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A model of the lateral line of fish for vortex sensing

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Abstract
In this paper, the lateral line trunk canal (LLTC) of a fish is modeled to investigate how it is affected by an external flow field. Potential flow theory is adopted to model the flow field around a fish’s body in the presence of a Karman vortex street. Karman and reverse Karman streets represent the flow patterns behind a bluff body and a traveling fish, respectively. An analytical solution is obtained for a flat body, while a fish-like body is modeled using a Joukowski transformation and the corresponding equations are solved numerically. The pressure distribution on the body surface is then computed employing Bernoulli’s equation. For a known external flow, the flow inside the LLTC is driven by the pressure gradient between a pair of consecutive pores, which can be solved analytically. Governing dimensionless parameters are obtained from this analytical solution, and the effects of these numbers on the amplitude or features of the velocity distribution inside the canal are studied. The results show that the main characteristics of a vortex street including the magnitude of vortices, their translational speed, their spacing, their distance from the fish’s body and the angle of the vortex street axis can all be recovered by measuring the velocity distribution along the canal and its changes with time. To this end, the proposed LLTC model could explain how a fish identifies the characteristics of a Karman vortex street shed by a nearby object or a traveling fish. It is also demonstrated that while this model captures the ac (alternating current) component of the external velocity signal, the dc (direct current) component of the signal is filtered out. Based on the results of our model, the role of the LLTC in a fish’s schooling and its evolutionary impact on fish sensing are discussed.

(Some figures may appear in colour only in the online journal)

Nomenclature

\(a\) Radius of cylinder
\(b\) Vertical distance between vortices
\(c, c^*\) Chord length of fish’s body, and its dimensionless form
\(c_0\) Coefficient of Joukowsky transformation
\(d, d^*\) Diameter of lateral line canal, and its dimensionless form
\(h, h^*\) Vertical distance between fish and vortex street, and its dimensionless form
\(l\) Horizontal distance between vortices
\(L\) Length of lateral line
\(P\) Total pressure at far field
\(P_0\) Total pressure at far field
\(P_L\) Pressure at left end of pipe
\(P_R\) Pressure at right end of pipe

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1. Introduction

Fish possess a mechanosensory lateral line system, which responds to the motion of the surrounding water relative to the fish’s skin. This sensing system could consist of a superficial neuromast (SN) and a canal neuromast (CN) subsystems. SNs are located on the outer surface of a fish, extending into the external fluid, while CNs are buried inside the lateral line canal (see figure 1). The basic sensing unit of the two is the fish’s hair cell. Such hair cells will deflect as the result of oncoming flow, thus allowing for detection. SNs are generally smaller in diameter but greater in height than CNs and contain fewer hair cells [1]. Research on the lateral line has shown that SNs respond to changes in external fluid velocity, whereas CNs respond to changes in external flow acceleration (associated with corresponding changes in external flow pressure) [2]. A simple interpretation would be that CNs sense information regarding pressure gradients, while SNs sense the velocity of the surrounding flow. From another perspective, SNs respond best to the dc (direct current) and low-frequency components (less than approximately 30 Hz) of the incoming flow, whereas CNs respond best to high-frequency components of the flow (approximately 30–150 Hz) [3]. The lateral line organ is capable of diverse sensing tasks, ranging from the detection of near field motion [4] produced by prey [5], predators [6] or during schooling [7] to the perception of water currents and static obstacles [8]. Although individual behavioral roles of SNs and CNs have not been well distinguished, it has been found that SNs control rheotaxis at low current velocities [9], whereas CNs control the orienting response of fish to the initial prey detection [3]. It is believed that CNs are important for fish schooling [10].

In hydrodynamic studies of the lateral line, models using potential flow theory have been proposed to find the pressure distribution on a fish’s skin in the presence of a stimulus. In potential flow theory, the fluid’s viscosity is neglected; thus, the boundary conditions around a solid body require only that the velocity normal to the wall be equal to the wall’s velocity. The potential flow model serves to approximate the pressure outside the boundary layer. This pressure essentially equals the pressure on the body’s surface, because the pressure across the thickness of a thin boundary layer can be accurately approximated as being constant [11]. Consequently, potential flow is a fair model as long as the study is confined to CNs that respond to pressure gradient. The potential dipole source flow equations have been used to model a vibrating sphere near a fish’s body [12]. The same equations were used by Kroese to compute the slip velocity along the fish’s skin caused by a nearby vibrating sphere [13]. Potential flow around a three-dimensional slender body with circular cross-section has been obtained by Handelsman and Keller [14]. Hassan adapted this solution to investigate the cases of fish moving in open water, gliding toward a plane’s surface [15], gliding alongside or above a plane’s surface [16], and near an oscillating sphere [17]. Various stimuli sensed by the lateral line have been the subject of many research investigations over the years. These stimuli usually involve a vibrating sphere, moving objects, an approaching wall and vortices [18–23]. In this paper, we specifically consider Karman vortex street sensing. We will develop a model of the lateral line sensing and show that
measurement in such a model provides enough information to
detect the most important parameters of such a vortex street,
namely the vortex strength, the lateral and flow wise spacings,
translational velocity and the orientation of the vortex street.

The ability of a fish to detect vortices by its lateral line
has been demonstrated in experiments [24, 25], Franosch et al
[26] followed the method used by Hassan [15–17] to model
the flow due to a three-dimensional vortex ring next to the
fish’s body. Their results showed that information about the
orientation of a vortex ring is encoded in the input of the lateral
line, and thus is accessible to the nervous system. Barbier has
studied numerically the two-dimensional flow field due to a
vortex street behind a prism [27]. He also investigated the
three-dimensional flow inside the LLTC in the presence of
the hair cells. However, he studied the flow in a subsegment
between two adjacent pores but did not consider this internal
flow over the entire length of the canal. Because of openings
along the canal, internal flow velocity is expected to be non-
uniform, which suggests that the distribution of velocity along
the LLTC may contain rich information on external flow. The
purpose of this study is to investigate the information captured
by the LLTC in the presence of a Karman vortex street and
to determine the relevance of this for schooling behavior.
Potential flow theory is used to create a two-dimensional
model of the external flow field created by a gliding fish,
and the pressure distribution along the fish’s surface is
calculated accordingly. The internal flow inside the canal is
also modeled analytically to calculate the velocity distribution
along the LLTC. Finally, this information is inverted in order
to determine how the original flow parameters may be gleamed
from the local data available to a fish. A preliminary result
from our group in this direction was presented in [28].

2. Theoretical modeling

In this section, a fish’s body is approximated by a flat plate,
and a two-dimensional potential flow model is presented
to determine the pressure field around this body next to a
vortex street, after which an analytical model is developed
to characterize the flow inside the LLTC. For simplicity, this
model is one-way coupled, meaning that the flow inside the
LLTC is driven by the external flow with no feedback from
internal to external flow. That is an acceptable assumption,
considering the higher viscosity of the fluid inside the canal
and small pore sizes which results in a very low Reynolds
number flow inside the canal.

2.1. External flow field modeling

In potential flow theory, an infinite single row of vortices with
separation l and strength \( \Gamma \) has a complex potential

\[
W(z) = \phi + \psi i = \frac{\Gamma}{2 \pi i} \sum_{n=-\infty}^{\infty} \ln (z - nl) = \frac{\Gamma}{2 \pi i} \ln \sin \left( \frac{\pi z}{l} \right).
\]

The wake of an obstacle or a swimming animal is generally
in the form of a Karman vortex street, which is a staggered
arrangement of two vortex rows with opposite orientations.

With the vertical spacing between two vortex rows denoted by
b, the stability analysis on this vortex street indicates that the
ratio b/l has a constant value of about 0.28. Accordingly, the
vertical distance between vortices is kept at b = 0.28l in
the present study. In a static fluid, such a double-layered vortex
street moves itself at an induced velocity \( u_0 \) which is a function
of the vortex street properties as given by

\[
u_0 = -\frac{\Gamma}{2l} \cot \left( \frac{\pi}{l} (l/2 + bi) \right) = \frac{\Gamma}{2l} \tan \left( \frac{\pi bi}{l} \right).
\]

Here, the sign of \( \Gamma \) is determined by the upper vortex street.
Superposing a free stream velocity, the total velocity of this
vortex street becomes

\[
u_z = U_\infty + u_0 = U_\infty - \frac{\Gamma}{2l} \cot \pi \left( 0.5 + 0.28i \right) = U_\infty + \frac{0.3536 \Gamma}{l}.
\]

At this point, we introduce a fish in this background flow.
We will consider two simplifying cases. In the first case, the
fish is modeled as a simple flat plate. While this is simplistic,
it is a reasonable model for fish with a low thickness-to-length
ratio. As will be shown shortly, the case of the flat plate in
a vortex street background flow can be solved analytically.
This is quite helpful in order to understand the role of each
parameter in the overall flow. For a more realistic presentation
of a fish, what we call a fish-like body in the following sections,
we will present a two-dimensional fish by a symmetrical
airfoil. As a boundary layer forms around the fish’s body
in the presence of a free stream, its surface approximates a
straight line parallel to the fish’s body [29]. As noted earlier,
the pressure predicted by the potential flow outside the boundary
layer is equal to the pressure inside the boundary layer. For
simplicity we assume that this flat body lies along the x axis
of the complex z-plane with a Karman vortex street parallel
to one side of the fish’s body; see figure 2. In order to make
the body surface streamlined, the method of images is used
by introducing another vortex street on the other side of the
body, so that the flow field is symmetric about the x axis. The
complex potential of such a flow field can be written as

\[
W(z, t) = U_\infty z + \frac{\Gamma}{2 \pi i} \left[ \ln \sin A - \ln \sin B - \ln \sin C + \ln \sin D \right].
\]
where
\[ A = \frac{\pi}{l} [z - u_t + 0.5l - i(h + 0.14l)] , \] (5)
\[ B = \frac{\pi}{l} [z - u_t + 0.5l + i(h + 0.14l)] , \] (6)
\[ C = \frac{\pi}{l} [z - u_t - i(h - 0.14l)] , \] (7)
\[ D = \frac{\pi}{l} [z - u_t + i(h - 0.14l)] . \] (8)

Here \( A, B, C \) and \( D \) terms indicate the locations of the four vortex trains. The complex velocity is then given by the derivative of the complex potential, \( W \), to be
\[ u - vi = \frac{\partial W}{\partial \zeta} = U_{\infty} + \frac{\Gamma}{2\pi i} [\cot A - \cot B - \cot C + \cot D] . \] (9)

On the body’s surface, where \( z = x + \Omega i \), the velocity normal to the body’s surface should be zero, namely \( v|_{z=\Omega} = 0 \). Thus, we have the surface velocity
\[ V_{\text{surf}} = u|_{z=\Omega} = U_{\infty} + \frac{\Gamma}{2\pi i} [\cot A - \cot B - \cot C + \cot D]|_{z=\Omega} . \] (10)

Owing to the symmetry of the flow field, the stream function on the body equals zero and this value does not change over time. As a result, the complex potential, \( W \), on the body is a real-valued function which equals the velocity potential, \( \phi \). Accordingly, the time derivative of the complex potential is also a real-valued function
\[ \frac{\partial W}{\partial t}|_{z=\Omega} = \frac{\partial \phi}{\partial t}|_{z=\Omega} = \frac{\Gamma u_t}{2\pi i} [-\cot A + \cot B + \cot C - \cot D]|_{z=\Omega} . \] (11)

Bernoulli’s equation can be employed in order to obtain a pressure distribution \( P \) along the fish’s body
\[ P_{\text{surf}} = \left[ P_0 - \frac{1}{2} \rho V_{\text{surf}}^2 - \mu \frac{\partial \phi}{\partial r} \right]|_{z=\Omega} . \] (12)

Here, \( P_0 \) is the total pressure at the far field. For convenience, we let \( P_0 = \frac{1}{2} \rho U_{\infty}^2 \). Substituting (3), (10) and (11) into (12), one obtains the following equation for the pressure distribution along the fish’s body:
\[ P_{\text{surf}} = \rho \frac{\Gamma^2}{8l^2} (\cot A - \cot B - \cot C + \cot D)^2 \]
\[ + \frac{0.3536 \rho \Gamma^2}{2l^4} (\cot A - \cot B - \cot C + \cot D) . \] (13)

The created pressure gradient along the body of the fish is the driving force behind the flow inside the LLTC. For simplicity, we assume that the pressure on the canal openings is determined only by the external flow and not affected by the flow inside the canal, meaning that the outside and inside flows are one-way coupled.

2.2. Model of the flow inside the canal

In the previous section, we obtained the pressure distribution on the body’s surface. In order to predict the response of hair cells inside the canal to this external pressure, a model of the flow inside the canal is required. In this study, the canal is modeled as a long, slender circular pipe with pores uniformly distributed along the surface of the pipe at certain intervals (see figure 3). For simplicity, the presence of hair cells inside the canal is not considered in the flow calculations. While the morphological characteristics of the trunk lateral line neuromasts vary across different species, averaged common values are used for several parameters in our model. Consequently, the diameter of the pipe, \( d \), is taken to be 250 \( \mu \text{m} \), the diameter of each pore, \( d_p \), to be 250 \( \mu \text{m} \) and the interval between two adjacent pores, \( \Delta L \), to be 4 mm [30]. Because the fluid inside the canal is more viscous than that outside [31] and the diameter of the canal is small, the Reynolds number for flow is much smaller than 1. As a result, the governing momentum equation of this flow simplifies to Stokes’ linear equation.

The flow inside the canal is driven by the pressure gradient between adjacent pores. The linearity of the governing equation allows one to consider the flow in each segment separately. Previous computational results indicate that the value of the pressure at a cross section of the canal located at the pore is essentially the external pressure at that opening [27]. Therefore, the pressure on the body’s surface where the canal openings are located determines the flow inside the canal.

For an infinitely long and periodic vortex street passing a fish, a periodic pressure wave is expected at each pore location. Accordingly, the pressure gradient between adjacent pores can be expanded in terms of a Fourier series. Here, the case of cosineloidal pressure difference is considered, namely
\[ P_L - P_R = |P_L - P_R| \cos \omega t , \] (14)
where \( P_L \) and \( P_R \) are the pressures at the left and right ends of the pipe, respectively, \( |P_L - P_R| \) and \( \omega \) are the wave amplitude and angular frequency, respectively.

Flow driven by a periodic pressure gradient inside a circular pipe of diameter \( d \) and length \( \Delta L \) has been investigated before by analytical techniques [32]. As the kinematic Reynolds number \( \omega |r| = \omega (d/2)^2/\nu < 4 \), where \( \nu \) is the kinematic viscosity of the fluid inside the pipe, the induced velocity in the pipe is in phase with the pressure oscillation. The flow velocity \( u_{\text{in}} \) of the fluid inside the pipe, written in cylindrical coordinates, is given by
\[ u_{\text{in}}(r) = \frac{K}{4\nu} [(d/2)^2 - r^2] \cos \omega t \]
\[ = \frac{1}{4\mu \Delta L} [(d/2)^2 - r^2] (P_L - P_R) , \] (15)
where \[ K = |P_L - P_R|/\rho \Delta L , \] (16)
and \( \rho \) and \( \mu \) are the density and dynamic viscosity of fluid inside the canal, respectively. We digress here to discuss the validation of this assumption. A small \( \omega |r| \) can be obtained,
as long as the oscillation frequency of the pressure gradient is low. Since the viscosity, $\nu$, of the fluid inside the canal is approximately five times that of water [31], the assumption of $\omega_0^2 < 4$ is satisfied as long as the frequency is below 200 Hz. The exact frequency limit for the validity of this assumption changes slightly with the canal diameter and fluid viscosity inside the canal which both vary from one species of fish to another. For the pressure gradient created by a vortex street, it can be shown that this stimulus contains only low-frequency components and hence the model applies for this specific case. For an arbitrary stimulus that changes in time, however, the model only applies to its low-frequency components. In other words, the model is valid for stimuli of which the leading components are within the frequency limitation. Nevertheless, the stimuli detected by fish are often in the low-frequency range [33].

According to equation (15), the amplitude of the velocity inside the canal is proportional to the pressure gradient between two adjacent pores, and the change of velocity is in phase with the pressure oscillation. For simplicity, here we assume that the spacing of adjacent pores is infinitely small so that the pressure gradient $dp/dx$ could be approximated by $\Delta p/\Delta L$. In this sense, the flow velocity distribution inside the canal is assumed to be driven by a continuous pressure gradient. The validation of this assumption and its produced errors will be discussed in a later section.

The local flow velocity inside the canal is characterized by the mean flow velocity over the cross-sectional area of the canal. By substituting (13) into (15) in the limit of $\Delta L \to 0$ and averaging over $r$, one obtains

$$V_m = \bar{u}_m = \frac{\pi d^2 \Gamma^2}{128 \nu l^3} \left[ (\cot^2 A - \cot^2 B + \cot^2 C - \cot^2 D) \times (\cot A - \cot B - \cot C + \cot D - 0.77072) \right].$$

(17)

For simplicity, no interactions between the inside flow and the neuromasts are considered here, but one can expect the local flow velocity to be detected by hair cells located between adjacent pores. As has been shown by van Netten, the combined frequency response of the canal and neuromast results in a nearly constant sensitivity to pressure gradient for frequencies up to approximately 100 Hz [34]. This enables us to neglect the frequency response of the system, since the problem considered here is in the low-frequency range.

Up to this point, we have derived an analytical expression of the inside flow velocity that is related to the external flow field. Yet this equation needs to be nondimensionalized in order to understand the effect of each variable.

2.3. Nondimensionalization

There are nine parameters involved in the velocity equation (17) inside the canal. These are

$$U_\infty, \quad V_m, \quad \Gamma, \quad x, \quad l, \quad h, \quad d, \quad t, \quad \nu \quad \left[ LT^{-1} \right], \left[ LT^{-1} \right], \left[ L^2 T^{-1} \right], \left[ L \right], \left[ L \right], \left[ L \right], \left[ T \right], \left[ L^2 T^{-1} \right].$$

(18)

The dimension of each variable is shown in brackets. Using the Buckingham $\Pi$ theorem, one can write the expected functional relationship as

$$\pi^* = \frac{U_\infty V_m \Gamma x l h d t \nu}{\Gamma 1^{[7]}}. \quad \left[ \frac{LT^{-1}}{\nu} \right] \quad \left[ \frac{LT^{-1}}{} \right] \quad \left[ L^2 T^{-1} \right] \quad \left[ L \right] \quad \left[ L \right] \quad \left[ L \right] \quad \left[ T \right] \quad \left[ L^2 T^{-1} \right].$$

(19)

Considering that the rank of the coefficient matrix is 2, one expects seven independent nondimensional numbers. In the following, we use $l$ and $U_\infty$ as the proper length and velocity scales for nondimensionalization. These seven nondimensional numbers are

$$x^* = \frac{x}{l}, \quad h^* = \frac{h}{l}, \quad d^* = \frac{d}{l}, \quad \Gamma^* = \frac{\Gamma}{U_\infty l}, \quad V^* = \frac{V_m}{U_\infty}, \quad t^* = \frac{\Gamma 1^{[7]}}{U_\infty}, \quad \nu^* = \frac{\nu}{U_\infty l}.$$

(20)

Accordingly, the dimensionless internal velocity equation is

$$V^* = \frac{\pi (d^* \Gamma^*)^2}{128 \nu l^3} \times \left( \cot A - \cot B - \cot C + \cot D - 0.77072 \right).$$

(21)

with

$$A^* = \pi [x^* + (0.7111^* - 1)t^* + 0.5 - 5(i(h^* + 0.14))],$$

(22)

$$B^* = \pi [x^* + (0.7111^* - 1)t^* + 0.5 + i(h^* + 0.14)),$$

(23)

$$C^* = \pi [x^* + (0.7111^* - 1)t^* - i(h^* - 0.14)],$$

(24)

$$D^* = \pi [x^* + (0.7111^* - 1)t^* + i(h^* - 0.14)].$$

(25)

As seen, $A^*$, $B^*$, $C^*$ and $D^*$ terms are functions of $\Gamma^*$, $x^*$, $h^*$ and $t^*$. At the initial time $t^* = 0$, these terms reduce to

$$A_0^* = \pi [x^* + 0.5 - i(h^* + 0.14)],$$

(26)

$$B_0^* = \pi [x^* + 0.5 + i(h^* + 0.14)],$$

(27)

$$C_0^* = \pi [x^* - i(h^* - 0.14)],$$

(28)

$$D_0^* = \pi [x^* + i(h^* - 0.14)].$$

(29)

We now recast equation (21) to its nondimensional form to read

$$\frac{128 \nu l^3 \pi}{\pi (d^* \Gamma^*)^2} \times \left( \cot A_0^* - \cot B_0^* - \cot C_0^* + \cot D_0^* \right) \times \left( \cot A_0^* - \cot B_0^* - \cot C_0^* + \cot D_0^* - 0.77072 \right).$$

(30)

Now defining a rescaled velocity

$$\Pi := \frac{128 \nu l^3}{\pi (d^* \Gamma^*)^2} \times \left( \cot A_0^* - \cot B_0^* - \cot C_0^* + \cot D_0^* - 0.77072 \right) \times \left( \cot A_0^* - \cot B_0^* - \cot C_0^* + \cot D_0^* \right).$$

(31)

equation (30) can be reduced to

$$\Pi = g_0(x^*, h^*).$$

(32)

From this equation, one can see that the contribution of $\Gamma$, $l$, and $d$ to the inside velocity is simply to scale its amplitude uniformly along the canal, while the general shape of the velocity distribution along the canal is given by $\Pi = g_0(x^*, h^*)$. This dimensionless function is plotted in figure 4. Since, for each $h^*$, the velocity profile is determined uniquely by (32) up to a scaling factor, one may think of $h^*$ as a parameter which controls the characteristic features (e.g., locations of peaks and troughs) of the velocity profile.

We now turn our attention to equation (21) in order to investigate the time dependence as the vortex street translates with respect to the fish. Since the effect of increasing $t^*$ is equivalent to increasing $x^*$, the shape of the velocity plot does not change as the vortex street moves in time, $t^* > 0$, and it moves downstream at the same speed as the vortex street $u_c$. 

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flow velocity is proportional to $h$ around $x^* = 0$. This section degenerates and eventually disappears with increasing vortex strength, $x^*$, the streamwise spacing between vortices, $h^*$ and $d$, while the dependence of the velocity on $x^*$ and $h^*$ is captured by the function $g_0(x^*, h^*)$. This function has a periodicity of 1, inherited by the periodicity of the problem.

As shown in figure 4, the features of $\Pi = g_0(x^*, h^*)$ plot along the $x^*$ axis change as the value of $h^*$ is varied. Consequently, one may expect to be able to calculate $h^*$ from these changes. For small values of $h^*$, a near-flat section is found between a positive peak and a negative trough. As $h^*$ increases, this section degenerates and eventually disappears around $h^* = 0.5$. Besides small shifts in the locations of the peaks and troughs, no visible qualitative change in features of the scaled velocity $\Pi$ is observed for constant $h^* > 0.5$. The distance between adjacent peaks and troughs is found to depend exclusively on $h^*$. This conclusion can also be verified by analyzing the analytical solution. The $x^*$ locations of the peaks and troughs of the function $g_0(x^*, h^*)$ can be determined by

$$\frac{\partial g_0(x^*, h^*)}{\partial x^*} = 0.$$  (33)

This calculation results in two roots, namely $x_L = h_1(h^*) + n$ and $x_R = h_2(h^*) + n (n = 1, 2, 3, \ldots)$, which represent the locations of the peaks and troughs, respectively. Therefore, the distance between an adjacent peak and trough can be calculated to be $x_L - x_R = h_2(h^*) - h_1(h^*)$. This relation is shown in figure 5. As the value of $h$ is easily determined from the mean velocity distribution inside the canal, the value of $h$ is simply obtained by $h = h^* l$.

Now, assume that a fish can measure the velocity magnitude $V_{in}$ inside the canal at the location of the hair cells. As discussed before, a fish is able to determine $h^*$ from the measured velocity inside the canal. It also has access to information on the dimensionless velocity $\Pi$ at the same location through the functional relation of $\Pi$ to $h^*$ and $x^*$,

Figure 4. (a) The surface plot of $\Pi = g_0(x^*, h^*)$. (b) The plot of $\Pi = g_0(x^*, h^*)$ as a function of $x^*$ for several values of $h^*$. Since the inside flow velocity is proportional to $\Pi$, the shape of the curve is essentially the velocity distribution along the entire canal.

2.4. Analytical results for a flat body

In the previous section, we investigated the qualitative dependence of each dimensionless variable to the velocity distribution along the canal. This velocity distribution is driven by the differential pressure at the pores which represent the external pressure outside a thin boundary layer. To this end, one might hypothesize that a fish senses the flow inside its lateral line canal without direct knowledge of the external velocity field. This is in contrast to SNs where it is believed that the external pressure outside a thin boundary layer. To this end, one might hypothesize that a fish senses the flow inside its lateral line canal without direct knowledge of the external velocity field. This is in contrast to SNs where it is believed that the external velocity field is directly measured. It is of interest to see how much information can potentially be retrieved by a fish if it is able to precisely measure the flow velocity inside the lateral line canal. In this section, we will study the possibility of recovering these variables once the velocity distribution inside the canal is measured.

The main parameters identifying a vortex street are the vortex strength, $\Gamma$, the streamwise spacing between vortices, $l$, the distance between the vortex street and the body, $h$, and the velocity of the vortex street, $u_v$. According to the analytical solution given by equation (17), the flow velocity inside the canal is a function of these variables. As previously seen in equations (31) and (32), the amplitude of the velocity inside the canal is controlled by $\Gamma$, $l$ and $d$, while the dependence of the velocity on $x^*$ and $h^*$ is captured by the function $g_0(x^*, h^*)$. This function has a periodicity of 1, inherited by the periodicity of the problem.

As shown in figure 4, the features of $\Pi = g_0(x^*, h^*)$ plot along the $x^*$ axis change as the value of $h^*$ is varied. Consequently, one may expect to be able to calculate $h^*$ from these changes. For small values of $h^*$, a near-flat section is found between a positive peak and a negative trough. As $h^*$ increases, this section degenerates and eventually disappears around $h^* = 0.5$. Besides small shifts in the locations of the peaks and troughs, no visible qualitative change in features of the scaled velocity $\Pi$ is observed for constant $h^* > 0.5$. The distance between adjacent peaks and troughs is found to depend exclusively on $h^*$. This conclusion can also be verified by analyzing the analytical solution. The $x^*$ locations of the peaks and troughs of the function $g_0(x^*, h^*)$ can be determined by

$$\frac{\partial g_0(x^*, h^*)}{\partial x^*} = 0.$$  (33)

This calculation results in two roots, namely $x_L = h_1(h^*) + n$ and $x_R = h_2(h^*) + n (n = 1, 2, 3, \ldots)$, which represent the locations of the peaks and troughs, respectively. Therefore, the distance between an adjacent peak and trough can be calculated to be $x_L - x_R = h_2(h^*) - h_1(h^*)$. This relation is shown in figure 5. As the value of $l$ is easily determined from the mean velocity distribution inside the canal, the value of $h$ is simply obtained by $h = h^* l$.

Now, assume that a fish can measure the velocity magnitude $V_{in}$ inside the canal at the location of the hair cells. As discussed before, a fish is able to determine $h^*$ from the measured velocity inside the canal. It also has access to information on the dimensionless velocity $\Pi$ at the same location through the functional relation of $\Pi$ to $h^*$ and $x^*$,
Therefore, a fish has enough information to calculate the strength of the vortices in a vortex street from
\[
\Gamma = \sqrt{\frac{\pi d^2 \Pi}{128 \nu V_{in} l^3}},
\]
(34)
where the canal diameter, \(d\), and the viscosity of fluid inside the canal, \(\nu\), are known for a given fish.

The translation of a vortex street outside the fish’s body is accompanied by a similar periodic variation in the measured velocity inside the canal. As a result, the velocity \(u_s\) of the vortex street is essentially the rate of motion of the sensed velocity peaks inside the canal. Therefore, one can imagine that a fish detects \(u_s\) by detecting the distance traveled by the sensed peak or trough velocity during a certain time period.

So far we have developed an analytical solution for the flow around a fish represented by a 2D flat surface in the presence of a vortex street. We were also able to show that there is enough sensed information in an LLTC to recover all of the main parameters of a vortex street. A summary of our discussion for detecting these parameters is shown in figure 6.

In order to extend this analytical analysis from a flat plate to a fish-like body in the presence of a vortex street, a few of the steps need to be conducted computationally. This will enable us to investigate the effect of body thickness on the sensing capabilities of an LLTC. This is presented in the following section.

**3. Numerical method for fish-like body**

**3.1. Joukowski transformation**

In this section, we approximate the shape of a fish by a symmetric airfoil. In order to take advantage of some of our theoretical findings from the previous section, we restrict this study to the realm of potential flow theory. The shape of a fish’s body can be obtained using a Joukowski transformation [35]. The flow over a cylinder of radius \(a\) located at a point \((c_0 - a, 0)\) can be mapped onto a symmetric airfoil (see figure 7), namely a fish-like body, by
\[
z = \xi + \frac{c_0}{\xi},
\]
(35)
The chord length, \(c\), and the thickness, \(\delta\), of the body are then determined by
\[
c = 2c_0 - \frac{(c_0 - 2a) + c_0^2}{(c_0 - 2a)},
\]
(36)
\[
\delta = \frac{3\sqrt{3}}{2} (a - c_0).
\]
(37)

The Karman vortex street is modeled by placing two rows of staggered, periodic 2D vortices on one side of the airfoil. We are not able to obtain a simple closed form analytic formulae for this case. As a result, we model the infinite array of vortices by including enough terms in the series so that the value of velocity potential difference at a specific surface point, after one cycle of the finite vortices moving downstream, is less than 0.5% of the initial potential value. In this sense, the resulting flow field near the body is a good approximation to the infinite vortex street case. The locations of the point vortices are first determined in the \(z\)-plane, and then their corresponding locations in the \(\xi\) plane are calculated according to the inverse Joukowski transformation. Solving the inverse Joukowski transformation (\(z\) to \(\xi\) coordinate mapping) in equation (35) for \(\xi\) results in a multi-valued solution
\[
\xi = \frac{1}{2} (z \pm \sqrt{z^2 - 4c_0^2}).
\]
(38)
The implication of this function is that there are two possible \(\xi\) coordinates in the cylinder frame (\(\xi\)-plane) for each \(z\) coordinate in the airfoil frame (\(z\)-plane). One of the \(\xi\) coordinates is outside the cylinder and the other is inside the cylinder. Therefore, in order to properly map a \(z\) coordinate to the appropriate \(\xi\) coordinate, the following condition is applied in order to ensure that the mapped area is outside the cylinder. The \(\xi\) coordinate corresponding to a given vortex located at a \(z\) coordinate will be the solution in equation (38) if the distance
potential in this case can be finally written as

$$W(\zeta) = U_\infty \left[ \zeta + \frac{2}{\zeta} \sum_{n=1}^{N} \left[ \ln(\zeta - \zeta_0) + \ln(\zeta - \zeta_n) - \ln(\zeta - \zeta_0 - \frac{a^2}{\zeta_n - \zeta_0}) \right] \right].$$

where $\zeta_0$ is the center of the cylinder and $\zeta_n$ is the coordinate of each vortex with its complex conjugate denoted by $\bar{\zeta}_n$. The real part of the above function is the velocity potential $\phi$. The flow velocity on the body surface can be determined as the gradient of this function on the fish’s body surface, that is,

$$U(z) = \sqrt{\left( \frac{\partial \phi}{\partial x} \right)^2 + \left( \frac{\partial \phi}{\partial y} \right)^2}.$$  \hspace{1cm} (40)

Assuming a quasi-steady flow, the velocity potential around the fish at any instant of time can be modeled by the 2D potential in equation (39). In order to obtain the pressure distribution around the fish’s body, $P(z)$, we employ the Bernoulli equation

$$\left[ \frac{P(z)}{\rho} + \frac{1}{2}U^2(z) + \frac{\partial \phi}{\partial r} \right]_{r=a} = P_0,$$  \hspace{1cm} (41)

where $P_0$ is the total pressure at the far field. Just as with our analytical model in the previous section, here we let $P_0 = \frac{1}{2}\rho U_\infty^2$ for convenience. Once we have computed the external pressure distribution, we may continue to use the same analytical model presented in the previous section to calculate the flow inside the canal. The results of this numerical model are discussed in the next sections in comparison with the results from our analytical calculations for a flat-body object.

### 3.2. Effects of body thickness

To evaluate the accuracy of the numerical method, we consider the flow around a flat-body object as a test case. A flat-body object could be considered as the limit of a fish-like body from the previous section when the body thickness approaches zero. In this case we will be able to compare our numerical procedure with the exact analytical result for a flat-body object. In order to evaluate the effect of body thickness, we define the thickness ratio, as illustrated in figure 8(a), to be body thickness over body length

$$\lambda = \delta/c.$$  \hspace{1cm} (42)

Figure 8(b) shows the variation in the pressure distribution as $\lambda$ is changed from a flat plate to a relatively rounded fish body with $\lambda = 0.08$. For larger $\lambda$ when the fish’s body is rather bluff, a sharp drop in pressure along the body is observed as one moves away from the stagnation point on the fish’s head. After that initial drop, pressure gradually increases along the body toward the tail. By increasing the thickness ratio, the pressure around the head region will be dominated by the stagnation pressure, which is dictated by the incoming free stream. The existence of a vortex street hardly alters the pressure distribution in this region. Since the induced velocity in the canal is proportional to the external pressure gradient, the inside velocity in this region does not contain much information on the vortex street. Also based on the fact that the LLTC often does not extend to the fish’s head, in this study we assume that the LLTC ranges from 0.1$c$ to 0.9$c$ along the fish’s body.

For convenience, the value of $l$ is measured in the cord length, $c$, units. The variable with a star is still scaled by $l$ as previously defined. The data shown in figure 9(a) use the same $\lambda$ values as before with $c^* = 2$, $h^* = 0.5$ and $\Gamma^* = 0.6$. For smaller values of $h^*$, where the vortex street moves closer to the fish’s body, a secondary positive peak-velocity appears on the left side of the absolute peak-velocity value by increasing the value of the thickness ratio $\lambda$. The adjacent locations of the peak and trough velocities move closer to each other by a small distance, and the amplitude evidently increases especially at the negative trough. It seems that the enhanced surface curvature in a fish-like body tends to amplify the peaks and troughs in the velocity curve. It should be noted that these features, created by the external vortices in the velocity distribution, are much more distinct than the features of the velocity induced by the pressure distribution due to the body curvature in a uniform flow. Therefore, one might consider the
The results of four bodies with thickness ratios of \( \lambda = 0.02, 0.04, 0.06 \) and 0.08 are shown in comparison with that of a flat body with \( \lambda = 0 \).

Figure 8. (a) Drawing of fish-like bodies with thickness ratios, \( \lambda \), ranging from 0 to 0.08. (b) The pressure distribution along the body surface. \( C_p \) is the pressure coefficient defined as \( C_p = P/\frac{1}{2} \rho U_\infty^2 \) with \( c^* = 2, h^* = 0.5 \) and \( \Gamma^* = 0.6 \). The dimensionless velocity inside the canal, \( \Pi \), along the LLTC. Four bodies with thickness ratios of \( \lambda = 0.02, 0.04, 0.06 \) and 0.08 are shown in comparison with a flat body. (a) \( c^* = 2, h^* = 0.5 \) and \( \Gamma^* = 0.6 \); (b) \( c^* = 2, h^* = 0.8 \) and \( \Gamma^* = 0.6 \).

The effect of the canal on the flow signal as of a differential filter. For larger values of \( h^* \), the features of the velocity distribution move slightly in the downstream direction for increasing body thickness values; see figure 9(b). Aside from this phase shift, the velocity plot maintains its main features, meaning that the relative locations of the velocity peaks and troughs are rather unchanged. For \( \lambda = 0.04 \) and higher, some perturbations in the velocity profile are observed around the fish’s head where the surface curvature is maximum. Because the fish’s posterior body is fairly similar to a flat plate, the features of the inside velocity distribution in this region are rather unchanged as compared to the flat-body case (see also figure 4(b)). Similar to the analytical results for a flat body, the value of \( l \) can be determined as the distance between two corresponding peaks or troughs as demonstrated in figure 10. In addition, the variations in \( l \) do not change the general features of the inside velocity distribution, as seen in figure 11.

Figure 12 demonstrates the effect of varying the vortex strength. The dimensionless number \( \Gamma^* \) does not change the overall features of the velocity distribution inside the canal, but it could change the amplitude of the velocity peaks. This observation also agrees with our analytical result in the previous section for a flat body. Inspired by the definition of \( \Pi \) in our analytical model, we further consider a dimensionless number \( \Pi \Gamma^{*2} \) instead of \( \Pi \). The result indicates that the amplitudes of \( \Pi \Gamma^{*2} \) at peaks and troughs fit a quadratic polynomial with respect to \( \Gamma^* \),

\[
\Pi \Gamma^{*2} = a_1 \Gamma^* + a_2 \Gamma^* + a_3,
\]

where \( a_1, a_2 \) and \( a_3 \) are functions of \( l \) and \( h \). Since \( l \) and \( h \) may be determined by sensed information available to a fish, these
coefficients can also be calculated from the sensed data. If a fish is able to measure the velocity inside the LLTC, Π, then it can deduct the strength of the vortices, $\Gamma^*$, in an incoming vortex street from the amplitude of the velocity peaks as shown in figure 12(b).

Having determined the effects of $\Gamma^*$ and $l$ on the velocity distribution inside the LLTC, one can conclude that, for a given fish’s body, $h^*$ had a dominant effect on changing the features of the velocity distribution inside the LLTC. The value of $h^*$ can again be evaluated by detecting the distance between consecutive maximum and minimum points in Π, similar to the flat-body case. It should be noted that as a vortex street is shifted away from the fish’s body, say that $h^*$ exceeds a threshold value $h^*_c$, the influence of the vortex street on the velocity inside the canal is diminished and the canal velocity will only be influenced by the pressure created by the free stream flow passing the body; see figure 13. On this view, the fish with a larger thickness ratio has less sensitivity to measure the vortex street at a greater distance, because its body induces a larger pressure gradient which could overwhelm information of other stimuli at a greater distance.

3.3. Vortex street on one side of the body

In our analytical model, we introduced vortex streets on both sides of the fish’s body in order to preserve the streamlined fish-like body in our theoretical solution. Yet, the case with a vortex street on one side of the body is actually more pertinent. This can easily be done by employing the numerical technique in the previous section. This result is shown in figure 14. As one can expect, there is no significant difference between vortices on only one side and vortices on both sides of the body, since the vortices on one side of the body have little influence on the near-body flow of the other side. Only the velocity near the body’s head and tail is altered. In this sense, all the conclusions drawn in the previous sections could be carried over to the case of a fish with a vortex street on one
distinct as a negative trough, which is near the body’s head, becomes more
angled with the fish’s body. To this end, this might provide enough information to a fish in order to determine the
rotated along with the vortex street. To this end, this might
results could be interpreted as if the whole velocity curve is
degenerates due to the departure of vortices. The presented
both sides of the body.

4. Discussion

In this investigation we considered modeling the LLTC of a fish
in a uniform background flow with a vortex street. A simplified
model of a vortex street is constructed if one knows the
following parameters: the vortex strength, $\Gamma$, the streamwise
spacing between vortices, $l$, the distance between the vortex
street and the body, $h$, and the velocity of the vortex street with
respect to the fish, $u_s$. While the velocity of the vortex
street, $u_s$, can be easily determined by monitoring the peak
or trough velocities inside an LLTC, sensing or calculation of
the other parameters of a vortex street requires information
on the velocity distribution throughout the entire lateral line
canal of the fish. To this end, accurate sensing of the velocity
inside the LLTC, information about the spacing between the
pores and the overall length of the canal are all quite important
for processing this information. Equation (17) for the velocity
inside the LLTC shows that $V_{in}$ is scaled with $d^2$ and $1/\nu$.
Therefore, one could expect that fish evolved in an environment
with a low level of hydrodynamic activity, which favors
-evolving a more sensitive sensing organ, will have a larger
canal diameter than those inhabiting a more hydrodynamically
active environment. This observation matches the measured
data from [37]. It should point out that, as noted earlier, it is
also desired to keep a small kinematic Reynolds number $\text{Re}_\nu$
for the flow inside the canal such that the velocity changes
in phase with the external pressure oscillation; otherwise the
inside flow velocity would exhibit a more complex profile,
which may impede the determination of the external pressure

An interesting observation from our LLTC modeling is
that while the sensory system of LLTC provides enough
information for detecting all the relevant parameters of an
incoming vortex street, it is not able to detect the mean
incoming velocity $U_\infty$. This is an interesting result and
consistent with observations by others [33] that an LLTC
filters out the dc (direct current) component of an incoming
velocity signal. To this end, it would be interesting to see
whether a fish evolved to have more lines of SNs shows the
same signal filtering characteristics. Furthermore, is there any
environmental reason for this lack of sensitivity to the dc
component of a velocity signal? Can a line of SNs be used
to detect the parameters of a vortex street?

Our LLTC model consists of two parts: (i) predicting the
-pressure gradient along the body of a fish and (ii) modeling
a sensed velocity inside the LLTC based on this pressure
gradient. While the second part (ii) is not applicable to a line of

Figure 13. The plot of the dimensionless velocity inside the canal, 
$\Pi$, for $h^+ = 1.5 > h^*$ with a flat body $\lambda = 0$ and a fish-like body
with $\lambda = 0.06$. Peaks and troughs indicating the vortex street are
overwhelmed by the background flow in the fish-like body case.

Figure 14. A comparison of the dimensionless velocity inside the
canal, $\Pi$, between a vortex street on one side and vortex streets on
both sides of the body.

It is also of interest to consider a more complex case
where the vortex street axis is not parallel to the fish’s body
axis. Assume that the angle between the vortex street axis
and the fish’s body is $\theta$. The velocity distribution inside the
LLTC for several values of $\theta$ is shown in figure 15. The first
negative trough, which is near the body’s head, becomes more
distinct as $\theta$ increases, while the secondary negative trough
degenerates due to the departure of vortices. The presented
results could be interpreted as if the whole velocity curve is
rotated along with the vortex street. To this end, this might
provide enough information to a fish in order to determine the
angle of an incident vortex street with the fish’s body.
SNs, the first part (i) is still valid and applicable. Therefore, one can still use equation (10) as a model to obtain an estimate for the surface flow velocity outside the boundary layer using a line of SNs. If the free stream velocity $U_\infty$ is much larger than the surface-induced velocity by the rest of the terms representing a vortex street, the response of the SNs will be dominated by this term and it will not be sensitive to perturbations by the vortex street. In contrast, the LLTC sensing system filters out this dc component and only processes the signal from the vortex street. As a result, our model predicts that a fish with an LLTC is more sensitive in detecting a vortex street and its parameters than a fish with a line of SNs.

In the previous sections, the flow velocity inside the LLTC was calculated to be proportional to the gradient of the external pressure along the fish’s body. This is based on the assumption that the pressure can be measured with an infinitely high spatial resolution. However, in reality the spatial resolution of this detection is dictated by the pore spacing $\Delta L$. Any pressure gradient changes over a distance smaller than this value may not be detected by the LLTC. In order to detect a vortex street, the spacing of the pores of an LLTC should be smaller than the characteristic length of the incident vortex street. Figure 16 shows the effect of the pore spacing, $\Delta L$, on the sensed velocity inside the LLTC. A typical pore spacing for the LLTC of fish is about 0.01 times of the body length. The pressure gradient recorded by such a canal has a fairly good spatial resolution. As pore spacing increases, the spatial resolution is decreased and as a result some of the more detailed features of the surrounding flow field could be lost during the measurement. This has also been observed by Fernandez et al [38] in their studies on a linear pressure sensor array for identifying small objects. They concluded that the pressure sensor array should have a sensor-to-sensor spacing of less than 0.03 times of the sensing body length. One can imagine that if the streamwise period of the Karman vortex street is less than the pore spacing, that is, $l < \Delta L$, a fish would have a very hard time in detecting a vortex street. On the other hand, the length of the LLTC also determines how much information regarding the vortex street can be obtained. It is evident that a longer LLTC would be helpful to collect more information on the surrounding flow field. An LLTC with small overall length, on the other hand, requires latency to experience the changes in the surrounding flow field as it passes over the fish. If the overall length of the LLTC is larger than the wavelength of the incident flow, the parameters of the incident vortex street could be detected from instantaneous measurements along the LLTC. On the other hand, detection of signals with wavelengths larger than the overall length of the LLTC, while possible, requires a longer detection sensing period and more processing. Therefore, an ideally fast LLTC sensing system should have its pore spacing much smaller than the characteristic length of the signal to be detected (e.g. its wavelength) and it should have a total length much smaller than the characteristic length of the signal to be detected (e.g. its wavelength). Thus, the ratios of $\Delta L/l$ and $L/l$ are of importance for the sensitivity of the LLTCs of fish. This result may account for the generally small pore spacings found in most fish, and for the fact that the LLTC stretches nearly all the way along a fish’s body.

The results of our study may also be used to explain some other fish behaviors, for instance, schooling behavior. It has been demonstrated that a blind fish is still able to school and to monitor the swimming speeds and directions of other fish in the school using its LLTC [7]. To this end, it is believed that a fish utilizes its LLTC sensing system to detect a vortex street wake by other fish in the school. Many fish schooling observations show that the preferred distance to the nearest neighbor for schooling fish is about one body length [10]. Considering the school structure, the vortex street shed by an anterior fish is 0.5 times the fish’s body length, that is, $h = 0.5c$. The vertical vortex spacing, $b$, could be approximated by the tail-beating amplitude, which is estimated by $0.1c—0.21c$ according to Wolfgang et al [39]. The downstream spacing of vortices $l$ is then determined to be around $0.4c—0.75c$ by the stability requirement of a Karman street $b/l \approx 0.28$. Accordingly, the dimensionless number $h^*$ is in the range of $0.7c—1.25c$, which is well below the critical distance $h^*$, meaning that the vortex street shed by anterior fish can be detected by the LLTC.

![Figure 16](https://via.placeholder.com/150)

**Figure 16.** The flow velocity inside an LLTC with different pore spacings. (a) $\Delta L = 0.01c$. (b) $\Delta L = 0.05c$. 

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If a fish in a school swims at a lower speed than that of its anterior neighbors, the relative speed of the vortex street with respect to the fish’s body, $u_s$, increases. Furthermore, it is expected that an accelerating anterior fish in a school creates a vortex street with larger vortex spacing and strength, marked by an increased $l$ and $\Gamma$. Another fish following this accelerating fish is then expected to sense these changes (increased $l$ and $\Gamma$) in the vortex street and thus to react appropriately to catch up to the speed of the school. On the other hand, if the anterior fish changes its direction, such a change will also alter the velocity distribution inside the LLTC, as discussed in the previous section. Again, the LLTC provides enough information to a fish in a school about the changes in the swimming direction for the anterior fish. In summary, our model demonstrates that an LLTC can provide enough information to a fish in a school in order to maintain its direction and speed with its neighboring fish.

A vortex street, shed by a large fish, is expected to have large distances between consecutive shed vortices. As discussed before, the LLTC of a smaller fish is not ideally suited for detecting such a vortex street instantly. On the other hand, if a vortex street is shed by a much smaller fish, the vortices’ strength is relatively small. Therefore, the detecting fish must be sufficiently close to the wake in order to be able to sense such a vortex street. This suggests that fish are especially sensitive to the wakes created by other fish of a similar length.

5. Conclusions

In this paper, we developed an analytical model for the flow field around a two-dimensional fish in proximity to a vortex street. This model is used to calculate the pressure distribution on the body surface. An analytical model for the LLTC of a fish is also developed to correlate the inside canal flow velocity to the external flow field. Nondimensional parameters governing the flow detection are also identified. Our model demonstrates the effects of each dimensionless number for a flat-bodied fish. The properties of a vortex street can be theoretically determined from the sensed velocity distribution inside the canal. A strategy is proposed to determine the properties of an outside vortex street, once a velocity distribution inside the LLTC is sensed. We also developed a numerical model to validate the result of our analytical model and to extend our investigation to the case of a fish-like body. The results of this numerical model show that the velocity distribution retains similar features to that of a fish with a small thickness ratio. Consequently, a similar technique to the one we developed for a flat-bodied fish is applicable here to identify the parameters of a vortex street from velocity measurements inside the LLTC. Therefore, our model provides a framework on how a fish might identify the parameters and characteristics of a wake including its speed, amplitude of vortices, spacing of vortices, angle of the vortex street axis, etc. Such information could be used by a fish to identify the size, speed and direction of fish traveling around it. Our model also provides possible explanations on functional adaptation of the LLTC’s morphology. These include the following: a lengthened canal favors sensing resolution, reduced pore spacing favors sensing resolution and widened canal favors sensing sensitivity. It is also argued that an LLTC could provide enough information to a fish about the direction for swimming or the accelerating/decelerating nature of the neighboring fish in a school.

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