

# Efficient Neighbor Search for Particle-based Fluids

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## Abstract

Lagrangian particle-based animation is a popular strategy for simulating complex phenomena as fluids. Due to its inherent mesh-less nature the set of neighbor particles within a specified range must be efficiently found.

In this paper we propose *Cell Indexing* a novel approach for searching approximate neighbor particles necessary for efficient fluid simulation using SPH. Instead of storing particles into a fixed 3D grid or hash map, we encode their coordinates and index into a to key. The list of keys is then sorted using linear time radix sort. A simple traversal using *H*-mask (see subsection (4.2)) can quickly accumulate approximate neighbors without problematic cache misses of Spatial Hashing, large memory requirements of full 3D grids or  $O(n \log n)$  time complexity of kd-trees. Furthermore we can achieve sub-cell precision<sup>1</sup> by using larger *H*-masks, while having only constant factor slowdown. Using *H*-mask can substantially increase the precision of Spatial Hashing or 3D grids, however more cache misses or larger memory requirements arise.

We have demonstrated our approach within a standard SPH based fluid simulation.

**Mathematics Subject Classification 2000:** I.3.5, I.3.7

**Additional Key Words and Phrases:** Neighbor Search, Cell Indexing, H-Mask, Multi-Phase Smoothed Particle Hydrodynamics

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## 1. INTRODUCTION

Physically based animation of complex natural phenomena is a very active topic among the whole computer graphics research. Simulation methods for rigid and deformable solids are been coupled with various fluid simulation methods. For both Eulerian [Guendelman et al. 2005; Carlson et al. 2002; Losasso et al. 2006] and Lagrangian approaches [Keiser et al. 2005; Solenthaler et al. 2007] unified solid-fluid simulation techniques has been proposed. Still a number of issues related to stability, accuracy, realistic boundary conditions, performance etc. arise in any fluid simulation approach. Recent graphics hardware allows huge parallelization of many time consuming problems, thus simulation algorithms has to be adapted.

Generally fluid simulation techniques solving full 3D Navier-Stokes equations can

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be categorized as *Eulerian* and *Lagrangian*. In Eulerian approaches governing equations are evaluated on a fixed mesh (usually a 3D grid), whereas Lagrangian methods can be both *mesh-based* or *mesh-less*. In both Lagrangian strategies mesh or particles are not fixed to the domain, but are advected with the fluid. This can simplify governing equations (see section (3)) and allows virtually unlimited simulation space. Beside these advantages, it is usually complex to extract smooth boundary representation of interfaces, correctly handle interface tension in small features as bubbles and droplets, perform complex remeshing (mesh-based techniques) or alternatively efficiently find neighbor (closest) particle pairs (mesh-less techniques). Beside full Navier-Stokes simulations, several "shallow water" techniques exist, where fluid interface is simulated using wave equations. However, in combination with Navier-Stokes equation they can achieve quite realistic results.[Thürey et al. 2006; Irving et al. 2006]

In this paper we focus on Lagrangian particle-base simulation of viscous fluids using *Smoothed Particle Hydrodynamics* (SPH). (see section (3)) It is a mesh-less approach, where all physical quantities are sampled on particle locations. The influence of each particle is only local<sup>2</sup> thus it is essential to find the set of neighbor particles. Since this highly affects further calculations of fluid dynamics it becomes usually the bottleneck of the overall animation. In section (4) we have extended the well known *Spatial Hashing* for varying particles support length (see subsection (4.1)) and developed a novel neighbor search algorithm fixing several issues of previous methods.

## 2. RELATED WORK

It is beyond the scope of this article to give an extensive overview of the fluid simulation problematics, therefore we focus here only to related Eulerian and mesh-less Lagrangian works.

### 2.1 Eulerian Grid-based Methods

Since introducing (by Foster and Metaxas) the simulation of full 3D Navier-Stokes equations to graphics community, an active research started to improve their famous Eulerian-based *MAC-grid* method[Foster and Metaxas 1996].<sup>3</sup> Stam proposed the popular extension of the basic MAC by using a semi-Lagrangian integration scheme and iterative solver of the pressure equation.[Stam 1999] Fedkiw et al. developed a *Particle Level Set* method for accurate interface tracking.[Enright et al. 2002; Enright et al. 2005]

Advanced simulation of melting and flowing of highly viscous, non-Newtonian fluids (e.g. wax) were presented in [Carlson et al. 2002]. Later Carlson et al. used distributed Lagrangian multipliers [Carlson et al. 2004] for animating the interplay between rigid bodies and viscous incompressible fluids. Direct two way coupling between mesh-based thin solids and Eulerian-based fluids was done in [Guendelman et al. 2005; Losasso et al. 2006].

Hong and Kim [Hong and Kim 2003] use a *Volume Of Fluid* (VOF) indicator function to simulate a two-phase fluid flow and bubbles with surface tension forces.

<sup>2</sup>Within a support distance from particles location

<sup>3</sup>Nevertheless a full 3D fluid simulation in CFD engineering has been already well established

Losasso et al. extended the particle Level Set Method interface tracking of multiple interacting fluids [Losasso et al. 2006] and presented a cutting-edge fluid simulator. A generalized solution to modeling hydraulic erosion is presented by Beneš et al. in [Benes et al. 2006]. Recently Chentanez [Chentanez 2007] developed a method for animating incompressible liquids with detailed free surfaces using Lattice-Based Tetrahedral Mesh.

Octrees has been used to speed-up grid-based fluid simulations [Losasso et al. 2004]. Further combining 2D height-fields with a full 3D Navier-Stokes simulation near the interface allowed efficient animation of large bodies of water.[Irving et al. 2006]. Recently controlling fluid motion while preserving details was successfully applied to both grid-based *Lattice-Boltzmann Method* (LBM) and Lagrangian *Smoothed Particle Hydrodynamics* (SPH).[Thürey et al. 2006]

## 2.2 Lagrangian Particle-based Methods

Particle-based simulations were introduced to graphics community by Reeves. These animations of simple fountains and sprays were represented by uncoupled particles, which was insufficient for realistic fluid simulation.

Later Desbrun et al. [Desbrun and Cani 1996] started using Smoothed Particle Hydrodynamics<sup>4</sup> (SPH) in computer graphics to couple particles and simulate animating highly deformable bodies including viscous fluids. Müller et al. further popularize SPH technique by animating a pouring water into a glass at interactive rates [Müller et al. 2003]. By averaging viscosity, they extended the model to multi-phase fluids. [Müller et al. 2005].

Since the pressure force is computed locally using explicit integration, their model is suitable only for compressible and near incompressible fluids. For stiff fluids incompressibility is achieved using very small time steps.<sup>5</sup> Premoze et al. gained incompressibility by solving the pressure globally (using Poisson equation) with an iterative linear equation solver. [Premoze et al. 2003] This technique is referred as *Moving Particle Semi-implicit* (MPS).

Clavet et al. describe a SPH based interactive technique for simulating visco-elastic and plastic fluids by connecting particles with temporary springs [Clavet et al. 2005]. Unified particle-based SPH model for coupling solids and fluids has been presented in [Keiser et al. 2005] and later in [Solenthaler et al. 2007]. Recently a successful approach of coupling SPH and Particle Level Set has been done by Losasso.[Losasso 2007]. Promising results are shown by Harada when moving the entire SPH calculation to GPU and precalculate the influence of solid boundary particles.<sup>6</sup> Their simulating of 60k particles run at 17 frames.[Harada et al. 2007a; 2007b].

Further simplifications of SPH has been proposed by Adams et. al [Adams et al. 2007] by adaptively sampling particles and allowing them to have various sizes. For sparse particles Kipfer et al. developed novel technique for computing neighbor particles [Kipfer and Westermann 2006].

<sup>4</sup>originally developed by Lucy and Monaghan

<sup>5</sup>Mainly due to stability issues

<sup>6</sup>Commonly solid-fluid boundary conditions are solved by fixing particles to the solid volume near the surface. This increase the number of particles and lower down performance

### 3. MULTI-PHASE SMOOTHED PARTICLE HYDRODYNAMICS

Inspired by Monte-Carlo integration *Smoothed Particle Hydrodynamics* (SPH) was initially developed by Lucy and Monaghan for simulating flow of interstellar gas within Astrophysics [Monaghan 2005]. Unlike traditional mesh-based methods (FEM, FDM) where physical quantities are evaluated on a fixed Mesh (grid), SPH belongs to mesh-less methods, where calculated values are approximated from neighboring points (particles). Particles have no fixed topology, thus are suitable for complex dynamic phenomena as fluids. Due to the particle-based Lagrangian nature of SPH mass conservation is trivially satisfied and convection of the substance is inherent.

#### 3.1 Smoothing with Particles

Given a set of particles (interpolation points)  $\mathbf{r}_i$  carrying values  $A_i = A(\mathbf{r}_i)$  of field quantity  $A$  we can approximate  $A(\mathbf{r})$  at arbitrary point  $\mathbf{r}$  by a convolution with a radial symmetric *smoothing kernel*  $W(\mathbf{r}, h)$  as [Monaghan 2005; Müller et al. 2005]

$$A(\mathbf{r}) \approx A \star W = \int_{\mathbf{r}'} A(\mathbf{r}') W(|\mathbf{r} - \mathbf{r}'|, h) d\mathbf{r}' \approx \sum_j V_j A_j W(\mathbf{r} - \mathbf{r}_j, h) \quad (1)$$

and further replaced by summation over neighbor particles having finite *volume*  $V_j = m_j/\rho_j$ . Mass of particles is a constant property, thus volume is defined as the ration between *mass*  $m_j$  and *density*  $\rho_j$ . The convolution smoothing kernel must have the following properties

$$\int_{\mathbf{r}} W(\mathbf{r}, h) d\mathbf{r} = 1 \quad \wedge \quad \lim_{h \rightarrow 0} W(\mathbf{r}, h) = \delta(\mathbf{r}) \quad (2)$$

where  $h$  is the kernels *support length*, i.e. the distance to which particle affects other particles. Notice in limit case  $h \leftarrow 0$  kernel  $W(\mathbf{r}, h)$  must be the Dirac delta function  $\delta(\mathbf{r})$ . Making the kernel second order differentiable we can further express gradient  $\nabla A(\mathbf{r})$  and laplacian  $\nabla^2 A(\mathbf{r})$  of field function  $A(\mathbf{r})$  at arbitrary point as

$$\begin{aligned} A(\mathbf{r}) &= \sum_j V_j A_j W(\mathbf{r} - \mathbf{r}_j, h) \\ \nabla A(\mathbf{r}) &= \sum_j V_j A_j \nabla W(\mathbf{r} - \mathbf{r}_j, h) \\ \nabla^2 A(\mathbf{r}) &= \sum_j V_j A_j \nabla^2 W(\mathbf{r} - \mathbf{r}_j, h) \end{aligned} \quad (3)$$

This properties greatly simplify further calculations of Lagrangian-based fluid dynamics.

#### 3.2 Evaluating Fluid Properties using SPH

The well known *Navier-Stokes* governing equations of a simple Newtonian isothermal and incompressible fluid can be expressed in Eulerian form by two conservation laws, i.e. the conservation of mass (*continuity equation*) and the conservation of momentum (*momentum equation*)

$$\begin{aligned} \frac{\partial \rho}{\partial t} &= -\rho \nabla \cdot \mathbf{v} & (\text{continuity eq.}) \\ \rho \left( \frac{\partial \mathbf{v}}{\partial t} + \mathbf{v} \cdot \nabla \mathbf{v} \right) &= -\nabla p + \mu \nabla^2 \mathbf{v} + \rho \mathbf{g} & (\text{momentum eq.}) \end{aligned} \quad (4)$$

Lagrangian (particle based) formulation can be obtained using the material derivative  $\frac{D\mathbf{q}}{Dt} = \frac{\partial\mathbf{q}}{\partial t} + \mathbf{v} \cdot \nabla\mathbf{q}$ .<sup>7</sup> as

$$\begin{aligned}\frac{D\rho}{Dt} &= -\rho\nabla \cdot \mathbf{v} \\ \rho\frac{D\mathbf{v}}{Dt} &= -\nabla P + \mu\nabla^2\mathbf{v} + \rho\mathbf{g} \quad (= \mathbf{F}^{\text{press}} + \mathbf{F}^{\text{visco}} + \mathbf{F}^{\text{ext}})\end{aligned}\quad (5)$$

Assuming constant mass of all particles total mass of fluid is always conserved. we can thus completely omit continuity equation from further calculations. Viewing the momentum equation as Newton's second law ( $\mathbf{f} = m\mathbf{a}$ ), i.e. we only need to calculate forces acting on particles. Notice the right hand side of the momentum equation can be thus expressed as a sum of *pressure*  $\mathbf{F}^{\text{press}}$ , *viscosity*  $\mathbf{F}^{\text{visco}}$  and *external*  $\mathbf{F}^{\text{ext}}$  force fields. This external force field further contains all other force fields acting on fluid (e.g. interface tension  $\mathbf{F}^{\text{int}}$  or gravity  $\mathbf{F}^{\text{grav}}$ )

Assuming a finite volume of particles, we get the relation between total force  $\mathbf{f}_i$  acting on  $i$ -th particle and respective force field. Simply by integrating the force field over the particles volume, setting  $\mathbf{F}_i = \mathbf{F}(\mathbf{r}_i)$  and using SPH approximation (equation (3)) we get

$$\mathbf{f}_i = \int_{\mathbf{r}} \mathbf{F}(\mathbf{r}) d\mathbf{r} \approx \int_{\mathbf{r}} V_i \mathbf{F}_i W(\mathbf{r} - \mathbf{r}_j, h) = V_i \mathbf{F}_i \int_{\mathbf{r}} W(\mathbf{r} - \mathbf{r}_i, h) = \mathbf{F}_i = V_i \mathbf{F}_i \quad (6)$$

*Fluid Properties.* Assuming  $\rho_j = m_j/V_j$  density can be approximated using SPH as well. Further pressure  $P_i$  occurred at  $i$ -th particle is described by Tait's equation [Becker and Teschner 2007]<sup>8</sup>

$$\rho(\mathbf{r}_i) = \sum_j m_j W(\mathbf{r}_i - \mathbf{r}_j, h) \quad P_i = k^{\text{gas}} \left( \left( \frac{\rho}{\rho_0} \right)^\gamma - 1 \right) \quad (7)$$

Using blindly the SPH approximation for pressure force  $\mathbf{f}_i^{\text{press}}$  and multiphase viscosity force  $\mathbf{f}_i^{\text{visco}}$  violates the action-reaction principle, since the forces are not symmetric ( $\mathbf{f}_i \neq -\mathbf{f}_j$ ). Various symmetrization approaches exists among the SPH literature (see [Ritchie and Thomas 2001; Monaghan 2005]), however we adapt here to simplified formulation by Müller [Müller et al. 2005],<sup>9</sup> where the pressure and viscosity is only averaged.<sup>10</sup> Notice particle volume  $V_i = m_i/\rho_i$  is using approximate density  $\rho_i$  computed in equation (7).

$$\mathbf{f}_i^{\text{press}} = V_i \mathbf{F}_i^{\text{press}} = - \sum_j V_i V_j \frac{P_i + P_j}{2} \nabla W^{\text{press}}(\mathbf{r}_i - \mathbf{r}_j, h) \quad (a)$$

$$\mathbf{f}_i^{\text{visco}} = V_i \mathbf{F}_i^{\text{visco}} = \sum_j V_i V_j \frac{\mu_i + \mu_j}{2} (v_j - v_i) \nabla^2 W^{\text{visco}}(\mathbf{r}_i - \mathbf{r}_j, h) \quad (b) \quad (8)$$

$$C_i^{\text{int}} = \sum_j V_j c_j^{\text{int}} \nabla^2 W^{\text{poly}}(\mathbf{r}_i - \mathbf{r}_j, h) \quad (c)$$

<sup>7</sup>Since particles move with the fluid the convective term  $\mathbf{v} \cdot \nabla\mathbf{q}$  is not present

<sup>8</sup>An simpler alternative is the ideal gas equation  $P_i = k(\rho - \rho_0)$

<sup>9</sup>See [Desbrun and Cani 1996] for alternative formulations

<sup>10</sup>This gives more stable simulation, with the cost of small energy dissipation

Interface tension force  $\mathbf{f}_i^{\text{int}}$  acts along the normalized gradient  $\mathbf{n} = \frac{\nabla C_i^{\text{int}}}{|\nabla C_i^{\text{int}}|}$  of the interface color field  $C_i^{\text{int}}(\mathbf{r})$  and is proportional to its curvature  $\kappa_i = \nabla^2 C_i^{\text{int}}$ . Interface color field values  $C_i^{\text{int}}$  at particle locations are approximated from particles interface color values  $c_i^{\text{int}}$  using SPH (see equation (8) (c)). Interface tension force is thus defined as

$$\mathbf{f}_i^{\text{int}} = -\sigma^{\text{int}} \kappa_i \mathbf{n}_i = -\sigma^{\text{int}} \nabla^2 C_i^{\text{int}} \frac{\nabla C_i^{\text{int}}}{|\nabla C_i^{\text{int}}|} \quad (9)$$

*Smoothing Kernels.* In our computations we use the following smoothing kernels adapted from [Müller et al. 2003] and its derivatives (see appendix (6)).

$$\begin{aligned} W^{\text{poly}}(\mathbf{r}, h) &= \frac{315}{64\pi h^9} \begin{cases} (h^2 - r^2)^3 & 0 \leq r \leq h \quad \wedge \quad r = |\mathbf{r}| \\ 0 & \text{otherwise} \end{cases} \\ W^{\text{press}}(\mathbf{r}, h) &= \frac{15}{\pi h^6} \begin{cases} (h - r)^3 & 0 \leq r \leq h \quad \wedge \quad r = |\mathbf{r}| \\ 0 & \text{otherwise} \end{cases} \\ W^{\text{visco}}(\mathbf{r}, h) &= \frac{15}{2\pi h^6} \begin{cases} -\frac{r^3}{2h^3} + \frac{r^2}{h^2} + \frac{h}{2r} - 1 & 0 \leq r \leq h \quad \wedge \quad r = |\mathbf{r}| \\ 0 & \text{otherwise} \end{cases} \end{aligned} \quad (10)$$

### 3.3 Simulation Overview

Our simulation loop contains three main steps, namely the *Neighbor Search*, *Force Computations* and the *Time Integration*. We use explicit time integration, thus the overall simulation can be categorized as explicit and force-based. (see algorithm (1)).

As shown in the algorithm searching for neighbor particles (i.e. building the list of close particle pairs) is done in `SEARCHNEIGHBORS(h, H)`, where  $h$  is support length and  $H$  is the subdivision factor (see section (4)). Next the force computation is done by iterating over all particles (lines 2-15). First density  $\rho_i$  is accumulated from neighbors (lines 4-6), then pressure  $P_i$  is calculated (line 7) and finally total force acting on particle is accumulated (lines 8-14). Time integration is performed using standard second order "Leap-frog" scheme (lines 16-19). This looks similar to the simple first-order explicit Euler scheme, however when the velocity is initialized<sup>11</sup> correctly as  $\mathbf{v}_i = \mathbf{v}_i(t_0 - \frac{1}{2}\Delta t)$  the code perfectly mimic the following "Leap-frog" update rules (see equation (11))

$$\begin{aligned} \mathbf{v}_i(t + \frac{1}{2}\Delta t) &\leftarrow \mathbf{v}_i(t - \frac{1}{2}\Delta t) + \Delta t \frac{\mathbf{f}_i(t)}{m_i} \\ \mathbf{r}_i(t + \Delta t) &\leftarrow \mathbf{r}_i(t - \Delta t) + \Delta t \mathbf{v}_i(t + \frac{1}{2}\Delta t) \end{aligned} \quad (11)$$

Due to the explicit nature of our method, only small time steps are allowed to maintain stability and accuracy. Formally,  $i$ -th particle can be integrated with a maximal time step  $\Delta t_i$  according to the *Courant condition* of convergence<sup>12</sup>

$$\Delta t_i \leq \Delta t^c = \alpha \frac{h}{c} \quad \Delta t_i = \min \left( \Delta t^c, \alpha \frac{h}{|\mathbf{v}_i|} \right) \quad (12)$$

<sup>11</sup>The initialization routine trivial and is omitted here

<sup>12</sup>Combating numerical errors time step should be even smaller.

```

In: support length  $h$ , subdivision factor  $H$  and delta time  $\Delta t$ 

function SPH( $h, H, \Delta t$ )
1:  NEIGHBOURS  $\leftarrow$  SEARCHNEIGHBORS( $h, H$ )
2:  foreach  $\mathcal{P}_i$  in PARTICLES do
3:     $\rho_i \leftarrow 0$ ;  $\nabla C_i \leftarrow \mathbf{0}$ ;  $\nabla^2 C_i \leftarrow 0$ ;  $\mathbf{f}_i \leftarrow \mathbf{f}_i^{\text{ext}}$  /* initialize */
4:    foreach  $\mathcal{P}_j$  in NEIGHBOURS( $\mathcal{P}_i$ ) do /* accumulate density */
5:       $\rho_i \leftarrow \rho_i + m_j W^{\text{poly}}(\mathbf{r}_i - \mathbf{r}_j, h)$ 
6:    end
7:     $P_i \leftarrow k^{\text{gas}} \left( \left( \frac{\rho}{\rho_0} \right)^\gamma - 1 \right)$  /* calculate pressure */
8:    foreach  $\mathcal{P}_j$  in NEIGHBOURS( $\mathcal{P}_i$ ) do /* accumulate forces */
9:       $\mathbf{f}_i \leftarrow \mathbf{f}_i - V_i V_j \frac{P_i + P_j}{2} \nabla W^{\text{press}}(\mathbf{r}_i - \mathbf{r}_j, h)$  /* ( $= \mathbf{f}_i^{\text{press}}$ ) */
10:      $\mathbf{f}_i \leftarrow \mathbf{f}_i + V_i V_j \frac{\mu_i + \mu_j}{2} (v_j - v_i) \nabla^2 W^{\text{visco}}(\mathbf{r}_i - \mathbf{r}_j, h)$  /* ( $= \mathbf{f}_i^{\text{visco}}$ ) */
11:      $\nabla C_i \leftarrow \nabla C_i + V_j c_j^{\text{int}} \nabla W^{\text{poly}}(\mathbf{r}_i - \mathbf{r}_j, h)$  /* ( $= \nabla C_i^{\text{int}}$ ) */
12:      $\nabla^2 C_i \leftarrow \nabla^2 C_i + V_j c_j^{\text{int}} \nabla^2 W^{\text{poly}}(\mathbf{r}_i - \mathbf{r}_j, h)$  /* ( $= \nabla^2 C_i^{\text{int}}$ ) */
13:   end
14:    $\mathbf{f}_i \leftarrow \mathbf{f}_i - \sigma^{\text{int}} \nabla^2 C_i^{\text{int}} \frac{\nabla C_i^{\text{int}}}{|\nabla C_i^{\text{int}}|}$  /* ( $= \mathbf{f}_i^{\text{int}}$ ) */
15: end
16: foreach  $\mathcal{P}_i$  in PARTICLES do /* Leap-Frog */
17:    $\mathbf{v}_i \leftarrow \mathbf{v}_i + \Delta t \frac{\mathbf{f}_i}{m_i}$ 
18:    $\mathbf{r}_i \leftarrow \mathbf{r}_i + \Delta t \mathbf{v}_i$ 
19: end
end

```

**Algorithm 1:** Our fluid simulation step using SPH

where  $c^{13}$  is speed of sound in the fluid and  $\alpha \approx 0.3$  is the Courant number. Stability (for fast moving particles) can be further increased by choosing  $\Delta t_i$  as in equation (12).

Choosing the global time step as  $\Delta t = \min_i \{\Delta t_i\}$  is safe, but obviously inefficient. Since usually only a few particles will need this minimal time step, we can rather integrate each particle using its own time step  $\Delta t_i$ . Similarly as Desbrun [Desbrun and Cani 1996] to synchronize integration of particles we choose a user defined simulation frame time  $\Delta t^{\text{max}}$  and find smallest positive number  $n_i$  satisfying  $\Delta t'_i = \Delta t^{\text{max}} / 2^{n_i} \leq \Delta t_i$  and set particles time step to  $\Delta t'_i$ . When the time step changes, we must further correct the position of the particle as

$$\mathbf{r}_i \leftarrow \mathbf{r}_i + \frac{(\Delta t'_{i,\text{new}})^2 - (\Delta t'_{i,\text{old}})^2}{8m_i} \mathbf{f}_i \quad (13)$$

<sup>13</sup>Material property, i.e. user defined constant

*Boundary Conditions.* In SPH context solid-fluid boundary conditions are usually modeled by simply attaching *ghost* particles<sup>14</sup> near the boundary inside the solid object. Since its complex deformations this is a natural choice for deformable and melting solids. However for pure rigid objects one can precompute the influence of ghost particles (with respect to bodies frame) to a distance field and further speed up simulation omitting from them further calculations. [Harada et al. 2007a].

*Interface Extraction.* Realistic looking visualizations can be achieved by extracting a smooth interface representation. Beside using Marching Cubes to extract the zero-level of the implicit density function  $\rho(\mathbf{r}) = 0$  several attempts has been done. We refer interested reader to [Adams et al. 2007] for further details.

#### 4. NEIGHBOR SEARCH

Similarly to n-body problem, searching for close particle pairs is a crucial problem in almost any particle-based fluid simulation. Since this usually becomes the bottleneck of the simulation time, efficient algorithms are necessary. In the SPH context each particle affects only a set of neighbor particles which lie within its support distance  $h$ . Formally we define for each particle  $p_i$  its *set of neighbor particle indices*  $N_i(h)$  as

$$N_i(h) = \{ j \mid |\mathbf{r}_i - \mathbf{r}_j| \leq h \} \quad (14)$$

Beside the naive and inefficient  $O(n^2)$  "all-pair-test", several algorithms usually based on spatial subdivision and fast approximate neighbor search has been proposed [Teschner et al. 2003; Kipfer and Westermann 2006; Keiser 2006; Cohen et al. 1995]. Here we extend the usual *Spatial Hashing* (subsection (4.1)) technique for faster SPH simulation and propose a novel approach *Cell Indexing* (subsection (4.2)) based on indexing non-empty cells in a virtual subdivision grid.

##### 4.1 Extended Spatial Hashing

A common approach to optimize the "all-pair-test" is to subdivide the smallest enclosing axis aligned bounding box (AABB) into a 3D grid of cells with size  $h$ . For each particle with position  $\mathbf{r}_i = (x, y, z)$  we use its relative coordinates to AABBs *minimal corner*  $\mathbf{c}_{\min} = (x_{\min}, y_{\min}, z_{\min})$  and calculate its (positive) cell coordinates  $\text{cell}(x, y, z, h) = (i, j, k)$  (see (15)). Into each cell  $(i, j, k)$  we could store indices of particles with the same cell coordinates. Depending on the AABB and cell size ratio, such voxelization obviously lead to huge memory consumption.<sup>15</sup>

Teschner et. al [Teschner et al. 2003] overcome this problem by introducing *spatial hash function*  $\text{hash}(i, j, k)$  which naturally maps particles cell coordinates to a bucket in a hash map. This enables grid size to be virtually unlimited and does not store empty cells. The hash function is defined as

$$\begin{aligned} \text{cell}(x, y, z, h) &= \left( \left\lfloor \frac{x - x_{\min}}{h} \right\rfloor, \left\lfloor \frac{y - y_{\min}}{h} \right\rfloor, \left\lfloor \frac{z - z_{\min}}{h} \right\rfloor \right) = (i, j, k) \\ \text{hash}(i, j, k) &= (i \cdot p_1 \text{ xor } j \cdot p_2 \text{ xor } k \cdot p_3) \bmod M \end{aligned} \quad (15)$$

<sup>14</sup>For static objects their positions are not integrated

<sup>15</sup>storing empty cells is not optimal



where  $p_1 = 73856093$ ,  $p_2 = 19349663$  and  $p_3 = 83492791$  are large prime numbers and  $M$  is the size of the hash map. Using cell and hash functions insertion of a particle is  $O(1)$  thus building the data structure for approximate neighbor search is linear. To find neighbors  $N_i(h)$  for  $i$ -th particle we need examine only particles from all (26 in 3D) surrounding cells. Assuming the particle distribution is approximately even, each cell contains only a constant number of particles and thus overall neighbor search is near linear.

However in the SPH framework particles can cluster<sup>16</sup>. This leads usually to larger bucket sizes<sup>17</sup> or slower hashing. To reduce this problem we can naturally hash particles into smaller cells but then we need to examine particles from more neighboring cells.

Formally let  $h' = h/(2H + 1)$  be the subdivided cell size and  $H$  the *subdivision factor*. When calculating approximate neighbors of particle in cell  $(i, j, k)$  we need to examine particles from all  $(i + s, j + t, k + u)$  cells, where  $(s, t, u) \in \mathcal{M}_H(h)$ . The set  $H$ -mask  $\mathcal{M}_H(h)$  contains relative coordinates of all neighbor cells which are not completely outside the support of processed cell. (see figure (1) (d))

$$\mathcal{M}_H(h) = \{(s, t, u) \mid |(h's, h't, h'u)| < h \quad \wedge \quad s, t, u \in \{-H, \dots, +H\}\} \quad (16)$$

Finally notice, that larger masks<sup>18</sup> completely contains smaller masks, i.e.  $G < H \Rightarrow \mathcal{M}_G(h) \subset \mathcal{M}_H(h)$ . This allows simulations where particles have different support length. Indeed, assuming cell size is  $h'$  and particles support is  $h$  we select  $H$ -mask for this particle according to  $H = \lfloor h/h' \rfloor + 1$ <sup>19</sup>.

Hashing naturally introduce cache misses, thus examining all  $\mathcal{M}_H(h)$  can even slow down the process<sup>20</sup>. With larger  $H$  size of hash map ( $M$ ) must increase to avoid numerous collisions. It can be estimated experimentally.

## 4.2 Cell Indexing

Inspired by the *staggered grids* [Kipfer and Westermann 2006] we have developed a novel approach for searching approximate neighbor particles in a linear time, trying to avoid several disadvantages of the spatial hashing. Similarly to other grid methods we first calculate the AABB of all particles and use further only their relative coordinates to AABBs minimal corner  $\mathbf{c}_{\min}$ . For each particle we define it's  $\text{key}(n, i, j, k)$  as

$$\text{key}(n, i, j, k) = n + 2^I i + 2^{I+J} j + 2^{I+J+K} k \quad \text{where} \quad \begin{aligned} 2^I &> N \\ 2^J &> \lfloor B_x/h \rfloor \\ 2^K &> \lfloor B_y/h \rfloor \\ 2^L &> \lfloor B_z/h \rfloor \end{aligned} \quad (17)$$

where  $n$  is index of particle,  $N$  is the number of particles,  $B_x$ ,  $B_y$  and  $B_z$  are dimensions of bounding box,  $(i, j, k) = \text{cell}(x, y, z)$  are cell coordinates and  $I, J, K$  and are arbitrary constants<sup>21</sup> depending on the size of the AABB and cell size  $h$ .

<sup>16</sup>simulating compressible (near incompressible) fluids with strong pressures

<sup>17</sup>We assume all buckets have fixed size as speed-up

<sup>18</sup>with larger subdivision factor  $H$

<sup>19</sup>Naturally this ration must be reasonable small

<sup>20</sup>Storing all indices of  $H$ -neighbor cells leads to more memory usage and slowdown

<sup>21</sup>usually 16 or 32

Notice, that function  $\text{key}(n, i, j, k)$  encode its parameters to a unique key. Thus given a key  $q$  we can compute unique  $(n, i, j, k)$ .<sup>22</sup>

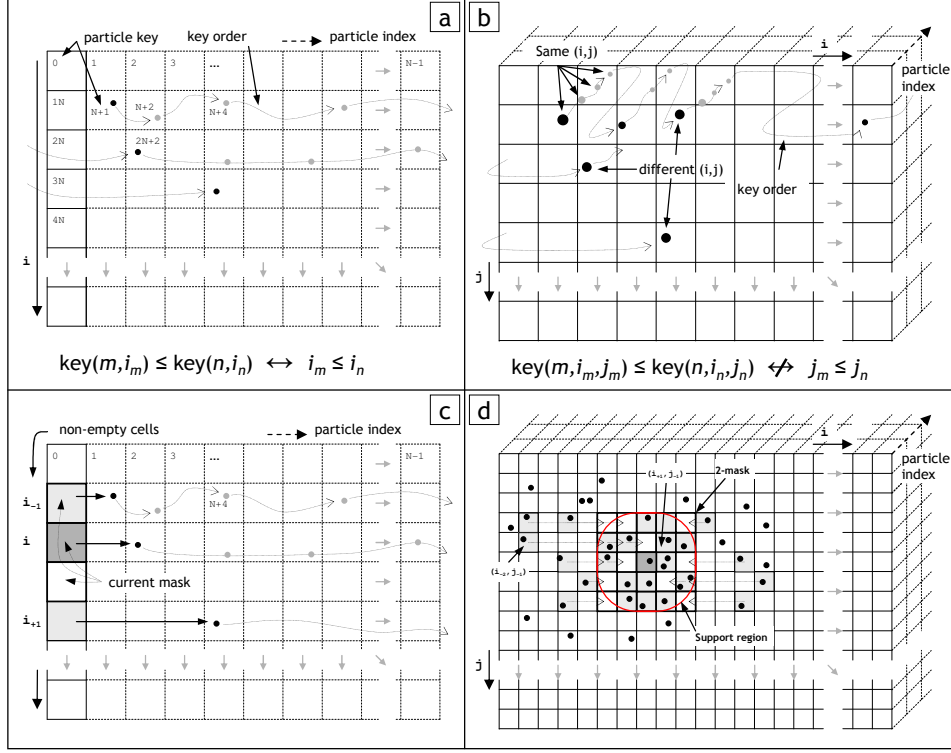


Fig. 1. Cell Indexing principles. In (a) 1D case is shown. The left most column represents 1D space divided into cells. Dashed columns symbolize indices of particles. Each row thus contain all particles with equal  $i$ -coordinate. Dotted lines show the ordering of sorted keys. In (b) 2D extension is illustrated. Notice the third dimension depicts particle index, thus has no geometrical meaning. In (c) 1D searching principle is shown. The search follows key ordering, thus (in 1D) the  $i$ -coordinate as well. Simple 1-mask (cells with thick outline) is show and respective non-empty cells  $i_{-1}, i, i_{+1}$  (light-gray cells). In (d)  $H$ -mask is extended to 2D showing its non-empty cells and their corresponding location in the mask. Since the presented 2-mask is (in 2D) equal to the full  $5 \times 5$  mask, the benefits of  $H$ -mask is not obvious from the picture, however for larger  $H$  (or in 3D) one can clearly see, that some "corner" cells of the full  $(2H+1) \times (2H+1)$  square mask will be omitted in respective "spherical"  $H$ -mask.

*1D Case.* Suppose a 1D case show in figure (1) (a), where the keys reduce to  $\text{key}(n, i) = n + 2^I i$ . Given two arbitrary particles  $m$  and  $n$  with keys  $\text{key}(m, i_m) \leq \text{key}(n, i_n)$  their cell coordinates also satisfy  $i_m \leq i_n$  and vice-versa. After sorting

<sup>22</sup>Particle index is  $n = q \bmod 2^I$ , cell coordinates are  $i = (q/2^I) \bmod 2^J, \dots$

all keys, we can thus efficiently access particles with increasing cell coordinates. Given a key  $q$  index of associated particle is  $m = q \bmod 2^I$ .

When building the neighbor set  $N(h)$  for particle  $p$  which lies in cell  $i$  we need to examine only particles in cells<sup>23</sup>  $i-1, i, i+1$ . This can be simply achieved by storing indices  $\mathcal{K}(i_{-1})$ ,  $\mathcal{K}(i)$  and  $\mathcal{K}(i_{+1})$ <sup>24</sup> of first<sup>25</sup> keys in *non-empty* cells  $i_{-1} < i < i_{+1}$ . We examine all keys (associated particles) in cell  $i_{-1}$  only when  $i_{-1} = i-1$  starting with key at index  $\mathcal{K}(i_{-1})$ . Similarly we examine cell  $i_{+1}$  only when  $i_{+1} = i+1$ . We are processing particles in  $i$ -th cell until the respective key maps coordinates of this cell, i.e. for each key  $q$  must hold  $\lfloor q/2^I \rfloor = \lfloor q_{\mathcal{K}(i)}/2^I \rfloor$ .

Assuming the maximum number of particles inside one cell is bound, this traversal is  $O(n)$ . However when simulating particles with SPH this upper bound is *not* guaranteed, but fortunately in average case there are only few particles in one cell. To overcome particle clustering problem<sup>26</sup> we can choose the cell size  $h' = h/(2H+1)$ , where  $H$  is a *subdivision factor* depending on the fluid properties. When searching for neighbors we need now to examine cells  $i_{-H} < i_{-H+1} < \dots < i \dots < i_{+H-1} < i_{+H}$  and thus store  $2H+1$  key indices  $\mathcal{K}(i_{-H}), \dots, \mathcal{K}(i), \dots, \mathcal{K}(i_{+H})$ , what slows down the traversal by a constant factor  $O(H)$  per iteration. The algorithm thus stays linear  $O(HN)$ .

*2D and 3D Case.* The extension to 2D and 3D is not obvious (see figure (1) (b)). In 2D when  $\text{key}(m, i_m, j_m) \leq \text{key}(n, i_n, j_n)$  then  $j_m \leq j_n$  while the order of  $i_m$  and  $i_n$  is non-decreasing only if  $j_m = j_n$  (j-coordinates are equal). When calculating neighbors for particle in cell  $(i, j)$  we need to store all 9 key indices  $\mathcal{K}(i_s, j_t) \mid s, t \in -1, 0, +1$  and examine non-empty cell  $(i_s, j_t)$  only when  $(i_s, j_t) = (i + s, j + t)$ . Furthermore in 3D we need 27 indices  $\mathcal{K}(i_s, j_t, k_u) \mid s, t, u \in -1, 0, +1$  and examine non-empty cell  $(i_s, j_t, k_u)$  only when  $(i_s, j_t, k_u) = (i + s, j + t, k + u)$ . We could generalize this approach for cell subdivision taking  $s, t, u \in -H, \dots, +H$  or even better using  $H$ -mask where  $(s, t, u) \in \mathcal{M}_H(h)$ . The traversal is therefore in average case linear with complexity  $O(|\mathcal{M}_H(h)|N)$ .

*Sorting Keys.* To make the overall neighbor search algorithm linear, we must sort keys in linear time. As pointed by Terdiman [Terdiman 2000] even a set of floating point numbers can be quickly sorted in  $O(n)$  using radix sort. Since we have encoded indices of particles into their keys<sup>27</sup> we can use radix sort and have correct indices after sorting.

## 5. IMPLEMENTATION AND RESULTS

As shown in figure (2) three different simulations of the classical Dam-break test has been performed. Increasing the stiffness and surface tension coefficient, fluid behaves less compressible thus more realistic. In all three scenarios we used 1600 particles and set the rest density  $\rho_o = 1000 \text{ kg/m}^3$ , mass  $m_i = 0.0012 \text{ kg}$ , support length  $h = 0.05m$  viscosity  $\mu_i = 50 \text{ N s/m}^2$  and used a fixed time stepping with

<sup>23</sup> Assuming size of cell is  $h$

<sup>24</sup>  $\mathcal{K}(i)$  is index of key in list of all keys

<sup>25</sup> key in cell  $i$  with the smallest particle index

<sup>26</sup> Too many particles in one cell

<sup>27</sup> Keys are usually 64-bit numbers, 16 bits for each  $(n, i, j, k)$  component

delta time  $\Delta t = 0.002s$ . The surface tension coefficient  $\sigma$  varied from 0.6 (top), 1.4 (mid) to 2.2 (bottom). We set stiffness  $k^{\text{gas}}$  to  $20Nm/g$  (top),  $40Nm/g$  (mid) and  $70Nm/g$  (bottom). Boundary conditions are handled by inelastic particle-plane collision resolution with wet friction. Simple particle-to-plane projection is used to prevent overshooting. State variables are integrated using explicit Leap-Frog integration scheme with fixed time stepping. The simulation has been performed on Mobile P4 1.7 GHz with GeForce 4 Go.

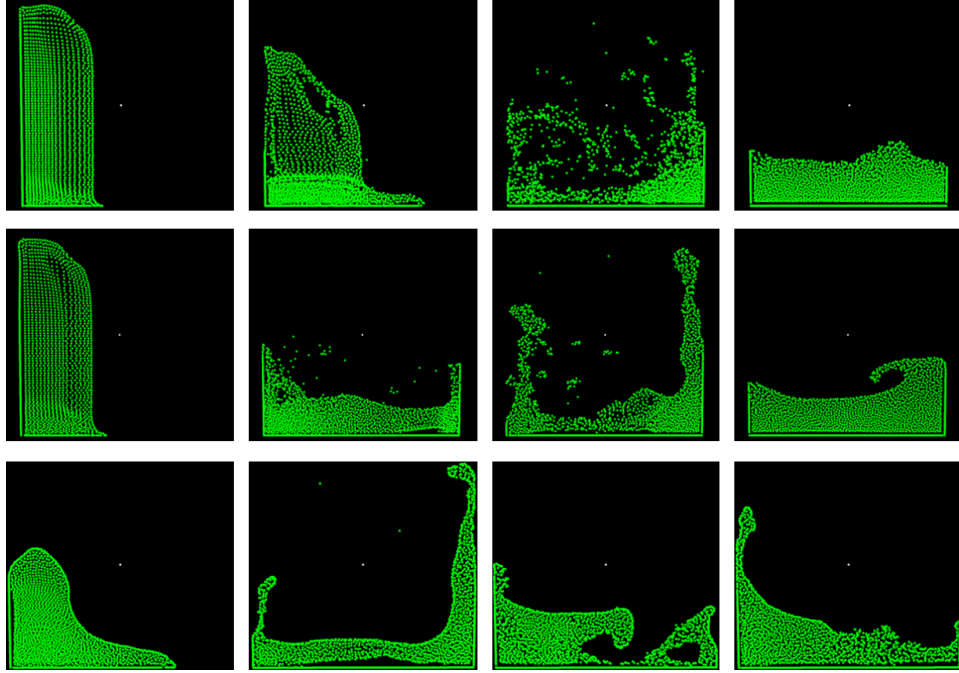


Fig. 2. The classical Dam-break test within our SPH simulation environment. The three different test simulations demonstrate various fluid behavior with respect to changing stiffness and surface tension parameters. Top row describes the behavior of more compressible fluid with less attraction between particles, middle row corresponds to higher constants and finally the last row represents a stiffer fluid with stronger surface tension. Notice the snapshots are not taken in equal time steps.

To test our approach we run each test case with three different neighbor search methods, namely the naive  $O(n^2)$  "all-pairs-test", *Spatial Hashing* and our *Cell Indexing*.<sup>28</sup> and measured the average time spent for searching neighbors and integration of one time step. These results are summarized in table (I).

<sup>28</sup>Other (tree) methods are left a the subject of future work

	All-Pairs	Spatial Hashing	Cell Indexing	Dynamics
Test Case 1	1,95 s	0,42 s	0,25 s	0,25 s
Test Case 2	1,97 s	0,45 s	0,27 s	0,24 s
Test Case 3	1,93 s	0,43 s	0,27 s	0,28 s

Table I. Dam-break tests. 1600 Particles. Average time spent in one simulation time step in each test case.

## 6. CONCLUSION AND FUTURE WORK

We have proposed and demonstrated Cell Indexing as a novel approach for searching approximate neighbor particles within a SPH base fluid simulation. Due to cache misses, Spatial Hashing has been slightly outperformed by our method, without large memory requirements of Full 3D voxelization. Our approach is inherently linear, thus will probably outperform other hierarchical methods<sup>29</sup> for larger data sets. We have introduced a  $H$ -mask to achieve speed-up by searching on a sub-cell resolution. This seamlessly fits into Cell Indexing and can even extend Spatial Hashing.<sup>30</sup>

Due to the immense computational power of current graphics hardware, we will investigate in the future the possibility of implementing our approach on GPU. Further we will try to involve spatial and temporal coherence into our method and allow it to use multiresolutional fluid as been done in [Keiser et al. 2006]. The inherent compressibility of SPH can be decreased by solving the pressure implicitly. Therefore we will explore methods as MPS [Premože et al. 2003] and solve the pressure equation iteratively.

### Appendix - Smoothing Kernel Derivatives

$$\begin{aligned}
\nabla W^{\text{poly}}(\mathbf{r}, h) &= -\frac{945}{32\pi h^9} \begin{cases} (h^2 - r^2)^2 \mathbf{r} & 0 \leq r \leq h \quad \wedge \quad r = |\mathbf{r}| \\ 0 & \text{otherwise} \end{cases} \\
\nabla^2 W^{\text{poly}}(\mathbf{r}, h) &= \frac{315}{64\pi h^9} \begin{cases} (h^2 - r^2)(7r^2 - 3h^2) & 0 \leq r \leq h \quad \wedge \quad r = |\mathbf{r}| \\ 0 & \text{otherwise} \end{cases} \\
\nabla W^{\text{press}}(\mathbf{r}, h) &= -\frac{45}{\pi h^6} \begin{cases} (h - r)^2 \frac{\mathbf{r}}{r} & 0 < r \leq h \quad \wedge \quad r = |\mathbf{r}| \\ 0 & \text{otherwise} \end{cases} \\
\nabla^2 W^{\text{visco}}(\mathbf{r}, h) &= \frac{45}{\pi h^6} \begin{cases} (h - r) & 0 \leq r \leq h \quad \wedge \quad r = |\mathbf{r}| \\ 0 & \text{otherwise} \end{cases}
\end{aligned} \tag{18}$$

<sup>29</sup>They usually need  $O(n \log n)$  to rebuild.

<sup>30</sup>However more cache misses or larger memory requirements arise

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