Hydrodynamic model of fish orientation in a channel flow

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- Abstract For over a century, scientists have sought to understand how fish orient against an 12 incoming flow, even without visual and flow cues. Here, we make an essential step to elucidate 13 the hydrodynamic underpinnings of rheotaxis through the study of the bidirectional coupling 14 between fish and the surrounding fluid. By modeling a fish as a vortex dipole in an infinite 15 channel with an imposed background flow, we establish a planar dynamical system for the 16 cross-stream coordinate and orientation. The system dynamics captures the existence of a 17 critical flow speed for fish to successfully orient while performing cross-stream, periodic 18 sweeping movements. Model predictions are validated against experimental observations in the 19 literature on the rheotactic behavior of fish deprived of visual and lateral line cues. The crucial 20 role of bidirectional hydrodynamic interactions unveiled by this model points at an overlooked 21
- ²² limitation of existing experimental paradigms to study rheotaxis in the laboratory.
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24 Introduction

²⁵ Swimming animals display a complex behavioral repertoire in response to flows (*Chapman et al.,*

- 26 2011). Particularly fascinating is the ability of several fish species to orient and swim against an
- ²⁷ incoming flow, a behavior known as rheotaxis. While intuition may suggest that vision is necessary
- ²⁸ for fish to determine the direction of the flow, several experimental studies of midwater species
- ²⁹ swimming in a channel have documented rheotaxis in the dark above a critical flow speed (*Coombs*
- *et al., 2020*). When deprived of vision, fish lose the ability to hold station and they may perform sweeping, cross-stream movements from one side of the channel to other (*Bak-Coleman et al.,*
- ³¹ sweeping, cross-stream movements from one side of the channel to other (*Bak-Cole* 2013; *Bak-Coleman and Coombs, 2014; Elder and Coombs, 2015*) (Fig. 1).

In addition to vision, fish may rely on an array of compensatory sensory modalities to navigate the flow. For example, fish could sense and actively respond to linear accelerations caused by the surrounding flow using their vestibular system (*Pavlov and Tjurjukov, 1995*). Similarly, with the help of tactile sensors on their body surface, fish could maintain their orientation against a current through momentary contacts with their surroundings (*Lyon, 1904; Arnold, 1969*). Several modern studies have unveiled the critical role of the lateral line system, an array of mechanosensory receptors located on the surface of fish body (*Montgomery and Baker, 2020*), in their ability to orient against a current (*Montgomery et al., 1997; Baker and Montgomery, 1999*), hinting at a

⁴¹ hydrodynamics-based rheotactic mechanism that has not been fully elucidated. When deprived

*For correspondence: mporfiri@nyu.edu (MP); peterson@uwaterloo.ca (SDP) of vision, can fish rely only on lateral line feedback to perform rheotaxis? Is there a possibility for

43 rheotaxis to be achieved through a purely passive hydrodynamic mechanism that does not need

any sensing?

Through experiments on zebrafish larvae swimming in a laminar flow in a straight tube. Oteiza 45 et al. (2017) have recently unveiled an elegant hydrodynamic mechanism for fish to actively per-46 form rheotaxis. Utilizing their mechanosensory lateral line, fish can sense the flow along different parts of their body, which is sufficient for them to deduce local velocity gradients in the flow and adjust their movements accordingly. As further elaborated upon by *Dabiri (2017)*, the insight of-40 fered by **Oteiza et al.** (2017) is grounded in the fundamental relationship between vorticity and 50 circulation given by the Kelvin-Stokes' theorem, so that fish movements will be informed by local 51 sampling of the vorticity field. While offering an elegant pathway to explain rheotaxis, the frame-52 work of Oteiza et al. (2017) does not include a way for rheotaxis to be performed in the absence of 53 information about the local vorticity field. Several experimental studies have shown that fish can 54 perform rheotaxis even when their lateral line is partially or completed ablated, provided that the 55 flow speed is sufficiently large (Bak-Coleman et al., 2013: Bak-Coleman and Coombs, 2014: Baker 56 and Montgomery, 1999; Elder and Coombs, 2015; Montgomery et al., 1997; Oteiza et al., 2017; 57 Van Trump and McHenry, 2013). 58 Mathematical modeling efforts seeking to clarify the mechanisms underlying rheotaxis are scant 59 (Oteiza et al., 2017: Burbano-L and Porfiri, 2021; Colvert and Kanso, 2016; Chicoli et al., 2015), de-60 spite experiments on rheotaxis dating back more than a century (Lyon, 1904). A common hypothe-61 sis of existing mathematical models is that the presence of the fish does not alter the flow physics 62

with respect to the background flow, thereby neglecting interactions between the fish and the

⁶⁴ walls of the channel. For example, the model by *Oteiza et al. (2017*) implements a random walk in ⁶⁵ a virtual flow, matching experimental measurements of the background flow in the absence of the

animal through particle image velocimetry. A similar line of approach was pursued by **Burbano-L** and **Porfiri (2021)** for the study of multisensory feedback control of adult zebrafish.

67 Thus, according to these models, the fish acts as a perfectly non-invasive sensor that probes 68 and reacts to the local flow environment without perturbing it. There are countless examples in 69 fluid mechanics that could question the validity of such an approximation, from coupled interactions between a fluid and a solid in vortex-induced vibrations (Williamson and Govardhan, 2004) 71 to laminar boundary layer response to environmental disturbances that range from simple decay 72 of the perturbation to bypass transition (Saric et al., 2002). We expect that accounting for bidirec-73 tional coupling between the fluid flow and the fish will help clarify many of the puzzling aspects of 74 rheotaxis. 75

To shed light on the physics of rheotaxis, we formulate a mathematical model based on the 76 paradigm of the finite-dipole, originally proposed by *Tchieu et al.* (2012a). Within this paradigm, a 77 fish is viewed as a pair of point vortices of equal and opposite strength separated by a constant 78 distance in a two-dimensional plane. The application of the finite-dipole has bestowed important 79 theoretical advancements in the study of hydrodynamic interactions between swimming animals 80 (Gazzola et al., 2016; Filella et al., 2018; Kanso and Tsang, 2014; Kanso and Michelin, 2019), upon 81 which we investigate the bidirectional coupling between a fish and the surrounding fluid flow in a 82 channel. Our work contributes to the recent literature on minimal models of fish swimming (Gaz-83 zola et al., 2014, 2015; Sánchez-Rodríguez et al., 2020) that builds on seminal work by Lighthill 84 (1975), Taylor (1952), and Wu (1971) to elucidate the fundamental physical underpinnings of loco-85 motion and inform the design of engineering systems. 86 We focus on an ideal condition, where fish are deprived of vision, vestibular system, and tactile 87

sensors. Their only potential ability to probe the environment is through their lateral line that
gives them access to information about the flow. Such flow information is coupled, however, to
the motion of the fish itself, which acts as an invasive sensor and perturbs the background flow.
Just as fish motion influences the local flow field, so too does the local flow field alter fish motion

⁹² through advection. Predictions from the proposed model are compared against existing empirical



Figure 1. Fish rheotaxis. (a) Illustration of the problem with notation, showing a fish swimming in a backgroud flow described by Eq. (4). (b) Schematic of the cross-stream sweeping movement of some fish species swimming without visual cues; snapshots of fish at earlier time instants are illustrated by lighter shading.

- observations on fish rheotaxis, compiled through a comprehensive literature review of published
- work since 1900. Data presented in the literature are used to validate the predicted dependence of
- ⁹⁵ rheotaxis performance on local flow characteristics, individual fish traits, and lateral line feedback.

96 Results

97 Model of the fluid flow

⁹⁸ Consider a single fish swimming in an infinitely long two-dimensional channel of width *h* (Fig. 1(a)).

Let one wall of the channel be at y = 0 and the other at y = h, with x pointing along the channel.

The fish position at time *t* is given by $\vec{r}_f(t) = x_f(t)\hat{i} + y_f(t)\hat{j}$, where \hat{i} and \hat{j} are the unit vectors in the *x* and *y* directions, respectively. The orientation of the fish with respect to the *x* axis is given by $\theta_f(t)$ (positive counter-clockwise) and its self-propulsion velocity is $\vec{v}_f = v_0(\cos\theta_f \hat{i} + \sin\theta_f \hat{j}) = v_0\hat{v}_f$, where

 v_0 is the constant speed of the fish and \hat{v}_f is a unit vector in the swimming direction. The flow is modeled as a potential flow, which is a close approximation of the realistic flow field around a fish. This simple linear fluid model is intended to capture the mean flow physics, thereby averaging any turbulence contribution. The fish is modeled as a finite-dipole (*Tchieu et al., 2012a*), the potential field of which at some location $\vec{r} = x\hat{i} + y\hat{j}$ is given by

$$\phi_f(\vec{r}, \vec{r}_f, \theta_f) = -r_0^2 \left(\frac{(\vec{r} - \vec{r}_f) \cdot \vec{v}_f}{||\vec{r} - \vec{r}_f||^2} \right),\tag{1}$$

where r_0 is the characteristic dipole length-scale (on the order of the amplitude of the fish tail beating), so that the circulation of each vortex is $2\pi r_0 v_0$. This potential field is constructed assuming a far-field view of the dipole (*Filella et al., 2018*), wherein r_0 is small in comparison with the characteristic flow length scale, which is satisfied for $\rho = r_0/h \ll 1$. The velocity field at \vec{r} due to the dipole (fish) is $\vec{u}_f = \nabla \phi_f$.

A major contribution of the proposed model is the treatment of the fish as an invasive sensor that both reacts to and influences the background flow, thereby establishing a coupled interaction between the fish and the surrounding environment. A fish swimming in the vicinity of a wall will induce rotational flow near the boundary. In the inviscid limit, this boundary layer is infinitesimally



Figure 2. Method of images. Schematic of the fish (black) in the channel (thick lines) and the set of images (gray) needed to generate the channel. The streamlines generated by the fish in an otherwise quiescent fluid are shown in the channel colored by local velocity magnitude (red: high; blue: low). Dashed and solid lines are mirroring planes for the method of images, the pattern for which continues *ad infinitum*.

thin and can be considered as wall-bounded vorticity (*Batchelor, 2000*). Employing the classical
method of images (*Newton, 2011*), the influence of the wall-bounded vorticity on the flow field is
equivalent to that of a fictitious fish (dipole) mirrored about the wall plane. For the case of a fish
in a channel, this results in an infinite number of image fish (dipoles) (Fig. 2), the position vectors
for which are

$$\vec{r}_{\leq n}^{+} = x_f \hat{i} + (y_f - 2(n+1)h)\hat{j},$$
(2a)

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$$\vec{r}_{<,n}^{-} = x_f \hat{i} + (-y_f - 2nh)\hat{j}, \tag{2b}$$

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 $\vec{r}_{>,n}^{+} = x_f \hat{i} + (y_f + 2(n+1)h)\hat{j},$ (2c)

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$$\vec{r}_{>n} = x_f \hat{i} + (-y_f + 2(n+1)h)\hat{j},$$
(2d)

where *n* is a non-negative integer representing the *n*-th set of images. Subscripts "<" and ">" correspond to position vectors of the images at y < 0 and y > h, respectively. Likewise, superscript " \pm " denotes the orientation of the image dipole as $\pm \theta_f$; that is, a position vector with superscript "+" indicates that the associated image has the same orientation as the fish.

The potential function for a given image is found by replacing \vec{r}_f in (1) with its position vector from (2) and adjusting the sign of θ_f in (1) to match the superscript of its vector. The potential field at \vec{r} due to the image dipoles is

$$\phi_{w}(\vec{r},\vec{r}_{f},\theta_{f}) = \sum_{n=0}^{\infty} \left(\phi_{f}(\vec{r},\vec{r}_{<,n}^{+},\theta_{f}) + \phi_{f}(\vec{r},\vec{r}_{<,n}^{-},-\theta_{f}) + \phi_{f}(\vec{r},\vec{r}_{>,n}^{+},\theta_{f}) + \phi_{f}(\vec{r},\vec{r}_{>,n}^{-},-\theta_{f}) \right).$$
(3)

Thus, the velocity field due to the wall is computed as $\vec{u}_w = \nabla \phi_w$, and the overall velocity field induced by the fish is $\vec{u}_f + \vec{u}_w$. (A closed-form expression for the series in terms of trigonometric and hyperbolic functions is presented in Appendix 1.) Overall, the presence of the walls distorts the flow generated by the dipole, both compressing the streamlines between the fish and the walls in its proximity and creating long-range swirling patterns in the channel (Fig. 2). The presence of a background flow in the channel is modelled by superimposing a weakly rotational flow,

$$\vec{u}_b(\vec{r}) = U_0 \left(1 - 4\epsilon \left(\frac{y}{h} - \frac{1}{2} \right)^2 \right) \hat{i},\tag{4}$$

which has speed U_0 at the channel centerline and $U_0(1 - \epsilon)$ at the walls, ϵ being a small positive parameter. As $\epsilon \to 0$, a uniform (irrotational) background flow is recovered: such a flow is indistinguishable from the one in Fig. 2, provided that the observer is moving with the background flow.

The overall fluid flow in the channel is ultimately computed as $\vec{u} = \vec{u}_f + \vec{u}_w + \vec{u}_b$.

The circulation in a region \mathcal{R} in the flow field centered at some location y is $\Gamma = \int_{\mathcal{R}} \omega \, dA$, where $\omega = (\nabla \times \vec{u}) \cdot \hat{k}$ is the local fluid vorticity ($\hat{k} = \hat{i} \times \hat{j}$). For the considered flow field, we determine

$$\omega(\vec{r}) = \frac{8U_0\epsilon}{h} \left(\frac{y}{h} - \frac{1}{2}\right),\tag{5}$$

whereby the irrotational component of the flow field does not contribute to the circulation, andthe circulation at a point (per unit area) is equivalent to the local vorticity.

147 Model of fish dynamics

From knowledge of the fluid flow in the channel, we compute the advective velocity $\vec{U}(\vec{r}_{f},\theta_{f})$ and

hydrodynamic turn rate $\Omega(\vec{r}_f, \theta_f)$ at the fish location, which encode the influence of the confining

walls and background flow on the translational and rotational motion of the fish, respectively. Neglecting the inertia of the fish so that it instantaneously responds to changes in the fluid flow, we

152 determine (Filella et al., 2018)

$$\dot{\vec{r}}_{f}(t) = \vec{U}(\vec{r}_{f}(t), \theta_{f}(t)) + \vec{v}_{f}(\theta_{f}(t)),$$
 (6a)

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$$\dot{\theta}_f(t) = \Omega(\vec{r}_f(t), \theta_f(t)) + \lambda(\vec{r}_f(t), \theta_f(t)), \tag{6b}$$

where λ is the feedback mechanism based on the circulation measurement through the lateral line.

The advective velocity is found by de-singularizing the total velocity field \vec{u} at $\vec{r} = \vec{r}_f$, which is equivalent to calculating the sum of the velocity due to the walls and the background flow in correspondence of the fish (*Milne-Thomson*, **1996**)

$$\vec{U}(\vec{r}_f, \theta_f) = \left(\vec{u}_w(\vec{r}, \vec{r}_f, \theta_f) + \vec{u}_b(\vec{r})\right)|_{\vec{r}=\vec{r}_f}$$

$$= -\frac{\pi^2 v_0 \rho^2}{12} \left[\left(1 + 3\csc^2\left(\frac{\pi y_f}{h}\right) \right) \cos\theta_f \hat{i} - \left(1 - 3\csc^2\left(\frac{\pi y_f}{h}\right) \right) \sin\theta_f \hat{j} \right]$$

$$+ U_0 \left(1 - 4\epsilon \left(\frac{y_f}{h} - \frac{1}{2}\right)^2 \right) \hat{i}. \quad (7)$$

Equation (7) indicates that the walls have a retarding effect on the swimming speed of the fish that increases in magnitude the closer the fish gets to either wall of the channel. A fish swimming with orientation $\theta_f = 0$ at the center of the channel, for example, will swim with velocity $\dot{r}_f(t) = v_0(1 - (\pi^2/3)\rho^2)\hat{i} + U_0\hat{i}$. This effect should not be mistaken as traditional viscous drag, which is not included

in potential flow theory; rather, it should be intended as the impact of nearby solid boundaries.

Hydrodynamic turn rate is incorporated by considering the difference in velocity experienced by the two constituent vortices comprising the dipole (*Filella et al., 2018*), namely,

$$\Omega(\vec{r}_f, \theta_f) = -\hat{v}_f \cdot \left[\nabla \left(\vec{u}_w(\vec{r}, \vec{r}_f, \theta_f) + \vec{u}_b(\vec{r}) \right) |_{\vec{r} = \vec{r}_f} \right] \hat{v}_f^{\perp} \\ = -\frac{\pi^3 \rho^2 v_0}{4h} \cot \left(\frac{\pi y_f}{h} \right) \csc^2 \left(\frac{\pi y_f}{h} \right) \cos \theta_f + \frac{8U_0 \epsilon}{h} \left(\frac{y_f}{h} - \frac{1}{2} \right) \cos^2 \theta_f, \quad (8)$$

where $\hat{v}_{f}^{\perp} = \hat{k} \times \hat{v}_{f}$; see Methods and Materials Section for the mathematical derivation. Equation (8) indicates that interaction with the walls causes the fish to turn towards the nearest wall;

for example, a fish at $y_{\ell} = 3/4h$, will experience a turn rate due to the wall of $(\pi^3 \rho^2 v_0)/(2h) \cos \theta_{\ell_1}$ 163 such that it will be rotated counter-clockwise if swimming downstream and clockwise if swimming 164 upstream. On the other hand, the turning direction imposed by the background flow is always 165 positive (counter-clockwise) in the right half of the channel and negative (clockwise) in the left half. 166 irrespective of fish orientation, so that a fish at $v_r = 3/4h$ will always be rotated counter-clockwise. 167 As a result, the fish may turn towards or away from a wall, depending on model parameters and 168 orientation 169 Based on experimental observations and theoretical insight (Burbano-L and Porfiri, 2021; Oteiza 170 et al., 2017), we hypothesize that hydrodynamic feedback, that is, lateral line measurements of the 171 surrounding fluid that fish can employ to navigate the flow, is related to the measurement of the 172 circulation in a region surrounding the fish. We consider a rectangular region \mathcal{R} of width r_0 along 173 the fish body length l. For simplicity, we assume a linear feedback mechanism, $\lambda(\vec{r}_{c}, \theta_{c}) = K\Gamma(\vec{r}_{c}, \theta_{c})$. 174 where we made evident that circulation is computed about the fish location and K is a non-negative 175 feedback gain. Assuming that the fish size is smaller than the characteristic length scale of the flow, 176

¹⁷⁷ we linearize the vorticity along the fish in (5) as $\omega(\vec{r}) \approx \omega(\vec{r}_f) + \nabla \omega(\vec{r}_f) \cdot \hat{v}_f \Delta I$. By computing the integral ¹⁷⁸ from $\Delta I = -I/2$ to I/2, we obtain

$$\lambda(\vec{r}_f, \theta_f) = Kr_0 l \frac{8U_0 \varepsilon}{h} \left(\frac{y_f}{h} - \frac{1}{2}\right).$$
(9)

Compared to established practice for modeling fish behavior in response to visual stimuli (Gau-179 trais et al., 2009; Calovi et al., 2014; Zienkiewicz et al., 2015b; Couzin et al., 2005), the proposed 180 model introduces rich nonlinear dynamics arising from the bidirectional coupling between the mo-181 tion of the fish and the flow physics in its surroundings. We note that the employed feedback in 182 (9) neglects additional potential sensing mechanisms, including vision (Lvon, 1904), acceleration 183 sensing through the vestibular system (Paylov and Tiuriukov, 1995), and pressure sensing through 184 sensory afferents in the fins (Hardy et al., 2016), which might enhance the ability of fish to navigate 185 the flow. 186

187 Analysis of the planar dynamical system

Given that the right hand side of equation set (6) is independent of the streamwise position of the fish, the equations for the cross-streamwise motion and the swimming direction can be separately studied, leading to an elegant nonlinear planar dynamical system. We center the cross-stream coordinate about the center of the channel and non-dimensionalize it with respect to *h*, introducing $\xi = y_f/h - 1/2$. The governing equations become

$$\dot{\xi} = \left[1 - \frac{\pi^2 \rho^2}{12} \left(3 \csc^2\left(\pi \left(\xi + \frac{1}{2}\right)\right) - 1\right)\right] \sin \theta_f, \tag{10a}$$

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$$\dot{\theta}_f = -\frac{\pi^3 \rho^2}{4} \cot\left(\pi \left(\xi + \frac{1}{2}\right)\right) \csc^2\left(\pi \left(\xi + \frac{1}{2}\right)\right) \cos\theta_f + 8\alpha\xi \left(\cos^2\theta_f + \kappa\right),\tag{10b}$$

where we non-dimensionalized by the time needed for the fish to traverse the channel in the absence of a background flow, that is, h/v_0 , and introduced $\alpha = U_0 \epsilon/v_0$ and $\kappa = Kr_0 l$ (see Methods and Materials Section for estimation of these parameters from experimental observations).

In search of the equilibria of the dynamical system, we note that swimming downstream or 197 upstream ($\theta_f = 0$ and π , respectively) solves (10a) for any choice of the cross-stream coordinate, 198 the value of which is determined from the solution of (10b) for the corresponding orientation θ_{r} . 199 In the case of downstream swimming, the only solution of the resulting transcendental equation 200 is $\xi = 0$. For upstream swimming, depending on the value of the parameter $\beta = (\alpha(1+\kappa))/\rho^2$, we 201 have one or three solutions: if $\beta < \beta^* = \pi^4/32$, the only solution is $\xi = 0$, otherwise, in addition to 202 $\xi = 0$, there are two solutions symmetrically located with respect to the centerline that approach 203 the walls as $\beta \to \infty$ (Fig. 3(a), see Methods and Materials Section for mathematical derivations). 204 Local stability of these equilibria is determined by studying the eigenvalues of the state matrix 205

²⁰⁶ of the corresponding linearized dynamics. For all the considered dynamics, the trace of the state



Figure 3. Qualitative dynamics of equation set (10). (a) Cross-stream equilibria for upstream swimming as a function of β . (b,c) Phase plot for downstream and upstream swimming in the case $\alpha = 0.1$, $\rho = 0.01$, and $\kappa = 1$, so that $\beta = 20$. In all panels, red refers to unstable equilibria and green to stable equilibria.

- ²⁰⁷ matrix is zero, so that the equilibria can be saddle points (unstable) or neutral centers (stable), if
- ²⁰⁸ the determinant is negative or positive, respectively (*Bakker, 1991*) (see Methods and Materials
- ²⁰⁹ Section for mathematical derivations). In the case of downstream swimming, the determinant is

always negative, such that the equilibrium ($\theta_f = 0, \xi = 0$) is a saddle point (Fig. 3(b)). For upstream

swimming, the equilibrium ($\theta_f = \pi, \xi = 0$) is stable if $\beta > \beta^*$, leading to periodic oscillations similar

to experimental observations (Bak-Coleman et al., 2013; Bak-Coleman and Coombs, 2014; Elder

and Coombs, 2015) (Fig. 1(b)); the other two equilibria located away from the centerline are always

unstable (Figs. 3(b,c)). Oscillations about the centerline during rheotaxis have a radian frequency

 $\omega_0 \simeq (\pi^2/2)\rho \sqrt{\beta/\beta^* - 1}$, such that the frequency increases with the square root of β and is zero at

 $_{216}$ β^* (see Methods and Materials Section for the mathematical derivation).

217 Validation against empirical observations

To support our proposed model with existing experimental data, we conducted an expansive literature review, identifying 11 relevant publications after applying a set of inclusion and exclusion criteria (see Methods and Materials Section). In particular, we only included experiments wherein fish performed rheotaxis in the absence of visual cues, in alignment with the assumption of a purely hydrodynamic feedback mechanism in (9).

The studies identified from the literature review are utilized to validate the proposed theoretical framework with respect to the rheotaxis stability threshold. We first express the stability threshold $\beta = \beta^*$ in dimensional form in terms of the rheotaxis threshold speed

$$U_c = \frac{\pi^4 r_0^2 v_0}{32h^2 (1 + K r_0 l)\epsilon},$$
(11)

such that $U_0 > U_c$ corresponds to the stable condition $\beta > \beta^*$, and vice versa. From most studies, the values of U_c can be identified and its confidence level can be inferred (see Methods and Materials Section).

As evidenced in (11), a series of parameters could influence the rheotaxis threshold speed. 229 including the lateral line feedback, flow gradient, swimming domain size, and fish body length. 230 Specifically, (11) predicts that increasing the lateral line feedback, flow gradient, and/or width of 231 swimming channel promotes rheotaxis at lower flow speeds, whereas increasing fish size will re-232 quire higher flow speeds to elicit rheotactic behavior. The effects of these parameters are validated 233 independently in Table 1, where we include experimental evidence garnered within each study 234 and, when possible, carry out a comparison, across them. We compare each of these empirical 235 observations to model predictions and assess if they support the model contradict the model or 236 are inconclusive. An observation is considered supportive of (contradictive to) our model if the 237 measured U exhibits with statistical significance the same (opposite) dependence on a certain pa-238 rameter (see Methods and Materials Section for details on the statistical analysis). Data that lack 239 statistical significance are considered inconclusive. 240

Several studies provide evidence in favor of the prediction of our model of the beneficial role of lateral line feedback, showing a significant reduction in rheotactic performance when the lateral line is compromised (*Kulpa et al., 2015; Oteiza et al., 2017; Suli et al., 2012*), see Table 1. In these studies, fish locomotion was measured in steady background flows, so that a fish holding station would experience minimal linear acceleration and marginally engage the vestibular system. Throughout these studies, fish were not observed to make contact with the swim channel, indicating that tactile senses played a negligible role in rheotaxis.

We identified two studies (*Lyon*, *1904*; *Oteiza et al.*, *2017*) that could validate the predicted effect of the flow velocity gradient on rheotaxis, as summarized in Table 1. In both studies, fish locomotion was recorded in flows with varying velocity gradients. In agreement with the proposed model, the rheotaxis performance of zebrafish larvae significantly improved with increasing gradient magnitudes (*Oteiza et al.*, *2017*). Similar observations were obtained by Lyon on blind Fundulus (*Lyon*, *1904*), where in a flow with a small gradient, fish performed rheotaxis only when tactile cues were

available, while in a jet flow with a large flow gradient, rheotaxis could be elicited solely by the 254 flow. Although qualitatively in line with our predictions, we conservatively considered this study as 255 inconclusive due to a lack of quantitative data for statistical tests.

To elucidate the role of the swimming domain size on rheotaxis threshold speed, we conducted 257 cross-study comparisons as shown in Table 1. As evidenced through comparisons between two ex-258 periments on zebrafish larvae (Oteiza et al., 2017: Peimani et al., 2017) in swim channels of dras-259 tically different sizes, rheotaxis was elicited at a higher threshold speed in a smaller flow channel. 260 which supports our model prediction. Our model is also gualitatively supported by comparisons 261 between Bak-Coleman and Coombs (2014) and Baker and Montgomery (1999), or Bak-Coleman 262 and Coombs (2014) and Van Trump and McHenry (2013), where experiments on blind cavefish of 263 comparable body sizes uncovered higher threshold speeds in smaller flow channels. In the ex-264 periments of Bak-Coleman and Coombs (2014), blind cavefish were observed to receive transient 265 tactile senses while swimming, which could have contributed to its lower rheotaxis threshold. As 266 a result. experimental data on blind cavefish were conservatively deemed to be inconclusive. 267

To validate the effect of fish body size on the rheotaxis threshold speed, we examined a pair 268 of studies by Bak-Coleman and Coombs (2014) and Elder and Coombs (2015), where experiments 269 were conducted on fish of the same species (Astyanax mexicanus) in swim tunnels of the same size. 270 and tested in flows at a similar range of speeds. The high flow speeds in both studies suggest that 271 the flow gradients in these experiments were small. We assume that the lateral line feedback were 272 equivalent in both studies, as the subjects were conspecific. Although the factile cues present in 273 the experiments by Bak-Coleman and Coombs (2014) hinder our ability to reach a definitive conclu-274 sion on the effect of fish body size, the higher threshold speed observed in larger fish qualitatively 275 supports our model prediction (see Methods and Materials Section). The paucity of data for valida-276 tion of the effect of fish size is a result of a lack of studies with matching experimental conditions. 277 including dimensions of the flow facilities, flow conditions, and functionality of the lateral line. 278 In summary, we identified a total of five sets of experiments in support of our model, and nine 279 sets of studies that offer inconclusive evidence. None of the data contradicted predictions from 280

the proposed model. 281

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Table 1. Results of the bibliographical research on fish rheotaxis in the absence of visual cues, used to validate the proposed model.

| Poforonco | Fish species | [†] Evidonco | Comparison with model | |
|------------------------|------------------------------|--|-----------------------|--------------|
| Reference | FISH Species | Evidence | Supportive | Inconclusive |
| Within studies | | | | |
| Effect of lateral line | | | | |
| Bak-Coleman | Giant danio (<i>Devario</i> | No significant difference in fish heading | | × |
| et al. (2013) | aequipinnatus) | angle against current was detected | | |
| | | between LL+ and LL– | | |
| Bak-Coleman | blind cavefish | Rheotaxis threshold speed was slightly (but | | × |
| and Coombs (2014) | (Astyanax mexicanus) | not significantly) lower in LL– condition | | |
| Baker and | blind cavefish | Rheotaxis threshold speed was significantly | | × |
| Montgomery | (Astyanax fasciatus) | higher in LL– condition; fish received | | |
| (1999) and | | intermittent tactile senses | | |
| Montgomery | | | | |
| et al. (1997) | | | | |
| Elder and | Mexican tetras | No significant influence of LL condition was | | × |
| Coombs (2015) | (Astyanax mexicanus) | detected on rheotactic performance | | |
| Kulpa et al. | blind cavefish | Significantly higher rheotaxis index in LL+ | × | |
| (2015) | (Astyanax mexicanus) | fish than LL– fish in jet stream | | |
| Oteiza et al. | zebrafish (Danio | Posterior lateral line ablation or chemical | × | |
| (2017) | <i>rerio</i>) larva 5–7 dpf | neuromast ablation severely reduced | | |
| | | rheotaxis | | |
| Suli et al. (2012) | zebrafish (Danio | LL hair cell damage led to a significant | × | |
| | <i>rerio</i>) larva 5 dpf | decrease in rheotaxis; regeneration of LL | | |
| | | hair cells restored rheotaxis | | |

| Poforonco | Fish species | [†] Evidonco | Comparison with model | |
|----------------------|------------------------------|--|-----------------------|--------------|
| Reference | | | Supportive | Inconclusive |
| Van Trump and | blind Mexican | In LL+ and LL–, fish exhibited statistically | | × |
| McHenry (2013) | cavefish (Astyanax | indistinguishable rheotaxis behavior | | |
| | fasciatus) | | | |
| Effect of flow gradi | ient | | | |
| Lyon (1904) | blind Fundulus | In a flow with small gradient, rheotaxis was | | × |
| | | elicited only when fish received tactile cues; | | |
| | | in jet flow with large gradient, rheotaxis | | |
| | | was elicited by flow without tactile cues. | | |
| | | Lack of data on statistical significance | | |
| Oteiza et al. | zebrafish (Danio | Rheotaxis of fish improved with increasing | × | |
| (2017) | <i>rerio</i>) larva 5–7 dpf | gradient magnitudes | | |
| Across studies | | | | |
| Effect of channel w | vidth | | | |
| Bak-Coleman | blind cavefish | Significantly different threshold speed for | | × |
| and Coombs | (Astyanax mexicanus); | LL+ fish: $0.90 \pm 0.137 \text{ cm/s}$ (mean \pm s.e.m.) in | | |
| (2014); Baker | blind cavefish | 25 cm wide tunnel; between 2 cm/s and | | |
| and Mont- | (Astyanax fasciatus) | 3 cm/s in 9 cm wide tunnel. Tactile cues | | |
| gomery (1999) | | available to fish in Bak-Coleman and | | |
| | | Coombs (2014) | | |
| Bak-Coleman | blind cavefish | Significantly different threshold speed for | | × |
| and Coombs | (Astyanax mexicanus); | LL+ fish: $0.90 \pm 0.137 \mathrm{cm/s}$ (mean \pm s.e.m.) in | | |
| (2014); | blind cavefish | 25 cm wide tunnel; between 2 cm/s and | | |
| Van Trump | (Astyanax fasciatus) | 4 cm/s in ~ 11 cm diameter tunnel. Tactile | | |
| and McHenry | | cues available to fish in Bak-Coleman and | | |
| (2013) | | Coombs (2014) | | |
| Oteiza et al. | zebrafish (Danio | Onset of rheotaxis in LL+ fish observed at | × | |
| (2017); Peimani | <i>rerio</i>) larva 5–7 dpf | flow speed 0.95 cm/s in 1.6 mm wide tunnel; | | |
| et al. (2017) | | rheotaxis observed in LL+ fish at flow | | |
| | | speed 0.2 cm/s in 2.22 cm diameter tunnel | | |
| Effect of body leng | th | | | |
| Bak-Coleman | blind cavefish | Significantly different threshold speed for | | × |
| and Coombs | (Astyanax mexicanus); | LL+ fish: $0.90 \pm 0.137 \text{ cm/s}$ (mean \pm s.e.m.) | | |
| (2014); Elder | Mexican tetras | for 4.2–5.0 cm long fish; 1.96 \pm 0.350 cm/s | | |
| and Coombs | (Astyanax mexicanus) | (mean \pm s.e.m.) for 8.3 cm long fish. Tactile | | |
| (2015) | | cues available to fish in Bak-Coleman and | | |
| | | Coombs (2014) | | |
| Total | | | 5 | 9 |

Table 1. Results of the bibliographical research on fish rheotaxis in the absence of visual cues, used to validate the proposed model.

282 [†] LL+: lateral line enabled; LL–: lateral line disabled

Discussion

There is overwhelming evidence that fish can negotiate complex flow environments by responding 284 to even small flow perturbations (Liao, 2007). However, seldom are these perturbations included 285 in mathematical models of fish behavior, which largely rely on vision cues (Gautrais et al., 2009; 286 Calovi et al., 2014: Zienkiewicz et al., 2015b: Couzin et al., 2005). In this paper, we proposed a 287 hydrodynamic model for the bidirectional coupling between fish swimming and fluid flow in the 288 absence of any sensory input but lateral line feedback. The model reduces to a nonlinear planar 289 dynamical system for the cross-stream coordinate and orientation, of the kind that are featured in 290 nonlinear dynamics textbooks for their elegance, analytical tractability, and broad physical interest 201 (Sastry, 2013). 292 The planar system anticipates several of the surprising features that have challenged our un-203 derstanding of rheotaxis for over a century. In agreement with experimental observations on fish 294 swimming in the absence of visual cues (see Table 1), we uncovered an equilibrium at the chan-295 nel centerline for upstream swimming whose stability is controlled by a single non-dimensional 296

- parameter that summarizes flow speed, flow gradient, lateral line feedback, fish size, and chan-
- nel width. Above a critical value of this parameter, the model predicts that rheotaxis is stable and

fish will begin periodic cross-stream sweeping movements whose amplitude can be as large as thechannel width.

The mathematical proof of the existence of a poptrivial threshold for β above which upstream 301 swimming in the absence of visual cues becomes stable is in agreement with experimental observa-302 tions on a number of species (see Table 2). Experiments have indicated the existence of a threshold 303 in the flow speed or flow gradient above which fish successfully perform rheotaxis, in the absence of visual stimuli. Importantly, the presence of the walls is necessary for the emergence of such a 305 threshold, since for $\rho \to 0, \beta \to \infty$, thereby automatically guaranteeing the stability of upstream 306 swimming, against experimental evidence. Based on our estimation of α and ρ , β can be as small 307 as 10^{-1} and exceed 10^2 , thereby encompassing the critical value $\beta^* \simeq 3$ (see Methods and Materials 308 Section for estimation of model parameters). 300

The determination of such a threshold resolves a long-standing dilemma about the role of hy-310 drodynamics on rheotaxis, laid bare by Lvon (1904) in 1904: "It is equally absurd to imagine a fish in 311 the Gulf Stream to be stimulated and oriented by a uniform forward motion of the water. Whether 312 orientation be a simple reflex or a conscious process, points of reference -i.e., points relatively 313 at rest – are necessary." Such points of reference could be visual stimuli (Davidson, 1949), static 314 objects near the bottom of the swimming channel (*Needham and Jones*, 1959), or hydrodynamic 315 points of reference consisting of flow regions with distinctive features from the background. It is 316 the gradient of the flow that creates hydrodynamic points of reference for a fish to undertake rheo-317 taxis in the dark, even without access to sensory information through the lateral line. The higher 318 is the speed of the flow with respect to the speed of the animal, the more the role of these hydro-319 dynamic reference points will be magnified until reaching the stability threshold for rheotaxis. 320

The model is successful in predicting the emergence of rheotaxis in the absence of sensory information from the lateral line and visual cues (see Table 1). Setting $\kappa = 0$ in our model eliminates hydrodynamic feedback, yet, the fish is able to perform rheotaxis at sufficiently large flow speeds and steep flow gradients. Increasing κ broadens the stable region, leading to more robust rheotaxis, in agreement with experimental observations of blind cavefish (*Kulpa et al., 2015*) and zebrafish larvae (*Oteiza et al., 2017; Suli et al., 2012*) with intact versus compromised lateral lines (see Table 1).

The model prediction on the influence of the environment, including the flow gradient and flow 328 channel size, on rheotaxis in the absence of visual cues is also supported by observations in the 329 literature. Consistent with observations on zebrafish larvae by Oteiza et al. (2017), increasing the 330 flow gradient ϵ enhances hydrodynamic feedback, resulting in improved rheotaxis. In contrast to 331 Oteizg et al. (2017), however, the proposed model treats the fish as an invasive sensor, accounting 332 for interactions between the fish and the walls of the channel. The model anticipates that wider 333 channels should promote rheotaxis, which is supported by experimental observations (Baker and 33/ Montgomerv. 1999: Van Trump and McHenrv. 2013: Bak-Coleman and Coombs. 2014: Oteiza et al., 335 2017: Peimani et al., 2017) (see Table 1). Similarly, in alignment with experimental observations 336 (Bak-Coleman and Coombs, 2014; Elder and Coombs, 2015), the model predicts a lower threshold 337 for longer fish, owing to a magnification of the hydrodynamic feedback received by a longer body. 338 Finally, the model anticipates the onset of periodic cross-stream sweeping, which has been 339 studied in some experiments on fish swimming in channels without vision (Coombs et al., 2020). 340 While there is not conclusive experimental evidence regarding the dependence of the frequency 341 of oscillations on flow conditions, the model is in qualitative agreement with experiments by *Elder* 342 and Coombs (2015), showing a sublinear dependence on the flow speed. Therein, it is shown that 343 the radian frequency has a weak positive tendency with respect to the flow speed for Mexican tetra 344 swimming with or without cues from the lateral line. Above 2 cm s^{-1} the animals can successfully 345 perform rheotaxis and display sweeping oscillations at about three cycles per minute and increase 346 to about four cycles per minute at $12 \,\mathrm{cm}\,\mathrm{s}^{-1}$. These correspond to a radian frequency on the order 347 of 0.1 rad s⁻¹, which is similar to what we would predict for β ranging from 10⁰ to 10¹ and ρ of the 348 order of 10^{-1} (recall that the time is scaled with respect to time required by the animal to traverse 349

the channel from wall to wall in the absence of a background flow).

Most experiments used in our model validation listed in Table 2 were conducted in the past 351 25 years, and only two studies date back to before 1970. This disparity is attributed to an evolu-352 tion of the methodologies for the study of rheotaxis over time. Among the earlier efforts, a large 353 portion relied on observations of fish behavior in the field (Arnold 1974) Although these studies 354 minimized the introduction of external stimuli stemming from human presence and unfamiliar 355 environments that could alter the behavior of fish in the wild a lack of flexibility in the design of controlled experiments in the field, together with an insufficient measurement resolution, has led 357 to only a limited number of works that could distinguish the impact of one sensory cue from an-358 other. As a result, a large number of earlier efforts do not meet our inclusion criteria (see Methods 350 and Materials Section). 360

Likely, the interest in fish rheotaxis was recently reignited owning to the advancements in tech-361 nologies that could selectively deactivate specific fish sensory organs, thereby allowing for the 362 targeted investigation of the role of each sensory cue in rheotaxis. For instance, pharmacologi-363 cal methods that could disable the lateral line led to studies (Montgomery et al., 1997: Baker and 364 Montgomery, 1999) challenging the long-standing perception that the lateral line could not mediate 365 rheotaxis. In addition, the development of high speed cameras with infra-red sensing capabilities 366 enabled precise measurements of fish behavior in the dark, allowing for the elimination of visual 367 cues from the study of lateral line functionality in rheotaxis. Some early experiments that have 368 been considered in the past as evidence against the role of the lateral line are not listed in Table 2 360 due to a lack of a controlled experimental design in the field setting. For example, some species of 370 salmonids, including salmon and trout, were observed to swim against the current in the day and 371 rest on the bottom of a stream at night (Davidson, 1949; Gibson, 1966; Edmundson et al., 1968). 372 leading to a conclusion that the lateral line played a minimal role in rheotaxis (Arnold, 1974). How-373 ever, we did not include these experiments for our model validation due to confounding factors 374 posed by the field settings, such as variations in water temperature (*Needham and Jones*, 1959: 375 Edmundson et al., 1968; Fraser et al., 1993) and current speeds at different hours of the day. Daily 376 fluctuations in the availability of food (Waters, 1962; Elliott, 1965) is another factor that could influ-377 ence the activity levels of fish at night, as observed in white bass (McNaught and Hasler, 1961) and 378 trout (*Elliott. 1965*). Another class of experiments that led to the previous rejection of lateral line 379 was the demonstration of the imperative role of vision in rheotaxis. Experiments on salmon (*Hour*, 380 1954) and herring (Brawn, 1960) showed a reduction in rheotaxis when vision was obscured in the 381 dark or in muddy water, conflating the role of visual cues in rheotaxis. Again, these observations 382 do not directly contradict the proposed model, which suggests that in the absence of visual cues. 383 rheotaxis could still manifest provided that the flow speed is sufficiently high. 384

Just as other minimal models of fish swimming have helped resolve open questions on scaling laws (*Gazzola et al., 2014*), gait (*Gazzola et al., 2015*), and drag (*Sánchez-Rodríguez et al., 2020*), the proposed effort addresses some of the baffling aspects of rheotaxis through a transparent and intuitive treatment of bidirectional hydrodynamic interactions between fish and their surroundings. The crucial role of these bidirectional interactions hints that active manipulation of their surroundings by fish offers them a pathway to overcome sensory deprivation when swimming in the dark or in the absence of feedback from their lateral line.

The proposed model is not free of limitations, which should be addressed in future research. 392 From a theoretical point of view, the current model neglects the elasticity and inertia of the fish. 303 which might reduce the accuracy in the prediction of rheotaxis, especially transient phenomena. 394 Future research should refine the dipole paradigm toward a dynamic model that accounts for 305 added mass effects and distributed elasticity, similar to those used in the study of swimming robots 396 (Sfakiotakis et al., 1999; Colgate and Lynch, 2004). Likewise, the current model does not describe 397 contact and impact with the walls of the channel, which could be important in further detailing 398 the onset of cross-sweeping motions that could involve stick-and-slip at the bottom of the chan-399 nel (Van Trump and McHenry, 2013). The model could also be expanded to account for additional 400

sensory modalities, such as vision, vestibular system, and tactile sensors on the fish body surface;

however, any of these extension shall require detailed experiments to tease out the contribution

⁴⁰³ of each of these sensory modalities (*Coombs et al., 2020*).

Even within the current scope of the model, further experiments could strengthen its predic-404 tive value. For example, the model assumes a linear hydrodynamic feedback mechanism, which 405 is yet to be validated experimentally; in this yein, future experiments should be designed to para-406 metrically vary the flow speed and quantify the activity level of lateral line nerve fibers through 407 neurophysiological recordings (*Mogdans, 2019*). We also see merit in experiments with robotic 408 fish (Zhang et al., 2016), mimicking the swimming gait of live animals and allowing to precisely 400 control sensory input. In this yein, we foresee experiments with robotic fish in a complete open-410 loop operation that does not utilize any sensory input. These experiments could bring conclusive 411 evidence to our model predictions of a purely passive hydrodynamic pathway to rheotaxis, which 412 can only be partially supported through live experiments where live fish may still have access and 413 utilize the vestibular system and tactile sensors. 414

Despite its limitations, the proposed minimalistic model is successful in anticipating many of 415 the puzzling aspects of rheotaxis and points at the possibility of attaining rheotaxis in a purely pas-416 sive manner, without any sensory input. Most importantly, the model brings forward a potential 417 methodological oversight of laboratory practice in the study of rheotaxis, caused by bidirectional 418 hydrodynamic interactions between the swimming fish and the fluid flow. To date, there is no 419 gold standard for the selection of the size of the swimming domain, which is ultimately chosen on 420 the basis of practical considerations, such as facilitating behavioral scoring and creating a laminar 421 background flow. The model demonstrates that the width of the channel has a modulatory effect 422 on the threshold speed for rheotaxis and the cross-stream swimming frequency, which challenges 423 the comparison of different experimental studies and confounds the precise quantification of the 424 role of individual sensory modalities on rheotaxis. Overall, our effort warrants reconsidering the 425

behavioral phenotype of rheotaxis, by viewing fish as an invasive sensor that modifies the encom-

⁴²⁷ passing flow and hydrodynamically responds to it.

428 Methods and Materials

429 Derivation of the turn rate equation for the fish dynamics

⁴³⁰ The expression for the turning moment in equation (8) is obtained from the original finite-dipole

model by *Tchieu et al.* (2012b), in the limit of small distances between the vortices in the pair $(r_0 \rightarrow 0)$.

433 Specifically, equation (2.11) from *Tchieu et al.* (*2012b*), adapted to the case of a single dipole 434 reads

$$\dot{\theta}_{f} = \operatorname{Re}\left[\frac{\left(\mathcal{U}(\vec{r}_{f,r}) - i\mathcal{V}(\vec{r}_{f,r})\right) - \left(\mathcal{U}(\vec{r}_{f,l}) - i\mathcal{V}(\vec{r}_{f,l})\right)}{r_{0}}e^{i\theta_{f}}\right],\tag{12}$$

where subscript *l* and *r* refer to the left and right vortices forming the pair and $\vec{U} = U\hat{i} + V\hat{j}$ is the advective velocity field acting on the dipole. The advective field consists of the interactions with the walls and the background flow, so that $\vec{U}(\vec{r}) = \vec{u}_w(\vec{r}, \vec{r}_f, \theta_f) + \vec{u}_b(\vec{r})$; in the case of **Tchieu et al. (2012b)**, such a field encompasses the velocity field induced by any other dipole in the plane. Left and right

vortices are defined so that $\vec{r}_{f,l} = \vec{r}_f + r_0 \hat{v}_f^{\perp}/2$ and $\vec{r}_{f,r} = \vec{r}_f - r_0 \hat{v}_f^{\perp}/2$, which yields $\vec{r}_{f,l} - \vec{r}_{f,r} = \hat{v}_f^{\perp} r_0$.

By carrying out the complex algebra in (12), we determine

$$\dot{\theta}_f = \left(\frac{-\vec{\mathcal{U}}(\vec{r}_{f,l}) + \vec{\mathcal{U}}(\vec{r}_{f,r})}{r_0}\right) \cdot \hat{v}_f,\tag{13}$$

which supports the intuition that the dipole will turn counter-clockwise if the right vortex would

experience a stronger velocity along the swimming direction. Upon linearizing the term in paren-

thesis in the neighborhood of \vec{r}_{f} , this expression becomes

$$\dot{\theta}_f = -\nabla \vec{\mathcal{U}}(\vec{r_f}) \hat{v}_f^{\perp} \cdot \hat{v}_f.$$
(14)



Figure 4. Visual illustration of the process of determining the roots of (15). (a) Plot of the function $\frac{\pi^3}{32} \cot \left(\pi \left(\xi + \frac{1}{2}\right)\right) \csc^2 \left(\pi \left(\xi + \frac{1}{2}\right)\right)$ (black), superimposed with three lines of different slope: 200 (red), –200 (dashed blue), and –2 (solid blue). (b) Zoomed-in view of the curves in (a) showing that the blue line can only intersect the black curve at the origin.

⁴⁴⁴ Determination of the equilibria of the planar dynamical system

By setting $\theta_f = 0$ or $\theta_f = \pi$ in equation set (10), we determine that ξ should be equal to some constant, which is a root of the following transcendental equation:

$$\frac{\pi^3}{32}\cot\left(\pi\left(\xi+\frac{1}{2}\right)\right)\csc^2\left(\pi\left(\xi+\frac{1}{2}\right)\right) = \pm\beta\xi,\tag{15}$$

where the positive sign corresponds to $\theta_f = 0$ and the negative sign to $\theta_f = \pi$. Here, $\beta = \alpha(1 + \kappa)/\rho^2$ as introduced from the main text.

As shown in Fig. 4, for $\theta_f = 0$, there is only one root of the equation ($\xi = 0$; see the intersection

between the solid red line and the solid black curve), while up to three roots can rise for $\theta_f = \pi$

depending on the value of β . For β smaller than a critical value β^* , only $\xi = 0$ is a solution (see the intersection between the solid blue line and the solid black curve), while for $\beta > \beta^*$ two additional

Intersection between the solid blue line and the solid black curve), while for $\beta > \beta^*$ two additional solutions, symmetrically located with respect to the origin emerge (see the intersections between

the dashed blue line and the solid black curve). The critical value β^* is identified by matching the

slope of the black curve at $\xi = 0$, so that $\beta^* = \pi^4/32$. Notably, the two solutions symmetrically

located with respect to the centerline approach the walls as $\beta \to \infty$.

457 Local stability analysis of the planar dynamical system

458 To examine the local stability of the equilibria of the planar dynamical system, we linearize equation

459 set (10). The state matrix of the linearized dynamics, A, describes the local behavior of the nonlinear

system when perturbed in the vicinity of the equilibrium, that is,

$$\dot{\delta \mathbf{q}}(t) = A \,\delta \mathbf{q}(t),\tag{16}$$

where $\delta \mathbf{q} = [\delta \xi, \delta \theta_f]^T$ is the variation about the equilibrium. The eigenvalues of the *A* are indicative of local stability about each equilibrium.

For $\theta_f = 0$ and $\xi = 0$, the state matrix is given by

$$A = \begin{bmatrix} 0 & 1 - \frac{\pi^2 \rho^2}{6} \\ 8(1+\kappa)\alpha + \frac{\pi^4 \rho^2}{4} & 0 \end{bmatrix}.$$
 (17)

- Given that the trace of the matrix is zero (tr A = 0), the analysis of the stability of the equilibrium
- resorts to ensuring the sign of the determinant to be positive (det A > 0). Specifically, if the deter-
- 466 minant is positive, the eigenvalues are imaginary and the equilibrium is a neutral center (stable,
- although not asymptotically stable), otherwise one of the eigenvalues is positive and the equilib-
- rium is a saddle point (unstable) (*Bakker, 1991*). Hence, stability requires that

$$\frac{1}{24} \left(-6 + \pi^2 \rho^2\right) \left(32\alpha (1+\kappa) + \pi^4 \rho^2\right) > 0.$$
(18)

- Since the first factor is always negative ($\rho \ll 1$) and the second is positive, the inequality is never
- fulfilled and the equilibrium is a saddle point (unstable) (Fig. 3(a,b)).
- For $\theta_f = \pi$ and $\xi = 0$, the state matrix is given by

$$A = \begin{bmatrix} 0 & -1 + \frac{\pi^2 \rho^2}{6} \\ 8(1+\kappa)\alpha - \frac{\pi^4 \rho^2}{4} & 0 \end{bmatrix}.$$
 (19)

⁴⁷² Similar to the previous case, stability requires that det A > 0, that is,

$$\frac{1}{24} \left(-6 + \pi^2 \rho^2\right) \left(-32\alpha (1+\kappa) + \pi^4 \rho^2\right) > 0.$$
⁽²⁰⁾

⁴⁷³ Due to the sign change in the first summand appearing in the second factor with respect to the ⁴⁷⁴ previous case, stability becomes possible. Specifically, the equilibrium is a neutral center (stable) ⁴⁷⁵ for $\beta > \beta^* = \pi^4/32$, which is also the necessary condition for the existence of the two equilibria ⁴⁷⁶ symmetrically located with respect to the channel centerline (Fig. 3(a,c)).

When $\beta > \beta^*$, we register the presence of two more equilibria at $\pm \xi \neq 0$. The state matrix takes the form

$$A = \begin{bmatrix} 0 & -1 - \frac{\pi^2 \rho^2}{12} + \frac{1}{4} \pi^2 \rho^2 \sec^2(\pi\xi) \\ 8(1+\kappa)\alpha - \frac{1}{4} \pi^4 \rho^2 (2 - \cos(2\pi\xi)) \sec^4(\pi\xi) & 0 \end{bmatrix},$$
 (21)

Also in this case, stability requires that det A > 0, that is,

$$\frac{1}{48} \left(-12 + 3\pi^2 \rho^2 \sec^2(\pi\xi) - \pi^2 \rho^2 \right) \left(-32\alpha(1+\kappa) + \pi^4 \rho^2(2 - \cos(2\pi\xi)) \sec^4(\pi\xi) \right) > 0$$
(22)

Once again, for $\rho \ll 1$, we can assume that the first factor in parenthesis is negative. (This assump-

tion is grounded upon (15), which yields that $(\xi \pm 1/2) = \mathcal{O}(\rho^{2/3})$; since $\cos(\pi\xi)^2 = \mathcal{O}((\xi \pm 1/2)^2)$, we have that $\rho^2 \sec^2(\pi\xi) \to 0$ as $\rho \to 0$.) Hence, we obtain

$$\beta > \frac{\pi^4}{32} (2 - \cos(2\pi\xi)) \sec^4(\pi\xi), \tag{23}$$

which is not satisfied for any choice of $\beta > \beta^*$. Thus, the two equilibria away from the channel

centerline, close to the walls are always saddle points (unstable) (Figs. \exists (a,c)).

485 Frequency of cross-stream sweeping

The linearized planar system about the stable focus in (19) is equivalent to a classical second-order

487 system in terms of the cross-stream coordinate, similar to a mass-spring model. Hence, the radian

resonance frequency of the system is

$$\omega_0 = \sqrt{\det A} \simeq \frac{\pi^2}{2} \rho \sqrt{\frac{\beta}{\beta^*} - 1}.$$
(24)

where the last approximation holds for $\rho \ll 1$. Equation (24) shows that, close to the threshold, the

frequency of oscillations is small and it increases with β and ρ .

491 Bibliographical survey

492 We surveyed over three hundred publications cited by Arnold (1974) and Coombs et al. (2020) – two

review papers on rheotaxis, with the former focusing on early investigations from 1900s to 1970s,

and the latter highlighting more recent works conducted between 1970s and 2020. Publications were selected through the following inclusion and exclusion criteria.

Inclusion criteria: We select studies where: i) the subject animals were fish; ii) fish demonstrated 496 rheotactic behavior; iii) no unsteady flow events were present in the swimming domain, such as 497 the wake structure of obstacles: iv) the sensory cues available to fish could be identified with some 498 confidence: v) fish behavior was not influenced by social interactions: vi) fish swam without visual 499 cues; and vii) the publication was written in English. Within criterion iii), we focused on experi-500 ments with steady flows where the flow gradient is consistent over time, thus excluding swimming 501 in random flow events. Criterion vi) was introduced to direct our search toward the effects of hy-502 drodynamic cues and lateral line sensing, which limited our search to experiments using blind fish 503 or experiments in the dark. 504

Exclusion criteria: Among studies identified through the selection criteria, we excluded experiments on pleuronectiform flatfishes, which swim on their side and generate propulsive undulations in a vertical plane (*Webb, 2002*). This locomotory pattern differs fundamentally from the current model, derived on the assumption the fish align their bodies vertically and undulate on a horizontal plane, which is the swimming strategy of the majority of fishes.

Table 2 presents data extracted from the selected studies, including the fish species, size of the 510 swimming domain, flow conditions, sensory cues available to the fish, and the measured rheotaxis 511 threshold speed. Swimming domains with rectangular cross-sections are defined by their length 612 (L), width (h), and depth (W), while swimming domains with circular cross-sections by their length 513 and diameter (D). Flow conditions are quantified through the flow speed and the flow gradient. If 514 information of the flow gradient was not measured in a study, we qualitatively estimated its value 515 through the Reynolds number of the flow, defined based on the width of the channel (or diameter 516 of the channel in case of a cylindrical domain) and the background flow speed as $Re = \frac{hU_0}{2}$ (or 517 $Re = \frac{DU_0}{2}$), where v is the kinematic viscosity of water at room temperature. For a sufficiently high 518 Re, the flow gradient near the center of the channel is expected to be low. 519

When possible, we made direct comparisons between studies, as detailed in Table 1. The con-520 fidence intervals of the measured U_{a} values were estimated to determine if U_{a} were significantly 521 different across studies. When the mean and standard error of the mean (s.e.m.) of U, were 522 provided. we estimated its 95% confidence interval as (mean - 1.96 s.e.m., mean + 1.96 s.e.m.). If the 523 confidence intervals of two U_{1} values did not overlap, we considered them significantly different. 524 For instance, in Bak-Colemon and Coombs (2014) and Elder and Coombs (2015), the confidence 525 intervals of U, were determined to be (0.63, 1.17) cm/s and (1.27, 2.64) cm/s, respectively, and thus 526 the U_{a} values were considered significantly different. 527

In several studies, such as **Baker and Montgomery** (1999) and **Van Trump and McHenry** (2013), the threshold speeds were only estimated as intervals, where fish swimming below a lower bound did not perform rheotaxis, while they exhibited rheotaxis above an upper bound. We treated this speed interval as the confidence interval for U_c in our statistical analysis.

The relationship between the threshold speed and fish body size is less straightforward, as the body size not only determines the value of *I*, but also influences r_0 , which is on the order of the fish tail beat amplitude. We assumed $r_0 = 0.2I$, which is a typical tail-beat-amplitude-to-body-length ratio (*Gazzola et al., 2014*). For fish with functional lateral lines that produce positive feedback, K > 0, we obtain $U_c \sim 1 - \frac{1}{1+0.2Kl^2}$. For fish with disabled lateral line, K = 0, we find $U_c \sim l^2$. In both cases, the model predicts that U_c is larger for fish with larger body length. **Table 2.** Relevant publications on fish rheotaxis in the absence of visual cues, identified through literature review.

| Poforonco | Fish | | Swimming | Flow properties | | †Sonson/ cuos | Rheotaxis |
|-----------------------------------|---|--------------|--|--|---|--|--|
| Reference | Species | Length | domain | Flow speed | Flow gradient | sensory cues | threshold sneed |
| Bak-Coleman et al. (2013) | Giant danio (Devario aequipinnatus) | 6.0 – 7.3 cm | Flow tank of $25 \times 25 \times 25$ cm $(L \times h \times W)$ | 0, 3, and 7 cm/s | Re ~ 7500 at LL+ threshold speed; flow gradient expected to be small near center of tank | LL+/LL- | ≤ 3 cm/s |
| Bak-Coleman and Coombs (2014) | blind cavefish (Astyanax mexicanus) | 4.2 - 5.0 cm | Flow tank of $25 \times 25 \times 25$ cm $(L \times h \times W)$ | 0, 1, 2, 3, 4, 7, and 8 cm/s | Re ~ 2000 at LL+ threshold speed; flow gradient expected to be small near center of tank | LL+/LL–; fish made transient contacts with substrate | LL+: 0.90 cm/s; LL-: 0.54 cm/s |
| *Baker and Mont- gomery (1999) | blind cavefish (Astyanax fasciatus) | 4 – 7 cm | Flow tank of $51 \times 9 \times 20 \text{ cm}$ $(L \times h \times W)$ | 0, 2, 3, 5, 9, 16 cm/s | Re ~ 2000 at LL+ threshold speed; flow gradient expected to be small near center of tank | LL+/LL–; tactile senses | LL+: 2-3 cm/s; LL-: 9-16 cm/s |
| Elder and Coombs (2015) | Mexican tetras (Astyanax mexicanus) | 8.3 cm | Flow tank of $25 \times 25 \times 25$ cm $(L \times h \times W)$ | 0, 1, 2, 4, 7, and 12 cm/s | Re ~ 5000 at threshold speed; flow gradient expected to be small near center of tank | LL+/LL- | ~ 2 cm/s for LL+ and LL– |
| Kulpa et al. (2015) | blind cavefish (Astyanax mexicanus) | 4.4 – 5.3 cm | Flow tank of $25 \times 25 \times 10$ cm $(L \times h \times W)$ | Maximum speed of 8 cm/s | Jet flow across center of tank; flow gradient expected to be large | LL+/LL- | $\leq 8 \mathrm{cm/s}$ |
| [‡] Lyon (1904) | blind Fundulus | unspecified | Trough with unspecified dimensions; tideway leading to pond | "not too strong current" in trough and current with "more or less eddy and irregularity" in tideway | Flow gradient expected to be small | LL+; some fish gained tactile senses | Not measured; rheotaxis elicited only by tactile cues |
| [‡] Lyon (1904) | blind Fundulus | unspecified | Trough with unspecified dimensions | flow "gushing rather violently" | Jet flow; flow gradient expected to be large | LL+ | Not measured; rheotaxis elicited by flow |
| *Montgomery et al. (1997) | blind cavefish (Astyanax fasciatus) | 4 – 7 cm | [§] Flow tank of 51 \times 9 \times 20 cm ($L \times h \times W$) | 0, 2, 3, 5, 9, and 16 cm/s | Re ~ 2000 at LL+ threshold speed; flow gradient expected to be small near center of tank | LL+/LL–; tactile senses | LL+: 2-3 cm/s; LL-: 9-16 cm/s |

Table 2. Relevant publications on fish rheotaxis in the absence of visual cues, identified through literature review.

| Reference - | Fish | | Swimming | Flow properties | | †Consorry guos | Rheotaxis |
|-----------------------|----------------|------------------------|----------------------------|-----------------------|---|----------------|-----------------|
| | Species | Length | domain | Flow speed | Flow gradient | - Sensory cues | threshold speed |
| Oteiza et al. (2017) | zebrafish | unspecified | 13 cm-long | 0.2-0.8 cm/s | Low to high flow gradients | LL+/LL- | LL+: rheotaxis |
| | (Danio rerio) | | circular tube | | identified through particle | | observed as low |
| | larva 5–7 days | | with diameter | | image velocimetry | | as 0.2 cm/s |
| | post | | 1.27-4.76 cm | | | | |
| | fertilization | | | | | | |
| | (dpf) | | | | | | |
| Peimani et al. (2017) | zebrafish | estimated | Flow channel | 0.95-3.8 cm/s | $\text{Re} \sim 10$ at threshold speed; | LL+ | 0.95 cm/s |
| | (Danio rerio) | $\sim 0.35\mathrm{cm}$ | of 63.3 × 1.6 × | | flow gradient expected to | | |
| | larva 5–7 dpf | | 0.55 mm | | be large | | |
| | | | $(L \times h \times W)$ | | | | |
| Suli et al. (2012) | zebrafish | ~ 0.33 cm | Flume of | 0.075, 0.15, 0.2 cm/s | Re < 75; flow gradient | LL+/LL- | Not quantified |
| | (Danio rerio) | | 110×3.7×2.8 cm | | expected to be large | | |
| | larva 5 dpf | | $(L \times h \times W)$ | | | | |
| Van Trump and | blind Mexican | 3 – 7 cm | Cylindrical | 0, 1, 2, 4, 6, 8, 10, | Re > 2000 at threshold | LL+/LL- | 2-4 cm/s |
| McHenry (2013) | cavefish | | channel of | 13, 16 cm/s | speed; flow gradient | | |
| | (Astyanax | | $150 \times 11 \text{ cm}$ | | expected to be small near | | |
| | fasciatus) | | $(L \times D)$ | | center of tank | | |

538 [†]LL+: lateral line enabled; LL–: lateral line disabled

539 * Data are extracted from the same set of experiments

 ${}^{$}$ Two experiments are considered from the same paper

541 § Data are from *Baker and Montgomery* (1999)

542 Estimation of model parameters

In a typical experimental setup on rheotaxis, the width of the channel, h_{i} is on the order of three 543 to ten times the body length of the animal, *l*. For example, experiments from *Elder and Coombs* 544 (2015) on Mexican tetras of $l = 8.3 \,\mathrm{cm}$ were conducted in a channel with $h = 25 \,\mathrm{cm}$. Similarly, in **5**/ **5** the experiments on adult zebrafish from **Burbano-L and Porfiri (2021)**, l = 3.6 cm and h = 13.8 cm. 546 and in the experiments on zebrafish larvae from **Oteizg et al.** (2017), l = 4.2 mm (inferred from the 547 animals' age) and h = 1.27 - 4.76 cm. The distance between the vortices simulating a fish, r_0 , should 548 be on the order of a tail beat, which has a typical value of 0.21 (Gazzola et al., 2014). As a result, it 549 is tenable to assume that ρ^2 is between 10^{-4} and 10^{-2} . 550 A safe estimation of the velocity of the animal in the absence of the background flow, v_0 , would 551 be on the order of few body lengths per second (*Gazzola et al., 2014*). The speed used for the 552 background flow across experiments, U_0 , tend to be of the same order as the magnitude of v_0 553 leaning toward values close to one body length per second (Coombs et al., 2020). For instance, 554 data on zebrafish from **Burbano-L and Porfiri (2021)** suggest $v_0 = 5.7 \,\mathrm{cm \, s^{-1}}$ and $U_0 = 3.2 \,\mathrm{cm \, s^{-1}}$. 555 The estimation of the non-dimensional parameter ϵ associated with the shear in the flow is more 556 difficult, since data on the velocity profiles are seldom reported. That being said, for channel flow 557 of sufficiently high Reynolds number, the velocity profile in the channel is expected to be blunt. approximating a uniform flow profile near the channel center (*White, 1974*). Thus, it is tenable to 559 treat ϵ as a small parameter, between 10^{-2} and 10^{-1} . For flow of low Reynolds number (*Oteiza et al.*, 560 **2017**) (Re < 100), the velocity gradient in the channel has been observed to be large, corresponding 561 to ϵ values in the range of 10^{-1} and 1. By combining these estimations, we propose that α ranges 562 between 0 and 1 563

An estimation of κ is difficult to offer, whereby feedback from the lateral line has only been 564 included in few studies (Oteiza et al., 2017; Burbano-L and Porfiri, 2021; Colvert and Kanso, 2016; 565 Chicoli et al., 2015). Using the data-driven model from Burbano-L and Porfiri (2021), it is tenable 566 to assume values on the order of 10^1 for individuals showing high rheotactic performance. This 567 gain can also be estimated by comparing the threshold speeds of fish with and without the lateral 568 line, through $\frac{U_c(LL-)}{U_c(LL+)} = 1 + \kappa$, according to (11). The significant increase in the threshold speed 569 following lateral line ablation in **Baker and Montgomery** (1999) indicates that $\kappa \in [2,7]$, while the 570 indistinguishable threshold speed between LL+ and LL– fish in a few other studies (Bak-Coleman 571 and Coombs, 2014; Elder and Coombs, 2015; Van Trump and McHenry, 2013) may suggest that 572 $\kappa \sim 0$. In Table 3, we summarize the model parameters identified from data in the experimental 573

studies comprising Tables 1 and 2.

| Reference | ρ | e | α | К | β |
|-------------------------|----------------|----------------------|------------------------|----------|----------|
| Bak-Coleman et al. | ~ 0.05 | $[10^{-2}, 10^{-1}]$ | [0, 0.17] | _ | _ |
| (2013) | | | | | |
| Bak-Coleman and | ~ 0.04 | $[10^{-2}, 10^{-1}]$ | [0, 0.16] | ~ 0 | [0, 100] |
| Coombs (2014) | | | | | |
| Baker and Montgomery | ~ 0.1 | $[10^{-2}, 10^{-1}]$ | [†] [0, 0.32] | [2,7] | [0,256] |
| (1999) and Montgomery | | | | | |
| et al. (1997) | | | | | |
| Elder and Coombs (2015) | ~ 0.066 | $[10^{-2}, 10^{-1}]$ | [†] [0, 0.24] | ~ 0 | [0, 55] |
| Kulpa et al. (2015) | ~ 0.04 | ~ 1 near center | ~ 1.3 | _ | _ |
| | | of jet | | | |
| Oteiza et al. (2017) | [0.018, 0.066] | [0.20, 0.82] | _ | _ | _ |
| Peimani et al. (2017) | ~ 0.044 | ~ 1 | _ | _ | _ |
| Suli et al. (2012) | ~ 0.018 | [0.1, 1] | _ | _ | _ |
| Van Trump and McHenry | [0.055, 0.127] | $[10^{-2}, 10^{-1}]$ | [†] [0, 0.32] | ~ 0 | [0, 106] |
| (2013) | | | | | |

Table 3. Estimation of model parameters from data in the literature.

 t_{a} tL+ cavefish swimming speed $v_0 \sim 5$ cm/s in zero background flow in **Bak-Coleman and Coombs (2014)** is used to estimate α

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Author Contributions

- 500 M.P. and S.D.P. conceived the study. M.P. and S.D.P. developed the theoretical model and per-
- formed data analysis. P.Z. conducted the literature review and performed the model validation.
- M.P. and S.D.P. wrote a first draft of the manuscript, which was consolidated in its present form by
- ⁵⁸³ all the authors.
- **584** Competing Interests
- ⁵⁰⁵ The authors declare that they have no competing financial interests.
- **Data and materials availability**
- ⁵⁸⁷ The authors declare that the data supporting the findings of this study are available within the pa-
- per. The Mathematica notebook used to derive the governing equations, study the planar dynami-
- cal, and generate associated figures are also available at https://github.com/dynamicalsystemslaboratory/
 Rheotaxis.
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793 Appendix 1

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Complete expression for the velocity field caused by image dipoles

The velocity field $\vec{u}_f = u_f \hat{i} + v_f \hat{j}$ at \vec{r} induced by the single dipole at \vec{r}_f , given by the potential function in equation (1), is

$$u_f(\vec{r}, \vec{r}_f, \theta_f) = r_0^2 v_0 \left(\frac{((x - x_f)^2 - (y - y_f)^2) \cos \theta_f + 2(x - x_f)(y - y_f) \sin \theta_f}{((x - x_f)^2 + (y - y_f)^2)^2} \right),$$
 (25a)

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$$v_f(\vec{r}, \vec{r}_f, \theta_f) = r_0^2 v_0 \left(\frac{-((x - x_f)^2 - (y - y_f)^2)\sin\theta_f + 2(x - x_f)(y - y_f)\cos\theta_f}{((x - x_f)^2 + (y - y_f)^2)^2} \right).$$
(25b)

The potential function describing the image vortex system for a dipole in a channel presented in equation (3) can be simplified using Mathematica, yielding

$$\phi_{w}(\vec{r},\vec{r}_{f},\theta_{f}) = \frac{r_{0}^{2}v_{0}}{4} \left[4 \frac{(x-x_{f})\cos\theta_{f} + (y-y_{f})\sin\theta_{f}}{(x-x_{f})^{2} + (y-y_{f})^{2}} - \frac{\pi e^{-i\theta_{f}}}{h} \left(e^{2i\theta_{f}} \left(\coth(\pi A) + \coth(\pi B^{*}) \right) + \coth(\pi A^{*}) + \coth(\pi B) \right) \right], \quad (26)$$

where $A = ((x - x_f) + i(y - y_f))/(2h)$, $B = ((x - x_f) + i(y + y_f))/(2h)$, $i = \sqrt{-1}$, and a superscript * indicates complex conjugate. The velocity field at \vec{r} due to the walls, $\vec{u}_w = u_w \hat{i} + v_w \hat{j}$, is

$$u_w = \frac{r_0^2 v_0}{4} \left[\frac{\pi^2 e^{-i\theta_f}}{2h^2} \left(e^{2i\theta_f} (\operatorname{csch}^2 \pi A + \operatorname{csch}^2 \pi B^*) + \operatorname{csch}^2 \pi A^* + \operatorname{csch}^2 \pi B \right) \right) \\ + \frac{4 \cos \theta_f}{(x - x_f)^2 + (y - y_f)^2} - \frac{8(x - x_f)((x - x_f)\cos \theta_f + (y - y_f)\sin \theta_f)}{((x - x_f)^2 + (y - y_f)^2)^2} \right], \quad (27a)$$

$$v_w = \frac{r_0^2 v_0}{4} \left[\frac{i\pi^2 e^{-i\theta_f}}{2h^2} \left(e^{2i\theta_f} (\operatorname{csch}^2 \pi A - \operatorname{csch}^2 \pi B^*) - \operatorname{csch}^2 \pi A^* + \operatorname{csch}^2 \pi B \right) \right) \\ + \frac{4\sin\theta_f}{(x - x_f)^2 + (y - y_f)^2} - \frac{8(y - y_f)((x - x_f)\cos\theta_f + (y - y_f)\sin\theta_f)}{((x - x_f)^2 + (y - y_f)^2)^2} \right].$$
(27b)

Superimposing the velocity fields from the dipole and its images and setting y = 0 (or y = h) yields $v_f + v_w = 0$, thereby confirming that the walls of the channel are streamlines.

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