Fast Algorithms for Intersection of Non-matching Grids Using Plücker Coordinates.

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Abstract

The XFEM and Mortar methods can be used in combination with non-matching or non-conforming grids to deal with problems on complex geometries. However the information about the mesh intersection must be provided. We present algorithms for intersections between 1d and 2d unstructured multi component meshes and their intersections with a background unstructured 3d mesh. A common algorithm based on the advancing front technique is used for the efficient selection of candidate pairs among simplicial elements. Bounding interval hierarchy (BIH) of axes aligned bounding boxes (AABB) of elements is used to initialize the front tracking algorithm. The family of element intersection algorithms is built upon a line-triangle intersection algorithm based on the Plücker coordinates. These algorithms combined with the advancing front technique can reuse the results of calculations performed on the neighbouring elements and reduce the number of arithmetic operations. Barycentric coordinates on each of the intersecting elements are provided for every intersection point. Benchmarks of the element intersection algorithms are presented and three variants of the global intersection algorithm are compared on the meshes raising from a hydrogeological application.

Keywords: non-matching grid, mesh intersections, mixed-dimensional mesh, Plücker coordinates, advancing front method

1. Introduction

The grid intersection algorithms are crucial for several techniques that try to overcome some limitations of the classical finite element method. The Chimera method [1], also called overset grid, and similar Nitche method [2] allow solution of the problems with changing geometry as in the fluid-structure problems. The Mortar method [3] allows domain decomposition, independent meshing of domains, and supports sliding boundaries. However our primal motivation is

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usage of XFEM methods [4] and non-matching meshes of mixed dimension in groundwater models.

The realistic models of groundwater processes including the transport processes and geomechanics have to deal with a complex nature of geological formations including fractures and wells. Although of a small scale, these features may have significant impact on the global behavior of the system and their representation in the numerical model is imperative. One possible approach is to model fractures and wells as lower dimensional objects and introduce their coupling with the surrounding continuum. The discretization then leads to the meshes of mixed dimensions, i.e. composed of elements of different dimension. This approach called mixed-dimensional analysis in the mechanics [5] is also studied in the groundwater context, see e.g. [6], [7], [8] and already adopted by some groundwater simulation software, e.g FeFlow [9] and Flow123d [10]. Nevertheless as the complexity of the geometry increase (e.g. when lots of fractures are randomly generated) the compatible meshing becomes painful or even impossible. In order to avoid these difficulties we may discretize the continuum and every fracture and well independently, getting a non-matching (or incompatible) mesh of mixed dimensions and then apply XFEM to represent jumps of the solution on the fractures or singularities around the wells. The prerequisite for such approach is a fast and robust algorithm for calculating intersections of individual meshes.

We consider a composed mesh \mathcal{T} consisting of simplicial meshes \mathcal{T}_i of dimensions $d_i \in \{1, 2, 3\}$, $i = 1, \ldots, N_{\mathcal{T}}$ in the 3d ambient space. We assume that every mesh \mathcal{T}_i is a connected set with no self intersection. Further we assume only single 3d mesh \mathcal{T}_1 . The mesh intersection problem is to find all pairs of elements $L \in \mathcal{T}_i$, $K \in \mathcal{T}_j$, $i \neq j$ that have non-empty intersection and to compute that intersection. The mesh intersection problem consists of the two parts: First, generate a set of candidate pairs (K, L). Second, compute the intersection of a particular pair.

According to our knowledge, there are lots of works using non-matching grids, yet only few of them discuss algorithms how to compute their intersections. Gander and Japhet [11] present the PANG algorithm for 2d-2d and 3d-3d intersections that can be used e.g. for mesh overlapping methods. They use the advancing front technique to get candidate pairs in linear time. The algorithm is part of the DUNE library [12]. Massing, Larson, and Logg [2] present an algorithm for 2d-3d intersections as part of their implementation of the Nitche method which is part of the Doplhin project [13]. They use axes aligned bounding boxes of elements (AABB) and bounding interval hierarchy (BIH) to get intersection candidate pairs of elements, the GTS library [14] is used for 2d-3d intersections. Finally, there is the work of A. Elsheikh and M. Elsheikh [15] presenting an algorithm for 2d-2d mesh union operation which includes calculation and imprinting of the intersection curves. They exploit binary space partitioning for search of initial intersection and the advancing front method for intersection curve tracking.

In this paper we present a new approach to intersection calculation of simplicial elements of different dimensions based on the Plücker coordinates, further developing the algorithm of Platis and Theoharis [16] for ray-tetrahedron intersections. Element intersections based on Plücker coordinates are combined with the advancing front method which allows us to reuse Plücker coordinates and their products between neighbouring elements and reduce the number of arithmetic operations.

The paper is organized as follows. In Section 2 the algorithms for 1d-2d, 1d-3d and 2d-3d intersections of simplices are described. In Section 3 we discuss our implementation of the advancing front technique and usage of AABB and BIH for its initialization. Finally, in Section 4, we present benchmarks and comparison of individual algorithms.

2. Element Intersections

In this section, we present algorithms for computing intersection of a pair of simplicial elements of a different dimension in the 3d ambient space. In particular we address intersection algorithms for 1d-2d, 1d-3d, 2d-3d pairs of elements. We have implemented the case 2d-2d as well, however the treatment of the special cases is quite technical and not fully completed yet. The fundamental idea is to compute intersection of 1d-2d simplices using the Plücker coordinates and reduce all other cases to this one.

We denote S_i a simplicial element with i+1 vertices (of dimension i). We call vertices, edges, faces and simplices itself the n-faces and we denote M_i the set of all n-faces of the simplex S_i . In general, an intersection can be a point, a line segment or a polygon called *intersection polygon* (IP) in common. The intersection polygon is represented as a list of its corners called *intersection corners* (IC). The IP data structure keeps also reference to the intersecting simplices. A data structure of a single IC consists of:

- the barycentric coordinate w_K of IC on K,
- the dimension d_K of the most specific n-face the IC lies on,
- the local index i_K of that n-face on K,

for each intersecting element K of the pair. The pair $\tau_K = (d_K, i_K)$ is called the topological position of the IC on K.

2.1. Plücker Coordinates

Plücker coordinates represent a line in 3d space. Considering a line p, given by a point \boldsymbol{A} and its directional vector \boldsymbol{u} , the Plücker coordinates of p are defined as

$$\pi_p = (\boldsymbol{u}_p, \boldsymbol{v}_p) = (\boldsymbol{u}_p, \boldsymbol{u}_p \times \boldsymbol{A}).$$

Further we use a permuted inner product

$$\pi_p \odot \pi_q = \boldsymbol{u}_p \cdot \boldsymbol{v}_q + \boldsymbol{u}_q \cdot \boldsymbol{v}_p.$$

The sign of the permuted inner product gives us the relative position of the two lines, see Figure 1.

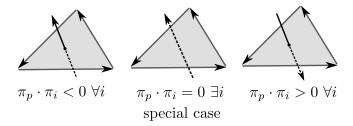


Figure 1: Sign of the permuted inner product is related to the relative position of the two oriented lines. Dashed line symbolizes that the line is in the back, the lines intersect in the middle case. **PE**: circle dot permuted inner product

2.2. Intersection Line-Triangle (1d-2d)

PE: I would stick to chosen notation - triangle S_2 , and possibly renamed side s_i ... Let us consider a line segment p with parametric equation

$$X = A + tu, \ t \in (0,1) \tag{1}$$

and a triangle T given by vertices $(\boldsymbol{V}_0, \boldsymbol{V}_1, \boldsymbol{V}_2)$ with oriented sides $s_i = (\boldsymbol{V}_j, \boldsymbol{V}_k)$, $j = (i+1) \mod 3$, $k = (i+2) \mod 3$.

Lemma 2.1. The permuted inner products $\pi_p \odot \pi_{s_i}$, i = 0, 1, 2 have the same non-zero sign if and only if there is an intersection point X on the p and inside the triangle T. The barycentric coordinates of X on T are

$$w_i = \frac{\pi_p \odot \pi_{s_i}}{\sum_{i=0}^2 \pi_p \odot \pi_{s_i}}.$$
 (2)

Proof. Using the barycentric coordinates the intersection point can be expressed as $\mathbf{X} = \mathbf{V}_0 + w_1 \mathbf{s}_2 - w_2 \mathbf{s}_1$. The line p has Plücker coordinates $(\mathbf{u}, \mathbf{u} \times \mathbf{X})$ since these are invariant to change of the initial point. Combining these two expressions we get

$$\pi_p \odot \pi_{s_1} = u \cdot (s_1 \times V_2) + s_1 \cdot (u \times [V_0 + w_1 s_2 - w_2 s_1]) = -w_1 u \cdot (s_1 \times s_2).$$

Since $s_0 + s_1 + s_2 = 0$ we have $s_1 \times s_2 = s_2 \times s_0 = s_0 \times s_1$ and thus

$$\pi_p \odot \pi_{s_i} = -w_i \boldsymbol{u} \cdot (\boldsymbol{v}_1 \times \boldsymbol{v}_2).$$

The point X is inside of T if and only if $w_i > 0$ for all i = 0, 1, 2.

Having the barycentric coordinates of X on T, we can compute also its local coordinate on p from its parametric form:

$$X_i = A_i + tu_i$$
, for $i = 1, 2, 3$ (3)

We use i with maximal $|u_i|$ for practical computation.

The calculation of the intersection proceeds as follows:

- 1. Compute or reuse Plücker coordinates and permuted inner products: π_p , π_i , $\pi_s \odot \pi_i$, for i = 1, 2, 3.
- 2. Compute barycentric coordinates w_i , i = 1, 2, 3 using (2).
- 3. If any w_i is less than ϵ , there is no intersection, return empty IP.
- 4. If all w_i are greater than ϵ , we set $\tau_T = (2,0)$ for the IC.
- 5. If one w_i is less than ϵ , intersection is on the edge s_i , we set $\tau_T = (1, i)$.
- 6. If two w_i are less than ϵ , intersection is at the vertex V_i , we set $\tau_T = (0, i)$.
- 7. If all w_i are less than ϵ , the line is coplanar with the triangle, both objects are projected to the plane $x_i = 0$ where i is the index of maximal component of the triangle's normal vector. Every pair p, s_i is checked for an intersection on T boundary either inside s_i or at a vertex V_i , setting the topological info τ_T to (1,i) or (0,i), respectively. At most two ICs are obtained.
- 8. For each IC the barycentric coordinates (1-t,t) on the line p are computed according to (3).
- 9. If $t \in (-\epsilon, \epsilon)$ or $t \in (1 \epsilon, 1 + \epsilon)$, we set $\tau_p = (0, 0)$ or $\tau_p = (0, 1)$, respectively.
- 10. If $t \notin (-\epsilon, 1+\epsilon)$, the IC is eliminated.

We shall call every IC, that was created in the coplanar case, degenerate since these will later special treatment.

Checking the same sign of the inner products can be viewed as a geometric predicate for the presence of the intersection and orientation of the line with respect to the triangle. Adaptive-precision evaluation of the geometric predicates was designed by Schewchuk [17] and used for 2d-2d mesh intersections in [15]. However, we rather apply a fixed tolerance check for the zero barycentric coordinates and consistently keep the topological positions in this and related algorithms. JB: Can we make the algorithm parsimonious in the spirit of the Fortune [18] quoted by Schewchuk? Seems that our problem is more local than the line example that was proven to be NP-hard.

The algorithms for 1d-3d and 2d-3d intersections use simpler version of the 1d-2d intersection algorithm, in particular the search for ICs in the coplanar case (item 7) is not necessary, and the test in the last point is not performed.

2.3. Intersection Line-Tetrahedron (1d-3d)

PE: degenerate not defined yet.. do you call it coplanar now? In this section we consider an intersection of a line segment p given by the parametric equation (1) with a tetrahedron S_3 . The used algorithm is based on the 1d-2d algorithm and closely follows [16]. Our modification takes into account intersection with the line segment and consistently propagates topological position of ICs.

Algorithm 1 first computes line-face intersections for every face of S_3 . Tetrahedron has six edges, so 7 Plücker coordinates and 6 permuted inner products are computed at most. Precomputed coordinates and products are passed into the 1d-2d algorithm which is performed for the whole line p (line 3). If no IC is found, or coplanar case occurs in line-face computation, we continue to the

Algorithm 1: 1d-3d intersection

```
Input: Tetrahedron S_3, line segment p.
   Output: List I of ICs sorted along p.
 1 I = \{\}
 2 for unmarked face f of S_3 do
      L = intersection(p, f)
 3
      if L is none or degenerate then continue
 4
      if L is inside the edge e then
 5
          set \tau_{S_3} = (1, e)
 6
          mark faces coincident with e
 7
      else if L is at the vertex v then
 8
          set \tau_{S_3} = (0, v)
 9
          mark faces coincident with v
10
      append L to I
11
      if |I| = 2 then break
13 if |I| = 1 and I is outside of p then erase I
  else if |I| = 2 then
      trim intersection with respect to the line segment p
15
```

next face. Note, ICs that would be created in coplanar case are to be found as ICs with the other faces, since they lie on edges. Next, IC can be on an edge or at a vertex; then we set the correct topological position and mark the adjacent faces to be skipped, since there cannot be another IC (and coplanar case has been checked already). Finally at the line 11, we append the IC to the result and check whether the maximal amount of ICs has been reached.

After collecting line-tetrahedron ICs, we do the line segment trimming from the line 13 further. If we have only one IC, we check that it actually lies inside the line segment (otherwise, we throw it away). If we have two ICs, and if both lie outside the line segment p, we eliminate both of them. If one of the ICs lies out of p, we use the closest end point of the line segment instead and interpolate barycentric coordinates of the IC on S_3 . The topological positions are updated as well. The result of the algorithm is 0, 1, or 2 ICs, sorted by the parameter t in the direction of the line p.

2.4. Intersection Triangle-Tetrahedron (2d-3d)

The intersection of a triangle S_2 and a tetrahedron S_3 is an n-side intersection polygon (IP), $n \leq 7$. The sides of the polygon lie either on sides of S_2 or on faces of S_3 . Thus each vertex (IC) of the polygon can arise either from side-face intersection, or from edge-triangle intersection, or be a vertex of S_2 . To get all ICs, we have to compute at most 12 side-face intersections and at most 6 edge-triangle intersections. However, to this end we only need to compute 9 Plücker coordinates (3 sides, 6 edges) and 18 permuted inner products, one for every

side-edge pair. Computation of IP consists of three stages: calculation of side-tetrahedron ICs (Section 2.4.1), calculation of edge-triangle ICs (Section 2.4.2), ordering of ICs (Section 2.4.4). The intersection corners are appended to the list I as they are computed, however their order on the polygon boundary is defined by the connection tables $F_g(:)$ and $F_p(:)$. Every side of the polygon that lies on n-face $x \in M_2 \cup M_3$ is followed by an IC given by $F_g[x]$ and every IC p is followed by the side that lies on $F_p[p] \in M_2 \cup M_3$ (see JB: possible figure). The vertices of the polygon are ordered in the same way as is the order and orientation of the sides of S_2 , that is counterclockwise around the interior with normal pointing to us. There are special cases when the IP is degenerated to a line or a point. PE: next sentence: how should I understand 'may possibly be'; 'an IC' instead of 'some IC' If some IC may possibly be part of the such degenerated IP we put it into the list J.

2.4.1. Intersections on sides of S_2

```
Algorithm 2: 2d-3d intersection, ICs on sides of S_2
   Input: input data
   Output: List of ICs on sorted output data
 1 F_g(:) = -1, F_p(:) = -1
                                                                // Unset links.
 2 for side \ s \ of \ S_2 do
       L = intersection(s, S_3)
       if |L| = 0 then continue
 4
 5
       if |L| = 1 then append p to J continue
       for p in L do
 6
 7
          p lies on x \in M_3
          if p lies at the vertex v of S_3 then x = v
 8
           if p is the first in L then
 9
              F_q[x] = p, F_p[p] = s
10
           else p is the last in L
11
              F_g[s] = p, \ F_p[p] = x
if F_g[x] = -1 then F_g[x] = p
12
13
```

Algorithm 2 computes all ICs on the boundary of S_2 . It passes through every side s of the triangle S_2 and computes the line-tetrahedron intersection L. If the side is just touching S_3 , we get L with a single IC. These ICs will be rediscovered again in Algorithm 3 with better topological information, however this is not the case if the touched edge e of S_3 is coplanar with S_2 and the IC is inside of e. To this end we save the IC into separate list J and skip filling of the connection tables. **PE:** here it is weird... there is no line appending the IC into resulting list.. Which is the resulting list?? I find the reference to 'line:one side loop' also incorrect.. In the regular case, we process each of the two ICs in L (loop on line 2). The IC p is added to the list I, vertices of S_2 added twice are merged at final stage. Then (line 7) we identify the n-face $x \in M_3$ the point

p lies on and set the tables F_g , F_p . For the IC at the vertex of S_2 we set x to that vertex regardless of its position on S_3 . The backward temporary link on the line 13 is used later in Algorithm 3 to allow proper continuation of the IC if it lies on edge or at vertex of S_3 . The condition at the same line deals with the case that two sides of S_3 intersect the same object $x \in M_3$, in such case the backlink is unnecessary and would overwrite the correct link set by previous IC.

2.4.2. Intersections on edges of S_3

```
Algorithm 3: 2d-3d intersection, ICs on edges of S_3
   Input: I with ICs on S_2 boundary, partially filled F
   Output: all ICs in I, complete F
 1 for edge e of S_3 do L[e] = intersection(e, S_2)
 2 for edge e of S_3 with non-empty L[e] do
 3
      p = L[e]
      if p is inside e then
 4
          (f_0, f_1) = \mathbf{edge} \ \mathbf{faces} \ (e)
 5
      else p at the vertex v of S_3
 6
 7
       (f_0, f_1) = vertex faces (v,L)
                                                                // Algorithm 4
      if p is on the boundary of S_2 then
 8
          p lies on edge or at vertex x \in M_3
 9
                                                // q is already computed p
           q = F_q[x]
10
          if F_p[q] = x then F_p[q] = f_1
11
          else F_g[f_0] = q
12
          F_q[x] = -1
                                                     // remove the backlink
13
      else
14
          append p to I
15
          F_{q}[f_{0}] = p, \ F_{p}[p] = f_{1}
                                                 // overwrite the backlink
16
```

Algorithm 3 uses the line-triangle intersection algorithm for the edges of S_3 (line 1). First, the intersection L[e] is evaluated for every edge e. The loop produces ICs in the interior of S_2 and possibly those ICs with special position at the vertex or on the edge of S_3 already computed in Algorithm 2. Then we pass through once again skipping the edges with none or degenerate L[e]. For every IC p = L[e] we first get its (generalized) faces that would appear before and after the IC on IP, assuming that p is inside S_2 .

The function edge faces returns the adjacent faces f_0 , f_1 to the edge e, on which the IC lies (see the situation in Figure 2). The faces are sorted using the sign of the permuted inner product in 1d-2d intersection. The order of faces matches the order of sides of IP if the sign is negative. If the sign is positive the function edge faces returns face pair (f_1, f_0) . If the IC is at the vertex v of S_3 , the function vertex faces described later (Algorithm 4) is used. It returns a pair of generalized faces (face or edge) adjacent to the IC L[e] at the vertex v of S_3 . If IC p is inside S_2 , we add it into I and set the connection tables (line

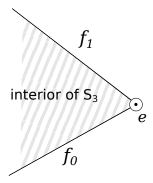


Figure 2: Order of faces adjacent to the oriented edge e pointing up.

16). However, if p is on boundary of S_2 it is already in I. Denoting x the n-face of S_3 the IC lies on, we use the back-link from x to IC $q \in I$ and if q points back to x we set the new successor f_1 to it (line 11). Otherwise f_0 is used as the predecessor of q (line 12).

2.4.3. Vertex Faces Algorithm

```
Algorithm 4: 2d-3d intersection, vertex faces
   Input: vertex v of S_3, L[:] intersection results for edges of S_3
   Output: (x_1, x_2), x_1, x_2 \in M_3, coincident with v and intersected by the
             plane of S_2
 1 e_0, e_1, e_2 edges coincident with v oriented out of v s[i] = L[e_i], for
    i = 0, 1, 2,
 2 if s[:] have 1 non-degenerate edge e then
     return pair of degenerate edges sorted according to edge faces (e)
 4 else if s have 1 degenerate edge e then
       f is face opposite to e
 5
      if other two edges e_a, e_b have different sign then
 6
          z = \mathbf{edge} \ \mathbf{faces}(e_a)
 7
          replace g \in z, g \neq f with e, return z
      else append IC of v to J, return anything
10 else if s have edge e with sign opposite to other two then
      return edge faces(e)
  else s have all signs same
      append IC of v to J, return anything
13
```

This function gets as a parameter IC p at the vertex v of S_3 which is a special case of anon-degenerate edge-triangle intersection. We use signs and degenerate indicator of ICs of the three edges coincident with v to return generalized faces

of S_3 preceding and succeeding p on the polygons boundary assuming p is at interior of S_2 . Possible cases are:

- Single degenerated IC (line 4). Let us denote e the edge with degenerated IC and f the face between the other two edges. The other two (non-degenerates) edges may have either the opposite sign (the plane is cutting S_3) or the same sign (the plane is touching S_3 at the edge e). In the first case, the call of edge faces for e returns (f_x, f) or (f, f_x) , then the vertex faces function returns (e, f) or (f, e), respectively. In the second case, we just return anything since there are at most two ICs in IP so the connection tables are not necessary. However, we add current IC into J.
- Two degenerated ICs (line 2). A face of S_3 lies in the plane of S_2 . Let e be the single non-degenerate edge. We treat the two degenerate edges as faces adjacent to e and return them sorted like the faces given by edge faces of edge e.
- Single IC has the opposite sign to the other two (line 10). Let e be the edge of the single IC with the different sign. The plane of S_2 separates e from the other two edges so it goes through the faces adjacent to e. The order is determined by the function $edge\ faces$ called for the edge e.
- All ICs have the same sign (line 12). Since S_2 is touching S_3 at the vertex v, the polygon degenerates into point and thus no connection information is necessary. We add IS to J and return any pair of faces.

2.4.4. Ordering of intersections

The final stage of the 2d-3d intersection is ordering of ICs. If there are less then 3 ICs in the list I we merge lists I and J remove duplicities and get at most 2 distinguish ICs forming a degenerate IP. Otherwise, we start with first IC p in I, put it into final list K, get its successor n-face $x = F_p(p)$ and its successor IC $q = F_q(x)$. Unless x is vertex of S_2 , we put q into K and repeat the process until we pass all ICs in I.

3. Global mesh intersection algorithm

Having the algorithms for element-element intersections we can proceed to the mesh intersection algorithm. We first compute all component-bulk mesh intersections that is (1d-3d and 2d-3d) and then the 1d-2d and 2d-2d intersections use the 3d mesh to get the intersection pair candidates. Element intersections for the particular component and bulk mesh are obtained in two phases: firstly, we look for the first two elements intersecting each other (initialization); secondly, we prolong the intersection by investigating neighbouring elements (front tracking). JB: Not sure that "prolong" is the right verb.

Initialization. We choose an arbitrary element of the component mesh and check it intersection with all elements of the bulk mesh. The process is accelerated by computing the axis aligned bounding box (AABB) for every element and use intersection of the bounding boxes as a fast indicator for possible intersection of the elements. This step takes O(N) time with respect to the number of elements of the bulk mesh N. For the small number of components k and if they are contained inside the bulk mesh the total time of initialization may be still linear O(kn). However, for more complex cases we organize the bounding boxes of the bulk mesh into the bounding interval hierarchy (BIH) [19] a data structure in principle equivalent to R-trees [20], [21]. The creation time is $O(N \log(N/n))$ and the lookup time $O(\log(N/n))$ where n is number of the bulk elements in the leaf nodes of the tree.

Advancing Front Method. The advancing front algorithm requires and employ the neigbouring information for the elements within the component as well as within the bulk mesh. It can be viewed as the breadth first search algorithm for a graph which vertices are intersection polygons and edges are sides shared by two polygons. Side every side of IP is on the boundary of either component or the bulk element we can distinguish bulk and component edges and correspondingly the bulk queue and the component queue.

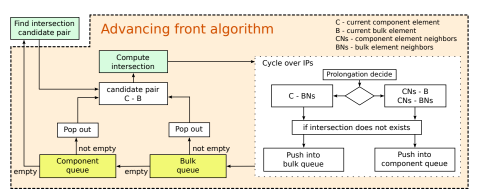


Figure 3: Advancing front algorithm for 1d-2d and 2d-3d intersections.

The scheme of the advancing front algorithm is displayed by the scheme in Figure 3. The main idea is to compute intersections for a component element with all possible bulk elements, and then move to a next neighboring component element. So the bulk queue is emptied before the component queue (see yellow boxes).

On the input we consider a candidate pair provided be the initialization (top-left corner). Now we look for new candidate pairs among the neighboring elements (the block Prolongation decide). Therefore, we iterate over the intersection corners and further exploit their topological position on the component element c and the bulk element b. There are 3 possible cases (applies both for 1d-3d and 2d-3d), how the intersection might be prolonged:

• IC on the boundary of c and inside b

We find all the sides S coincident with the n-face of c on which IC lies. Then we get all component elements C' neighbouring with c over any sides $\in S$. And finally, we push all pairs (c',b), $c' \in C'$ into the component queue. Note, that c can have more then one neighbour component elements over the single side s, i.e. branches are allowed.

\bullet IC on the boundary of both c and b

We find all the faces F coincident with the n-face of b on which IC lies. Then we get all neighbours B' through all faces in F as in the previous case. We push pairs (c, b'), $b' \in B'$ into the bulk queue and pairs (c', b'), $c' \in C'$, $b' \in B'$ into the component queue.

• IC inside of c and on the boundary of b)

We proceed as in previous case, and push pairs (c, b'), $b' \in B'$ into the bulk queue.

JB: Can we just use two (not excluded) cases one for IB on boundary of c second for CI on boundary of b?

We also see that the candidate pairs are of three types: (c, b'), (c', b), (c', b') from which only the first one goes into the bulk queue, trying to cover current component element c before we proceed to the next one.

When the pair is taken from either of the queues we check if the pair was computed and possibly skip it. To this end every component element keeps a hash of the bulk elements with which we have called the intersection test.

4. Benchmarks

JB: We compare just against NGH which we should briefly describe. Neither of the algorithms is optimized. The new one deals correctly with special cases and provides barycentric coordinates (so more work for less time). **JB:** Also write little about Flow123d here. The language etc.

In this section, we present numerical results on several benchmark problems. In the beginning, we provide some software development related information. Then we compare the performance of our algorithms for element intersections with other approaches. Next we shall compare our algorithms with different initialization phase (candidate pairs search), and using the advancing front method or not. We shall show the results both on a mesh of a real locality and an artificial mesh.

4.1. Software Flow123d

The intersection algorithms are implemented as a part of the software Flow123d [10]. The software is developed with an open source licence at the Technical University in Liberec, the implementation is based mostly on C++ language. Flow123d provides models for (un)saturated groundwater flow, transport of solutes possibly coupled with first order reactions and heat transfer in fractured porous

media. The speciality of the software is that it enables computations on complex meshes consisting of simplicial elements of different dimensions. However, non-matching meshes are at the moment available only in experimental 1d-2d flow model, using mortar like coupling.

PE: check NGH.. how to refer to NGH later? Currently, there is also another implementation of the element intersections in the software (we shall refer to it later as NGH). It uses standard means of analytic geometry for computation of intersections line-line and line-plane. This leads to solving a large amount of small linear systems of equations. This algorithm also does not provide us barycentric coordinates and topological position of ICs. At last, the algorithm cannot handle all of the special cases. We must note, that neither the old algorithms nor the new ones are optimised at the moment.

Note, that we use Armadillo [22], a C++ linear algebra library, for linear algebra operations. Most importantly, all coordinates are represented as Armadillo vectors, so we can use dot and cross product and further vector and matrix operations provided by the library.

4.2. Element Intersection Algorithms

The first benchmark focuses on the 1d-3d and 2d-3d element intersections. We randomly generated 100000 element pairs inside a unit cube, from which approximately 65% have nonempty intersection. In some of the empty cases, the bounding boxes did not collide, so the actual intersection algorithms were skipped. A single element pair was computed 100 times to obtain reasonable computational time.

algorithm, 2d-3d	FLOPs
	estimate
parametric plane	585
normal plane (reuse)	765
Moller and Trumbore	783
Plücker (reuse)	306

Table 1: Estimation of floating point operations count and comparison of different approaches. Only 2d-3d case considered.

PE: comment on different strategies **JB:** Rather put into text. I shall comment that.

We see the benchmark results in the Figure 4. Green values correspond to the presented algorithms using Plücker coordinates, red values to NGH code. The gained speed up factor is approximately 5 in 1d-3d case and 2 in 2d-3d. Further we provide estimated count of floating point operations for different approaches in the Table 1. **PE:** *Improve and complete...*

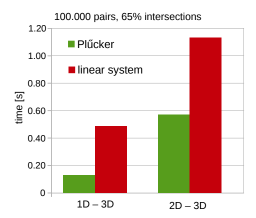


Figure 4: Efficiency of Plücker approach compared to the legacy NGH code.

4.3. General algorithms

In previous sections, we have suggested an intersection algorithm that uses axes aligned bounding boxes, BIH to find initial intersection and advancing front method (shortly AFront) to prolong the intersection. From now on, we shall refer to the algorithm made of these components BIHsearch. In this section, we shall compare two additional algorithms BBsearch and BIHonly to see the effects of BIH and AFront (see Table 2). BBsearch does not use BIH, but searches for the initial intersection using only AABB, and then follows AFront. The version BIHonly computes AABB together with BIH, but does not use AFront.

BIHsearch	BBsearch	BIHonly
BIH(AABB)	AABB	BIH(AABB)
\mathbf{AFront}	AFront	

Table 2: Structure of the general algorithms. 'AFront' stands for the proposed advancing front method. **JB:** Just in the text. Possibly use different name for variants. **PE:** jsem vzal z posledni prezentace.. prislo mi to prehledny.. vim jak jsem se v diplomce VF hrozne krkolome vracel od grafu k popisu algoritmu..

Let us now start with the problem on an artificial mesh.

Next, we study the performance of the intersection algorithms on a mesh of a real problem, see Figure 7. The mesh represents a mountain ridge in the Jizera mountains which includes a system of geological fractures (Figure 7a). We also used this model to create a less real mesh, in which the fractures extend through the bulk surface (Figure 7b). However, this situation might rather occur in another application than approus media problem. In total, 28 fractures are in the mesh.

The results for both meshes can be seen in the Figure 8, pay attention to the different time scales in the graph. In the first case, we notice that algorithms using AFront are nearly twice as fast as *BIHonly*. The speed up corresponds

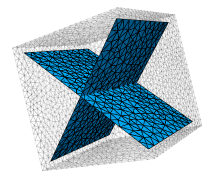


Figure 5: Artificial mesh – a cube with two perpendicular planes placed on the diagonals of the cube. The planes are also nonmatching, therefore can be seen as two independent components.

Dependence on mesh refinement

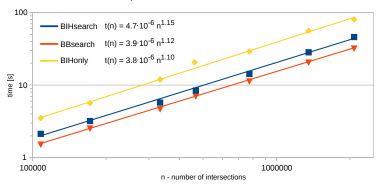


Figure 6: Dependence of the algorithms performance on number of intersection (and mesh refinement).

to the time saved by using AFront instead of searching the whole mesh for candidate pairs. Computation of BIH in BIHsearch pays off and the algorithm performs better than BBsearch.

In the second case, we observe large blow up for *BBsearch*. It is caused by the exterior elements, for which all bulk elements bounding boxes are iterated, so we can be sure there is no intersection. This of course is much better performed using BIH.

In contrast to the artificial case on the fractured cube, BIH construction is worth due to the presence of more components. Having more fractures inside the bulk would result to more dramatic difference.

5. Conclusions

TODO: - line intersection tracking for accelerate 2d-2d intersections - better handling of special cases in paarticular in relation to prolongations - better

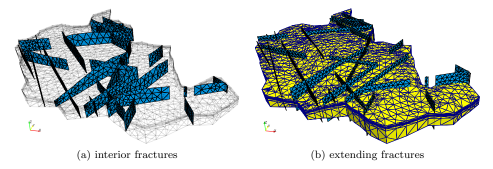


Figure 7: A mesh of the real locality of Bedřichov in the Jizera mountains. We see fractures inside the bulk mesh in the left figure, fractures are extending the bulk mesh.

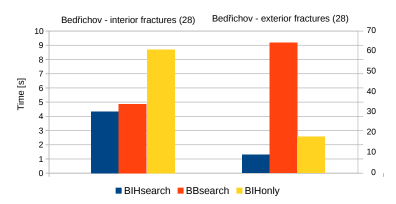


Figure 8: Comparison of the algorithms on meshes of Bedřichov locality – interior fractures on the left, extending fractures on the right.

calculation reuse (pass with prolongations) - optimisation of element intersection - skip unnecessary calculations

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