MLS Pressure Extrapolation for the Boundary Handling in Divergence-Free SPH

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Abstract

We propose a novel method to predict pressure values at boundary particles in incompressible divergence-free SPH simulations (DFSPH). Our approach employs Moving Least Squares (MLS) to predict the pressure at boundary particles. Therefore, MLS computes hyperplanes that approximate the pressure field at the interface between fluid and boundary particles. We compare this approach with two previous techniques. One previous technique mirrors the pressure from fluid to boundary particles, but uses a gradient that is computed with Smoothed Particle Hydrodynamics (SPH). We motivate that gradient-based extrapolation is more accurate than mirroring. We further motivate that our proposed MLS gradient is less error prone than the SPH gradient at the boundary. In our experiments, we indicate artifacts in previous approaches. We show that these artifacts are significantly reduced with our approach resulting in simulation steps that can be twice as large compared to previous methods. We further present challenging and complex scenarios to illustrate the capabilities of the proposed boundary handling.

CCS Concepts

• Computing methodologies → Physical simulation; Massively parallel and high-performance simulations;

1. Introduction

Iterative pressure solvers such as PCISPH [SP09], IISPH [ICS*14] or DFSPH [BK17] compute a pressure field \( p \) and apply pressure accelerations of the form
\[
a_f^p = -\sum_j m_j \left( \frac{p_f}{\rho_f^2} + \frac{p_b}{\rho_b^2} \right) \nabla W_{ij}
\]
to particles \( i \). The sum considers all neighboring particles \( j \) of particle \( i \). The variables \( m, p, \nabla W \) denote mass, density and the gradient of the SPH kernel function \( W \), respectively. There exist minor variations, but the solvers follow the same concept (see Section 4).

Iterative solvers typically compute pressure \( p_f \) only at fluid particles \( f \). Pressure \( p_b \) at boundary particles \( b \) is not computed, but approximated if needed. This is, e.g., the case in the computation of the pressure acceleration \( a_f^p \) at fluid particles \( f \) close to the boundary. Here, the respective SPH sum iterates over fluid neighbors with known pressure, but also over boundary neighbors with unknown pressure. I.e.,
\[
a_f^p = -\sum_{f_j} \frac{m_{f_j}}{\rho_{f_j}} \left( \frac{p_f}{\rho_f^2} + \frac{p_{f_j}}{\rho_{f_j}^2} \right) \nabla W_{f_j f} - \sum_{b_j} \frac{m_{b_j}}{\rho_{b_j}} \left( \frac{p_f}{\rho_f^2} + \frac{p_{b_j}}{\rho_{b_j}^2} \right) \nabla W_{f_j b}
\]
with \( f_f \) and \( f_b \) denoting fluid and boundary neighbors of fluid par-
ticle \( f \), respectively. Equation (1) requires a notion of mass \( m_b \), density \( \rho_b \), and pressure \( p_b \) at boundary particles \( b \). The density \( \rho_b \) is typically set to the rest density of the adjacent fluid and the mass \( m_b \) can be geometrically motivated from the boundary particle volume, see e.g. [AIA*12].

In terms of the pressure \( p_b \), there exist different options. E.g., Akinci et al. [AIA*12] propose to mirror known pressure from fluid particles to adjacent rigid particles. Alternatively, Adami et al. [AHA12] propose to extrapolate pressure from the fluid to boundary particles using an SPH approximation of the pressure gradient.

**Our contribution:** We propose a novel variant to predict unknown pressure \( p_b \) at boundary particles \( b \). In contrast to Akinci et al. [AIA*12], we compute one unique pressure value per boundary particle. We further extrapolate the pressure using the pressure gradient instead of copying pressure from the fluid to the boundary. This improves the quality of the pressure gradient of fluid particles near the boundary as illustrated in Fig. 2. In contrast to Adami et al. [AHA12], where pressure is extrapolated using SPH, we propose to use MLS [KK56,Nea04]. MLS is more accurate than SPH in case of particle deficiency which can particularly be the case near the boundary. We illustrate artifacts when using the boundary handling of Akinci et al. [AIA*12] and Adami et al. [AHA12]. We further show that these artifacts can be reduced with the proposed MLS extrapolation of pressure values. We have implemented our boundary handling in a DFSPH framework. Capabilities of the approach are illustrated for scenarios with challenging boundary setups, e.g. Fig. 1. We particularly show performance gain factors of up to two compared to [AIA*12].

**Organization:** The remainder of this paper is organized as follows. The following Section 2 describes existing approaches related to SPH fluid simulation and the handling of solid boundaries. In Section 3, we discuss the proposed pressure extrapolation concept whereas implementation details are described in Section 4. In Section 5, we show simulations employing our method and compare it to the boundary handling schemes of Akinci et al. [AIA*12] and Adami et al. [AHA12]. Finally, we conclude in Section 6.

## 2. Related Work

SPH is a popular choice for Lagrangian simulations in computer graphics [IOS*14]. First used by Stam and Fiume [SF95] to simulate gaseous phenomena and by Desbrun and Cani [DG96] for deformable objects, Müller et al. [MCG03] employed SPH to simulate compressible fluids. From that time on, research has focused on practical formulations of incompressible fluids with recent improvements in volume preservation [ICS*14, BK17, TDNL16], multiphase simulation [MSKG05, SP08, RLY*14, ATO17], highly viscous fluids [PICT15, TDF*15, PT17, BGFAO17, WKBB18] and deformable objects [KAG*05, SSP07, PGBT18]. Incompressibility can be enforced in various ways. Unlike non-iterative state equation solvers, e.g. [Mon04, MCG03, APKG07, BT07], iterative SPH pressure solvers, such as PCISPH [SP09], IISPH [ICS*14] and DFSPH [BK17], compute a pressure field \( p \) by solving a pressure Poisson equation (PPE) of the form \( \nabla^2 p = \mathbf{s} \), c.f. [Cho68].

Thereby, \( \mathbf{s} \) is a source term that either encodes a predicted density deviation [SL03, SP09, ICS*14], the divergence of a velocity field [CR99, PDC*03] or a combination of both [HA07, BK17]. Computing the pressure field from a global formulation seems to improve the stability of the simulation. Rather large time steps can typically be used compared to the aforementioned non-iterative state equation solvers.

This paper focuses on the optimization of the boundary handling. Therefore, we briefly discuss related works regarding the modeling of solid boundaries in the next section.

### Boundary Handling in SPH

As particle-based representations are very flexible and can handle arbitrarily shaped geometries, representing solid boundaries with particles is a popular choice for SPH fluid simulations, e.g. [Mon05, IAGT10, ICS*14, BK17, TDNL16]. One popular technique for handling fluid-boundary contact is to apply penalty forces between two particles as soon as they are within a certain distance, e.g. [Mon05, MST*04, MK09]. Penalty forces should prevent fluid particles from penetrating the boundary. Therefore, the magnitude of the penalty force is determined based on a penetration measure between the particles. As the results are sensitive to the stiffness parameter of the penalty force, small time steps are typically required to produce a smooth pressure field.

In order to achieve larger time steps, the direct forcing method of Becker et al. [BT09] uses a predictor-corrector scheme to compute control forces and velocities. This method guarantees non-penetration. However, due to an incomplete support domain at the boundary, stacking of fluid particles can occur.

Another technique to treat boundaries are ghost particles [CL03, YRS09, SB12]. For fluid particles that are located at a certain dis-
tance to the boundary, a narrow layer of ghost particle is generated. Those ghost particles mirror the hydrodynamic quantities of their associated fluid particle, i.e. they have the same viscosity, mass, density and pressure. However, for complex geometries, generating such ghost particles is challenging. Furthermore, ghost particles have to be re-generated per simulation step.

Using only one layer of pre-generated boundary particles, Akinci et al. [AIA∗12] treat irregular samplings by computing volume contributions, c.f. [OS03, SP08], and by mirroring the quantities of a fluid particle onto its neighboring boundary particles. While adhering to the SPH concept, this approach is efficient to compute and allows a versatile coupling of fluids and solid objects [ACAT13]. Band et al. [BGT17] propose an extension of the boundary handling scheme of Akinci et al. [AIA∗12] by employing MLS. To improve the accuracy of the density estimate and normal computation in planar regions, they locally reconstruct the surface of the true boundary by fitting boundary particles to a plane. This approach results in a smooth representation of the boundary. However, it is only applicable to planar boundaries. Furthermore, Band et al. [BGT17] do neither compute unique pressure values nor perform any pressure extrapolation at boundary particles. Instead, they use the mirroring scheme of Akinci et al. [AIA∗12]. Yet, MLS techniques have been successfully applied in many research areas, e.g. [Di99, ABCO∗03, MKN∗04, BRHN11].

Instead of mirroring fluid particle quantities onto the boundary, Adami et al. [AHA12] propose to use pre-generated dummy boundary particles. Thereby, fluid particles at the boundary interact with dummy particles according to the overlap of the kernel function. This has the advantage that, even for complex geometries, the boundary is well-described through-out the simulation. Furthermore, by extrapolating the pressure of boundary particles from the surrounding fluid particles, this method allows an accurate approximation of a fluid particle’s pressure gradient near the boundary.

As an alternative to particles, boundaries are also representable with triangle meshes [HEW15, FM15]. Yet, as stated in [AIA∗12], handling discontinuous surface normals and non-manifold structures that cause spatial and temporal discontinuities of the fluid properties is challenging for triangulated boundaries. Another alternative is an implicit representation of the boundary as proposed by Koschier and Bender [KB17]. Based on a pre-computed density map, this approach allows to efficiently evaluate the density and pressure gradient of fluid particles at the boundary.

3. Method

We first discuss the previous concepts of Akinci et al. [AIA∗12] in Section 3.1 and Adami et al. [AHA12] in Section 3.2. Our approach is introduced in Section 3.3. This section focuses on the concepts. The combination of the boundary handling with DFSPH is described in Section 4.

3.1. Pressure mirroring

Pressure mirroring is motivated by its simple and efficient implementation. When a pressure acceleration is computed at a fluid particle \( f \) that has a boundary particle \( b \) with unknown pressure in its neighborhood, the pressure at the boundary particle is simply set to the pressure of the fluid particle, i.e. \( p_{fb} = p_f \). The computation in Eq. (1) slightly changes to

\[
\mathbf{a}_f = -\sum_{j \in f} m_{fj} \left( \frac{p_f}{r_{fj}^2} + \frac{\rho_f}{\rho_{fj}} \right) \nabla W_{ff} \nabla
\]

\[
-\sum_{j \in b} \rho_j \nabla V_{fb} \left( \frac{p_f}{p_j} + \frac{p_j}{p_f} \right) \nabla W_{ff} \nabla .
\]

Compared to Eq. (1), the mass of a boundary neighbor \( m_{fb} \), i.e. its contribution in the SPH sum, is computed as \( m_{fb} = \rho_{fb} V_{fb} \) and the density of a boundary neighbor is set to \( \rho_{fb} = \rho_f \). Please refer to [AIA∗12] for a motivation of these choices and for a discussion how to compute the volume \( V_{fb} \).

While density and mass in Eq. (2) play an important role for the accurate weighting of a boundary particle in the SPH sum, the employed pressure approximation \( p_{fb} = p_f \) negatively affects the accuracy of the pressure gradient computation, i.e. the computation of the pressure acceleration. As illustrated in Fig. 2, it would be more appropriate to extrapolate the pressure from the fluid to the boundary.

Pressure mirroring does not require to iterate over boundary particles or to store pressure values at boundary particles. If a boundary pressure is required, it is simply set to the pressure of the currently processed fluid particle. While this efficiency is positive, it results in inconsistent pressure values at boundary particles. If two fluid particles \( f_1 \) and \( f_2 \) with different pressure values \( p_{f1} \) and \( p_{f2} \) share the same boundary particle \( b \), the gradient computations at both fluid particles work with different pressure values. Fluid particle \( f_1 \) works with \( p_b = p_{f1} \) in Eq. (2), while the other fluid particle \( f_2 \) uses \( p_b = p_{f2} \) at the same boundary particle \( b \).

3.2. Pressure extrapolation with SPH

Pressure extrapolation can be motivated by Pascal’s law for hydrostatic pressure which states that the pressure difference at two fluid points is proportional to their height difference. For a boundary particle \( b \) and an adjacent fluid particle \( f \), this can be written as \( p_b = p_f + \rho_b \mathbf{g} \cdot \mathbf{x}_{bb} \) with gravity \( \mathbf{g} \) and distance \( \mathbf{x}_{bb} = \mathbf{x}_b - \mathbf{x}_f \). In order to handle the interaction of one boundary particle with several neighboring fluid particles, the respective contributions are weighted with the kernel function \( W_{bb} \), summed up and normalized as proposed by Adami et al. [AHA12]:

\[
p_b = \frac{\sum_{b_j} p_{b_j} W_{bb_j} + \mathbf{g} \cdot \mathbf{x}_{bb_j} W_{bb_j}}{\sum_{b_j} W_{bb_j}}.
\]

In contrast to the pressure mirroring in [AIA∗12], this approach requires an additional loop over boundary particles to compute the pressure which is also stored at boundary particles. On the other hand, boundary pressures are not inconsistent as in [AIA∗12]. Instead, each boundary particle is attributed a unique pressure value. If the pressure \( p_b \) is computed for all boundary particles \( b \), Eq. (1) can be used to compute the pressure acceleration at fluid particles.

Although the computation in Eq. (3) is normalized, it nevertheless
We propose to resolve the particle deficiency issue by using MLS (which corresponds to). Therefore, we fit hyperplanes that approximate the pressure field through the pressure field. The hyperplane parameters \( b \) are estimated from pressure values at adjacent fluid particles (blue). This is conceptually different to [BGT17] where MLS is used to fit planes through boundary sample positions. In our case, MLS is performed from the perspective of a boundary particle. It employs all pressure values \( p_{bf} \) of fluid neighbors \( b \) adjacent to this boundary particle \( b \) to fit a hyperplane through the pressure field. The hyperplane parameters \( c_b = (\alpha_b, \beta_b, \gamma_b, \delta_b)^T \) at boundary particle \( b \) are estimated by minimizing \( \sum_{bf}(b_{bf} \cdot c_b - p_{bf})^2V_{bf}W_{bbf} \) with \( b_{bf} = (1, x_{bf}, y_{bf}, z_{bf})^T \). Considering the fact that the partial derivatives of this term with respect to the hyperplane parameters should be zero at the minimum, we get the following system that has to be solved:

\[
\sum_{bf}b_{bf}(\mathbf{b}_{bf} \cdot \mathbf{c}_b - p_{bf})V_{bf}W_{bbf} = \mathbf{0}. \quad (4)
\]

In order to solve for the unknown hyperplane parameters \( c_b \), we propose to rewrite Eq. (4) as

\[
\sum_{bf}(\mathbf{b}_{bf} \otimes \mathbf{b}_{bf})c_{bf}V_{bf}W_{bbf} = \sum_{bf}b_{bf}p_{bf}V_{bf}W_{bbf}, \quad (5)
\]

which corresponds to

\[
\sum_{bf} \begin{bmatrix}
1 & x_{bf} & y_{bf} & z_{bf} \\
1 & x_{bf} & x_{bf}y_{bf} & x_{bf}z_{bf} \\
1 & y_{bf} & y_{bf}y_{bf} & y_{bf}z_{bf} \\
1 & z_{bf} & x_{bf}z_{bf} & y_{bf}z_{bf} \\
1 & x_{bf} & y_{bf} & z_{bf}
\end{bmatrix} \begin{bmatrix}
\alpha_b \\
\beta_b \\
\gamma_b \\
\delta_b \\
p_{bf}V_{bf}W_{bbf}
\end{bmatrix} = \sum_{bf} \begin{bmatrix}
1 \\
0 \\
0 \\
0 \\
1
\end{bmatrix} p_{bf}V_{bf}W_{bbf}. \quad (6)
\]

This \( 4 \times 4 \) system can be rewritten such that \( \alpha_b \) can be directly computed and the parameters \( \beta_b, \gamma_b, \delta_b \) can be computed by solving a \( 3 \times 3 \) system. We therefore perform a basis transform: all considered positions \( \mathbf{x}_b \) and also \( \mathbf{x}_b \) are translated to positions \( \tilde{\mathbf{x}}_b \) and \( \tilde{\mathbf{x}}_b \) by

\[
\mathbf{d}_b = \frac{\sum_{bf} \mathbf{x}_{bf}V_{bf}W_{bbf}}{\sum_{bf} V_{bf}W_{bbf}}. \quad (7)
\]

i.e.

\[
\tilde{\mathbf{x}}_b = (\tilde{x}_b, \tilde{y}_b, \tilde{z}_b)^T = \mathbf{x}_b - \mathbf{d}_b. \quad (8)
\]

\[
\mathbf{d}_b = \frac{\sum_{bf} \mathbf{x}_{bf}V_{bf}W_{bbf}}{\sum_{bf} V_{bf}W_{bbf}}. \quad (9)
\]

This basis transform, i.e. the translation of all incorporated particle positions by the same vector \( \mathbf{d}_b \), does not affect the parameters of the hyperplane. Thus, instead of solving Eq. (6), we now consider the following system with the same solution:

\[
\sum_{bf} \begin{bmatrix}
1 \\
1 \\
1 \\
1
\end{bmatrix} \begin{bmatrix}
\tilde{x}_b \\
\tilde{y}_b \\
\tilde{z}_b \\
\mathbf{p}_{bf}V_{bf}W_{bbf}
\end{bmatrix} = \sum_{bf} \begin{bmatrix}
\alpha_b \\
\beta_b \\
\gamma_b \\
\delta_b
\end{bmatrix} \begin{bmatrix}
\mathbf{p}_{bf}V_{bf}W_{bbf} \\
V_{bf}W_{bbf}
\end{bmatrix}. \quad (10)
\]

Please note that \( W_{bbf} \), in Eq. (10) depends on the distance between \( \mathbf{x}_b \) and \( \mathbf{x}_{bf} \). Now, the elements \( \tilde{x}_b, \tilde{y}_b, \) and \( \tilde{z}_b \) can be replaced by zero based on the following observation:

\[
\sum_{bf} \tilde{x}_bV_{bf}W_{bbf} = \sum_{bf} (\mathbf{x}_b - \mathbf{d}_b)V_{bf}W_{bbf}
= \sum_{bf} \mathbf{x}_bV_{bf}W_{bbf} - \mathbf{d}_bV_{bf}W_{bbf}
= \mathbf{d}_b\sum_{bf} \mathbf{x}_bV_{bf}W_{bbf} - \mathbf{d}_bV_{bf}W_{bbf}
= \mathbf{0}. \quad (11)
\]

Therefore, \( \sum_{bf} \tilde{x}_bV_{bf}W_{bbf} = \mathbf{p}_{bf} \sum_{bf} \tilde{y}_bV_{bf}W_{bbf} \), but \( \mathbf{p}_b \sum_{bf} \tilde{z}_bV_{bf}W_{bbf} \) in Eq. (10) can be written as \( \mathbf{p}_{bf} \sum_{bf} \tilde{y}_bV_{bf}W_{bbf} \). In the same way, the coefficients \( \tilde{y}_b \) and \( \tilde{z}_b \) in the matrix in Eq. (10) can be replaced by zeros, resulting in the following form:

\[
\sum_{bf} \begin{bmatrix}
1 & 0 & 0 & 0 \\
0 & \tilde{x}_b & \tilde{y}_b & \tilde{z}_b \\
0 & \tilde{x}_b & \tilde{y}_b & \tilde{z}_b \\
0 & \tilde{x}_b & \tilde{y}_b & \tilde{z}_b
\end{bmatrix} \begin{bmatrix}
\alpha_b \\
\beta_b \\
\gamma_b \\
\delta_b
\end{bmatrix} \begin{bmatrix}
\mathbf{p}_{bf}V_{bf}W_{bbf} \\
V_{bf}W_{bbf}
\end{bmatrix} = \sum_{bf} \begin{bmatrix}
\tilde{x}_b \\
\tilde{y}_b \\
\tilde{z}_b \\
\mathbf{p}_{bf}V_{bf}W_{bbf}
\end{bmatrix}. \quad (12)
\]

Now, we can solve for \( \alpha_b \):

\[
\alpha_b = \frac{\sum_{bf} \mathbf{p}_{bf}V_{bf}W_{bbf}}{\sum_{bf} \mathbf{p}_{bf}V_{bf}W_{bbf}}. \quad (13)
\]
The other hyperplane parameters $\beta_b$, $\gamma_b$, and $\delta_b$ are obtained by solving

$$
\begin{pmatrix}
\beta_b \\
\gamma_b \\
\delta_b
\end{pmatrix} = \left( \sum_{b_j} \begin{bmatrix}
\delta_{b_j} & \gamma_{b_j} & \delta_{b_j}
\end{bmatrix} \begin{bmatrix}
\delta_{b_j} & \gamma_{b_j} & \delta_{b_j}
\end{bmatrix} W_{bb_j} \right)^{-1} \begin{bmatrix}
\sum_{b_j} \begin{bmatrix}
\delta_{b_j} & \gamma_{b_j} & \delta_{b_j}
\end{bmatrix} V_{b_j} W_{bb_j} \\
\sum_{b_j} \begin{bmatrix}
\delta_{b_j} & \gamma_{b_j} & \delta_{b_j}
\end{bmatrix} V_{b_j} W_{bb_j}
\end{bmatrix}.
$$

Finally, the pressure $p_b$ at boundary particle $b$ is computed as

$$
p_b = (1, \bar{x}_b, \bar{y}_b, \bar{z}_b)^T \cdot c_b.
$$

If the matrix in Eq. (14) is not invertible, we set $\beta_b$, $\gamma_b$, and $\delta_b$ to zero resulting in

$$
p_b = (1, \bar{x}_b, \bar{y}_b, \bar{z}_b)^T \cdot (\alpha_b, 0, 0, 0)^T = \alpha_b = \frac{\sum_{b_j} p_{b_j} V_{b_j} W_{bb_j}}{\sum_{b_j} V_{b_j} W_{bb_j}}.
$$

This pressure corresponds to the weighted average of the pressure values of fluid particles $b_j$ adjacent to boundary particle $b$. In our experiments, however, we only experienced issues with a singular matrix in Eq. (14) for boundary particles with a single fluid neighbor and for boundary particles whose fluid neighbors lie exactly on a line or plane. In these cases, employing the proposed basis transform sets many coefficients of the matrix to zero.

Similar to the SPH pressure extrapolation, our approach performs a loop over boundary particles to compute and store pressure at boundary particles. Each boundary particle has a unique pressure value. In contrast to the SPH extrapolation, our MLS approach does not suffer from the particle deficiency issue.

### 4. Implementation

We have combined our proposed boundary handling with a slightly modified DFSPH solver [BK17] that is outlined in Algorithm 1. We follow the idea of combining two solvers, one for the velocity divergence and one for the density invariance. As a minor notation change to DFSPH, our two solvers compute pressure $p$ instead of stiffness parameter $\kappa$. DFSPH implicitly introduces $\kappa$ with the relation $\nabla p_f = \sum_f m_f k_f \nabla W_{ff}$, with $f$ denoting a fluid or a boundary neighbor of $f$. From the SPH formulation $\nabla p_f = \sum_f \frac{m_f}{\rho_f} \nabla W_{ff}$, it follows that $p_f = k_f \rho_f$. So, we use the DFSPH solver, but multiply all $\kappa$ values with the density $\rho$ to get pressure values.

The functions CORRECTDIVERGENCEERROR and CORRECTDENSITYERROR in Algorithm 1 compute pressure at fluid particles, but they are also responsible for the pressure computation at boundary particles. In case of pressure mirroring, the pressure is not explicitly computed, but just considered in the computation of $a^*$ and $a^{**}$ by using Eq. (2) instead of Eq. (1). The DFSPH extrapolation and our proposed MLS extrapolation loop over the boundary particles to compute and store pressure. Then, Eq. (1) is used to compute the pressure accelerations $a^*$ and $a^{**}$. Finally, we update the pressure values $p_f$ of fluid particles $f$ in a Jacobi step. In our experiments, we set the relaxation coefficient $\omega = 0.5$.

![Algorithm 1 DFSPH with MLS pressure extrapolation](image)

```
procedure PERFORM SIMULATION
    for each fluid particle $f$
        find neighborhoods $N_f(t)$
    for each fluid particle $f$
        compute density $\rho_f(t)$
        compute factor $\alpha_f \leftarrow \left| \sum_{\ell} m_{f,\ell} \nabla W_{f\ell} \right|^2 \sum_{\ell} m_{f,\ell} \left| \nabla W_{f\ell} \right|^2 - \left( \rho_f(t) \right)^2$ 
        $a^* \leftarrow$ CORRECT DIVERGENCE ERROR $\triangleright$ divergence-free
        predict velocity $v_f^* \leftarrow v_f(t) + \Delta t a^*_f$
        for each fluid particle $f$
            do
                predict velocity $v_f^{**} \leftarrow v_f^* + \Delta t a^{**}_f$
                update position $x_f(t + \Delta t) \leftarrow x_f(t) + \Delta t v_f(t + \Delta t)$
            done
        return $a^*$

procedure CORRECT DENSITY ERROR
    for each fluid particle $f$
        compute source term $s_f \leftarrow -\frac{1}{\Delta t} \nabla \cdot v_f(t)$
        initialize pressure $p_f \leftarrow 0$
        while not converged do
            for each boundary particle $b$
                compute pressure $p_b$ using MLS $\triangleright$ Eq. (15)
            for each fluid particle $f$
                compute pressure acceleration $a^*_f$ $\triangleright$ Eq. (1)
            for each fluid particle $f$
                set pressure $p_f \leftarrow p_f + \frac{\alpha}{\Delta t} \left( s_f - \nabla \cdot a^*_f \right)$
            return $a^{**}$

procedure CORRECT DENSITY ERROR
    for each fluid particle $f$
        compute source term $s_f \leftarrow -\frac{1}{\Delta t} \nabla \cdot v_f^*(t)$
        initialize pressure $p_f \leftarrow 0$
        while not converged do
            for each boundary particle $b$
                compute pressure $p_b$ using MLS $\triangleright$ Eq. (15)
            for each fluid particle $f$
                compute pressure acceleration $a^{**}_f$ $\triangleright$ Eq. (1)
            for each fluid particle $f$
                set pressure $p_f \leftarrow \max\left( 0, p_f + \frac{\alpha}{\Delta t} \left( s_f - \nabla \cdot a^{**}_f \right) \right)$
            return $a^{**}$

```
which causes an incorrect movement of the fluid particles. In contrast with a support of two times the particle spacing. The rest density of Threading Building Blocks \[ \text{Phe08} \]. We use \[ \text{FIF18} \] to reconstruct To reduce the loss in turbulent details, we use a micropolar material viscosity at the boundary. The computation time for the pressure 5.1. Rotating Sphere a radius of 3 m and is rotating slowly at7 revolutions per minute. We have a particle spacing of 5 cm. The scenario consists of 417 k fluid and 445 k boundary particles and is simulated for ten seconds with a fixed time step of 1 ms. The number of density invariant iterations of our DFSPH solver was fixed to ten while the divergence-free error is 0.037 %. In this experiment, the boundary particles should not influence the fluid velocities and the fluid should rest inside the sphere. However, as shown in Fig. 4, this is not the case for \[ \text{AIA}^* \text{A12} \] and \[ \text{AHA12} \]. Both boundary handling schemes introduce an artificial viscosity, which causes an incorrect movement of the fluid particles. In contrast to this, our MLS pressure extrapolation does not suffer from artificial viscosity at the boundary. The computation time for the pressure field is very similar for all approaches (our: 64.29 ms, \[ \text{AIA}^* \text{A12} \]: 63.12 ms, \[ \text{AHA12} \]: 63.44 ms).

5.2. Breaking Dam In order to compare the solver iteration counts of our approach with \[ \text{AIA}^* \text{A12} \] and \[ \text{AHA12} \], we simulate a breaking dam scenario inside a cylindric-shaped domain of size 3 m × 3 m × 0.5 m with a particle spacing of 8 mm. Thus, making a total of 2.95 million fluid and 182.2 k boundary particles. The scenario is illustrated in Fig. 5. Furthermore, we use different fixed time step sizes. Table 1 summarizes the iteration measurements and computation times for a simulation over ten seconds.

In our experiment, our MLS pressure extrapolation approach always requires the minimum number of iterations per simulation step. For larger time step sizes, our approach outperforms \[ \text{AHA12} \] due to more accurate pressure gradients at the boundary. Computing unique pressure values for boundary particles is not expensive. For \[ \text{AHA12} \], we measured an average computation time for the boundary pressures of 1.12 ms per iteration. For our approach, as we have to iterate twice over fluid neighbors of the boundary particles, the computation time slightly increased to 1.87 ms.

5.3. Vase As shown in \[ \text{BGI}^* \text{18} \], computing unique pressure values for boundary particles can be beneficial for scenarios with a high water depth. In order to show that our approach can also handle such challenging scenarios, we simulate a vase of height 10 m that is filled with water over a duration of eighteen seconds. The scene is illustrated in Fig. 6. The particle spacing is 2 cm and the adaptive time step \[ \text{IAGT10} \] is 0.47 ms on average. The scene consists of up to 13.33 million fluid and 826 k boundary particles. The total computation time per simulation step is 3.70 s on average with MLS pressure extrapolation, 3.72 s for pressure mirroring and 3.84 s for SPH pressure extrapolation. The reduced computation time of our approach is the result of a reduced solver iteration count.

5.4. Teacup In order to demonstrate the applicability of our approach to two-way coupled dynamic objects, we integrated the Bullet physics
Table 1: Comparison of the different boundary handling schemes using different time steps for the Breaking Dam scenario.

<table>
<thead>
<tr>
<th>$\Delta t$</th>
<th>Mirroring</th>
<th>SPH extrapolation</th>
<th>MLS extrapolation</th>
<th>Mirroring</th>
<th>SPH extrapolation</th>
<th>MLS extrapolation</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.25 ms</td>
<td>4.0</td>
<td>4.0</td>
<td>4.0</td>
<td>116 ms</td>
<td>120 ms</td>
<td>123 ms</td>
</tr>
<tr>
<td>0.50 ms</td>
<td>5.7</td>
<td>5.7</td>
<td>5.6</td>
<td>186 ms</td>
<td>192 ms</td>
<td>192 ms</td>
</tr>
<tr>
<td>1.00 ms</td>
<td>12.0</td>
<td>12.2</td>
<td>11.8</td>
<td>448 ms</td>
<td>467 ms</td>
<td>462 ms</td>
</tr>
<tr>
<td>1.50 ms</td>
<td>14.9</td>
<td>16.0</td>
<td>14.9</td>
<td>559 ms</td>
<td>618 ms</td>
<td>587 ms</td>
</tr>
</tbody>
</table>

5.5. Washing Machine

Our proposed method is particularly appropriate for fast-moving and complex boundaries. This is indicated in Fig. 7 where we simulate a washing machine that contains seven two-way coupled rigid spheres with different radii. The washing drum is animated and contains holes, i.e. the fluid drains. The particle spacing is 2 cm and the scene consists of up to 2.04 million fluid particles and 1.18 million boundary particles. We use an adaptive time step with an average of 0.5 ms. Overall, with our approach the average computation time is 409 ms per simulation step whereof computing the boundary pressures takes 17.6 ms. The average iteration count is 4.42. Due to instabilities at the fast-moving boundary, pressure mirroring requires a time step that is half as large compared to our MLS extrapolation. This results in a speed-up of factor 1.8 compared to Akinci et al. [AIA*12]. This speed-up factor is particularly remarkable considering the fact that [AIA*12] typically works for time steps that correspond to rather large CFL numbers.

6. Conclusion and Future Work

MLS pressure extrapolation at boundaries reduces artifacts at fluid-solid interfaces which can improve the performance of the pressure computation in iterative solvers. We have shown that pressure mirroring and SPH extrapolation suffer from artificial viscosity which is not the case for the proposed MLS extrapolation. We have also shown that the reduced velocity artifacts in our boundary handling can positively influence the pressure computation time for challenging scenarios. As one of the next steps, we plan to investigate properties of the MLS gradient estimation for other purposes, e.g. the computation of the pressure acceleration at fluid particles.
Acknowledgments

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Figure 7: Animated washing machine with two-way coupled rigid spheres.

References


