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Article in *Journal of Graphics Tools* · January 2009

DOI: 10.1080/2151237X.2009.10129281 · Source: DBLP

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# A hierarchical hashing scheme for Near-est Neighbour Search and Broad-Phase Collision Detection

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**Abstract.** Increasing computational power allows computer graphics researchers to model spectacular phenomena such as fluids and their interactions with deformable objects and structures. Particle-based (or Lagrangian) fluid and solid simulations are commonly managed separately and mixed together for the collision detection phase. We present a unified dynamic acceleration model to be used for particle neighbourhood queries and broad-phase collision detection, based on a hierarchical hash table data structure. Our method is able to significantly reduce computations in large, empty areas, and thus gives better results than existing acceleration techniques such as multilevel hashing schemes or KD-trees in most situations.

## 1. Introduction

Fluid simulation has been a major focus in the Computer Graphics community for the last two decades. The difficulty of capturing the complex motion of a fluid encouraged researchers to focus on physically-based simulations. Particle-based (or Lagrangian) simulations are widely used, mainly because particle systems are easy to implement. This type of simulation has proved to give excellent results, such as the well-known Smoothed Particle Hydrodynamics (SPH) method described in [Müller et al. 03] for water simulation.

When considering interactions between a set of objects and a fluid modelled

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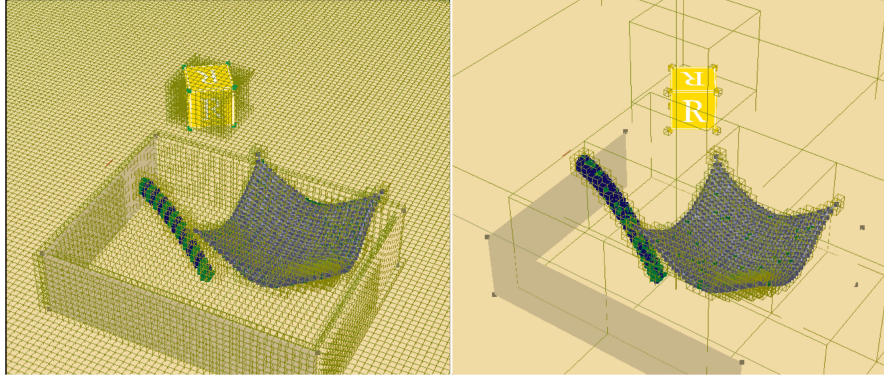
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by a set of particles, we rely on neighbourhood computations in two main cases:

1. for each particle, we must determine a set of neighbouring particles when computing the fluid dynamics (a fluid particle acts upon the surrounding fluid particles). This problem is known as **Nearest Neighbours Search** (NNS);
2. we must also check if a particle collides with objects in the scene in order to compute correct fluid-object coupling. This collision detection is a two-phase process: the **broad-phase** consists in finding a set of candidate objects, then in the narrow-phase all these candidate objects are checked for collision using exact arithmetic computations. It is therefore critical that the broad-phase is implemented efficiently to get the smallest possible set of candidates, and that all the situations where the particle has no chance of colliding with an object can be quickly discarded.

Since particles and objects are in motion, neighbourhood information must be updated at every integration step, which is the most computationally expensive part of the whole simulation process. Therefore, we must provide an acceleration method in order to efficiently determine the neighbourhood. Particle-based fluid simulations usually rely either on space-partitioning data structures such as KD-trees [Adabala and Manohar 00, Adams et al. 07] or hash tables [Teschner et al. 03, Mirtich 96, Eitz and Lixu 07] that store particles and objects and are dynamically updated at each time step. As stated in [Keiser 06], KD-trees supposedly show better computational performances when dealing with a high number of particles, but hash tables have other advantages: they represent an unbounded, implicit grid which does not limit the spatial extent of the simulation, and they are also very easy to implement.

Our method extends the multilevel hashing scheme described in [Eitz and Lixu 07] in order to accelerate broad-phase collision detections and NNS queries, by storing explicit relationships between hash cells and subcells using a *hierarchical* hashing. Since it is capable of efficiently pruning large, empty areas in the scene, our approach combines both the benefits of hash tables and hierarchical structures such as KD-trees. Test results show significant improvements over existing methods, especially for broad-phase collision detection.



**Figure 1.** A scene “rasterized” with a unique cell size (*left*) and using a multilevel model (*right*).

## 2. Background

### 2.1. Spatially Uniform Hashing for NNS

In particle-based simulations, spatially uniform hash tables are well-adapted for the NNS problem. Since fluid particles usually interact with all particles lying in a sphere of radius  $r$  [Müller et al. 03], this value can be used to efficiently store the particles in a hash table. One of the most efficient hash functions is based on the XOR operation, noted as  $\oplus$ :

$$h(\mathbf{p}) = h(p_x, p_y, p_z) = \left( \left( \left( \alpha \left\lfloor \frac{p_x}{r} \right\rfloor \right) \oplus \beta \left\lfloor \frac{p_y}{r} \right\rfloor \right) \oplus \gamma \left\lfloor \frac{p_z}{r} \right\rfloor \right) \bmod n \quad (1)$$

where  $\alpha, \beta, \gamma$  are big prime numbers,  $n$  is the hash table’s length and  $p_x, p_y, p_z$  are the  $x, y, z$  axis components of point  $\mathbf{p}$  [Eitz and Lixu 07].

If a particle  $\mathbf{p}$  is hashed to a given cell  $h(\mathbf{p})$ , then neighbouring particles lying within radius  $r$  are found in the same cell and its 26 neighbours (or 8 neighbours in 2D); the hashkeys of these neighbouring cells are obtained by  $h(p_x + r, p_y + r, p_z + r), h(p_x - r, p_y + r, p_z + r), \dots$

### 2.2. Multilevel Hashing for Broad-Phase Collision Detection

Unfortunately, the use of a unique hash size  $r$  is computationally expensive if used simultaneously for NNS and broad-phase collision detection since fluid particles may collide with objects of size  $b \gg r$ . Consequently, the computational complexity of a uniform hashing scheme is proportional to  $(\frac{b}{r})^3$ , and very large objects may cover thousands of cells as shown on Figure 1 (left). Depending on the configuration of the 3D scene, this is the bottleneck of

simulations that involve fluid-solids interactions.

An efficient and elegant solution presented in [Eitz and Lixu 07] consists in defining multiple hash sizes, so that no object can cover more than 8 cells. Each object is hashed using the most appropriate size, as shown on Figure 1 (right), thus the number of hash cells is significantly reduced. Hash sizes are defined as powers of 2, and a cell may contain 8 subcells like in a recursive octree structure. But only cells of defined sizes need to be stored, which reduces memory consumption, especially when there are big differences in objects sizes (e.g.  $2^{-2}, 2^4, 2^{15}$ ).

The hash size corresponding to an object must be at least the size of the longest edge  $l$  of its AABB (Axis-Aligned Bounding Box). This size  $s$  is computed by converting  $l$  into the nearest but greater power of 2:

$$s = 2^{\lceil \log_2(l) \rceil} \quad (2)$$

This computation is performed for each object during a preprocessing step at the beginning of the simulation, yielding a set of different *classes* associated with different sizes. We consider in the following that a given class can be identified either by its number  $c$  or its size  $s = 2^c$ .

This hashing scheme can handle deformable objects, i.e. whose size can change through time. At each integration step, if the class  $c$  corresponding to an object was not computed in the preprocessing step, then  $c$  is given by the nearest greater class. The object is thus hashed in the smallest possible class by considering the 8 endpoints of its AABB: for each point  $\mathbf{p}$ , the object’s id is added in the hash table at the position given by  $h^c(\mathbf{p})$ , defined by extending equation 1:

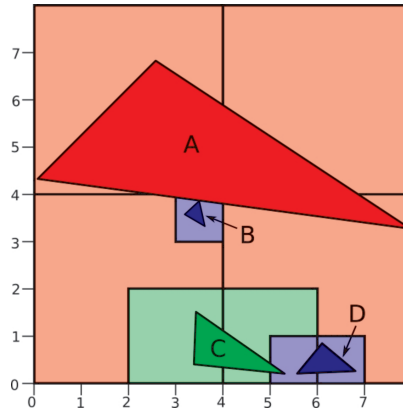
$$h^c(\mathbf{p}) = h^c(p_x, p_y, p_z) = \left( \left( \left( \alpha \left\lfloor \frac{p_x}{s} \right\rfloor \right) \oplus \beta \left\lfloor \frac{p_y}{s} \right\rfloor \right) \oplus \gamma \left\lfloor \frac{p_z}{s} \right\rfloor \right) \bmod n \quad (3)$$

An example of the hash structure obtained with this method is presented on Figure 2 (left). For the sake of clarity, we use left-bottom coordinates to identify cells, and hashkeys are computed without modulo operations (i.e.  $n = +\infty$ ). A unique hashtable, or alternatively one hashtable per class  $c$ , is sufficient to hash objects with minimal *hash collisions* (which should not be confused with collision detection), i.e. distant objects will unlikely have the same hashkey, although this also depends on the hashtable’s length  $n$ .

Using this multilevel hashing scheme, the broad-phase collision detection between a given object and the objects stored in the hash structure consists in considering each class, and check whether one of its cells overlaps the object’s AABB; if positive, ids stored in this cell are retrieved for the narrow-phase. Since a reduced number of cells has to be checked, this process is far more efficient than if a unique cell size was used.

As noted in [Eitz and Lixu 07], another advantage is the ease of implementation, since the multilevel hashing scheme adapts itself to any scene setup,

## A hierarchical hashing scheme for NNS and Broad-Phase Collision Detection



**Figure 2.** **Top:** a 2D example where objects with different sizes are hashed to corresponding classes 0, 1 or 2 (with resp. sizes 1, 2 and 4). Bounding boxes are not represented. **Bottom left:** Non-empty hash cells obtained with multilevel hashing [Eitz and Lixu 07], using Eq. 3. **Bottom right:** Non-empty hash cells obtained with our method using Eq. 4.

including moving and deforming objects over time as shown in Fig. 3. However, if there are no constraints on the deformations of the objects, an AABB can be larger than the largest class: in that case it must be subdivided and hashed to several cells.

### 3. Hierarchical Hashing

#### 3.1. Top-Down Hashing

We extend the hash function defined in Equation 3 with a hierarchical scheme: an element (object or particle) calls this hash function from the largest class  $k$  to the class  $c$  corresponding to its size (top-down), i.e. from  $[k \dots c]$ , where  $c \in [0, k]$ :

$$\tilde{h}^c(\mathbf{p}) = (h^k(\mathbf{p}) + h^{k-1}(\mathbf{p}) + \dots + h^c(\mathbf{p})) \bmod n \quad (4)$$

Then the element may be inserted in the cell given by the key  $\tilde{h}^c$ . A boolean flag is also stored in the hash table to indicate whether the cell has at least one child, i.e. if a cell exists in a smaller class that overlaps  $\tilde{h}^c$  and contains at least one element. Figure 2 (right) shows the hash structure obtained with our method compared to the multilevel hashing scheme of [Eitz and Lixu 07].

Our hash structure is built upon two main data structures:

- a hashmap `hm`: for each hash key, it stores overlapping objects’ ids, and a boolean `hasChildren`
- a set `classSizes`, initialized in a preprocessing step as in section 2.2 and sorted in decreasing order. The minimal size stored in `classSizes` should be the value  $r$  of the interaction radius for fluid particles. Since fluid simulations usually require that this value is the smallest possible, it is safe to assume that no object in the scene will be smaller than  $r$ .

Although multiple calls to the hash function  $h$  are necessary, the additive hash function described by Eq. 4 helps to prevent useless future and expensive computations by reducing *hash collisions*. Consider two 3D points  $\mathbf{p}_1, \mathbf{p}_2$  that are not in the same  $c+1$  class spatial cell, i.e.  $h^{c+1}(\mathbf{p}_1) \neq h^{c+1}(\mathbf{p}_2)$ . Then suppose that a hash collision occurs when hashing these points into class  $c$ , i.e.  $h^c(\mathbf{p}_1) = h^c(\mathbf{p}_2)$ . This hash collision is avoided using Eq. 4 since  $h^c(\mathbf{p}_1) + h^{c+1}(\mathbf{p}_1) \neq h^c(\mathbf{p}_2) + h^{c+1}(\mathbf{p}_2)$ . The offset must be the same for all the points coming from the same parent’s class, which is true in our case because we make use of hierarchical coherence. Hash collisions may still happen because nothing guarantees that the new cell is free, nonetheless this hashing scheme will obviously give better results than a unique hashing. We point out that increasing the hash table size also reduces the probability of hash collisions.

#### 3.2. Particles and Objects Insertions

We consider in Listing 1 the insertion of a single point  $\mathbf{p}$  in the structure, which belongs to an element referenced by its `id`, into a given class `size`. The

hash function  $h$  is implemented as in Equation 3.

**Listing 1.** Top-down point hashing

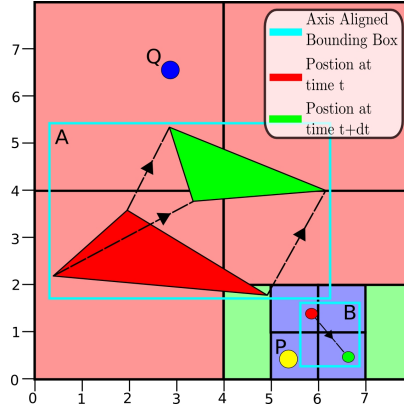
```
void insert(point p, int id, int size) {
    int hashKey = 0;
    for each (int currSize in classSizes) {
        hashKey += h(p, currSize);
        if (size == currSize) {
            hm[hashKey].insert(id); break;
        }
        else if (!hm[hashKey].hasChildren)
            hm[hashKey].hasChildren = true;
    }
}
```

Relying on Listing 1, the insertion of a particle is straightforward: we simply use its position and id as input values, and the interaction radius  $r$  as the hash size. Since  $r$  is also the minimal size in `classSizes`, each of its elements will be considered. The insertion of a larger object is also rather simple. First, we compute its AABB and determine the hash size  $s$  in which it should be hashed as in section 2.2. Then top-down hashing is achieved for each of the 8 endpoints of the AABB using Listing 1, with the object’s id and size  $s$  as input values.

This approach can be used to insert either static or moving elements. Assuming linear motion during one integration step, we simply consider the AABB of its entire path - e.g. the path of a moving triangle is a prism, as shown on Figure 3 - and insert the object as described above using Listing 1. In this case, the top-down hashing stops as soon as the size corresponding to the AABB is met; since we assume in Listing 1 that `size` belongs to the pre-computed `classSizes` set, it may be necessary to choose the nearest greater size as in section 2.2. Unlike some other approaches [Kondoh et al. 04], this method is capable of inserting moving elements independently of the distance covered during a time step.

### 3.3. Hash Structure Queries

Any query in the hash structure will take advantage of the top-down hashing of static or moving objects described in the previous section. As shown on Figure 3 (right), finding possible candidates that may collide or interact with a single point consists in iteratively hashing this point for each class, starting from the largest class, and retrieving the ids stored in the corresponding cell as candidates. The process ends if the cell has no child or if the smallest class is



Hashkey	Objects	hasChildren	Point $P$
$\tilde{h}^2(0, 0)$	A	false	$\downarrow$
$\tilde{h}^2(4, 0)$	A	true	$\tilde{h}^2(4, 0)$ : retrieve object A
$\tilde{h}^2(0, 4)$	A	false	$\downarrow$
$\tilde{h}^2(4, 4)$	A	false	$\tilde{h}^1(4, 0)$
$\tilde{h}^1(4, 0)$		true	$\downarrow$
$\tilde{h}^1(6, 0)$		true	$\tilde{h}^0(5, O)$ : retrieve object B
$\tilde{h}^0(5, 1)$	B		
$\tilde{h}^0(5, 0)$	B		Point $Q$
$\tilde{h}^0(6, 1)$	B		$\downarrow$
$\tilde{h}^0(6, 0)$	B		$\tilde{h}^2(0, 4)$ : retrieve object A

**Figure 3.** **Top:** Hashing moving objects. **Bottom left:** Corresponding, non-empty hash cells obtained with our method. **Right:** Top-down broad-phase collision detection for points  $P$  and  $Q$  (marked as colored circles)

reached. Unfortunately, the problem is more challenging for both broad-phase collision detection and NNS. This simple method is thus extended below to handle more complex queries.

### 3.3.1. Broad-phase Collision Detection

We are interested in checking if a given AABB overlaps any cell in the hash structure that may contain possible candidates for the narrow-phase collision detection. This broad-phase algorithm is described on Listing 2. In the context of fluid-solid interactions, the AABB encloses the path of a particle. In a more general context, it corresponds to a static or moving object that may collide with other objects in the scene as in Figure 3.

The main advantage of our method is the use of the boolean `hasChildren`, which quickly ends the process if not necessary by emptying the `points` set. Since we deal with an AABB `a` of arbitrary size, we may have to call `a.subdivide(t)` to subdivide the box  $t$  times as in an octree decomposition, in order to ensure that we do not miss hash cells smaller than `a.size()`. A more complex implementation (not presented here) could also optimize the process by ensuring that a value `p.hashKey` is not processed multiple times, which happens if `a.size()` is very small since several endpoints can be hashed in the same cell.

**Listing 2.** Get all the objects that may collide with a given AABB

```
set<int> broadPhase(AABB a) {
    set<int> candidateObjects;
    // Build a set with the 8 endpoints
    // (the initial hash key for these points is set to 0)
    vector<point> points = a.getEndpoints();

    for each (int currSize in classSizes) {
        int delta = a.size() - currSize;
        // Subdivide AABB if necessary
        if (delta > 0) points.insert(a.subdivide(delta));
        // Find possible candidates in corresponding cells
        for each (point p in points) {
            p.hashKey += h(p, currSize);
            candidateObjects.insert(hm[p.hashKey].retrieveIds());
            // Remove this point if the cell has no child
            if (!hm[p.hashKey].hasChildren) points.remove(p);
        }
    }
    return candidateObjects;
}
```

### 3.3.2. NNS

As stated in section 2.1, if a particle **p** is hashed to a given cell, then neighbouring particles lying within radius  $r$  are simply found in the same cell and its 26 neighbours. The hash keys of these neighbouring cells are simply given by  $\tilde{h}^r(p_x + r, p_y + r, p_z + r)$ ,  $\tilde{h}^r(p_x - r, p_y + r, p_z + r)$ , ...

Unfortunately, since  $r$  is the smallest class, this simple approach does not perform well with our method because computing neighbouring hash keys requires to consider all classes stored in `classSizes`. We thus propose to determine neighbouring particles directly when a particle is inserted. We add a new attribute in the hash structure called `neighbours` that stores, for a given

particle, the references of its neighbours. Then each particle is processed using the modified insertion procedure described on Listing 3, which makes use of the interaction radius  $r$  and of a method `get27Points(p)` that produces the set  $p, (p_x + r, p_y + r, p_z + r), (p_x - r, p_y + r, p_z + r), \dots$ ,

**Listing 3.** Particle insertion and NNS computation

```

void insert(particle P) {
    set<int> candidateParticles;
    // Build a set with P and 26 neighbouring points
    // (the initial hash key for these points is set to 0)
    vector<point> points = get27Points(P.position);

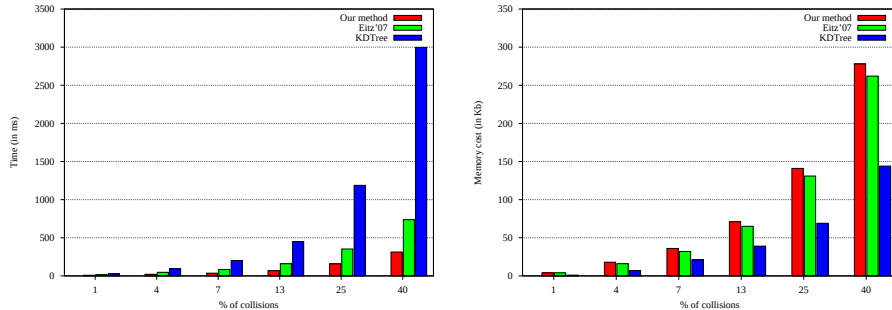
    for each (int currSize in classSizes)
        // Retrieve possible candidates if size r is reached
        for each (point p in points) {
            p.hashKey += h(p, currSize);
            if (currSize == r)
                candidateParticles.insert(hm[p.hashKey].retrieveIds());
            if (!hm[p.hashKey].hasChildren) points.remove(p);
        }
        // Update the 'neighbours' attribute if the distance
        // between particles is lower than r
        for each (int id in candidateParticles)
            if (distance(P.id, id) <= r) {
                neighbours[P.id].insert(id);
                neighbours[id].insert(P.id);
            }
        // Insert the particle (finally !)
        insert(P.position, P.id, r);
    }
}

```

This *on-the-fly* NNS computation gives better results than the uniform hash scheme described in section 2.1, again thanks to the boolean flag `hasChildren` that discards empty areas. When the whole set of particles has been processed, `neighbours[id]` stores the neighbours of a particle `id` which can be retrieved for fluid dynamics computations.

#### 4. Results and Comparisons

The results presented below are performance comparisons of our method against several other acceleration structures on random distributions, conducted on a single i686 processor running at 2.66 GHz, in order to validate



**Figure 4.** **Left:** Timings comparisons for broad-phase collision detection. **Right:** Memory consumption comparisons.

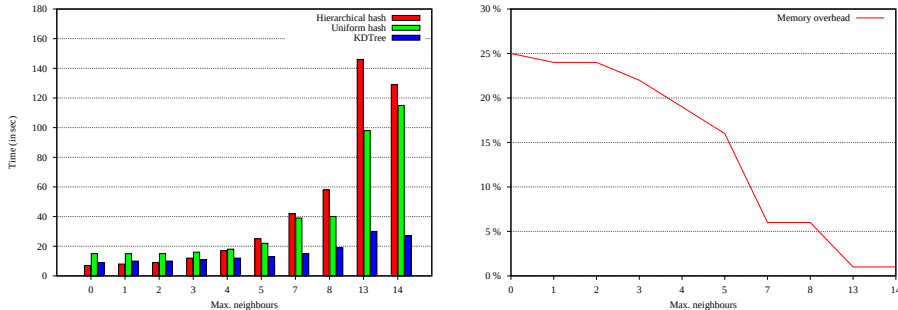
our model on typical particle-based simulations.

#### 4.1. Comparisons on 2D Random Distributions

Figure 4 shows the result on broad-phase collision detection tests, using a KD-tree, the multilevel hashing scheme of [Eitz and Lixu 07] and our method. A distribution of  $N$  objects is generated randomly in a 2D square of size  $2^{13}$  and inserted in the acceleration structure using each object’s AABB. Then a distribution of  $1M$  segments is computed, which accounts for moving particles in the scene; for each segment we query the structure to determine which objects it may intersect. The objects’ size distribution is implemented in a way that approx. 10% of the objects have an average size of  $2^{10}$ , 25% have an avg. size of  $2^6$ , and the remaining objects have an avg. size of  $2^3$ , whereas the length of each particle’s motion vector does not exceed  $2^4$ . The computation times are depicted as a function of the ratio of particles that collide with at least one of  $N$  objects; this ratio increases according to  $N$ . Our method improves the timings obtained with Eitz and Lixu’s method by at least a factor 2 in all situations, whereas the memory overhead induced by our hierarchical structure does not exceed 10%. Moreover, our tests indicate that hash-based methods in general seem a better choice for broad-phase collision detection over KD-trees, although more memory-consuming.

The second tests aim at evaluating the computational cost of our model for NNS, compared to a KD-tree and a uniform hashing scheme<sup>1</sup>. A distribution of  $1M$  points is generated randomly in a 2D square of size  $2^{20}$ , and inserted in the acceleration structure; then for each point we query its neighbours located in a circle of radius  $r$ . The computation times are depicted on Figure

<sup>1</sup>In that case, a uniform hashing scheme [Teschner et al. 03] can be considered as a restriction of Eitz and Lixu’s hashing scheme with only one level.



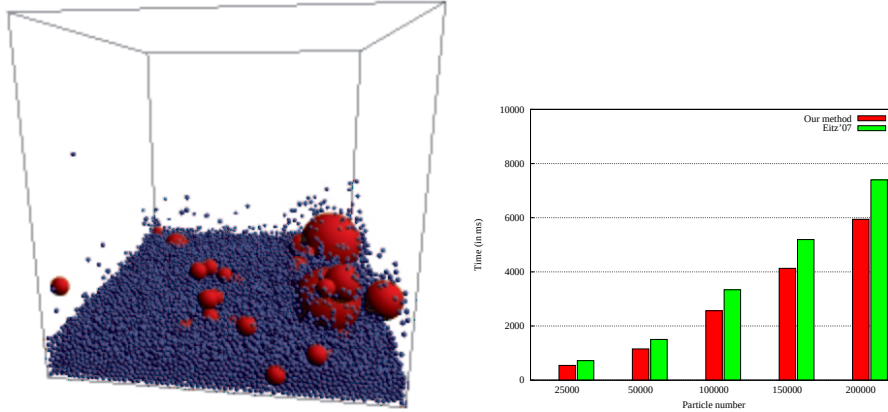
**Figure 5.** **Left:** Timings comparisons for Nearest-Neighbour Search. **Right:** Memory overhead induced by our method.

5 (left) as a function of the maximal number of neighbours lying within an increasing radius  $r$ . We can note that our model is competitive over uniform hashing when particles are far from each other, i.e. when maximal neighbours count is below 5; however the memory overhead in that case ranges from 10% to 25%. For more dense points distributions, the KD-tree exhibits the best performances, partly because the objects to be inserted all have the same size in the NNS case. Implementations of particle-based simulations that involve high neighbour count should thus prefer KD-tree acceleration over hash tables for NNS.

#### 4.2. Comparisons on a 3D fluid simulation

Our 3D test scene is presented on Figure 6 (left): a particle-based fluid (in blue) falls and bounces over a set of 2000 static, solid spheres (in red), and eventually stabilizes around its equilibrium position at the bottom of the simulation box. We have chosen a particle-based model with simple collision response for the fluid simulation, which was conducted over 1000 successive iterations, but other Lagrangian models such as SPH could be used [Müller et al. 03]. Dynamic particles and static spheres are initially located randomly in the box; static spheres’ sizes range from  $2^{-4}$  to  $2^{-2}$ , whereas particles are  $2^{-6}$  in size.

Timings are given as the time required to compute the whole collision detection process per frame: hashing objects (although spheres and walls are static in this example), hashing moving particles, retrieving possible candidates for each particle, and finally computing exact collisions. A uniform hashing scheme was used for Nearest-Neighbour search. As shown on Figure 6 (right), our method outperforms Eitz and Lixu’s by approximately 25%; memory measurements (not presented here) showed that memory overhead



**Figure 6.** **Left:** Our 3D test scene. **Right:** Performance comparisons for collision detection (in millisecond per simulation frame)

needed to store hierarchical relationships never exceeds 20%, as in the 2D case. One can notice that, due to the progressive aggregation of particles at the bottom of the box, the number of isolated particles tends to decrease over time. This explains why the acceleration brought by our method is not as spectacular as in the 2D case, but remains significant nonetheless.

#### 4.3. Discussion and Improvements

The 2D tests presented in this paper were implemented using the STL’s `hash_multimap` container [Musser et al. 01]. This simple implementation avoids using a fixed length for our hash structure, which means we can discard the modulo operation from Eqs. 1 and 4, and consider  $n = +\infty$ . Although this comes at the price of logarithmic time complexity to find objects associated with a given key, it also demonstrates that, without optimizations, our data structure already provides nice results even for 1M particles.

A more optimized implementation was used for our 3D experiments, based on a fixed-length vector. Again, satisfying results were obtained compared to Eitz and Lixu’s approach, but in that case a specific value for  $n$  must be chosen. This problem has a significant impact on memory consumption if too high; on the other hand, a lower value may imply more hash collisions, which in turn increase the number of candidate objects obtained after the broad-phase. Our experiments showed that the overhead computations induced by these ‘false’ candidates is negligible if  $n$  is set to the approximate number of

polygons in the scene multiplied by 100.

In the specific case of dynamic simulations, we could make use of a timestamp, i.e. a parameter that distinguishes two time steps [Teschner et al. 03]. If a new object is to be inserted in a hash cell and the timestamps of the cell and the object differ, then the object does not necessarily need a new memory allocation to be stored, since it can reuse the memory allocated for another object at the previous time step. The timestamp is thus used to reduce overall memory allocation when hash list insertions are achieved. By using this scheme, the hash table is never entirely swept between successive time steps. In STL-based implementations this approach is readily obtained since the `clear` operation does not necessarily free its memory when called at each time step. Performances also depend on simulation parameters such as the time step: the larger the time step is, the bigger the distance an element will cover and consequently a longer distance will have to be hashed.

One can point out that the complexity of our hash function  $\tilde{h}^c$  is proportional to the number of different classes, whereas the multilevel hashing scheme requires a single call to the hash function  $h^c$ . Nevertheless, it is not necessary to check all different classes to find candidate objects, thanks to the boolean flag `hasChildren` that indicates whether smaller objects could be found in a given area; this explains the good performances of our method, especially for collision detection.

**Acknowledgments.** The authors thank C. Rousselle for her corrections, and the reviewers who suggested numerous improvements.

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Received [DATE]; accepted [DATE].