Realistic Animation of Liquids

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http://www.cis.upenn.edu/~fostern/liquids.html

Abstract

We present a comprehensive methodology for realistically animating liquid phenomena. Physically accurate 3D motion is achieved by performing a two-stage calculation over an arbitrary environment of static obstacles surrounded by fluid. A finite difference approximation to the Navier-Stokes equations is first applied to a low resolution, voxelized representation of the scene. The resulting velocity and pressure fields describe the gross transport of liquid, including effects such as splashing, vorticity and overturning. Local fluid velocity is then used to drive a height field equation or to convect massless marker particles. The position of any free surface can thus be determined to a significantly higher resolution than that of the Navier-Stokes calculation. In addition, the pressure field, together with the Lagrange equations of motion, is used to simulate dynamic buoyant objects. Typical disadvantages to volumetric methods such as poor scalability and lack of control are addressed by assuming that stationary obstacles align with grid cells during the finite difference discretization, and by appending driving functions to the Navier-Stokes equations. The output from our system is suitable for many of the water rendering algorithms presented by researchers in recent years.

Keywords: Fluid Simulations, Navier-Stokes Equations, Physics-Based Modeling, Free-Surface Flow.

1. Introduction

Some of the most breathtaking animations in recent years have been generated by modeling the interaction between light and water. Effects such as caustic shading, reflection, refraction, and internal scattering have been addressed in some detail, with realistic results. One characteristic of this work however, has been that the motion of the water surface is approximated by a non physics-based function. Suggested methods have included parametric functions and sinusoidal phase functions. Two exceptions to this are the papers by Kass and Miller, and Chen and Lobo. Kass and Miller use a fast approximation to the two-dimensional shallow water equations to simulate surface waves in water of varying depth. Their model allows for the reflection and refraction of waves, and takes account of mass transport, but it does not address the full range of three-dimensional motion found in a liquid. Such motion includes rotational and pressure based effects responsible for the much of a fluid’s characteristic behavior. They also cannot easily incorporate dynamic objects or buoyant effects into the model, because the velocity of the fluid is known only on the surface, and internal pressure is not calculated at all. Chen and Lobo go further towards a physics-based solution by solving a simplified form of the Navier-Stokes equations in two dimensions. However, they assume that the fluid has zero depth, and calculate the elevation of the surface solely from the instantaneous pressure. This allows them to perform some interaction between moving objects and the flow field, but restricts the class of problems that can be solved using the method. Notably, obstacle geometry must be two-dimensional, and although the surface height is varied for animation, they treat the fluid as being completely flat during the calculation. Therefore, convective wave effects, mass transport, and submerged obstacles are not covered by their technique.

Comprehensive models of fluid motion do exist, and there are a variety of tools for solving them in the field of Computational Fluid Dynamics (CFD). These methods generally involve direct simulation techniques to get accurate fluid motion. Unfortunately, in any direct simulation technique the temporal resolution is strongly coupled to the spatial resolution. Thus, if the spatial resolution doubles, the temporal resolution must also be doubled so that the solution does not move more than one spatial sample per time step. This gives running times proportional to the fourth power of the resolution, so most of these techniques will scale poorly. Furthermore, an animator needs a fairly clear understanding of the system of equations being solved so that he or she can set initial and boundary conditions to get the desired results. An ideal fluid simulator for graphics applications would apply the correct conditions automatically based on the underlying geometry. CFD methods also resist external control, making it difficult to force a particular motion from a fluid, unless it is a natural consequence of the system. These restrictions are an inherent part of the fluid modeling problem. The question arises whether it is possible to accurately model realistic fluid motion while keeping within acceptable efficiency bounds for Computer Graphics.

In this paper we present a solution to the Navier-Stokes equations for modeling liquid motion, that satisfies many of an animator’s needs. Realism is provided through a finite difference approximation to the incompressible Navier-Stokes equations. This gives rise to a complete pressure and velocity profile of the simulated environment. This profile is then used to determine the behavior of free surfaces, and is loosely coupled to the Lagrange equations of motion to include buoyant rigid objects into a scene. The range of behaviors accounted for include wave effects such as refraction, reflection and diffraction, together with

This research is partially supported by ARPA DAMD17-94-J-4486; DMS0 DAAH04-94-G-0402; National Library of Medicine N01LM-43551 and ARO DURIP DAAH04-95-1-0023.
rotational motion such as eddies and vorticity. Furthermore, velocity and pressure are strongly coupled within the model. This means that even the simplest animation exhibits subtle realistic behavior not available using previous computer-graphics fluid models.

Usability has also been a strong motivation for this paper. The Navier-Stokes equations are solved over a coarse, rectangular mesh containing an arbitrary distribution of submerged or semi-submerged obstacles. Boundary conditions for the mesh are generated automatically by constraining the free variables at an obstacle-fluid or air-fluid boundary. This low resolution calculation together with homogeneous boundary conditions leads to a relatively efficient determination of fluid velocity and internal pressure. Detail is achieved by using the velocity field to concentrate attention on regions of interest, i.e., the fluid surface. The surface is represented as either a chain of massless marker particles, or a height field. The markers are carried around the mesh by convection, and can have arbitrary connectivity, accounting for multiple colliding surfaces in a scene.

Consideration is also given to controlling the overall behavior of the fluid. Liquid sources or sinks (known as inflow and outflow boundaries) can be included anywhere in the environment. They allow liquid to flow (or be forced) into a scene, or flow out at a natural rate. A time dependent pressure field may also be applied to the fluid surface. Thus, the effects of a strong wind can be simulated and initial waves be driven realistically. The output from the system exhibits subtle realistic behavior not available using previous equations. They are derived from Newton’s Second Law due to the local pressure gradient, the kinematic viscosity, \( \nu \), or thickness of the fluid. Together with appropriate boundary conditions and the constraint that not only momentum, but also mass should be conserved (see Section 3.1), the Navier-Stokes equations can be used to accurately simulate fluid phenomena.

\[ \begin{align*}
\frac{\partial u}{\partial t} + u \frac{\partial u}{\partial x} + v \frac{\partial u}{\partial y} + w \frac{\partial u}{\partial z} &= -\frac{\partial p}{\partial x} + \frac{\nu}{\rho} \left( \frac{\partial^2 u}{\partial x^2} + \frac{\partial^2 u}{\partial y^2} + \frac{\partial^2 u}{\partial z^2} \right) \\
\frac{\partial v}{\partial t} + u \frac{\partial v}{\partial x} + v \frac{\partial v}{\partial y} + w \frac{\partial v}{\partial z} &= -\frac{\partial p}{\partial y} + \frac{\nu}{\rho} \left( \frac{\partial^2 v}{\partial x^2} + \frac{\partial^2 v}{\partial y^2} + \frac{\partial^2 v}{\partial z^2} \right) \\
\frac{\partial w}{\partial t} + u \frac{\partial w}{\partial x} + v \frac{\partial w}{\partial y} + w \frac{\partial w}{\partial z} &= -\frac{\partial p}{\partial z} + \frac{\nu}{\rho} \left( \frac{\partial^2 w}{\partial x^2} + \frac{\partial^2 w}{\partial y^2} + \frac{\partial^2 w}{\partial z^2} \right) + g z, \tag{1}
\end{align*} \]

where \( u, v, w \) are velocities in the \( x, y, z \) directions respectively, \( p \) is the local pressure, \( g \) is the acceleration due to gravity (or any body force) \( \gamma \), and \( \rho \) is the density of the fluid. Together with appropriate boundary conditions and the constraint that not only momentum, but also mass should be conserved (see Section 3.1), the Navier-Stokes equations can be used to accurately simulate fluid phenomena.

3. Solving the Navier-Stokes equations

Despite the complexity of a system of equations such as (1), it is possible to solve it in an intuitive way, using standard analysis tools. The first step is to discretize both the equations and the environment that we want to model. There are a number of ways to do this, but it is important to keep four things in mind:

- In a typical graphics application involving liquids, there are many boundaries between the liquid and other objects, and between the liquid and the surrounding medium. Computation cost can be minimized if such interfaces are homogeneously incorporated into the model instead of being treated as special cases.

- It must be possible to apply some external control to the system so that the animator can accurately specify how the liquid will behave.

- The range of motion that can be animated using the technique should include the set of effects available with existing computer-graphics methods, and extend it by adding new, interesting, and useful behavior.

With some thought, a good discretization that provides a solution to the first of these constraints also provides solutions to the other three. In the following sections we present a numerical solution to the Navier-Stokes equations. This technique combines a low resolution 3D calculation to determine velocity and pressure fields within the liquid, with a height field equation that is used to precisely track the position of a free surface. At all times during the computation, boundary conditions due to solid obstacles and the fluid surface are homogeneous, and their application is transparent to the user.

2. Navier-Stokes Equations

The motion of a fluid at any point within a flow is completely described by a set of non-linear equations known as the momentum or Navier-Stokes equations. In three dimensions, for an incompressible fluid such as water, these equations can be written as

\[ \begin{align*}
\frac{\partial u}{\partial t} + u \frac{\partial u}{\partial x} + v \frac{\partial u}{\partial y} + w \frac{\partial u}{\partial z} &= -\frac{\partial p}{\partial x} + \frac{\nu}{\rho} \left( \frac{\partial^2 u}{\partial x^2} + \frac{\partial^2 u}{\partial y^2} + \frac{\partial^2 u}{\partial z^2} \right) \\
\frac{\partial v}{\partial t} + u \frac{\partial v}{\partial x} + v \frac{\partial v}{\partial y} + w \frac{\partial v}{\partial z} &= -\frac{\partial p}{\partial y} + \frac{\nu}{\rho} \left( \frac{\partial^2 v}{\partial x^2} + \frac{\partial^2 v}{\partial y^2} + \frac{\partial^2 v}{\partial z^2} \right) \\
\frac{\partial w}{\partial t} + u \frac{\partial w}{\partial x} + v \frac{\partial w}{\partial y} + w \frac{\partial w}{\partial z} &= -\frac{\partial p}{\partial z} + \frac{\nu}{\rho} \left( \frac{\partial^2 w}{\partial x^2} + \frac{\partial^2 w}{\partial y^2} + \frac{\partial^2 w}{\partial z^2} \right) + g z, \tag{1}
\end{align*} \]
3.1. Discretization

We solve (1) across the entire environment. Solid obstacles and the atmosphere are treated as fluid, but with special properties that remain constrained throughout the calculation. The computation domain is first divided into a fixed rectangular grid aligned with a Cartesian coordinate system. The u, v, and w velocities are defined at the centers of each face of a cell and referenced locally (see Figure 1) while pressure, \( p \), is defined in the center of each cell. Note from the figure that \( v_{i, j-1/2, k} \equiv v_{i, j-1} \). At the start of the calculation the contents of each cell are determined. A cell may either contain a solid obstacle, be Full of fluid, be a Surface cell on the boundary between the liquid and surrounding medium, or be Empty. In all four cases, the velocity and pressure fields are defined everywhere.

This discretization leads to an explicit finite difference approximation of (1) in the form

\[
\begin{align*}
\frac{\partial u_{i+1/2,j,k}}{\partial t} & = u_{i+1/2,j,k} + \frac{1}{\sqrt{\varepsilon}} \left[ \frac{1}{\varepsilon} \left( (u_{i,j,k})^2 - (u_{i+1,j,k})^2 \right) 
+ \frac{1}{\varepsilon} \left( (w_{i+1,j+1/2,k} - w_{i+1,j-1/2,k}) + \frac{1}{\sqrt{\varepsilon}} \left( (u_{i+1,j+1/2,k})^2 - (u_{i+1,j-1/2,k})^2 \right) 
+ \frac{1}{\varepsilon} \left( (p_{i+1,j,k} - p_{i+1,j+1/2,k}) + \frac{1}{\sqrt{\varepsilon}} \left( (u_{i+1,j+1/2,k})^2 - (u_{i+1,j-1/2,k})^2 \right) \right. 
- 2u_{i+1/2,j,k} + u_{i-1/2,j,k} + \frac{1}{\varepsilon^2} \left( (u_{i+1,j+1/2,k})^2 + (u_{i+1,j-1/2,k})^2 \right) 
- 2u_{i+1/2,j,k} + u_{i+1/2,j+1/2,k} + \frac{1}{\varepsilon^2} \left( (u_{i+1,j+1/2,k})^2 + (u_{i+1,j-1/2,k})^2 \right) \right) 
- 2u_{i+1/2,j,k} + u_{i+1/2,j-1/2,k} + \frac{1}{\varepsilon^2} \left( (u_{i+1,j+1/2,k})^2 + (u_{i+1,j-1/2,k})^2 \right) \right] \right],
\end{align*}
\]

for each velocity component \( u, v, \) and \( w \) of cell \( i, j, k \). Although this system of equations is complex, the solution process is straightforward. To move the solution ahead in time, velocities and pressures from the previous iteration are taken directly from individual cells and plugged into (2) to give the new velocities for the current iteration \([\tilde{u}, \tilde{v}, \tilde{w}]\). In some cases, velocities are required that do not lie on cell faces, in which case they are averaged over the nearest available values, e.g., \( \tilde{u}_{i,j,k} = \frac{1}{3} (u_{i+1/2,j,k} + u_{i-1/2,j,k}) \), and the square of a quantity, e.g., \( u^2 \) at \((i, j, k)\), is the square of the average, \( \tilde{u}_{i,j,k} \).

The new velocities are labeled with a tilde because the direct application of (2) does not ensure that (3) is satisfied. Due to the discretization of the environment, individual cell may not explicitly satisfy the criteria that mass be conserved and that the fluid is incompressible. Also, the new pressure field needs to be determined. These constraints are satisfied simultaneously by solving the mass conservation, or continuity equation, as

\[
\frac{\partial u}{\partial x} + \frac{\partial v}{\partial y} + \frac{\partial w}{\partial z} = 0, \tag{3}
\]

which essentially says that the net fluid flow into or out of a cell is zero.

Consider a cell \( i, j, k \). The divergence of fluid within the cell, or "missing mass", is given by

\[
D_{i,j,k} = -((1/\varepsilon)(u_{i+1/2,j+1/2,k} - u_{i-1/2,j+1/2,k}) + (1/\varepsilon)(v_{i+1/2,j+1/2,k} - v_{i+1/2,j-1/2,k}) + (1/\varepsilon)(w_{i+1/2,j+1/2,k} - w_{i+1/2,j-1/2,k})), \tag{4}
\]

Notice that this is a finite difference approximation to the continuity equation. A positive \( D \) therefore, represents an influx of fluid, and in the real world would correspond to an increase in cell pressure and subsequent increase in fluid outflow from the cell. Similarly, a negative \( D \) lowers internal pressure and increases inflow from neighboring cells. Thus if the change in cell pressure is scaled according to the divergence in the cell, and the face cell velocities are adjusted according to the change in pressure, the cell can be made to satisfy (3). The change in pressure for a cell is

\[
\delta p = \beta D, \tag{5}
\]

where \( \beta \) is given by

\[
\beta = \beta_0/2(\frac{1}{\varepsilon^2} + \frac{1}{\varepsilon^2} + \frac{1}{\varepsilon^2}), \tag{6}
\]

and \( \beta_0 \) is a relaxation coefficient within the range \([1, 2]\). The cell face velocities are then updated according to \( \delta p \) such that

\[
\begin{align*}
u_{i+1,j+1/2,k} &= u_{i+1,j+1/2,k} + (\delta t/\varepsilon)\delta p, \\
u_{i-1,j+1/2,k} &= u_{i-1,j+1/2,k} - (\delta t/\varepsilon)\delta p, \\
v_{i,j+1/2,k} &= v_{i,j+1/2,k} + (\delta t/\varepsilon)\delta p, \\
v_{i,j-1/2,k} &= v_{i,j-1/2,k} - (\delta t/\varepsilon)\delta p, \\
w_{i,j+1/2,k} &= w_{i,j+1/2,k} + (\delta t/\varepsilon)\delta p, \\
w_{i,j-1/2,k} &= w_{i,j-1/2,k} - (\delta t/\varepsilon)\delta p, \\
\end{align*}
\]

and the cell pressure is updated according to

\[
\tilde{p}_{i,j,k} = \tilde{p}_{i,j,k} + \delta p. \tag{7}
\]

Use of the above equations satisfies (3) for a single cell, but neighboring cells may now have a non-zero divergence. In order for the whole mesh to simultaneously satisfy (3), the pressure and velocities are first adjusted using (5)–(7) for every cell in the grid. This procedure is then repeated until all cells in the flow field have a divergence less than some prescribed small \(\epsilon\). With a \( \beta_0 \) of 1.7 and \( \epsilon = 0.0001 \), the examples shown in this paper converged in 2–6 sweeps on average. Once convergence is achieved, the fluid is considered to be locally incompressible and the velocity and pressure fields are complete for buoyant object inclusion and the start of the next cycle.

3.2. Boundary Conditions

The boundary conditions for our model are set automatically once the contents of each cell in the mesh have been determined. They are also homogeneous. That means that once they have been set, the Navier-Stokes equations can be applied blindly without determining exactly where surfaces or obstacles lie. This makes for cheap computation.
because boundary conditions need only be checked once at the beginning of an iteration, rather than for the velocity component calculation of each cell. A boundary is an interface between the fluid and a solid obstacle, or between the fluid and atmosphere, or a point at which fluid flows into or out of the system. In all cases, generalizing assumptions about the shape of static obstacles, and the position of free surfaces can greatly reduce the amount of work that we have to do, without compromising accuracy or realism.

3.2.1. Stationary Obstacles

Consider Figure 2, which shows an obstacle and a free surface. We assume that the walls of an obstacle are always co-incident with the face of a computational cell. It then becomes a trivial process to set correct solid obstacle boundary conditions; That is, velocity and pressure for use in the finite difference expressions. For example, the component of fluid velocity normal to the face of a non-permeable obstacle is zero. Because obstacle and cell faces are coincident, the normal velocities are set directly (\(w_0 = 0\) in the figure). In the case of a non-slip obstacle which exerts a drag on the fluid, the tangential velocity at the boundary is also zero. This is set indirectly by making the tangential cell face velocity inside the boundary cell equal and opposite to that outside in the fluid (\(w_0 = -w_1\)). Finally, the pressure in the boundary cell, which is also needed for the finite difference calculation, is set equal to the pressure in the adjacent fluid cell, preventing any acceleration across the boundary.

Another useful type of obstacle is a free-slip boundary. The treatment of pressure and velocity is the same as for a non-slip boundary except that the inner tangential velocity is set equal to that outside in the fluid (\(w_0 = w_1\)). A free-slip boundary can be thought of as a plane of symmetry for motion tangential to it, thus it provides a convenient way to bound a flow field.

3.2.2. Inflow and Outflow

Fluid can easily flow into or out of the system by virtue of inflow or outflow boundary cells. For inflow, the required input velocity is set on the cell faces and held fixed throughout the calculation. In the case of an outflow boundary, velocities are initially set equal to the tentative velocity field in adjacent fluid cells and then allowed to relax without constraint during the pressure iteration step. This ensures that fluid can flow freely out of the system without causing any upstream artifacts.

3.2.3. Free Surface

Boundary conditions also need to be set on the free surface. When (2) is applied to a surface cell, velocities and pressures are needed from adjacent empty cells. We assume that for most applications, if the wavelength of any disturbance is longer than a few inches, forces due to surface tension will be negligible. We then relax the constraint that we need to know exactly where in a cell the surface lies. Thus, if any part of a free surface passes through a cell, that cell is labeled as a Surface cell, and the equation of continuity (3) is used to set boundary velocities. Consider a two dimensional surface cell which is surrounded on three sides by cells containing fluid. The velocity on the remaining surface side is set so that the divergence \(D\) of the fluid in the cell is zero. So referring to Figure 2,

\[
\begin{align*}
   u_{i,j,k} + u_{i,j,k-1} - u_{i,j,k+1} - (\delta \delta x) (w_{i+1,j,k} + w_{i-1,j,k} - w_{i,j,k}) = 0.
\end{align*}
\]

If the cell had two sides which face an empty cell, we require that \(\partial u/\partial x\) and \(\partial w/\partial z\) both vanish separately, that is that each open side velocity equals the velocity of the side of the cell opposite it. This also satisfies (3). Finally, for the case in which three sides are open, the side opposite the fluid carries the velocity of that side, while the remaining two sides follow freely the effects of the body force and do not otherwise change. A three dimensional surface cell has velocity components set in an analogous fashion, leading to 64 distinct Empty-Fluid configurations. The pressure in a surface cell is set to the applied atmospheric pressure or forcing pressure function (see Section 7.2).

4. Tracking fluid position

We have described a method for solving the full Navier-Stokes equations over a finite difference mesh. From the mesh we want to generate a smooth and accurate representation of the actual fluid surface position. We also want to track the motion of such a surface over time, so that we can adjust the contents of the mesh accordingly (i.e. Fluid, Surface, or Empty). Finally, to avoid aliasing, the resolution of the surface should not be restricted by the coarse resolution of the mesh. With these goals in mind, three methods of surface identification have been developed, each of which is useful for a particular class of liquid phenomena.

4.1. Marker Particles

The simplest and most functional way to track fluid position in 2D is to convect massless marker particles with local fluid velocity. In this way particles are continuously introduced at inflow boundaries and removed if they cross an outflow boundary, and can splash and flow freely. A particle’s new position is found using an area weighting interpolation over the four nearest cell velocities (See Figure 3) and multiplying the resultant velocity by the current timestep. The finite difference mesh is then labeled as follows:

- A cell containing no particles is Empty.
- A cell containing at least one particle that is adjacent to an Empty cell is a Surface cell.
- A cell containing at least one particle that is not a Surface cell is a Full cell.
Both of them and connect their neighbors. If two particles become too close together, delete them. Lowed to change dynamically as the position of the surface between them. This ensures that the surface always remains continuous and that colliding surfaces are smoothly connected. Two dimensions, this method is particularly useful because it is fast and can easily account for multiple surfaces.

4.2. Free Surface Particles

Marker particles can also be used to precisely delineate any free-surfaces in a scene. Instead of appearing within every cell containing fluid, a grid of markers is placed along the boundaries between fluid and obstacles or air. This grid is connected with local velocity as before. However, the number, distribution, and connectivity of particles are allowed to change dynamically as the position of the surface changes. The rules for removing and adding particles are simple. If two particles become too close together, delete both of them and connect their neighbors. If two particles become too far apart, insert a new particle on the link between them. This ensures that the surface always remains continuous and that colliding surfaces are smoothly connected. In two dimensions, this method is particularly useful because it is fast and can easily account for multiple surfaces.

4.3. Height Field

Liquid in the real world often has a surface that is single valued. Examples of this are puddles, rivers, or the ocean (as long as there are no overturning waves). For such cases the position of the surface can be calculated without using marker particles because we no longer need to track the complex geometry caused by overturning. We define the surface height along the y axis, at the center of each vertical column of cells in the three-dimensional mesh. The change in local surface elevation at each timestep is determined by the local fluid velocity, that is, by the vertical component of the fluid motion plus the horizontal convection of the surface elevation from adjacent cell columns,

\[
\frac{\partial h}{\partial t} = w - u \left( \frac{\partial h}{\partial y} \right) - v \left( \frac{\partial h}{\partial x} \right),
\]

where \( h \) is the surface height. This equation can be approximated by a finite difference expression \(^{11}\)

\[
h_{i,j}^{t+1} = \left( h_{i,j}^t + \Delta t \left( \frac{h_{i,j}^t - h_{i,j+1}^t}{4\Delta x} + \frac{h_{i,j+1}^t - h_{i,j+2}^t}{4\Delta x} + \frac{h_{i,j-1}^t - h_{i,j}^t}{4\Delta y} + \frac{h_{i,j+1}^t - h_{i,j+2}^t}{4\Delta y} \right) \right)
\]

This expression is used to update the position of the height field once the velocity and pressure fields have been calculated. It is important to note that despite superficial similarities to the method used by Kass et al. \(^5\), the height field equation is very different. Here, surface elevation is driven by the underlying fluid velocity. Therefore, velocity or pressure disturbances anywhere in the fluid volume can affect the surface (see Examples). Cell configuration for the height field approach is trivial. Cells crossed by the height field are Surface cells while those above it are Empty cells and those below it are Full.

For dramatic effects such as crashing waves or splashing, the height field can be combined with the marker particles. Whenever the vertical velocity of the surface is greater than some positive threshold, a set of particles are introduced just below the surface and the local fluid velocity is used to set their initial velocity. The particles are then removed from the Navier-Stokes calculation and affected only by gravity, wind, and air resistance. There is interesting discussion of topics related to the use of particle systems for fluid animation in \(^4, 5, 10\).

5. Buoyancy

Rigid dynamic objects can be included in a scene using the velocities and pressures calculated using the Navier-Stokes equations. Specifically, we assume that each rigid object is discretized and consists of a set of nodes \( n \). For each model surface node \( n \) which is within the fluid, the force acting on this node is calculated based on the following formula

\[
f_{ni} = - \nabla p_i dV_i + m_i g_i,
\]

where \( dV_i \) is a volume associated with the submerged node of the object and \( \nabla p_i \) is the gradient vector of the pressure. Each component of \( \nabla p_i \) is computed in discrete form as

\[
(\nabla p_i)_{x_j} = \frac{p_{n_i} - p_{n_{j+1}}}{\Delta s_{x_j}}, \quad j = 1, 2, 3,
\]

where \( p_{n_i} \) is the pressure in the cell containing \( n_i \), and \( p_{n_{j+1}} \) is the pressure in the previous cell in the \( x_j \) direction. Also, \( g \) is the gravitational acceleration, and \( m_i \) is the nodal mass assuming lumped masses. The total force on the object due to the fluid motion and gravity is given in discrete form by

\[
f_{\text{total}} = \sum_i f_{ni}.
\]

Based on the total force acting on each node, we compute the generalized external forces \( f_\delta \) (total force and torque)
acting on the object) as demonstrated in \(^9\) and we compute its motion based on the Lagrange equations of motion

\[
M \ddot{\mathbf{q}} + D \dot{\mathbf{q}} = f_\mathbf{q} + \mathbf{g}_\mathbf{q},
\]

where \(M\) and \(D\) are the object’s generalized mass and damping matrices, \(\mathbf{q}\) are the model translational and rotational degrees of freedom, and \(\mathbf{g}_\mathbf{q}\) are the generalized coriolis and centrifugal forces. The mass matrix, \(M\), is derived directly from the object in question \(^9\), and is unaffected by the fluid model. The damping matrix, \(D\), also has the same form as in \(^9\), but with the damping coefficients adjusted proportional to the relative velocity between a node, \(n_i\), and the local fluid.

In order to handle collisions of the floating objects with static obstacles, we also apply the techniques developed in \(^9\) for collision detection and collision force computation.

The floating objects that we used in the examples are small compared to the mesh size and therefore it is possible to make the simplifying assumption that they do not affect the water flow. Thus, they act like large marker particles moving and rotating according to local forces. For the objects to influence the motion of the fluid, more sophisticated techniques need to be employed.

6. Summary of the Navier-Stokes Algorithm

The complete algorithm for solving the Navier-Stokes equations and tracking the fluid surface can be summarized in the following steps;

1. Define obstacles and starting fluid configuration, and place dynamic objects.
2. Set initial pressure and velocity conditions.
3. Determine cell contents depending on the method used to track the surface.
4. Set up boundary conditions for the free surface and obstacle cells.
5. Compute \(\mathbf{\ddot{u}}, \mathbf{\ddot{v}}, \mathbf{\ddot{w}}\) for all Full cells.
6. Perform the pressure iteration for all Full cells.
7. Re-calculate boundary velocities for Surface cells.
8. Update the position of the surface and objects.
9. Go to step 3.

7. Control

An important part of the animation process is specifying how objects in a scene will move. Doing this for a fluid surface is difficult because the governing equations (1) are strongly coupled and non-linear. Large scale behavior of the system can be controlled by altering various constants such as gravity and viscosity, but it is difficult to specify a motion then solve backwards to find the correct boundary conditions to cause it. However, there are two places in our algorithm where coercion can be applied to the fluid. These can easily be exploited to yield effective methods for controlling the fluid surface.

7.1. Inflow and outflow velocities

A time dependent function can be used to determine the rate at which fluid is pumped into a scene or the rate at which it is allowed to exit, producing a variety of effects. For example, a broken dam initially generates a high input rate, but tails off exponentially as the water level drops. Also, for animating a river scene, a varying inflow and outflow rate will simulate different classes (speed, turbulence) of water flow without requiring any changes to the environment model.

7.2. Surface pressure history

Perhaps the most natural way to specify surface behavior is to model nature. As wind blows across a liquid surface, small, low pressure vortices induce a local change in surface elevation. This in turn, disturbs the airflow over the surface, changing the pressure. Gravity then provides a restoration force for the initial perturbation which results in oscillation. Over time, this process is amplified and a wind driven wave is born. A similar effect can be achieved in a shorter time by applying a forcing pressure history to the free surface during the Navier-Stokes computation. This may be constant, time dependent, or depend on the present height of the surface. For example, in two dimensions, interesting waves can be developed using the forcing function

\[
p_{\text{applied}}(z) = \frac{A + B \cos(Cz - \omega t)}{\delta t}.
\]
Figure 5: Deriving constants for an applied pressure function.

Figure 6: Starting configuration for 2D marble soup animation.

where $p_{\text{applied}}$ is the pressure within a Surface cell, $A/\delta t$ is the mean pressure, and $B$ and $C$ are constants derived from the desired wave motion. From Figure 5, if $2L$ is the wavelength of the oscillation and $D$ is the mean depth of the fluid, then,

$$B = a \sqrt{\frac{2D}{C}}, \quad (17)$$

and

$$C = \frac{\pi}{L}, \quad (18)$$

where $a$ is the wave amplitude, and $g$ is gravity. Such a function is used to set the applied pressure boundary condition on the free surface (See section 3.2.3).

8. Examples

We present a number of examples to show different aspects of the system described in this paper. Running times are given for a Silicon Graphics Crimson R4000. They do not vary linearly with the size of each problem because other factors, such as the total number of Full cells present, or the speed of the flow, make a larger contribution to the amount of CPU time required.

The first example (Figure 4(a-d)), is a two dimensional animation of a water jet splashing into a concrete tank. The water motion was calculated over a 30x40 grid of cells, and marker particles were used to delineate fluid position. Two input rates were specified; water inflow and particle inflow. The jet had a velocity of 0.8 m s$^{-1}$ and new particles were introduced at the inflow boundary at a rate of 500 particles per second. It is important to note that the only overhead associated with the marker particles is the cost of moving and displaying them. A relatively sparse distribution of particles was used in this case to clearly show that the model can account for colliding surfaces, overturning waves, and arbitrary splashing. A later frame from the animation shows that the small vortex in the tank slow down and the surface starts to settle (Figure 4(d)). This animation ran for 4500 iterations in just over sixteen minutes. The same grid size (30x40) was used again to animate a splashing drop (Figure 4(e-h)). Figures 4(e) and (f) show the starting configuration of the drop and its initial impact with the surface. The waves caused by the collision travel out to the sides of the drop (Figure 4(g)), and are reflected back to give the characteristic fluid rebound at the epicenter of the splash. Particle density was set at 25 per cell. This animation ran for 2500 iterations in twelve minutes, slightly slower than the water jet example above because the average number of Full cells per iteration was higher.

If the scene geometry is rotationally symmetric, computationally cheap two dimensional calculations can be made using linked chains of markers. Figure 8 shows two frames from an animation of a rigid marble dropping into a bowl of thick soup. The actual calculation was performed in two dimensions by setting the $z$ axis resolution to 1. The curved side of the bowl was approximated as a series of steps, and a semicircular drop of liquid was aligned along the $y$ axis (see Figure 6) to represent the marble. The drop was given an initial velocity of 0.2 m s$^{-1}$, and the viscosity of the soup was set relatively high (0.003). Two chains of particles were used to represent the free surfaces in the scene, 50 for the drop and 150 for the soup. The calculation was run for two thousand iterations at a resolution of 15x30x1 taking twelve minutes. The scene was rendered directly from the positions of the markers. Each chain was used to define the profile of a surface of revolution, which was smoothed using a series of bicubic splines. Finally, the marble and other objects were added, and the whole scene rendered using Pixar's PhotoRealistic RenderMan. The liquid surface was colored using a straightforward environment map, taking account of Fresnel's law to calculate the fraction of light reflected toward the camera, or transmitted to the bottom of the bowl.

The soup example clearly shows some of the advantages of our model. The liquid drop for the calculation is the same size as the marble object, so after impact the mean surface level has risen correctly. Also, the coupling between pressure and velocity develops as a non-linear oscillation which continues long after the wave due to the collision has subsided. Previous computer graphics fluid models would have accounted for the surface wave, but not for the accompanying pressure wave which is responsible for most of the final motion.

The first full 3D example is an animation titled Moonlight Cove (Figure 9). A 50x51x40 mesh was used to finely resolve the effect of two large ocean waves crashing into a shallow cove. Submerged rocks, and an irregular sea bottom, focus the waves into the center of the cove, causing a number of interesting features on the water surface. The wave becomes steeper as the water depth decreases, and eddies and pressure waves appear to the left of, and behind the initial obstacle (Figure 9(b)).

Setting up the scene was straightforward and proceeded in two stages. First, a voxel based editor was used to define the initial distribution of rocks and water (Figure 7). The last plane of cells opposite the cove were then designated as inflow cells, with inflow velocity defined as

$$u = u + u_{\text{inflow}} \cos \omega t, \quad (19)$$

...
where \( a \) was the desired wave amplitude and \( \omega \) the desired wave frequency. The calculation was run for \( 2/\omega \) seconds, then the inflow cells were changed to outflow and water allowed to leave the system at its natural rate. This approach resulted in two full waves while allowing the added water volume to flow back out of the scene once the waves had been reflected. The animation took two and a half hours to complete and ran for 20,000 iterations.

RenderMan was also used to render this example. Two spline meshes were used; one generated from the surface height field, and another from the distribution of boundary cells. The water surface was rendered as a glass-like object with small disturbances generated using the long crested wave model suggested in 14. Detail in the rocks was provided using a displacement map and suitable noise function on the spline surface.

The frames in Figure 10 show screen shots from an animation involving buoyant objects. Water flows into a closed container carrying soda cans along with it. When the flow is turned off, the cans gather at the far corner of the container because the walls in this example were set as non-slip so the tangential fluid velocity is zero. This simulates the effect that objects tend to gather in stagnant parts of a flow. The water motion was precomputed in thirty minutes over a \( 30 \times 10 \times 20 \) grid. The soda cans were added later using an interactive editor which takes a pre-computed velocity and pressure field, and calculates the forces on an object within the mesh. In this way, many different shapes and sizes of object can be experimented with, without having to re-do the fluid calculation.

9. Conclusions

We have presented a comprehensive method for animating fluid phenomena. A direct simulation technique is used to solve the Navier-Stokes equations in two or three dimensions yielding a range of behavior unavailable with previous computer graphics fluids models. The method does come with a computational cost, which, like other volumetric techniques, scales proportional to the fourth power of the spatial resolution. However, by careful discretization of the environment, the most expensive part of the computation can be made at a low resolution. Detail generation is then achieved by directly calculating the position of a height field representing the fluid surface, or tracking the changing connectivity of surface marker particles convected with local fluid velocity. The model allows for some novel control techniques that can be used to generate a variety of interesting effects, and is suitable as a front end to many of the more inspiring water rendering algorithms available.

References

Figure 8: A marble dropping into a bowl of thick soup. Initial collision (a). Oscillation due to coupling between pressure and velocity (b).

Figure 9: Moonlight Cove. Two ocean waves crash into a shallow cove. Pressure and velocity effects throughout the water volume manifest themselves at the surface (a, b).

Figure 10: Dynamic objects. Soda cans are carried along with the incoming water, colliding with obstacles (a), and getting caught in local eddies (b).