et al. (2004) summarized the Vogel number measured for different plant species, which ranged from $\nu = -1.3$ to $-0.32$ to $-0.2$, suggesting that the reconfiguration of leaves played a dominant role in the reduction of plant-scale drag. In addition, Västilä and

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Järvelä (2014) observed that the force per leaf area decreased as the number of leaves increased (increasing leaf area to stem area), suggesting that leaf sheltering was also contributing to the reduction in plant-scale drag. The contribution of leaf sheltering to drag reduction has not been explicitly included in models predicting the impact of reconfiguration on drag. The goal of this paper was to consider this mechanism.

Many previous studies have examined the drag force on leaves and its contribution to the total drag on a branch or tree. For example, individual broad leaves, especially those with long petioles and bilaterally lobed or heart-shaped bases, reconfigure into cones under strong wind (10–20 m/s), which significantly reduces their drag (Vogel 1989). Leaves can contribute 10–90% of the total drag on trees and individual branches (based on flume tests reported in Xavier et al. 2010; Jalonen and Järvelä 2013; Whittaker et al. 2013; Västilä and Järvelä 2014). Because of leaf reconfiguration, the fraction of leaf drag decreases with increasing velocity. For example, Figure 6 in Jalonen and Järvelä (2013) shows that leaves contribute up to 80% of the total drag on a tree dragged through water at low speed (0.1 m/s), but only 40% when the speed exceeded 0.8 m/s. Whittaker et al. (2013) observed that the drag contribution from leaves varied with species, with foliage contributing 75% of the total drag on Alnus glutinosa, but only 30% of the total drag on Salix alba. In contrast to tree species with significant leaf reconfiguration, the present study considered submerged plant species for which the leaves exhibit little reconfiguration, even as stem reconfiguration changes the overall plant posture. In this case drag reduction is not associated with leaf reconfiguration, but instead with the activation of leaf sheltering as the plant reconfigures.

Several previous studies have developed scaling laws predicting the Vogel exponent for simple structures such as beams, rods, and plates, and an excellent review of these models is provided in Gosselin (2019). For example, at sufficiently high velocity, flat plates, filaments (Gosselin et al. 2010; Gosselin and De Langre 2011), and rod-shaped fibers (Alben et al. 2002) exhibit 2-D reconfiguration, which produces a Vogel number \( \nu = -2/3 \). 3-D reconfiguration, such as the rolling up of tuliptree leaf (Vogel 1989) or a Irisidae flaccida blade (Gaylord et al. 1994), has been studied by Schouveiler and Boudaoud (2006) using a disk cut along its radius, such that the disk rolls up into an increasingly acute cone as the flow speed increases. This 3-D reconfiguration produces a Vogel number \( \nu = -4/3 \). Luhar and Nepf (2011) expanded the 2-D reconfiguration model of Gosselin et al. (2010) by including buoyancy. The influence of drag, buoyancy, and elastic restoring force on a simple 2-D structure (e.g., flat blade or cylindrical stem) can be characterized by two dimensionless parameters, the Buoyancy number \( B \) (representing the ratio of restoring force due to buoyancy and restoring force due to stiffness) and the Cauchy number (representing the relative magnitude of hydrodynamic drag and the restoring force due to stiffness). The existing definition of Cauchy number will be denoted as \( Ca' \) (Eq. 2), so that we can later introduce a modified Cauchy number (\( Ca \)) specific to the more complex morphology considered in this study.

\[
B = \frac{\Delta \rho g V_p l^2}{EI}
\]

\[
Ca' = \frac{1}{2} \frac{C_D bU^2 l^3}{EI}
\]

Here, \( \Delta \rho \) is the density difference between water (\( \rho \)) and plant (\( \rho_p \)). \( g \) is the gravitational acceleration. \( V_p \) is the total volume of the plant, which equals \( bt_1 t_2 l/4 \) for a flat blade or cylindrical stem, respectively, with \( b \), \( t \), \( D \), and \( l \) the width, thickness, diameter, and length, respectively. \( E \) is the elastic modulus, and \( I \) is the second moment of area. For a flat blade, \( I = bt^3/12 \). For a cylindrical stem, \( I = \pi D^4/64 \). \( C_D \) is the drag coefficient. \( U \) is the velocity.

To describe the reduction in drag associated with blade reconfiguration, Luhar and Nepf (2011) introduced the effective blade length, \( l_o \), defined as the length of a rigid, vertical blade that generates the same drag as a flexible blade with length \( l \). The effective length is related to the Vogel number, \( l/l_o - U^* \) (see discussion in Pan et al. 2014). Using theory and simulation, Luhar and Nepf (2011) derived the functional dependence of \( l \) on \( Ca' \) and \( B \). When buoyancy is negligible, \( l/l_o - U^{-2/3} - Ca'^{-1/3} \), consistent with 2-D bending (\( \nu = -2/3 \), e.g., Gosselin 2019). Luhar and Nepf (2013) predicted the flow within and above a submerged meadow of flexible blades using this drag model for a single blade, demonstrating that a physical description of an individual plant can be extended to predict flow-vegetation interaction at the canopy scale.

The model proposed by Luhar and Nepf (2011) assumes a blade morphology, representing the seagrass Zostera marina (Fig. 1) and the freshwater grass Vallisneria natans. For a simple blade morphology, in which the bending element is the same as the drag generating element, \( l/l_o - Ca'^{-1/3} \) (\( \nu = -2/3 \), e.g., Alben et al. 2002; Gosselin et al. 2010; Luhar and Nepf 2011). Plants with a rod geometry, such as Equisetum hyemale (Horsetail rush) also follows the 2-D bending model, with \( l/l_o - Ca'^{-1/3} \) (\( \nu = -2/3 \), Hassani et al. 2016). Other aquatic grasses are composed of a slender stem with multiple leaves. The leaves have many shapes: needle, feather, round, short, and long. In addition, the leaves may attach to the stem directly (e.g., Elodea nuttallii) or be linked through thin stalks called petioles (e.g., Ludwigia repens). When multiple leaves are attached to a stem by petioles, both the stem and petioles may reconfigure, which approximates a 3-D reconfiguration. Consistent with this, Puijalon et al. (2005) measured the drag on Mentha aquatica, which has leaves attached by petioles, and found a Vogel number of \( \nu = -1.2 \) to \( -0.75 \), which is higher than the 2-D reconfiguration value (\( \nu = -2/3 \) and approaching the value of 3-D reconfiguration (\( \nu = -4/3 \)). When leaves are attached to the stem directly (without petioles), the leaves are less prone to reconfiguration. This is particularly true when the leaf is short.
and wraps around the stem (e.g., *Rotala bonsai* in Fig. 1), forcing the leaf to arch, which makes it more resistant to bending. In this case, the stem reconfigures to a much greater degree than the leaves, which has been observed for *R. bonsai* and *Bacopa caroliniana* (as discussed in this paper). For this morphology, the leaves provide most of the drag, while the stem does most of the bending. With this disconnection between the drag and bending elements, a new model must be developed to predict the plant posture and drag. Further, for plants with many leaves distributed along the stem (Fig. 1), as the stem bends the upstream leaves may shelter downstream leaves, providing additional drag reduction. This paper uses laboratory experiments with plastic and live plants to guide the extension of previous models (e.g., Gosselin et al. 2010; Luhar and Nepf 2011) to plants with leaves, specifically accounting for the sheltering between leaves.

**Theoretical model to predict drag and posture of buoyant, flexible vegetation with multiple leaves**

A model to describe the reconfiguration and drag for a plant with evenly distributed leaves (Fig. 2), such as *R. bonsai* (Fig. 1) and *B. caroliniana*, was developed with the following assumptions. (1) The material density of the stem and leaves is uniform and constant ($\rho_p$). (2) The leaves are distributed uniformly along the stem and each leaf has the same size (length $l_b$, width $w$, and area $A_l$). (3) The leaves do not deform and the angle between the leaf and stem ($\alpha$) remains unchanged as the stem bends. (4) The stem diameter ($D$) and elastic modulus $E$ are constant. (5) The lift force is assumed to be negligible. Finally, (6) the drag associated with the leaves is much larger than the drag associated with the stem, such that the drag on the stem is neglected. $A_r$

A curved coordinate system is used to describe the stem reconfiguration (Fig. 2). The coordinate along the stem is $s$, with $s = 0$ at the base and $s = l$ at the top of stem, $l$ is the height of the plant when fully erect. $\theta$ is the local angle between the stem and vertical. For simplicity, the forces exerted on the leaves are assumed to be distributed uniformly along the stem, such that the buoyancy force ($f_B$) per stem length is:

$$f_B = \frac{\Delta \rho g V_p}{l} = \frac{(\rho - \rho_p) g V_p}{l}$$

(3)

For a rigid plant, the total drag force can be expressed as:

$$F_r = \frac{1}{2} \rho C_D A_r U^2$$

(4)

in which $A_r$ is the total projected area of a rigid plant onto the vertical. The subscript $r$ denotes the conditions for the rigid, erect plant. The velocity ($U$) is assumed to be steady and uniform over depth. Since the drag on the stem was neglected, $A_r$
is the sum of projected area of each leaf, \( A_r = NA_l|\cos \alpha | \) with \( A_l \) the area of a single leaf and \( N \) the number of leaves. For simplicity, the leaves are assumed to emerge in pairs and to be distributed equally at the left and right side of the stem (see Fig. 2). When the plant is deflected, the stem has angle \( \theta \) to vertical, changing the angle between the leaves and vertical to \( |\theta - \alpha | \) and \( |\theta + \alpha | \) for the leaves on the upstream (left) and downstream (right) sides of the stem, respectively. Consequently, the drag force per stem length, \( f_D \), is a function of \( \theta \) and varies along the stem. Finally, for some stem angles, upstream leaves may shelter downstream leaves, reducing the drag on downstream leaves. To account for this, a sheltering coefficient, \( C_s \), is included in the model. The form drag per stem length can then be described as the sum of drag on the upstream leaves and downstream leaves (first and second term in the numerator, respectively):

\[
f_D = \frac{1}{2} \rho CD C_s \left( \frac{2}{l} A_l |\cos(\theta - \alpha)| + \frac{2}{l} A_l |\cos(\theta + \alpha)| \right) U^2 \tag{5}
\]

Since \( A_r = NA_l|\cos \alpha | \), \( f_D \) can be expressed in terms of \( A_r \):

\[
f_D = \frac{1}{2} \rho CD C_s \frac{A_r |\cos(\theta - \alpha)| + |\cos(\theta + \alpha)|}{2|\cos \alpha|} U^2 \tag{6}
\]

Note that Eq. 6 neglects skin friction and stem drag, because the leaf form drag dominates for most leaf orientation. An exception occurs for an initially vertical plant with horizontal leaves (\( \alpha = 90^\circ, \theta = 0 \)), for which Eq. 6 predicts zero drag, such that the model plant will not reconfigure, which is unrealistic.

When the stem reconfigures, a stem-normal restoring force \( (V) \) is generated, with magnitude equal to the derivative of internal bending moment \( (M = EI \frac{d^2 \theta}{ds^2}) \), i.e.,

\[
V = -EI \frac{d^2 \theta}{ds^2} \tag{7}
\]

The stem-normal force balance for stem section \( s > s^* \), where \( s^* \) is an arbitrary position along the stem, can be expressed as:

\[
- \frac{d^2 \theta}{ds^2} \big| s^* + \Delta \rho g V_p / l (1 - s^*) \big| \sin \theta = \frac{1}{2} \rho CD C_s \frac{A_r |\cos(\theta - \alpha)| + |\cos(\theta + \alpha)|}{2|\cos \alpha|} \cos \theta ds \tag{8}
\]

In dimensionless form, Eq. 8 is:

\[
- \frac{d^2 \theta}{ds^2} \big| s^* + B (1 - s^*) \big| \sin \theta = Ca \int_{s^*}^{s} C_s \frac{|\cos(\theta - \alpha)| + |\cos(\theta + \alpha)|}{2|\cos \alpha|} \cos \theta ds \tag{9}
\]

in which \( s = s/l, B \) was defined in Eq. 1, but the Cauchy number, \( Ca \), has been modified to reflect the more complex plant geometry considered here. Specifically,

\[
Ca = \frac{1}{2} \rho CD A_r U^2 \left( \frac{1}{EI/l^2} \right) \frac{\text{drag on leaves in rigid upright posture}}{\text{restoring force in stem}} \tag{10}
\]

This modified Cauchy number reflects the fact that the drag is associated with the projected area of the leaves, \( A_r \), and the rigidity is associated with the stem, represented by stem length scale \( l \). Finally, two boundary conditions are needed to use Eq. 9 to predict the plant posture and drag. As in previous studies (e.g., Alben et al. 2002), the following two boundary conditions were applied: \( \theta = 0 \) at \( s = 0 \) and \( d\theta/ds = 0 \) at \( s = l \). The total drag on the deflected plant, \( F_r \), can be then obtained by integrating \( f_D \) (Eq. 6) along the stem.

\[
F_r = \int_0^l f_D ds = \int_0^l \frac{1}{2} \rho CD C_s \frac{A_r |\cos(\theta - \alpha)| + |\cos(\theta + \alpha)|}{2|\cos \alpha|} U^2 ds = \frac{1}{2} \rho CD A_r U^2 \tag{11}
\]

The right-most expression defines an effective area, \( A_e \). Similar to the effective length \( (l_e) \) defined in Luhar and Nepf (2011), \( A_e \) is the area of rigid plant that provides the same drag as a flexible plant. With this definition, the ratio of drag on a flexible plant to the drag on a rigid plant can be represented as a ratio of areas, specifically, \( A_e/A_r \) (Eq. 12).

\[
\frac{F_r/F_e}{A_r/A_e} = \int_0^l C_s \frac{|\cos(\theta - \alpha)| + |\cos(\theta + \alpha)|}{2|\cos \alpha|} ds \tag{12}
\]

These ratios are linked to the Vogel number which was defined within the context of quadratic dependence of the drag force, \( F_r \), with the velocity, i.e., \( F_r \sim U^2 + \psi \) (Vogel 1984). For a rigid plant \( F_r \sim U^2 \), such that \( F_r/F_e = A_e/A_r \sim U^\psi \) (see more discussion in Pan et al. 2014).

The sheltering coefficient, \( C_s \), describes the drag reduction associated with the sheltering between leaves. The sheltering of elements within an array has been studied in both air (e.g., Wang and Takele 1997; Btsuamlak et al. 2010; Wu et al. 2010) and water (Barsu et al. 2016; Jin et al. 2018). Based on these studies, \( C_s \) is expected to depend on the leaf width \( (l_w) \) and spacing \( (d) \), the angle between leaf and stem \( (\alpha) \), and the angle of the stem to vertical \( (\theta) \). For simplicity, consider a linear stem that tilts relative to the vertical (Fig. 3). For a linear stem, the angle \( \theta \) is related to the ratio of deflected height, \( h \), and stem length, \( l \), specifically, \( \cos \theta = h/l \). As the stem tilts, sheltering between leaves will occur when two conditions are met. First, part of the upstream leaf must be in-line with the downstream leaf. Second, the spacing ratio \( (d/l_w) \) must be small enough that the downstream leaf falls within the wake of the upstream leaf. These two conditions are discussed in more detail below.
First, sheltering can only occur if part of an upstream leaf falls in front of a downstream leaf. Consider the plant illustrated in Fig. 3, as the plant deflects from its vertical position, at some degree of deflection the top of the upstream leaf will just pass the plane of the downstream leaf. This position will be denoted as the critical deflected height, \( h_c \), illustrated in Fig. 3b. The critical posture can also be described by the critical tilting angle \( \theta_c \), which is geometrically related to the critical deflected height, i.e., \( \cos \theta_c = h_c/l \). The critical angle, \( \theta_c \), is defined by geometry, as expressed in Eq. 13.

\[
\delta_1 l |\cos(\theta_c - \alpha)| + \delta_2 l |\cos(\theta_c + \alpha)| - d |\cos \theta_c| = 0 \tag{13}
\]

in which \( \delta_1 = 0 \) when \( \theta_c - \alpha < - \pi/2 \) (may occur when \( \alpha > \pi/2 \)), other wise, \( \delta_1 = 1 \). Similarly, \( \delta_2 = 0 \) when \( \theta_c + \alpha < \pi/2 \), other wise, \( \delta_2 = 1 \). Note that values of \( \theta \) range from 0 to \( \pi/2 \) and \( \alpha \) range from 0 to \( \pi \). The sheltering coefficient \( C_s = 1 \) when \( \theta < \theta_c \), i.e., \( h > h_c \). \( C_s \) decreases as \( \theta \) increases from \( \theta_c \) to \( \pi/2 \) (\( h \) decreasing from \( h_c \) to 0) at which point the plant reaches the maximum reconfiguration with a horizontal posture (Fig. 3c). At this posture, the sheltering coefficient reaches a minimum value, defined as \( C_{s0} \).

Second, the degree of sheltering in the final horizontal posture will depend mainly on leaf spacing \( (d) \) and leaf width \( (l_w) \), the smaller dimension of the leaf). Results from Barsu et al. (2016) provide insight into the dependence of sheltering on leaf spacing and leaf width. Specifically, Barsu et al. (2016) measured the drag on a row of blades considering different blade width and different spacing between blades. In comparison to the drag measured for an isolated blade, they showed that the drag is reduced due to sheltering when the spacing, \( d \), is less than four times \( l_w \). Further, Figure 6b in Barsu et al. (2016) shows that the drag coefficient decreases linearly with decreasing leaf spacing for \( d_c < 4l_{\omega} \). Based on this, we assume that the critical spacing for sheltering to occur is \( d_c = 4l_w \), and for \( d \geq d_c \), \( C_{s0} = 1 \). The critical value \( d_c = 4l_w \) makes sense because the recirculation zone developed behind an isolated flat plate is approximately 4 time the plate width (Barsu et al. 2016). For \( d < d_c \), Figure 6b in Barsu et al. (2016) suggests the following linear relationship.

\[
C_{s0} = d/(4l_{\omega}) \tag{14}
\]

Combining the above considerations and assuming \( C_s \) decreases linearly with deflected plant height, \( h/l \), we have:

\[
C_s = \left\{ \begin{array}{ll}
1 & d \geq d_c \text{ or } h \geq h_c \\
\frac{1-C_{s0}h}{h_c/l} + C_{s0} & d < d_c \text{ and } h < h_c
\end{array} \right. \tag{15}
\]

Laboratory experiments were conducted to first verify Eq. 14. Next, measured values of plant posture and drag were used to verify the prediction of plant posture and drag using Eq. 9 and Eq. 12, which incorporated the sheltering coefficient (Eq. 15). Because \( C_s \) and \( h/l \) are mutually dependent, the following iterative method was needed to predict drag and plant posture. An initial sheltering coefficient \( C_{s1} = 1 \) was applied to Eq. 9 to calculate \( h_{i1}/l \). Next, \( C_{s2} \) was calculated from \( h_{i1}/l \) and Eq. 15. After this, \( C_{s2} \) was applied to Eq. 9 to calculate \( h_{i2}/l \). This was repeated until step \( i \), for which the difference between \( h_{i}/l \) and \( h_{i-1}/l \) was less than 1%.

**Laboratory experiments**

Three types of experiments were conducted using plastic and live plants. First, in Experiment 1, the limiting sheltering coefficient \( C_{s0} \) (Eq. 14), was studied using plastic R. bonsai leaves distributed along a thin rigid wire (with 0.9 mm diameter), held rigidly by a small post at stem angle \( \theta = 90^\circ \) representing the most extreme stem reconfiguration (Fig. 4a). The number of leaf pairs was varied from 1 to 20, and the spacing between leaves ranged from infinite to 0.5 cm. Two types of
leaf distributions were tested, type 1: pairs of leaves aligned along the stem, as assumed in the theoretic model (see Fig. 4a) and type 2: pairs of leaves alternating by 90° rotation along the wire, representative of many real plants (e.g., Fig. 4c). The drag force on 1 leaf and on N leaves was measured and subtracted. Assuming all leaves have equal area, \( C_{D_{AR}} = \frac{F_{DN}}{(N F_{D1})} \).

Second, in Experiment 2, rigid model plants were used to establish the baseline drag on a rigid plant without reconfiguration (Fig. 4b). The stem was made rigid by attaching a thin steel wire (1 mm diameter) along the back of the stem. To simplify the analysis, Eq. 4 was rewritten as a function of \( k \). The coefficient \( k \) was estimated for a 10-cm plant exposed to velocity \( U = 5–26 \) cm/s (Fig. 4b). For a rigid vertical plant, the total drag is proportional to stem length, such that the value of \( k \) measured for the 10-cm plant could be used to estimate the drag on a longer rigid plant. Specifically, \( F_r = k U^2 \), in which \( n = 1, 2, 3, \) and 4 for plants with length \( l = 10, 20, 30, \) and 40 cm, respectively. Applying, \( F_r = k U^2 \) to Eq. 10, the Cauchy number becomes,

\[
Ca = \frac{nkU^2l^2}{EI}
\]

For the live \( B. \ caroliniana \), there was no way to attach a rigid steel support without breaking the leaves. As an alternate method, the forces measured for the shortest \( B. \ caroliniana \) at the lowest three velocities \( (1.75–7.2) \) cm/s were used to define the coefficient \( k \) and \( Ca \). For this plant and over these three velocities the drag varied quadratically with velocity.

Third, in Experiment 3, flexible model \( R. \ bonsai \) (plastic plant, \( \rho_p = 914 \) kg/m\(^3\), \( E = 1.11 \times 10^8 \) Pa, \( I = 3.79 \times 10^{-13} \) m\(^4\)) and live \( B. \ caroliniana \) \( (\rho = 748 \) kg/m\(^3\), \( E = 7.51 \times 10^6 \) Pa, \( I = 2.02 \times 10^{-12} \) m\(^4\)) were used to validate the prediction of plant posture and plant drag force. To capture a wide range of Cauchy number, four different lengths of plastic \( R. \ bonsai \) (10, 20, 30, and 40 cm) and three different lengths of live \( B. \ caroliniana \) (9, 17, and 28 cm) were studied (Table 1). The stem rigidity, \( EI \), was measured with the cantilever beam method using a 5-cm stem specimen (leaves removed). Based on five replicate tests on the plastic sample the uncertainty was \( 2\% \). With the velocity \( U = 2–62 \) cm/s, the Cauchy number ranged from \( Ca = 0.01–3000 \). The uncertainty in the drag force was estimated to be \( 1\% \) based on repeated measurements in Experiment 2. Along with the uncertainty in \( 1\% \) in stem length and \( 2\% \) in stem rigidity \( EI \), the uncertainty of Cauchy number was approximately \( 10\% \).

All the experiments were conducted in a 24-m-long, 38-cm-wide, 60-cm-depth recirculating flume. Velocity was adjusted using two variable speed pumps. To reduce wave generation at the inflow, two layers of rubberized coconut fiber (80 cm length, 38 cm width, 10 cm thickness) were placed just below the water surface at the channel inlet (see Fig. 5).

In each experiment, a single live or model plant was attached to a 2.5-cm tall stainless steel post with a 2-mm diameter, which was connected to a submersible force sensor (Futek LSB210 100 g, see Fig. 5). The sensor was calibrated by National Instrument (NI-USB 9237), and force measurements were logged to a computer using a data acquisition module (Lei and Nepf 2019). The force measurements had an accuracy of \( 1\% \). The force sensor was mounted beneath two layers of rubberized coconut fiber to avoid interaction between flow and the sensor. For each flow condition, the drag force was measured for 3 min at a sampling rate of 1600 Hz.
Experiments with plastic plants were repeated four times while live plants were tested one time to keep the plants in good condition. A separate force measurement was made on the post alone (without plant) at each velocity, and the force on the post was subtracted from the force measured with the plant and post together. During each force measurement, digital images were captured by a smart cellphone (MIX 2S) camera with 12 million pixels in each photo. The camera was fixed to a tripod by a self-stick holder. The camera captured 100 photos in 10 s. This sampling frequency (10 Hz) was much higher than the estimated 0.1–0.8 Hz stem natural frequency (Table 8.1 in Blevins 1979) and 0.2–1 Hz vortex shedding frequency from the plant leaves (e.g., Blevins 1990), and the 10-s duration included a minimum of 10 cycles of possible oscillation. Twenty photos spaced 0.5 s apart were selected from each burst to obtain the range of plant posture and the deflected plant height. A photo with a ruler beside the plant (Fig. 4c) was used to calculate the scale (pixels/cm).

Three water depths, 16, 22, and 32 cm (above the ramp) were selected to achieve the desired velocity and maintain complete submergence of the plant. The Froude number ranged from 0.01 to 0.5. After the force measurements, the plant, post, and force sensor were removed, and an acoustic Doppler velocimeter (ADV, Nortek Vectrino) was used to measure velocity profiles at the middle of the ramp where the plant had been located. The vertical resolution of the velocity profile was 1 cm to 3 cm. At each measurement point, the Vectrino recorded a 3-min record with a sampling frequency of 200 Hz. The post elevated the plant above the boundary layer on the ramp, so that the plant experienced a nearly uniform velocity over depth (see Figure 3 in Luhar and Nepf 2011). Each velocity measurement was quality checked by the signal to noise ratio and de-spiked with the following method. First, the acceleration thresholding method identified and removed spikes in the velocity time series if the instantaneous acceleration was higher than the acceleration of gravity \( g \) (Nikora and Goring 2000). Second, the time-mean velocity \( \bar{u} \) and standard deviation \( \sigma \) of the remaining data were calculated and the velocity component \( u_i \) was removed if \( |u_i - \bar{u}| > 3\sigma \) (Goring and Nikora 2002). Finally, the depth-averaged velocity \( U \) was estimated as the average of time-mean velocity \( \bar{u} \) between 3 cm above ramp and flow surface.

**Results**

**Sheltering coefficient**

In Experiment 1 (Fig. 4a), force measurements made on a horizontal stem with leaves of varied spacing \( d \) were used to estimate the limiting sheltering coefficient, \( C_{s0} \), representing the condition with maximum sheltering. The measured values of \( C_{s0} \) are plotted as a function of the ratio of spacing \( d \) to leaf width \( l_w \) in Fig. 6. There was no systematic difference between the cases with aligned leaves (type 1) and alternating leaves (type 2), indicating the theoretic model assumption of aligned leaves was reasonable. At the maximum spacing \( d/l_w = 7.5 \), \( C_{s0} = 1 \), indicating that no sheltering impacted
the drag on individual leaves. For spacing less than $d/l_w = 5$, the sheltering coefficient decreased (indicating increasing sheltering) as $d/l_w$ decreased. This trend agreed with the observations in Barsu et al. (2016), despite the different shape in model leaves. Specifically, they noted that $C_{sd}$ decreased linearly from 1 for $d/l_w < 4$, as shown with a solid black line in Fig. 6, corresponding to $C_{sd} = d/(4l_w)$, i.e., Eq. 14. Note that for $d/l_w \leq 2$, $C_{sd} = d/(4l_w)$ underestimated $C_{sd}$, i.e., overestimated the sheltering effect because the first leaf will never be sheltered, such that the drag cannot actually reach zero. The same underestimation of $C_{sd}$ by Eq. 14 (overestimation of sheltering) would also occur for a small number ($< 4$) of leaves, because the drag on the first leaf, which is never sheltered, would be a large fraction of the total leaf drag.

Drag force on rigid plant

As expected, the drag force measured for the rigid plastic plant, $F_r$, followed a quadratic dependence on velocity (e.g., Fig. 7). The slope of $F_r$ vs. $U^2$ was denoted as $k = 1/2 \rho C_f A_r$. For the 10-cm long rigid plant, $k = 0.488 \pm 0.009 \text{ Nm}^{-2} \text{ s}^{-2}$ (Fig. 7a). The smallest specimen ($l = 9 \text{ cm}$) of live B. caroliniana exhibited limited reconfiguration at the three lowest velocity conditions, such that the drag increased linearly with velocity squared (Fig. 7b), providing an estimation of the rigid drag factor, $k = 0.97 \pm 0.02 \text{ Nm}^{-2} \text{ s}^{-2}$ (Fig. 7b).

Reconfiguration of and drag on flexible model plant with leaves

This section describes the measured deflected stem height and drag force and makes comparison to the model prediction. Consistent with the model assumption, the leaves in the experiment did not deform, even as the stem was deflected horizontally (see images in Fig. 8). The drag reduction was framed in terms of the effective area ratio, defined as the ratio of effective frontal area to rigid frontal area, $A_e/A_r$, which was estimated from the ratio of drag force on a flexible plant $F_l$ to drag force on a rigid upright plant $F_r$, as defined in Eq. 12. Figure 8 shows the measured and modeled values of stem deflected height, $h/l$, and effective area ratio, $A_e/A_r$, for the plastic model of R. bonsai (with $l_w = 1.2 \text{ cm}$, $d = 1.2 \text{ cm}$, $a = 48^\circ$, and from Eq. 14 $C_{so} = 0.25$). The critical posture was $\theta_c = 24^\circ$ ($h/l = 0.91$) as calculated from Eq. 13. See the summary of parameters in Table 1. To capture the range of Buoyancy number ($B = 0.4–14$), model predictions were made for $B = 0$ and $20$. Images of plant posture at different Cauchy numbers are included at the bottom of Fig. 8. Note that the plant was not perfectly straight, and image 1 represents the fully erect posture, which was the same as that observed for zero current. The red lines in the plant images are the postures predicted from the model (Eq. 9). The model captured the trends in both deflected height ($h/l$) and effective area ratio ($A_e/A_r$) over the entire Cauchy number range. The maximum error between the modeled and measured values of $h/l$ and
Fig. 8. (a) Normalized deflected height, $h/l$, vs. Cauchy number defined in Eq. 16. (b) Normalized effective frontal area, $A_e/A_r$, vs. Cauchy number. The symbols are measured values, and the lines are smoothed curves based on individual model predictions at 55 Cauchy numbers. The black arrow indicates the model predicted limiting value $A_e/A_r = 0.28$ from Eq. 12. The error bars indicate the standard deviation among four repeated measurements of force and 20 photos of postures for $A_e/A_r$ and $h/l$, respectively. The bold numbers correspond to the images shown at the bottom, which correspond to $Ca = 0.4, 5, 48, 540,$ and $1930$. The red lines in the images are the model predicted postures.

$A_e/A_r$ were $\pm 0.04$ and $\pm 0.1$, respectively. The good agreement between prediction and measurement supported our assumptions, including the neglect of lift force. A negligible lift contribution may be attributed to the leaves being distributed in symmetric pairs (see Figs. 2, 3), so that the lift force on each leaf in a pair was roughly equal in magnitude but opposite in sign, such that the pair contributed negligible lift. However, we caution that this may not be true for all leaf morphology, because at large angles of attack the lift force is not necessarily symmetric. In addition, the influence of sheltering on the lift experienced by downstream leaves has not been defined and requires further study.

The ratio $A_e/A_r$ exhibited four regions of behavior. First, for $Ca < 1$ the plant was not deflected ($h/l = 1$, Fig. 8a) and consistent with this $A_e/A_r$ was close to 1 (Fig. 8b). For $Ca = 1–10$, the plant began to deflect ($h/l < 1$), but the drag ratio remained close to 1. In this range $\nu = -0.20 \pm 0.04 \ (A_e/A_r = (1.02 \pm 0.03)Ca^{-0.10 \pm 0.02})$ and $\nu = -0.04 \pm 0.02 \ (A_e/A_r = (1.01 \pm 0.02)Ca^{-0.02 \pm 0.01})$, based on the model prediction of $B = 0$ and $B = 20$, respectively. The Vogel number estimated from the measured drag fell in between these values, with $\nu = -0.06 \pm 0.04 \ (A_e/A_r = (1.0 \pm 0.1)Ca^{-0.03 \pm 0.02})$. The fact that drag remained high even as the plant deflected can be attributed to the leaves, which exhibit negligible reconfiguration. That is, for small stem deflections, the total area of leaves remained roughly constant, even as the projected areas of leaves on the upstream and downstream face of the stem changed. As the stem deflected, the angle between the leaves on the upstream face and the vertical first decreased until the leaves were in the vertical plane and then increased, such that the projected leaf area initially increased and then decreased. However, the projected area of the leaves on the downstream side of the stem first decreased until the leaves were in the horizontal plane and then increased. These competing trends resulted in little change in total frontal area when the deflection was small (image 1 and image 2 in Fig. 8). In addition, at small stem deflection leaf sheltering was not important.

For $10 < Ca < 1000$, the deflected plant height decreased rapidly with the increasing $Ca$ (image 3 and image 4 in Fig. 8). Over this range of deflection, sheltering became increasingly important. The sheltering was predicted to begin at $h/l = 0.91$ (Table 1), which occurred at $Ca = 5$. For $Ca = 10–1000$, the Vogel number was $\nu = -0.46 \pm 0.04 \ (A_e/A_r = (1.6 \pm 0.1)Ca^{-0.23 \pm 0.02})$, based on the measurements). This transition from a weaker to a stronger dependence on $Ca$ (from $\nu = -0.06$ for $1 < Ca < 10$ to $\nu = -0.46$ for $10 < Ca < 1000$) was also presented in Figure 4b in Harder et al. (2004). Note that the peak magnitude of $\nu = -0.46$ was smaller than the value for simple 2-D bending ($\nu = -2/3$) (Alben et al. 2002; Gosselin et al. 2010; Gosselin and De Langre 2011; Luhar and Nepf 2011) and 3-D bending ($\nu = -4/3$) of simple structures (Gosselin et al. 2010), and this difference likely reflects the role of the leaves.

Finally, for $Ca > 1000$ (Fig. 8 image 5), the plant posture was horizontal, such that the sheltering coefficient $C_s$ approached its limiting value $C_{s0}$. For perfectly horizontal stem posture, the model predicted a limiting value of $A_e/A_r = 0.25\left[\frac{\cos(x/2 - a)}{2\cos^2(a)} + \frac{\cos(x/2 + a)}{2\cos(a)}\right]$ = 0.25 tan 48° $\approx 0.28$ (Eq. 13,
shown with a horizontal arrow in Fig. 8b). A limiting value of finite magnitude is not predicted by the previous 2-D and 3-D scaling laws, but it is likely an important aspect in all real plant morphology.

**Reconfiguration of and drag on live plant B. caroliniana**

The predictions of plant posture \(h/l\) and drag were also validated with live samples of *B. caroliniana* (Fig. 9), for which \(\alpha = 112^\circ\), \(h/l = 0.99\), and \(C_{A_0} = 0.125\) (Table 1). The plant remained upright when \(Ca < 1\). For \(Ca > 1\), the deflection was dependent on plant buoyancy, which was higher for the longer plant samples. Plant buoyancy has been observed to delay the reconfiguration of a single blade until \(Ca^* > B\) (see discussion in Luhar and Nepf 2011). A similar tendency was seen here. Specifically, for the shorter plants with \(l = 9\) cm and 17 cm, \(B = 1.4\) and 8.6, respectively, and the strong reconfiguration (rapidly declining \(h/l\) in Fig. 9a) was delayed to \(Ca > 5\). For the longer plant \((l = 28\) cm), \(B = 37\), and the strong reconfiguration was delayed until \(Ca > 20\). Importantly, at the start of plant reconfiguration, the reconfiguration was accompanied by an increase in drag (see \(Ca = 5–100\) in Fig. 9b), and this behavior was captured by the model prediction (Fig. 9b). The greatest increase in drag (effective area) was 29\% for both the measured and predicted results. This phenomenon has also been observed for tree branches (Vogel 1984) and a poroelastic ball (Gosselin and De Langre 2011). The magnitude of drag increase in our study was comparable to that of the poroelastic ball (18\%, Figure 9 in Gosselin and De Langre 2011). The new model can explain this unusual trend. The *B. caroliniana* used in our experiment had a large leaf angle \(\alpha = 112^\circ\), so that the angle between the leaves and the flow was small when stem deflection was small (see image 1 in Fig. 9). For small stem reconfiguration, the angle between the leaves and incoming flow initially increased, producing a larger projected area and hence a larger drag force. Note that leaf sheltering was not significant at these low Cauchy numbers, because the plant deflected height was higher than or close to the critical plant deflected height \((h_*=0.99)\) at which sheltering began. A similar progression can be described for the poroelastic ball. With the introduction of flow, filaments that initially point upstream become more perpendicular to the flow, with an associated increase in drag, but as velocity (Cauchy number) increases further, these filaments eventually bend downstream, with an associated decrease in drag (Gosselin and De Langre 2011).

Three images are shown at the bottom of Fig. 9 corresponding to the posture of the plant at \(Ca = 1\), 20, and 2900. At the maximum Cauchy number 2900, the plant was almost horizontal, and \(A_e/A_r\) approached the model predicted limit \(A_e/A_r = 0.31\) (estimated by Eq. 13 with \(\theta = \pi/2\) and shown with the black arrow in Fig. 9b). At this high Cauchy number, slight vibration in the leaves and stem were observed. However, the leaves exhibited very little deformation, even at high Cauchy number, consistent with the model assumptions. The red lines indicate the predicted stem postures and illustrate a reasonable agreement between the observed and modeled postures. The maximum deviations between model and measurements in \(h/l\) and \(A_e/A_r\) are \(\pm 0.12\) and \(\pm 0.2\), respectively. Over all, the model made a quite good prediction in both plant deflection and drag force reduction considering the complexity of the real plant morphology.
Discussion

The sheltering effect on drag reduction

This study developed a new sheltering-reconfiguration model, which was validated with measurements of both plastic R. bonsai and live B. caroliniana. The model can be used to explore how different plant structure, specifically, leaf spacing, leaf width, and leaf angle $\alpha$, affect sheltering and impact drag. To focus on the influence of sheltering, the plant material density was assumed to be comparable to water so that $B = 0$.

Several species of live plant can be represented by the model, which assumes a single flexible stem and multiple leaves distributed along the stem, with leaves that undergo negligible reconfiguration (Table 2). For real plants, the leaf angle $\alpha$ ranges from 30° to 110°, and the ratio of leaf spacing to leaf width $d/l_w$ ranges from 0.5 to 3.5 (estimated from web images of live plant with the links given in Supporting Information Table A1.).

First, the impact of leaf spacing on stem deflection and drag was studied with $d/l_w = 0.5, 1, 2, 3,$ and 4 at a fixed leaf angle $\alpha = 48^\circ$ (the same as R. bonsai). Second, the impact of leaf angle on stem deflection and drag was considered for $d/l_w = 1$ and for a range of leaf angles, summarized in Table 3. Note that $A_l \mid \cos(\pi - \alpha) = A_l \mid \cos(\alpha)$, such that $90^\circ < \alpha < 180^\circ$ has the same result as $\pi - \alpha$. For each model run, Eqs. 13, 14, and 15 were used to define the limiting sheltering coefficient, $C_s$, and the corresponding equation of $C_s$ for each plant model, and these are listed in Table 3.

With a constant leaf angle ($\alpha = 48^\circ$), variation in leaf spacing had little impact on the plant posture (Fig. 10a), but a significant impact on plant drag (Fig. 10b). The horizontal arrows at the right hand axis of Fig. 10b indicate the minimum values of $A_e/A_r$, which were calculated by Eq. 13 with $\theta = \pi/2$, i.e., with horizontal stem posture. In each case, the effective frontal area approached the minimum value between $Ca = 1000$ and $10,000$. At the point of maximum stem deflection, leaf sheltering produced a great variation in effective frontal area, spanning $A_e/A_r = 1.1$ for leaf spacing $d/l_w = 4$ with

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no sheltering to \( A_e/A_r = 0.14 \) for leaf spacing \( d/l_w = 0.5 \). Note that for the leaf spacing \( d/l_w = 4 \), the drag increased with increased stem reconfiguration, which reflected the absence of leaf sheltering. This trend has not been previously reported. Also note that none of the plants listed in Table 2 exhibit this spacing. Most plants have leaf spacing 0.5–2, for which leaf sheltering contributes to a reduction in drag (\( A_e/A_r \)) during reconfiguration (Fig. 10b). Therefore, the model results suggest that the commonly observed leaf spacing (Table 2) is selected because it facilitates leaf sheltering. However, as shown in the next example, for larger leaf angles, enhanced drag with reconfiguration can occur for smaller spacing between leaves.

For a constant leaf spacing (\( d/l_w = 1 \)), variation in leaf angle, \( \alpha \), impacted the Cauchy number dependence of both deflected height and drag (Fig. 11). For small \( Ca \), the stem deflected height followed the same dependence observed for a stem without leaves (represented by the 2-D bending model shown with the dashed gray line in Fig. 11a). However, at higher Cauchy number the stem with leaves pronated more rapidly (smaller \( h/l \)) than the stem without leaves, and the deviation from simple 2-D bending (dashed curve) occurred sooner (at smaller \( Ca \)) for greater leaf angle. After deviating from the 2-D bending curve, the plants with leaves all followed the same dependence, \( h/l-Ca^{-0.48 \pm 0.01} \) (Fig. 11a). Despite the similar scale dependence observed in stem deflection, the variation in drag force with \( Ca \) exhibited great variation with the leaf angle \( \alpha \) (Fig. 11b). As expected, for \( Ca \leq 1 \), the plant did not deflect (\( h/l = 1 \)) and \( A_e/A_r = 1 \). For \( 1 < Ca < 10 \), drag force increased for \( \alpha = 60^\circ \) and \( 80^\circ \), but decreased for all smaller angles. For example, when \( \alpha = 80^\circ \) and \( Ca = 2 \), the drag force increased to three times the drag generated by a rigid plant. For \( 10 < Ca < 1000 \), drag declined with increasing \( Ca \) for all leaf angles, but the rate of decline increased with decreasing \( \alpha \), and the greatest decrease occurred for \( \alpha = 10^\circ \). For \( Ca > 1000 \), \( A_e/A_r \) approached the limiting value for horizontal posture, indicated with horizontal arrows, and this minimum drag decreased with decreasing leaf angle.

### A range of drag reduction for flexible plants

The way that reconfiguration impacts drag varies with plant morphology and rigidity. The range of behavior is illustrated in Fig. 12, which shows the ratio of drag on a flexible plant, \( F_r \), normalized by the drag on a rigid plant of the same

![Fig. 12](image-url)
morphology, \( F_r \). Rigid plants, such as mangrove roots and tree trunks, do not reconfigure and follow the quadratic drag law, such that \( F/F_r = 1 \) (black line in Fig. 12). A seagrass blade or simple cylindrical stem exhibits 2-D reconfiguration, follows the scaling \( F/F_r \sim \text{Ca}^{-13/2} (\alpha = -2/3) \) for \( \text{Ca} > 10 \) (gray curve in Fig. 12). A circular plastic sheet (Schouveiler and Boudaoud 2006), representative of some leaves, reconfigures in both the lateral and vertical dimensions, and follows the scaling for 3-D reconfiguration, \( F/F_r \sim \text{Ca}^{-2/3} (\alpha = -4/3) \) (Schouveiler and Boudaoud 2006; Gosselin et al. 2010; Gosselin et al. 2019) for \( \text{Ca} > 1 \) (dashed line in Fig. 12). Finally, plants with multiple leaves distributed along a flexible stem (Table 2) are influenced by both stem reconfiguration and leaf sheltering. For these plants, the dependence of drag on Cauchy number depends on plant structure, and in particular on the stem configuration, \( \alpha \). For example, when \( B. \, caroliniana \) \((d/l_w = 0.5, \alpha = 112^\circ) \) begins to reconfigure, the drag force increases over the range \( 1 < \text{Ca} < 30 \), and then decreases for \( \text{Ca} > 30 \) following \( F/F_r \sim \text{Ca}^{-0.22} (\alpha = -0.44) \) (thick black curve in Fig. 12). As listed in Table 2, leaf spacing most commonly falls between \( d/l_w = 0.5 \) and 2 and leaf angle between \( \alpha = 20^\circ \) and \( 80^\circ \), for which the model developed in this paper predicts a wide range of force variation with \( \text{Ca} \) represented by the light gray shading (Fig. 12).

Previous studies have considered Cauchy numbers up to the order of 1000, and focused on the trend of decreasing drag with increasing reconfiguration (Gosselin and De Langre 2011; Luhar and Nepf 2011; Li et al. 2018). However, the present study has highlighted that there is a limit in drag reduction, and the necessarily existence of a maximum deflection with a finite, minimum drag (e.g., Figs 3, 8). Luhar and Nepf (2011) noted that even as the form drag on a seagrass blade decreases with reconfiguration, the frictional drag will eventually become important when the majority of the blade is aligned with the flow. Figure 5 in Gosselin et al. 2010 also shows that the drag on a flexible plate tends away from the \(-4/3\) slope and levels off to a constant for Cauchy greater than 1000. The existence of a finite minimum drag is important to consider when modeling real vegetated systems.

**Conclusion**

This paper extended previous models (Gosselin et al. 2010; Gosselin and De Langre 2011; Luhar and Nepf 2011) describing the impact of plant reconfiguration on plant drag by considering the more complex morphology of a flexible stem with multiple leaves. The new model considered the sheltering between leaves using a sheltering coefficient, \( C_s \) (Eq. 15). The new model was validated by successfully predicting the reconfiguration and drag for the live plant \( B. \, caroliniana \). The sheltering of leaves provided an additional mechanism through which drag can be reduced during plant reconfiguration. The impact of sheltering was shown to depend on leaf size, spacing, angle of leaf to the stem, and the stem deflection. Importantly, a new regime was revealed for leaves attached to the stem at an angle close to \( 90^\circ \), in which the drag increased for an initial degree of stem reconfiguration, before subsequently decreasing with greater reconfiguration. In addition, new observations and modeling illustrated the existence of a finite minimum drag, which occurs when the stem reaches a horizontal posture, and which must be considered to properly represent real vegetation. This limit was not predicted by previous models.

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Conflict of Interest

None declared.

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Appendix A.

Table A1. List of links of webpage photos used to estimate $\alpha$ and $d/l_w$

| Name                  | Link                                                                 |
|-----------------------|-----------------------------------------------------------------------|
| Bacopa araguaia       | http://shuicao.cc/baike/view-1821.html                                |
| Bacopa colorata       | http://202.39.251.199/encyclopaedia/plant?plantID=207                 |
| Bacopa salzmannii     | http://202.39.251.199/encyclopaedia/plant?plantID=992                 |
| Bacopa monnieri       | http://shuicao.cc/baike/view-402.html                                 |
| Cuphea anagalloidea   | http://www.denatuurinhuis.nl/zen/index.php?album=Planten&image=cuphea-anagalloidea.jpg |
| Elatine orientaris    | http://202.39.251.199/encyclopaedia/plant?plantID=343                 |
| Hedyotis salzmannii   | https://pearlingplants.com/products/hedyotis-salzmannii-live-aquarium-plants |
| Limnophila fragrans   | https://www.aquagreen.com.au/plant_data/Limnophila_fragrans.html       |
| Lindernia grandiflora| https://www.aquaticplantcentral.com/forumapc/plantfinder/details.php?id=340 |
| Lindernia parviflora  | http://www.staraqua.ru/content/lindernia-parviflora-variegated         |
| Micranthemum umbrosum| https://aqualabaquaria.com/products/micranthemum-umbrosum             |
| Nesaea icosandra      | http://202.39.251.199/encyclopaedia/plant?plantID=499                 |
| Nesaea triflora       | http://www.staraqua.ru/content/nesaea-triflora                         |
|                       | http://202.39.251.199/encyclopaedia/plant?plantID=974                 |

(Continues)
Table A1. Continued

| Name                        | Link                                              |
|-----------------------------|---------------------------------------------------|
| Oldenlandia salzmannii      | http://shuicao.cc/baike/view-1881.html             |
| Rotala indica               | https://www.aquariumplantsfactory.com/products/rotala-macrandra?variant=28553780428861&currency=USD&utm_campaign=gs-2018-12-09&utm_source=google&utm_medium=smart_campaign |
| Rotala macrandra            | http://202.39.251.199/encyclopaedia/plant?plantID=242 |
| “green”                     | http://shuicao.cc/baike/view-1913.html             |
| Rotala macrandra variegata  | http://shuicao.cc/baike/view-3520.html             |
| Rotala sp. “indian”         | http://202.39.251.199/encyclopaedia/plant?plantID=249 |
| “mini type 1”               |                                                   |