Stirring by swimming bodies

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A controversial proposition:

- There are many regions of the ocean that are relatively quiescent, especially in the depths (1 hairdryer/ km$^3$);
- Yet mixing occurs: nutrients eventually get dredged up to the surface somehow;
- What if organisms swimming through the ocean made a significant contribution to this?
- There could be a *local* impact, especially with respect to feeding and schooling;
- Also relevant in suspensions of microorganisms (Stokes regime).
Munk’s Idea

Though it had been mentioned earlier, the first to seriously consider the role of biomixing was Walter Munk (1966):

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Abyssal recipes

WALTER H. MUNK

(Received 31 January 1966)

Abstract—Vertical distributions in the interior Pacific (excluding the top and bottom kilometer) are not inconsistent with a simple model involving a constant upward vertical velocity \( w \approx 1 \cdot 2 \text{ cm day}^{-1} \) and eddy diffusivity \( \kappa \approx 1 \cdot 3 \text{ cm}^2 \text{ sec}^{-1} \). Thus temperature and salinity can be fitted by exponential-like solutions to \( \kappa \cdot d^2/dz^2 \equiv -w \cdot d/dz \) \( T, S = 0 \), with \( \kappa/w \approx 1 \text{ km} \) the appropriate “scale height.” For Carbon 14 a decay term must be included, \( \kappa \cdot d^2/dz^2 \equiv 14\text{C} = \mu 14\text{C} \); a fitting of the solution to the observed 14C distribution yields \( \kappa/w^2 \approx 200 \text{ years} \) for the appropriate “scale time,” and permits \( w \) and
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“...I have attempted, without much success, to interpret [the eddy diffusivity] from a variety of viewpoints: from mixing along the ocean boundaries, from thermodynamic and biological processes, and from internal tides.”
Basic claims

The idea lay dormant for almost 40 years; then

- Huntley & Zhou (2004) analyzed the swimming of 100 (!) species, ranging from bacteria to blue whales. Turbulent energy production is $\sim 10^{-5} \text{ W kg}^{-1}$ for 11 representative species.
- Total is comparable to energy dissipation by major storms!
- Another estimate comes from the solar energy captured: 63 TeraW, something like 1% of which ends up as mechanical energy (Dewar et al., 2006).
- Kunze et al. (2006) find that turbulence levels during the day in an inlet were 2 to 3 orders of magnitude greater than at night, due to swimming krill.
Rain on the parade

Visser (2007) debunks these claims:

Let the turbulence be generated at a scale $L$, with a rate of turbulent energy dissipation $\varepsilon$.

The **buoyancy frequency** $N$ is defined as

$$N^2 = -\frac{g}{\rho} \frac{d\rho}{dz}$$

where $g$ is the gravitational acceleration and $\rho(z)$ is the density.

The **buoyancy length scale** is

$$B = \left(\frac{\varepsilon}{N^3}\right)^{1/2}$$
The mixing efficiency is defined as

$$\Gamma = \frac{\text{change in potential energy}}{\text{work done}}$$

so $0 \leq \Gamma \leq 1$.

Visser’s point is that $\Gamma$ depends strongly on $L/B$.

For krill $L = 1.5$ cm, $B = 3$ to $10$ m, so $L/B = .005$ to .0015.

Conclude: $\Gamma = 10^{-4}$ to $10^{-3}$: almost none of the turbulent energy goes into mixing.

(from Visser (2007))
But it’s not over... 

Katija & Dabiri (2009) looked at jellyfish:

[movie 1]
Displacement by a moving body

(from Darwin (1953))

[movie 2] (movie from Katija & Dabiri (2009))
A sequence of kicks

The age-old paradigm for calculating an effective diffusivity consists of assuming a test particle undergoes uncorrelated “kicks”: if a test particle initially at $x(0) = 0$ undergoes $N$ encounters with axially-symmetric swimming bodies, its position is

$$x(t) = \sum_{k=1}^{N} \Delta(a_k) \hat{r}_k$$

where $\Delta(a)$ is the displacement, $a_k$ is the impact parameter, and $\hat{r}_k$ is a direction vector.

After squaring and averaging, assuming isotropy:

$$\langle |x|^2 \rangle = N \langle \Delta^2(a) \rangle$$

where $a$ is treated as a random variable.
Assuming the swimmers move in a straight line at speed $U$, the number that will hit an “interaction disk” of radius $R$ in time $t$ is $2RUnt$, where $n$ is the number density.

The approach from infinity means that $a$ is distributed as $da/R$. Putting this together,

$$\langle |x|^2 \rangle = 2Unt \int_0^\infty \Delta^2(a) \, da = 4\kappa t, \quad \text{2D}$$

which defines the effective diffusivity $\kappa$.

In 3D, the factors are modified slightly:

$$\langle |x|^2 \rangle = 2\pi Unt \int_0^\infty a \Delta^2(a) \, da = 6\kappa t, \quad \text{3D}$$
Numerical simulation

- Validate theory using simple simple simulations;
- Periodic box of size $L$;
- $N$ swimmers (spheres of radius 1), initially at random positions, swimming in random direction with constant speed $U = 1$;
- Target particle initially at origin advected by the swimmers;
- Since dilute, superimpose velocities;
- Integrate for some time, compute $|\mathbf{x}(t)|^2$, repeat for a large number $N_{\text{real}}$ of realizations and average.
A ‘gas’ of swimmers

[movie 3] \( N = 100 \) swimmers, \( L = 1000 \)
How well does the dilute theory work?

\[ \frac{\langle |x|^2 \rangle}{2nU^3} \]

- \( n = 10^{-3} \)
- \( n = 5 \times 10^{-4} \)
- \( n = 10^{-4} \)
- theory
Diffusion is dominated by rare events

2 × 10^6 realizations of N = 10 cylinders, with L = 1000
Contribution to displacement

Small $a$: $\Delta \sim -\log a$, large $a$: $\Delta \sim a^{-3}$ (Darwin, 1953)

\[
\log a = -5
\]

\[
\log a = 0
\]

\[
\log a = 2
\]

\[
\int_0^1 \Delta^2(a)da \simeq 2.31, \text{ whilst } \int_1^{\infty} \Delta^2(a)da \simeq .06.
\]

\[\Longrightarrow 97\% \text{ dominated by “head-on” collisions}\]
Origin of the singularity

At the leading and trailing ‘edges’ of a body, there is a hyperbolic point. Locally,

\[ \dot{x} = -\lambda x, \quad \dot{y} = \lambda y \]

so that \( y(t) = y_0 \exp(\lambda t) \). The time it takes to go from \( y_0 = a \) to \( y > a \) is

\[ t = \lambda^{-1} \log(y/a) \sim -\lambda^{-1} \log a \]

which is the source of the logarithmic divergence of the displacement:

\[ \Delta \sim -2U\lambda^{-1} \log a, \quad a \ll 1 \]

The factor of 2 is for leading±trailing edges.
Four cases

For a wedge half-angle $\beta$, $\Delta$ has four possible behaviors as $a \rightarrow 0$:

- $\pi > \beta > \frac{\pi}{2}$: $a^\gamma$, nonsingular
- $\beta = \frac{\pi}{2}$: $-\log a$, integrable
- $\frac{\pi}{2} > \beta > \frac{\pi}{4}$: $a^{-\gamma}$, integrable
- $\frac{\pi}{4} > \beta > 0$: $a^{-\gamma}$, non-integrable
Sphere in Stokes flow

A natural question is what happens in the presence of viscosity, which greatly increases the “sticking” to the swimmer’s surface?

(from Camassa et al., Sphere Passing Through Corn Syrup)
Considerable literature on transport due to microorganisms: Wu & Libchaber (2000); Hernandez-Ortiz et al. (2006); Saintillian & Shelley (2007); Ishikawa & Pedley (2007); Underhill et al. (2008); Ishikawa (2009)
Stokesian Spheres: Transport

The transport is not diffusive, but is nearly **ballistic**!

*(short-time result)*

\[ \langle |x|^2 \rangle^{1/2} \]

exponent = 0.992

prefac = 0.013
Squirmers

One problem with the Stokesian spheres is that they are an awful model for swimming: there is a net force on the fluid. It’s as if the spheres are pulled by invisible threads.

Lighthill (1952), Blake (1971), and more recently Ishikawa et al. (2006) have considered squirmers:

- Sphere in Stokes flow;
- Steady velocity specified at surface, to mimic cilia;
- Steady swimming condition imposed (no net force on fluid).

(Drescher et al., 2009)  (Ishikawa et al., 2006)
3D axisymmetric streamfunction for a typical squirmer, in cylindrical coordinates \((\rho, z)\):

\[
\psi(\rho, z) = -\frac{1}{2} \rho^2 + \frac{1}{2r^3} \rho^2 + \frac{3\beta}{4r^3} \rho^2 z \left(\frac{1}{r^2} - 1\right)
\]

where \(r = \sqrt{\rho^2 + z^2}\), \(U = 1\), radius of squirmer = 1.

Note that \(\beta = 0\) is the sphere in potential flow!

We will use \(\beta = 5\) for most of the remainder.
Particle motion for squirmer

A particle near the squirmer’s swimming axis initially (blue) moves towards the squirmer.

After the squirmer has passed the particle follows in the squirmer’s wake.

(The squirmer moves from bottom to top.)

[movie 4]
Small $a$ asymptotics for squirmer

Four stagnation points for the squirmer ($B$ is a “ring” around the squirmer). A particle coming close to the axis from $z = \infty$ will encounter $A$, $B$, $C$ in turn, but never come near the trailing edge stagnation point $D$.

The relative contribution of each point is proportional to $-\lambda^{-1} \log a$, where $\lambda$ is the coefficient of the linearized flow:

\[
\lambda_A^{-1} = \frac{2}{3(\beta + 1)} \approx 0.1111
\]

\[
\lambda_B^{-1} = \frac{4\beta}{3(\beta^2 - 1)} \approx 0.2778
\]

\[
\lambda_C^{-1} = (\text{mess}) \approx 3.0095
\]
Displacement for squirmer

\[ \log \Delta = \log a = -4 \]
\[ \log \Delta = \log a = 0 \]
\[ \log \Delta = \log a = 2 \]

\[ \rightarrow 81\% \text{ dominated by “head-on” collisions,} \]
\[ \text{or 92\% if we use the \textit{wake radius}, 1.96.} \]
Squirmers: Transport

\[ \langle |x|^2 \rangle \]
The short-time transport is anomalous. Exponent is consistent with Wu & Libchaber (2000).
Long-range correlations

Because the net displacement decays faster than total extent of trajectory, sensitive to perturbations.
Conclusions

- Biomixing: no verdict yet;
- Simple dilute model works well, at least for potential flow;
- Potential flow dominated by “sticking”;
- Stokes flow dominated by long-range effects;

Future work:
- Wake models and turbulence;
- PDF of scalar concentration;
- Buoyancy effects;
- Schooling: longer length scale?
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