# The PSurface Library 

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#### Abstract

We describe PSurface, a C++ library that allows to store and access piecewise linear mappings between simplicial surfaces in $\mathbb{R}^{2}$ and $\mathbb{R}^{3}$. These mappings are stored in a graph data structure and can be constructed explicitly, by projection, or by surface simplification. Piecewise linear maps can be used, e.g., to construct boundary approximations for finite element grids, and grid intersections for domain decomposition methods. In computer graphics the mappings allow to build level-of-detail representations as well as texture- and bump maps. We document the data structures and algorithms used and show how PSurface is used in the numerical analysis framework Dune and the visualization software Amira.


## 1 Introduction

Let $S_{1}$ and $S_{2}$ be $d$-dimensional simplicial surfaces in $\mathbb{R}^{d+1}$, where $d \in\{1,2\}$. A map $\Xi: D \subseteq S_{1} \rightarrow S_{2}$ will be called piecewise linear, if for each pair of simplices $T_{1} \in S_{1}$ and $T_{2} \in S_{2}$ the restriction of $\Xi$ to $T_{1} \cap \Xi^{-1}\left(T_{2}\right)$ is an affine function. We assume that all maps we consider are continuous, injective, and such that $\Xi\left(S_{1}\right)$ is a simplicial subsurface of $S_{2}$. However, we do not require that simplices of $S_{1}$ be mapped onto simplices of $S_{2}$. If, additionally, $S_{1}$ and $S_{2}$ are homeomorphic and $\Xi: S_{1} \rightarrow S_{2}$ is a homeomorphism, we call $\Xi$ a piecewise linear parametrization of $S_{2}$ over $S_{1}$.

It is occasionally useful to be able to handle such piecewise linear maps in a computer algorithm. In the numerical simulation of two-body contact problems in linear elasticity, e.g., regions of potential contact on the boundaries of the objects need to be identified with a homeomorphism. This homeomorphism needs to be evaluated repeatedly in the course of the simulation [19]. Also, when a partial differential equation is to be solved on a domain with a curved or highly detailed boundary, its approximation by a grid may discard important geometric information. To make this information available at the time of grid refinement, a parametrization of the domain boundary over the grid boundary is necessary. In computer graphics, parametrizations can be used, e.g., to generate textureand bump maps [6, 17]. Atlas-based methods for the automatic segmentation of medical image data use parametrizations to make the different models in an atlas comparable 15].

[^0]

Figure 1: Piecewise linear map from a subset $D$ of a one-dimensional simplicial surface $S_{1}$ to a second simplicial surface $S_{2}$.

PSURFACE is a $\mathrm{C}++$ library to handle continuous piecewise linear maps between simplicial surfaces. Its core is a data structure that can hold the simplicial surfaces $S_{1}, S_{2}$ and a piecewise linear map $\Xi: D \subset S_{1} \rightarrow S_{2}$. The surfaces can be one- or two-dimensional and are expected to be embedded in a Euclidean space of one dimension higher. $S_{1}$ and $S_{2}$ are not assumed to be manifolds but can, e.g., contain edges with more than two adjacent triangles. The mapping $\Xi$ itself is stored as a graph $\mathcal{G}$ on the simplices of $S_{1}$. This graph is the image of the edges of $\Xi\left(S_{1}\right) \subset S_{2}$ under $\Xi^{-1}$. If $d=2$ we obtain a planar graph on each two-dimensional simplex of $S_{1}$. If $d=1$ the graph consists of nodes along the one-dimensional simplices of $S_{1}$ and graph edges between them (Figure 1). For any point $x \in S_{1}$, the image $\Xi(x) \in S_{2}$ can then be evaluated by a point-location in $\mathcal{G}$ and subsequent linear interpolation.

Several ways of constructing mappings $\Xi$ have been implemented. The most simple way is by direct construction. A special factory class allows to set up maps by explicitly prescribing how vertices of $S_{1}$ are mapped onto $S_{2}$, what points of $S_{1}$ are mapped onto the vertices of $S_{2}$, and how the images of the edges of $S_{1}$ cross edges of $S_{2}$. Given two surfaces, a mapping can also be generated by projecting $S_{1}$ onto $S_{2}$ in the direction of a given vector field on $S_{1}$. Finally, a parametrization of a given fine surface $S_{2}$ over a coarser surface $S_{1}$ can be constructed by initially setting $S_{1}=S_{2}, \Xi=\mathrm{Id}$, and successively removing points from $S_{1}$. A valid $\Xi: S_{1} \rightarrow S_{2}$ is kept at each step, and hence this creates the coarse surface $S_{1}$ along with the parametrization function $\Xi$.

To the knowledge of the author, PSURFACE is the only library for the general handling of piecewise linear mappings between simplicial surfaces currently available. Very similar functionality exists in various places, e.g., the code used for atlas-based image segmentation in [15] or the LGM domain manager for piecewise linear boundary descriptions in the UG finite element software [3]. However, these implementations are all hardwired to their respective applications and cannot be used separately.

This article is intended to describe and document the data structures and algorithms in PSurface. Information about PSurface has appeared implicitly in various publications [13, 14, 18, 19], but this is the first document that describes PSURFACE explicitly, exclusively, and exhaustively. It is not a reference manual. As PSurface is still under development the actual method names and signatures can still be subject to change, and mention of them here would outdate quickly. Up-to-date information is available in form of a doxygengenerated documentation distributed with the library. The library itself is


Figure 2: Piecewise linear map from a subset of a sphere $S_{1}$ to another sphere $S_{2}$. The preimages of the edges of $S_{2}$ form a graph on $S_{1}$.
currently available under a free license at http://numerik.mi.fu-berlin.de/ dune/psurface.

## 2 The data structure

This section describes the data structure for continuous piecewise linear maps between simplicial surfaces, which is the heart of PSurface. We concentrate on the case $d=2$, because it is considerably more complicated than $d=1$. Remarks concerning maps between one-dimensional surfaces will be given where appropriate.

### 2.1 Simplicial surfaces

To fix ideas and notation we briefly define what we mean by 'simplicial surface'.
Definition 2.1. A simplicial $k$-complex $\mathcal{K}$ is a set of closed simplices of dimension less than or equal to $k$, that satisfies the following conditions:

1. Any face of a simplex from $\mathcal{K}$ is also in $\mathcal{K}$.
2. The intersection of any two simplices $\sigma_{1}, \sigma_{2} \in \mathcal{K}$ is a face of both $\sigma_{1}$ and $\sigma_{2}$, or empty.

A simplicial $k$-complex $\mathcal{K}$ is called homogeneous if every simplex of dimension less than $k$ is the face of some simplex $\sigma \in \mathcal{K}$ of dimension exactly $k$.

We call $d$-dimensional simplicial surface a homogeneous simplicial $d$-complex in a $d+1$-dimensional Euclidean space. Note that this definition does not restrict the setting to discrete approximations of manifolds. Indeed, situations like three triangles meeting at a common edge are easily handled by PSurface.

Let $S$ be a $d$-dimensional simplicial surface with $d \geq 1$. We call the 0 dimensional simplices of $S$ the vertices, and the 1-dimensional simplices the edges of $S$. The $d$-dimensional simplices are called elements. We denote the set of vertices by $\mathcal{V}$, the set of edges by $\mathcal{E}$, and the set of elements by $\mathcal{T}$.

### 2.2 A data structure for piecewise linear maps

The foundation of the data structure for piecewise linear maps $\Xi: D \subset S_{1} \rightarrow S_{2}$ are the data structures for the two surfaces $S_{1}$ and $S_{2}$. There are separate data structures for each surface. These are conceptually very similar, but not identical, because more functionality is needed for $S_{1}$. Both data structures store an array with the vertex positions and, for each $k, 1 \leq k \leq d$, an array of $k+1$-tuples of vertex indices to store the $k$-dimensional simplices.

Before describing the data structure for piecewise linear maps between simplicial surfaces we define these maps formally.
Definition 2.2. Let $S_{1}, S_{2}$ be simplicial surfaces. A function $\Xi: D \subset S_{1} \rightarrow S_{2}$ is called piecewise linear if for each pair of triangles $T_{1} \in S_{1}$ and $T_{2} \in S_{2}$ the restriction of $\Xi$ to $T_{1} \cap \Xi^{-1}\left(T_{2}\right)$ is an affine function.

We make the following additional assumption.
Assumption 2.1. For each element $T$ of $S_{2}$, the set $T \cap \Xi\left(S_{1}\right)$ is a simplex of $S_{2}$ or empty.

By restricting our attention to a subsurface of $S_{2}$ we even assume that $\Xi$ is surjective, i.e., the image surface $\Xi\left(S_{1}\right) \subset S_{2}$ is equal to $S_{2}$. However, this restriction is not imposed in the implementation.

The following special case will appear frequently.
Definition 2.3. If $S_{1}$ and $S_{2}$ are homeomorphic, and $\Xi: S_{1} \rightarrow S_{2}$ is defined on all of $S_{1}$ and is a homeomorphism, then $\Xi$ is called a parametrization of $S_{2}$ over $S_{1}$.

The edges and vertices of $S_{2}$ form a graph with a geometric realization, which is mapped by the inverse function $\Xi^{-1}$ onto $S_{1}$. By linearity, it is enough to know how $\Xi$ acts on this graph image, and this is how PSURFACE is implemented. Remember that a graph $\mathcal{G}$ is a finite set of vertices $V$ together with a set $E$ of unordered pairs of vertices which are called edges. A graph can be given a geometry in $\mathbb{R}^{d}$ by associating each vertex $v \in V$ with a position $p(v) \in \mathbb{R}^{d}$, and each edge $e=\left(v_{0}, v_{1}\right)$ with the line segment from $p\left(v_{0}\right)$ to $p\left(v_{1}\right)$. If $d \in\{1,2\}$ and none of these segments intersect except at vertices then the graph together with the embedding into $\mathbb{R}^{d}$ is called a straight-line plane graph 7]. Each plane graph divides $\mathbb{R}^{d}$ into a set of $d$-dimensional regions. If each region except for the unbounded one is a simplex, $\mathcal{G}$ is called a triangulation. More generally we can define embeddings of graphs into simplicial surfaces.

Definition 2.4. Let $\mathcal{G}=(V, E)$ be a graph and $S$ a simplicial surface. With each vertex $v \in V$ associate a position $p(v) \in S$, and with each edge $e=$ $\left(v_{0}, v_{1}\right) \in E$ associate a set of open line segments $\eta_{e}=\left\{\left(e_{0}, f_{0}\right), \ldots,\left(e_{n_{e}}, f_{n_{e}}\right)\right\}$ with $\left(e_{i}, f_{i}\right) \subset S$ for all $i=0 \ldots, n_{e}$. If

- $p\left(v_{0}\right)=e_{0}, p\left(v_{1}\right)=f_{n_{e}}, f_{i}=e_{i+1}$,
- for each $\left(e_{i}, f_{i}\right)$ there exists an element $T$ of $S$ such that $\left(e_{i}, f_{i}\right) \subset T$,
- for any two segments $\left(e_{i}, f_{i}\right)$ and $\left(e_{j}^{\prime}, f_{j}^{\prime}\right)$ we have $\left(e_{i}, f_{i}\right) \cap\left(e_{j}^{\prime}, f_{j}^{\prime}\right)=\emptyset$,
then $(p, \eta)$ with $\eta=\left\{\eta_{e} \mid e \in E\right\}$ is called a piecewise straight embedding of $\mathcal{G}$ in $S$. A piecewise straight embedding is called minimal, if for each triangle $T \in S$ and each edge $e \in E$ there is at most one segment $\left(e_{i}, f_{i}\right) \in \eta_{e}$ with $\left(e_{i}, f_{i}\right) \subset T$.


Figure 3: Implementation of a piecewise linear mapping $\Xi$ as a graph on $S_{1}$. Graph node types are corner $(\bullet)$, touching $(\square)$, intersection $(\square)$, ghost $(\bigcirc)$, and interior (no symbol).

A graph embedded into a triangulated surface $S$ subdivides $S$ into regions. In general these regions do not coincide with the triangles of $S$ (Figure 3).

Piecewise linear mappings induce graphs on the domain surface $S_{1}$. For a formal statement of this, we first define the preimage.
Definition 2.5. Let $\Xi: S_{1} \rightarrow S_{2}$ be piecewise linear and $A \subseteq S_{2}$. We define the preimage of $A$ under $\Xi$ as

$$
\Xi^{-1}(A)=\left\{x \in S_{1} \mid \Xi(x) \in A\right\}
$$

The vertices $\mathcal{V}_{2}$ and the edges $\mathcal{E}_{2}$ of the surface $S_{2}$ form the edge graph $\mathcal{G}_{2}$ of $S_{2}$. Its preimage $\mathcal{G}_{\Xi}=\Xi^{-1}\left(\mathcal{G}_{2}\right)$ under the piecewise linear map $\Xi: D \subset S_{1} \rightarrow S_{2}$ is a minimal piecewise straight embedding of $\mathcal{G}_{2}$ in $S_{1}$ (Figures 11 and 2).

Theorem 2.1. Let $S_{1}, S_{2}$ be two simplicial surfaces and let $\Xi: D \subset S_{1} \rightarrow S_{2}$ be a surjective, continuous, piecewise linear map. Denote by $\mathcal{G}_{2}=\left(\mathcal{V}_{2}, \mathcal{E}_{2}\right)$ the edge graph of $S_{2}$. Then $\Xi^{-1}\left(\mathcal{G}_{2}\right)$ is a minimal piecewise straight embedding of $\mathcal{G}_{2}$ in $S_{1}$, with

$$
p(v)=\Xi^{-1}(v)
$$

for all vertices $v \in \mathcal{V}_{2}$, and

$$
\eta_{e}=\left\{T \cap \Xi^{-1}(e) \mid T \in \mathcal{T}_{1}, T \cap \Xi^{-1}(e) \neq \emptyset\right\}
$$

for all edges $e \in \mathcal{E}_{2}$.
This graph embedding contains all relevant information about $\Xi$. Therefore, the mapping $\Xi: D \subset S_{1} \rightarrow S_{2}$ can be stored using a data structure for graphs on $S_{1}$. This is the approach taken by PSurface, and we will now describe how this data structure looks like.

To reduce complexity, the data structure for $\mathcal{G}_{\Xi}$ on $S_{1}$ consists of data structures for plane graphs on each element of $S_{1}$. Some additional bookkeeping then allows to reconstruct the complete graph $\mathcal{G}_{\Xi}$ from its restrictions to the individual elements. For each element $T \in \mathcal{T}_{1}$ we store a set of nodes, each storing its local position on $T$, its position on $S_{2}$, and a list of its neighbors in
cyclic order. We introduce five different types of graph nodes (Figure 3). The first three correspond to vertices of $S_{2}$.

Interior nodes: Nodes in the interior of an element $T$.
Corner nodes: Nodes on the corners of $T$
Touching nodes: Nodes on an edge $k \subset \partial T$, but not on a corner of $T$.
Nodes on edges or corners have copies on all elements that share the common edge or corner. The same holds for the following two node types which are needed to make the restriction of $\mathcal{G}_{\Xi}$ on each element a graph in its own right, and to allow the evaluation of $\Xi$ on any point of an element without having to use information from other elements.

Intersection nodes: For any edge $e=\left(v_{0}, v_{1}\right) \in \mathcal{E}_{2}$, the corresponding preimage $\Xi^{-1}(e)$ is generally not contained in a single element of $S_{1}$. Instead, from Theorem 2.1 we know that it consists of a sequence of segments $\left(e_{i}, f_{i}\right), 0 \leq i \leq n_{e}$, with each segment $\left(e_{i}, f_{i}\right)$ being contained in a single element. With the exception of $e_{0}$ and $f_{n_{e}}$, the points $e_{i}, f_{i}$ do not correspond to vertices of the edge graph $\mathcal{G}_{2}$. Still, the data structure needs to keep them as vertices anyways, in order to have a consistent graph data structure on each element of $S_{1}$. Since, by Theorem 2.1, the embedding of $\mathcal{G}_{2}$ into $S_{1}$ is minimal, intersection nodes cannot occur in the interior of triangles.

Ghost nodes: If a corner $c$ of an element $T \in \mathcal{T}_{1}$ does not get mapped onto a vertex of $S_{2}$, a ghost node without any neighbors is added at $c$.

Each node $p$ stores its image $\Xi(p) \in S_{2}$ by storing the index of a triangle $T \in \mathcal{T}_{2}$ with $\Xi(p) \in T$ and local coordinates of $\Xi(p)$ with respect to $T$. Finally, each triangle $T \in \mathcal{T}_{1}$ keeps three arrays containing the nodes on the three triangle edges in cyclic order, and each node on an edge knows its index in the corresponding array. That way, corresponding nodes on adjacent triangles are identified, and it is possible, in a given graph data structure, to efficiently track preimages of edges of $S_{2}$ across multiple triangles of $S_{1}$.
Remark 2.1. While globally $\Xi^{-1}\left(\mathcal{G}_{2}\right)$ is a triangulation of its domain of definition, its restrictions to individual triangles of $\mathcal{T}_{1}$ need not be (Figure 3). However, for efficient point-location it is necessary that the graph on each triangle be a triangulation (see [5] and Section 2.3). Therefore, we add additional graph edges to the data structure such that $\Xi^{-1}\left(\mathcal{G}_{2}\right)$ is a triangulation on each $T \in \mathcal{T}_{1}$. We call this the triangular closure.

The data structure simplifies considerably if $d=1$. Then the elements of $S_{1}$ and $S_{2}$ are simply line segments connecting pairs of vertices in $\mathbb{R}^{2}$. On each segment of $S_{1}$, the graph $\mathcal{G}_{\Xi}$ is merely a linear sequence of nodes ordered by local position $s \in[0,1]$ and with adjacent nodes (possibly) connected by a graph edge. The simpler structure also limits the set of node types that can occur in $\mathcal{G}_{2}$. As there are no edges in $S_{1}$, node types that live on edges do not occur. Only interior, corner, and ghost nodes are needed. For the same reason, the arrays storing the nodes on surface edges in cyclic order do not appear in the one-dimensional data structure.

### 2.3 Evaluating a piecewise linear map

Given a consistent data structure as described above and a point $p \in S_{1}$ specified by a triangle $T \in \mathcal{T}_{1}$ and local coordinates $\xi$ on $T$, the map $\Xi$ can be evaluated at $p$ in two steps. If $p$ is contained in the domain of $\Xi$, then it must be contained in a region $r$ of the graph $\mathcal{G}_{\Xi}=\Xi^{-1}\left(\mathcal{G}_{2}\right)$. First, this region $r$ (with corners $c(r)$ ) is determined using a point-location algorithm, and the barycentric coordinates $\zeta_{c(r)}$ of $p$ with respect to $r$ are computed. If $d=1$ the element $T$ is a line segment, and the graph data structure on $T$ has been prepared for point-location by sorting the graph nodes in increasing order of their local coordinates. Then the region $r=\left[v_{i}, v_{i+1}\right]$ containing $p$ is found in logarithmic time by a bisection search.

If $d=2$ we need a point location in a two-dimensional plane graph. Remember that we added the additional graph edges to make all regions of $\left.\mathcal{G}_{\Xi}\right|_{T}$ triangles (Remark 2.1). For point-location in triangulations we use the randomized version of the algorithm presented by Brown and Faigle [5]. It is simple to implement and its expected run-time is in $O\left(\sqrt{n_{e}}\right)$, with $n_{e}$ the number of edges in the triangulation.

After the point-location step we have a triangular region $r$ of $\mathcal{G}_{\Xi}$ on $T$ with $p \in r$, barycentric coordinates $\zeta$ of $p$ with respect to $r$ and the set of corners $c(r)$ of $r$. By construction, the images of all corners of $r$ are on the same element $T_{2}$ of $S_{2}$, and their local positions of their images $\Xi(c(r))$ with respect to $T_{2}$ are stored in the node data structure. Hence the local position of $\Xi(p)$ with respect to $T_{2}$ can be computed by linear interpolation

$$
\Xi(p)=\sum_{c(r)} \zeta_{c(r)} \Xi(c(r)) .
$$

By this algorithm we have proved the following result.
Theorem 2.2. $\Xi$ is completely determined by the presented data structure.

## 3 Constructing a piecewise linear mapping

There are various ways to set up a piecewise linear mapping, and these are related to different applications of PSurface. You can either directly construct a PSurface object by means of certain primitives, construct it by projection of $S_{1}$ onto $S_{2}$; or by starting with two identical surfaces $S_{1}=S_{2}$ and the identity map, and then coarsening $S_{1}$.

### 3.1 Direct construction

PSURFACE allows to set up a map $\Xi$ from scratch by explicitly specifying the vertices and triangles of $S_{1}$ and $S_{2}$, and their relations under $\Xi$. This can be used, for example, to read mappings provided in some file format 1 Also, it provides a clear interface to PSURFAcE creation for those wishing to implement their own construction algorithms.

[^1]Direct creation of a PSurface object is implemented by means of a factory class. The PSurfaceFactory is used internally by all other construction methods. It accepts the specification of surfaces $S_{1}$ and $S_{2}$, and a graph on $S_{1}$ describing a piecewise linear map. Upon a call to the method createPSurface, the factory hands back a pointer to the newly created object.

Creating the surfaces $S_{1}$ and $S_{2}$ is done by providing positions for the vertices, and $d+1$-tuples of vertex indices for the elements. Since the data structure for $S_{2}$ does not contain information about $\Xi$ it is also possible to build it outside of the factory. A pointer to $S_{2}$ is then handed to the PSurfaceFactory. At this point, the graph on $S_{1}$ is the empty graph, i.e., there is no mapping defined anywhere on $S_{1}$.

The actual mapping is specified in a separate step. In the simplest case, $S_{2}$ is a logical refinement of $S_{1}$ under $\Xi$, i.e., for each element $T_{2} \in S_{2}$, the preimage $\Xi^{-1}\left(T_{2}\right)$ is entirely contained in a single element $T_{1}$ of $S_{1}$. In this case, vertices of $S_{1}$ in the domain $D \subset S_{1}$ of $\Xi$ are mapped to vertices of $S_{2}$, and preimages of edges of $S_{2}$ do not cross edges of $S_{1}$. Graph nodes of $\mathcal{G}_{\Xi}$ can only be of corner, touching, or interior type, and the factory class provides a method to enter such nodes into the graph data structure on $S_{1}$. Interior nodes get entered once for the element they reside on. Corner and touching nodes get entered automatically on each element that borders the vertex and edge, respectively, that the node resides on. The method returns a handle to the newly created set of nodes. A second method allows to insert edges into the graph. Since, by assumption, an edge of $\mathcal{G}_{\Xi}$ is contained in a single element it can be inserted simply by giving the element number and handles to its two nodes. A convenience method allows to insert elements of $S_{2}$, which amounts to inserting their edges, if not already present.

If $S_{2}$ is not a logical refinement of $S_{1}$, more information is needed. First of all, ghost nodes appear at those vertices of $S_{1}$ in $D$ that do not get mapped to a vertex of $S_{2}$. Ghost nodes can be added to the factory by calling the method insertGhostNode. The arguments are the vertex number on $S_{1}$, and an element number and local coordinates on $S_{2}$. If the vertex of $S_{1}$ is to be mapped onto an edge of $S_{2}$ any one of the adjacent triangles will do. Secondly, an edge $e=(p, q)$ of $S_{2}$ may now connect vertices whose preimages are not on the same element of $S_{1}$. In this case, all crossings of $\Xi^{-1}(e)$ with edges of $S_{1}$ need to be specified. This leads to the insertion of intersection nodes at the triangle boundaries. The preimage of an edge is inserted using the method insertEdge. Its arguments are the nodes on $S_{1}$ corresponding to $p$ and $q$, an array of the crossed edges of $S_{1}$, and local coordinates of the intersection nodes on these edges. If $\Xi^{-1}(e)$ crosses a vertex of $S_{1}$ (on one-dimensional surfaces, this is even a certainty for edges visiting more than a single element of $S_{1}$ ), a ghost node is automatically inserted at the vertex instead of an intersection node, unless that ghost node has already been inserted previously by hand.

After all nodes and edges have been inserted, a finalization method creates the actual PSurface object. Among other things it computes the triangular closure (Remark 2.1), sorts the edges of each node in cyclic order 2 and builds up the arrays that store the nodes on the element edges.

[^2]

Figure 4: Backward normal projection from a point $q$ onto a triangle (left), and from an edge ( $\tilde{q}_{0}, \tilde{q}_{1}$ ) onto an edge ( $p_{0}, p_{1}$ ) (right).

### 3.2 Projection

For two given simplicial surfaces $S_{1}$ and $S_{2}$ a mapping can also be constructed algorithmically by projecting $S_{1}$ onto $S_{2}$ along a piecewise linear vector field on $S_{1}$ given by directions at the vertices of $S_{1}$. This is an important ingredient, e.g., of algorithms for computing contact problems in continuum mechanics (see Section 4.2). Frequently, the projection direction is chosen as a piecewise linear averaged surface normal of $S_{1}$, but other sufficiently well-behaved vector fields on $S_{1}$ can also be used.

More formally, let $\boldsymbol{\nu}: \mathcal{V}_{1} \rightarrow \mathbb{R}^{d+1}$ be a set of vectors associated to the vertices of $S_{1}$. We extend $\boldsymbol{\nu}$ by linear interpolation to a continuous vector field on $S_{1}$, which we denote again by $\boldsymbol{\nu}$. Our aim is to construct a set $D \subseteq S_{1}$ and a mapping $\Phi: D \rightarrow S_{2}$ such that $D$ is as large as possible and

$$
\begin{equation*}
\Phi(p)-p=\mu \boldsymbol{\nu}(p) \quad \text { for all } p \in D \tag{1}
\end{equation*}
$$

with $p$ and $\Phi(p)$ interpreted as points in $\mathbb{R}^{d+1}$ and $\mu \in \mathbb{R}_{0}^{+}$depending on $p$.
Assume that surfaces $S_{1}$ and $S_{2}$ and the vector field $\boldsymbol{\nu}$ are given. The construction of the set $D$ and the mapping $\Phi$ consists of three steps.

1. Computing $\Phi^{-1}(q)$ for all vertices $q \in \mathcal{V}_{2}$

For each $q \in \mathcal{V}_{2}$ we have to find a $p \in S_{1}$ such that

$$
\begin{equation*}
q-p=\mu \boldsymbol{\nu}(p) \tag{2}
\end{equation*}
$$

We can then define $\Phi^{-1}(q):=p$. For each $q \in \mathcal{V}_{2}$ and each element $T$ of $S_{1}$,(2) is a nonlinear system of equations for the barycentric coordinates $\lambda=\left(\lambda_{0}, \ldots, \lambda_{d+1}\right)$ of $p$ on $T$ and the distance $\mu$. For $d=2$ it reads

$$
\begin{equation*}
0=p_{2}-q+\lambda_{0}\left(p_{0}-p_{2}\right)+\lambda_{1}\left(p_{1}-p_{2}\right)+\mu \lambda_{0}\left(\boldsymbol{\nu}_{0}-\boldsymbol{\nu}_{2}\right)+\mu \lambda_{1}\left(\boldsymbol{\nu}_{1}-\boldsymbol{\nu}_{2}\right)+\mu \boldsymbol{\nu}_{2}, \tag{3}
\end{equation*}
$$

where $p_{0}, \ldots, p_{d}$ are the corners of $T$ and $\boldsymbol{\nu}_{0}, \ldots, \boldsymbol{\nu}_{d}$ are the directions at these corners. A point $p$ with (1) exists on $T$ if (3) has a solution with $\lambda_{0}, \lambda_{1}, \mu \geq 0$ and $\lambda_{0}+\lambda_{1} \leq 1$. This system can be solved conveniently using a Newton solver.
To find preimages for all vertices of $S_{2}$ we loop over the elements of $S_{1}$. For each element $T \in \mathcal{T}_{1}$ we use an octree to efficiently find all vertices of


Figure 5: In certain cases the edge projection algorithm fails: the edge preimage may leave $S_{1}$ (left), or the projection may not be continuous (right).
$S_{2}$ reachable in the direction of $\boldsymbol{\nu}$ from $T$. A vertex $v$ has a preimage $p$ on $T$ if (1) has a solution for $p$ in $T$ and there is no other solution $(\tilde{p}, \tilde{\mu})$ on an element $\tilde{T}$ with $\tilde{\mu}<\mu$. To increase robustness we additionally compare the normal $\tilde{\boldsymbol{\nu}}$ of $S_{2}$ at $q$ and the segment $w=q-p$, and accept a preimage only if $\langle w, \tilde{\boldsymbol{\nu}}\rangle<0$.
2. Computing $\Phi(v)$ for all vertices $v \in \mathcal{V}_{1}$

At this stage all vertices of $S_{2}$ in the range of $\Phi$ appear as nodes in the graph on $S_{1}$. We have to add additional ghost nodes at those vertices of $S_{1}$ that are not mapped onto vertices of $S_{2}$. This is comparatively easy as it does not involve solving nonlinear equations. Given a vertex $v \in \mathcal{V}_{1}$, its image on $S_{2}$ can be found by considering the ray $r$ in direction $\boldsymbol{\nu}(v)$ beginning in $v$, and looking for intersections of $r$ with elements of $S_{2}$. Potential candidate elements for intersections are produced efficiently using an octree. If $r$ intersects more than one triangle of $S_{1}$ the intersection closest to $v$ is chosen. If no intersection is found then $\Phi(v)$ will be left undefined by modifying $D$ such that $v \notin D$. If $v$ is mapped onto a vertex of $S_{2}$ then it has already been treated in Step 1 and the data structure already contains a node for it. Otherwise a ghost node is inserted.
3. Adding the graph edges

In order to enter an edge $\tilde{e}=\left(\tilde{q}_{0}, \tilde{q}_{1}\right)$ of $S_{2}$ into the graph on $S_{1}$ we try to 'walk' on $S_{1}$ along $\Phi^{-1}(\tilde{e})$ from $q_{0}=\Phi^{-1}\left(\tilde{q}_{0}\right)$ to $q_{1}=\Phi^{-1}\left(\tilde{q}_{1}\right)$. Since $q_{0}$ and $q_{1}$ will generally not be on the same triangle of $S_{1}$, we have to find the points where the path from $q_{0}$ to $q_{1}$ crosses edges of $S_{1}$. Let $T \in \mathcal{T}_{1}$ be the current triangle in this walking process. For an edge $e=\left(p_{0}, p_{1}\right)$ of $T$ we have to check whether there are points $p \in e$ and $q \in \tilde{e}$ with $q-p$ collinear to $\boldsymbol{\nu}(p)$ (Figure [4, right). This can be formulated as a nonlinear system of equations

$$
\begin{equation*}
p(\lambda)+\mu \boldsymbol{\nu}(\lambda)=q(\eta) \tag{4}
\end{equation*}
$$

for three variables $\lambda, \mu, \eta \in \mathbb{R}$ and
$p(\lambda)=p_{0}+\lambda\left(p_{1}-p_{0}\right), \quad \boldsymbol{\nu}(\lambda)=\boldsymbol{\nu}_{0}+\lambda\left(\boldsymbol{\nu}_{1}-\boldsymbol{\nu}_{0}\right), \quad q(\eta)=\tilde{q}_{0}+\eta\left(\tilde{q}_{1}-\tilde{q}_{0}\right)$,
which can be solved with a damped Newton algorithm. We have found an intersection if (4) has a solution with $0 \leq \lambda, \eta \leq 1$ and $0 \leq \mu$. This intersection is then inserted as an intersection node and the procedure is continued on the triangle which borders $T$ on $e$.


Figure 6: The point-removal primitive

There are various corner cases where this approach fails. For example, even though $\Phi^{-1}(\tilde{e})$ intersects $e$, there may not be another triangle that borders $T$ across $e$ (Figure 5, left). Alternatively, there may be such a triangle, but it may be hidden from $S_{2}$ by $T$ (Figure 5, right). In both cases, the edge insertion algorithm currently simply aborts and does not insert the edge $\tilde{e}$ at all. For the future one may consider adding smarter algorithms that can handle the first case. In the second case, the projection in the direction of $\boldsymbol{\nu}$ is not continuous, and hence storing it is beyond the scope of the PSURFACE data structure.

Assuming that the Newton solvers for (3) and (4) terminate after a constant number of iterations and that each edge of $S_{2}$ crosses only a constant number of elements from $S_{1}$, the projection algorithm described above requires an expected $O\left(N_{\max } \log N_{\max }\right)$ time, with $N_{\max }=\max \left(\left|\mathcal{V}_{1}\right|,\left|\mathcal{V}_{2}\right|\right)$. In the worst case the time needed is quadratic. This is optimal, because there are configurations where (almost) every edge of $S_{1}$ gets mapped onto (almost) every edge of $S_{2}$, leading to a quadratic number of intersection nodes.

### 3.3 Constructing parametrizations by surface simplification

In this section we give an algorithm that constructs piecewise linear mappings by surface simplification. It appeared originally in [18]. Given a not too coarse triangulated surface $S_{2}$, the result is a coarser surface $S_{1}$ and a parametrization $\Xi$ of $S_{2}$ over $S_{1}$, i.e., a homeomorphism from $S_{1}$ to $S_{2}$. The parametrization should be of good quality, a criterion to be specified more formally in Section 3.3.2.

Parametrizations of a simplicial surface over a coarser copy of itself are useful for the treatment of domain boundaries in finite element problems. These boundaries are sometimes given as triangulated surfaces of very high resolution, and need to be coarsened before being able to serve as input for a mesh generator [18]. The geometric information lost by coarsening can be regained later during mesh refinement if a parametrization of the fine boundary over the coarse boundary is available.

The idea of constructing a parametrization function by surface simplification originates in computer graphics [16], where it is used for a variety of problems, such as surface remeshing and texture map generation [17]. Our algorithm consists of two steps:

1. construct the base domain surface $S_{1}$ and a parametrization function $\Xi$ : $S_{1} \rightarrow S_{2}$ by means of a simplification algorithm (Section 3.3.1),
2. smooth $\Xi$ to obtain refined surfaces of optimal quality (Section 3.3.2).


Figure 7: The polar map: The star of $p$ is first cut along the edge $\left(p, p_{1}\right)$ and flattened out into the complex plane. The conformal map $z \rightarrow z^{a}, a=$ $2 \pi / \sum_{i} \theta_{i}$, then stretches it out so that it covers one full circle.

We describe the algorithm assuming that $S_{2}$ is a piecewise linear manifold. The algorithm also works well in the non-manifold case. See [18] for details.

### 3.3.1 Surface simplification

Let $S_{2}$ be a simplicial surface of sufficiently high resolution. We initialize the algorithm by setting $S_{1}^{0}=S_{2}$ and $\Xi^{0}: S_{1}^{0} \rightarrow S_{2}, \Xi^{0}=\mathrm{Id}$. The algorithm will construct a sequence of simplicial surfaces $S_{1}^{i}$ and functions $\Xi^{i}: S_{1}^{i} \rightarrow S_{2}$ such that $S_{1}^{i}$ contains less elements than $S_{1}^{i-1}$ and $\Xi^{i}\left(S_{1}^{i}\right)=S_{2}$ for all $0 \leq i \leq N$.

A surface simplification algorithm consists of two parts. The first is a geometric operation that allows the controlled local reduction of surface complexity. We chose point removal since it does not create new vertices and is easily extendible to non-manifold surfaces. Point removal works by repeatedly removing a vertex and its neighboring triangles, and retriangulating the resulting hole (Figure 6).

The second ingredient is a scalar oracle that allows to rank the different possible simplification steps according to the error they would introduce. Many different strategies have been described [12, 18]. We combine the following aspects:

- monitor a modified Hausdorff distance between different simplification stages to control geometric error,
- favor steps that remove triangles with a high aspect ratio and introduce triangles with a low aspect ratio, to maintain a high-quality triangulation,
- reject simplification steps that lead to surface intersections,
- penalize long edges, to obtain surfaces of uniform element size.

Let $i \in \mathbb{N}$ be the current simplification step and assume that $v_{i}$ is the current best vertex according to the error oracle. We define St $v_{i}$, the star of $v_{i}$, as the set of all edges and elements that contain $v_{i}$. Speaking first strictly in terms of the domain surface $S_{1}^{i}$, a simplification step consists of removing $v_{i}$ and its star from $S_{1}^{i}$. The resulting hole is then filled by a constrained Delaunay triangulation with a set of elements $P_{i}$. The resulting surface $S_{1}^{i+1}$ has one vertex and two elements less than $S_{1}^{i}$.


Figure 8: The flattened star of a vertex is merged into a single polygon.


Figure 9: The polygon is cut along the new triangulation.

At the same time we have to make sure that the mapping defined on St $v_{i}$ is correctly transferred to $P_{i}$. This means that the graphs defined on the elements of St $v_{i}$ must be transferred to the elements of $P_{i}$. For this, after having removed St $v_{i}$ from $S_{1}^{i}$, we flatten it out into $\mathbb{R}^{d}$. This is easy if $d=1$. If $d=2$ we use the polar map introduced in [8]. It maps St $v_{i}$ bijectively and conformally onto a star-shaped polygon in $\mathbb{R}^{2}$ (Figure [7). We then merge the graphs on the flattened elements of St $v_{i}$ into a single plane graph on the polygon consisting of the union of the elements of $\operatorname{St} v_{i}$ (Figure 8).

This polygon implements that part of the parametrization function $\Xi$ that had formerly been implemented by the triangles in St $v_{i}$. We then repartition the graph according to the new triangulation $P_{i}$ obtained by Delaunay triangulation (Figure 9). In the process, edges are cut and intersection nodes are added, and interior nodes may turn into touching nodes. The result is a set of graphs for the triangles that are inserted into the domain surface.

Denoting by $n$ the number of vertices of $S_{2}$, the surface simplification algorithm needs only $O(n \log n)$ steps to remove a constant fraction of the vertices. However, due to the overhead for maintaining a consistent parametrization at each step the constant is fairly large. The space requirements remain linear in the number of surface vertices.

### 3.3.2 Parametrization smoothing

The mappings constructed by the simplification algorithm are frequently of poor quality. By this we mean that the maximal relative condition number

$$
\kappa_{\Xi}=\max _{A \in \mathcal{A}_{\Xi}}\|A\|_{2}\left\|A^{-1}\right\|_{2}
$$

can get very large. Here, $A$ is the matrix corresponding to one linear piece of a piecewise linear map $\Xi$, and $\mathcal{A}_{\Xi}$ is the set of all these matrices. As a consequence, surfaces obtained by refining $S_{1}$ and using $\Xi$ to place the newly


Figure 10: The edge relaxation algorithm significantly increases the quality of the refined mesh. The figure shows a surface constructed by simplification of a sphere (left) and a refined surface without (center) and with (right) smoothing.
inserted vertices can have large triangle aspect ratios (see Figure 10). In the context of finite element methods this is very undesirable. We will now shortly describe an algorithm that considerably improves the quality of the refined grids.

Assume that we have a method that computes high-quality parametrizations of surface patches over convex polygons. Various algorithms for this have been proposed in the literature. It follows from Gauß' Theorema Egregium that such a parametrization can in general not be isometric, i.e., distortion-free. We use the shape-preserving parametrization introduced by Floater [9]. Inspired by planar graph-drawing, it places the vertices in the polygon such that each vertex is a convex combination of its neighbors. The weights are chosen such that if there is an affine mapping of the surface onto the polygon, then this mapping is reproduced by the parametrization algorithm. Floater [9] showed that such mappings always exist, and that they can be computed by solving a sparse non-symmetric system of equations. This type of mapping is widely used in surface parametrization algorithms [10, 11].

We can use this method to recompute the parametrization on each element of $S_{1}$, while keeping the values of $\Xi$ on the edges of $S_{1}$ fixed. This will already greatly improve the parametrization quality. A further improvement can be achieved if the domains are enlarged. In PSurface we have implemented smoothing on quadrilaterals. Let $T_{1}, T_{2}$ be two triangles of $S_{1}$ that share a common edge. Consider them to be separated from the remaining surface. They can then be 'folded' isometrically to lie in a single plane. The common edge is removed and the graphs on $T_{1}$ and $T_{2}$ are merged into a single graph on the quadrilateral formed by $T_{1}$ and $T_{2}$. The Floater parametrization method can now be applied to this quadrilateral. Afterwards the common edge is reintroduced and the graph is cut along this edge. Merging and cutting of graphs was already a part of the surface simplification algorithm of Section 3.3.1] and the code can be reused.

To achieve a global improvement of the parametrization quality we now loop over all edges of $S_{1}$ with two adjacent triangles. For each such edge we smooth the quadrilateral formed by the two adjacent triangles. The result after one iteration is a marked improvement of the parametrization quality. Since the smoothing across edges introduces a global coupling, it pays to loop several times over all edges. In practice, a satisfying parametrization quality is reached in less than ten iterations. See [18] for more information.

## 4 Integration in Dune

Several features of PSURFACE are useful in the context of the numerical analysis of partial differential equations (PDEs). For these cases, PSurface can be used by the Distributed and Unified Numerics Environment (Dune). Dune is a set of C++ libraries for grid-based methods for solving PDEs [4]. It has been designed with flexibility as the main goal, and consists of interface definitions to core components such as grids, linear algebra, or shape functions. These components can then be implemented in various ways, allowing the user to always choose the most appropriate implementation for the task at hand. PSURFACE is currently used by the Dune modules dune-grid and dune-grid-glue. The dune-grid module provides the interface to finite element grids, together with various implementations of this interface. dune-grid-glue provides infrastructure for the coupling of several grids.

### 4.1 Parametrized boundaries with dune-grid

In the numerical solution of PDEs, the shape of the domain may be given as a high-resolution triangulated surface. Such a surface may need to be coarsened before grid generation is possible. The geometric information lost should be regained when adaptively refining the grid. For this, the construction and handling of a mapping from the coarse grid boundary to the original boundary is necessary. This is provided by the PSurface library.

Several of the DUNE grid implementations can handle the refinement of grids with a parametrized boundary, provided the geometric information is given. dune-grid contains code that, given that the PSurface and AmiraMesh libraries are installed on the system, loads a PSurface from an AmiraMesh file into a PSurfaceBoundary object. This object then hands out information about the number of boundary segments, their types and vertices. Most importantly, for each boundary segment it hands out an object of a class derived from Dune: : BoundarySegment. These objects implement mappings $f_{i}: \mathbb{R}^{d_{b}} \rightarrow \mathbb{R}^{d_{w}}$, with $d_{b}$ the dimension of the grid boundary and $d_{w}$ the dimension of the world space, and can be handed directly to the grid factory class at grid creation time ${ }^{3}$ Grid managers that are able to will then honor the geometric information for grid refinement (see Figure 12).

As a variation of this idea, it is also possible to parametrize the elements of certain grids themselves. Consider a two-dimensional grid embedded in a three-dimensional space. As a finite element grid it will be piecewise linear (or piecewise polynomial at most), but in many cases it will be intended to approximate a smooth surface. Hence, upon grid refinement one would expect the grid to approximate the smooth surface better. This is only possible if the shape of the actual surface is known as a function on the coarsest grid. If the target surface is not given analytically but by a second triangulated surface, then this function can be implemented using PSurface. Unlike the previous case, the parametrizing function is now defined on the entire grid instead of only on the grid boundary.

[^3]

Figure 11: Two domains $\Omega_{1}$ and $\Omega_{2}$ that meet at a common interface $\Gamma$ (left). The restrictions of the two grids on $\Gamma$ (center). Together they form the set of remote intersections $G_{M}$ (right).

### 4.2 Domain decomposition methods with dune-grid-glue

A different use of PSurface occurs in the treatment of domain decomposition problems. In the simplest case, a partial differential equation is to be solved on two subdomains $\Omega_{1}, \Omega_{2} \subset \mathbb{R}^{d}$, connected through suitable coupling conditions on a common interface $\Gamma$. This interface is assumed to be a $d$-1-dimensional manifold. Two grids $G_{1}, G_{2}$ on $\Omega_{1}$ and $\Omega_{2}$, respectively, induce trace grids $G_{\Gamma, 1}$, $G_{\Gamma, 2}$ on $\Gamma$, which are not necessarily related to each other in any way. Various domain decomposition methods, in particular the popular mortar method, require the evaluation of integrals of the form

$$
m(v, w)=\int_{\Gamma} v w d x
$$

where $v$ and $w$ are finite element functions with respect to $G_{\Gamma, 1}$ and $G_{\Gamma, 2}$, respectively [22]. To compute these integrals exactly, the domain of integration $\Gamma$ needs to be split up in intersections $I_{i j} \subset \Gamma$ such that each $I_{i j}$ is the set intersection of two elements $T_{i} \in G_{\Gamma, 1}, T_{j} \in G_{\Gamma, 2}$, (Fig. 11). If the grid elements are convex polyhedra, then each intersection is a convex polyhedron itself and can be decomposed in simplices. On each such simplex, the product $v w$ is a polynomial and a quadrature rule is available, hence $v w$ can be integrated exactly. The total integral is given as the sum of the integrals over the individual intersections $I_{i j}$.

In more general cases, the two domains may be separated by a positive gap. In this case, the coupling boundaries $\Gamma_{1} \subset \partial \Omega_{1}$ and $\Gamma_{2} \subset \partial \Omega_{2}$ are identified through a homeomorphism $\Phi: \Gamma_{1} \rightarrow \Gamma_{2}$ (Fig. 12 left). The integrals used in the mortar method take the form

$$
\begin{equation*}
m_{\Phi}(v, w)=\int_{\Gamma_{1}} v(x) w(\Phi(x)) d x \tag{5}
\end{equation*}
$$

Note that this degenerates to the previous simpler case if $\Gamma_{1}=\Gamma_{2}$ and $\Phi=\mathrm{id}$. The intersections $I_{i j}$ are now defined as

$$
I_{i j}=T_{i} \cap \Phi^{-1}\left(T_{j}\right)
$$

In Dune, the task of computing the intersections $I_{i j}$ from the coupling boundary grids $G_{\Gamma, 1}$ and $G_{\Gamma, 2}$ and the function $\Phi$ is handled by the module
dune-grid-glue [2]. The set of intersections can be accessed by stl-style iterators. Each intersection provides information such as its embedding in $T_{i}$ and $T_{j}$, its embedding in the world space $\mathbb{R}^{d}$, normal vectors, etc.

The general problem of computing intersections between two grids is too diverse to be handled efficiently by a single implementation. A very general implementation will not be efficient in simple cases, while an efficient implementation cannot be sufficiently general. The software design of dune-grid-glue therefore follows the general Dune philosophy of declaring interfaces and allowing the user to choose between various implementations of this interface.

One such implementation uses the PSurface library. During setup, the coupling grids $G_{\Gamma, 1}$ and $G_{\Gamma, 2}$ are extracted and a new PSurface object is created with $S_{1}=G_{\Gamma, 1}$ and $S_{2}=G_{\Gamma, 2}$. If $G_{\Gamma, 1}$ or $G_{\Gamma, 2}$ contain elements that are not simplicial, these are triangulated to obtain valid input surfaces for PSURFACE. A mapping $\Phi: S_{1} \rightarrow S_{2}$ is then constructed using the projection algorithm of Section 3.2. The default projection direction is the field of surface vertex normals of $S_{1}$, but user-specified directions are also possible. Note that this yields the identity mapping when $\Gamma_{1}=\Gamma_{2}$. The intersections $I_{i j}$ can then be read off as the regions of the graph $\mathcal{G}_{\Xi}$ on $S_{1}$ implementing $\Phi$. Since PSurface triangulates all regions of $\mathcal{G}_{\Xi}$ to facilitate point location queries, all intersections are simplices. dune-grid-glue then implements the various iterators over the intersections.

### 4.3 Example application: two-body contact

The possibilities of PSurface in Dune are demonstrated by the following short example. Consider a two-body contact problem occurring as part of a biomechanical application. We simulate the left distal femur and proximal tibia from the Visible Human data set 1]. The data was segmented and a high-resolution boundary surface was extracted. The femur surface consisted of 14468 triangles, and the tibia surface of 14902 triangles. They were simplified as described in Section 3.3 to yield coarse surfaces with 532 triangles for the femur and 444 triangles for the tibia. The Amira 20] grid generator produced two tetrahedral grids with 1328 and 1044 elements, respectively (Figure 12).

We modeled bone with an isotropic, homogeneous, linear elastic material. The bottom section of the proximal tibia was clamped and a downward displacement of 6 mm was prescribed on the upper section of the femur. The parts of the bones usually covered with articular cartilage were marked as the coupling boundaries $\Gamma_{1}$ and $\Gamma_{2}$, but the actual coupling boundary was smaller, because the normal projection $\Phi$ could only be constructed on a part of the prescribed coupling boundary.

Mutual nonpenetration of the two objects was modeled by a linearized inequality constraint. Let $\mathbf{u}_{i}: \Gamma_{i} \rightarrow \mathbb{R}^{3}, i=1,2$, be the traces of the displacement functions on the contact boundaries. We required

$$
\begin{equation*}
\left\langle\mathbf{u}_{1}-\mathbf{u}_{2} \circ \Phi, \mathbf{n}\right\rangle \leq g, \tag{6}
\end{equation*}
$$

where $\mathbf{n}$ is the outer unit normal on $\Gamma_{1}$ and $g: \Gamma_{1} \rightarrow \mathbb{R}$ is the normal distance in the undeformed state. The mortar discretization of (6) is

$$
\begin{equation*}
\int_{\Gamma_{1}}\left\langle\mathbf{u}_{1}-\mathbf{u}_{2} \circ \Phi, \mathbf{n}\right\rangle \mu d s \leq \int_{\Gamma_{1}} g \mu d s \tag{7}
\end{equation*}
$$



Figure 12: Two-body contact problem. Note the stress peaks on the condyles due to the contact, and how the refined grid boundary approximates the smooth domain boundary.
for a set of test functions $\mu$ from a suitable test function space $M$. Expressing this algebraically in terms of basis functions leads to terms of the form (5). See [19] for the details. Figure 12 show a result of the computation after three steps of adaptive refinement. It can be seen that the refined grid boundary approaches the original finer boundary. The effect of the weak nonpenetration condition (17) can be seen by the stress peaks at the condyles.

## 5 Integration in Amira

Amira is a software for the visualization of scientific data. Originally developed at the Zuse Institute Berlin (ZIB), Amira is now distributed commercially 4 Amira is written in $\mathrm{C}++$ and has a plug-in infrastructure. Plug-ins have access to most internal data structures, and visualization and data processing algorithms. Conversely, users can control the plug-ins with the Amira GUI and the tcl scripting language integrated into Amira.

The development of PSurface started when the author was employed at the visualization department of ZIB. Therefore, the original code consisted entirely of Amira plug-ins. Later, the core data structures and algorithms were moved into a generic C++ library which could be compiled and used independently from Amira. However, some useful functionality still exists in form of Amira code. It is collected in the plug-in hxpsurface.

The core of the PSurface Amira integration is a class HxPSurface which encapsulates a PSurface and makes it available as an Amira data object. Using this wrapper, an Amira viewer module allows to visualize the PSurface. Both surfaces $S_{1}$ and $S_{2}$ are shown as triangulated surfaces, and the mapping is visualized as the edge graph of $S_{2}$ on $S_{1}$ (Figure 2). Such a representation gives useful insight into the structure and quality of the parametrization, among other things for debugging purposes. Additionally, the display module can visualize the graph nodes with a color code according to the node type, and the images

[^4]of the edges of $S_{1}$ under $\Xi$. The regularity of these edges is another indicator of the parametrization quality.

Besides the direct visualization of PSurface objects, the Amira plug-in offers several other useful features. In particular, parts of the surface simplification algorithm of Section 3.3 are contained in the plug-in. While PSurface itself contains the code for removing single vertices, retriangulating the hole and transferring the preimage graphs, the error criteria and ranking algorithms reside in hxpsurface. That way, the simplification can be controlled conveniently using the Amira GUI. Similarly, the code to smooth a parametrization across an edge is contained in PSurface, while the global loop and a GUI are in the plug-in. A general PSURFACE editor, modeled after the Amira surface editor, allows to interactively modify the domain surface $S_{1}$. Among other things it is possible to delete single vertices, and flip or split edges, while always maintaining a valid parametrization. Also, various quality measures of $S_{1}$ such as the triangle aspect ratio can be computed.

Another useful feature is the remesher, which allows to visualize a uniformly refined $S_{1}$, where all new vertices have been moved to the positions prescribed by a parametrization $\Xi$. For a parametrization intended for use as a finite element grid boundary, this allows to visually inspect the quality of the resulting mesh boundary.

As a conclusion, using PSurface as an Amira plug-in offers a simple way to implement and experiment with computer graphics and visualization algorithms involving parametrized surfaces. For example, the construction of texture and bump maps and their visualization would be straightforward. As another example, we show the linear morphing of one triangulated surface into another. This example originally appeared in [18].

### 5.1 Morphing

Morphing is the continuous deformation of a surface into another one. Given two homeomorphic surfaces $S$ and $S^{\prime}$, we are looking for a one-parameter family of surfaces $\mathcal{S}=\{S(\rho) \mid \rho \in[0,1]\}$ with $S(0)=S, S(1)=S^{\prime}$ and such that the map $\rho \rightarrow S(\rho) \in \mathcal{S}$ is continuous. The user is usually requested to mark sets of feature points on the surfaces that will be required to correspond via the morphing. These points will generally have some geometric significance. For example, when morphing two heads, one might choose the tip of the nose, the chin, the corners of the mouth and the like. The algorithm should then construct a surface family under those constraints.

A basic algorithm looks like this: given two triangulated surfaces $S$ and $S^{\prime}$ of identical topological type, and an equal number of feature points on each of them. On each surface, the points are then connected by paths that triangulate each surface such that the two triangulations are combinatorially equivalent. This can be done manually or automatically. From these triangulations coarse surfaces $B$ and $B^{\prime}$ are constructed. The vertices of $B$ and $B^{\prime}$ are the feature points of $S$ and $S^{\prime}$, respectively, and two vertices in $B\left(B^{\prime}\right)$ are connected by an edge if there is a corresponding path in $S\left(S^{\prime}\right)$. The triangulated surfaces are then parametrized over these base surfaces $B, B^{\prime}$ : We map points on patch boundaries onto base grid edges via an arc-length parametrization and we use the Floater scheme for all other vertices. The quality of the parametrization can be improved further by applying the relaxation scheme described in


Figure 13: Constructing a parametrization of ' $\mathrm{Al}^{\prime}$ ', the cartoon figure, that is suitable for morphing. i) the surface with marked points and a rough triangulation; ii) the corresponding base grid; iii) the base grid after the application of edge-relaxation; iv) the original surface showing the relaxed triangulation.


Figure 14: A morphing sequence from the cartoon figure into a dinosaur head showing five equidistant steps.

Chapter 3.3.2. This sequence of steps is shown for the cartoon figure head in Figure 13.

Since the two triangulations were required to be equivalent, we now have the two surfaces $S$ and $S^{\prime}$ parametrized over two isomorphic base grids $B$ and $B^{\prime}$ using the respective parametrization functions $\Xi$ and $\Xi^{\prime}$. Thus, there exists a bijection $\theta$ from $B$ to $B^{\prime}$, which in turn creates a homeomorphism $\Phi$ from $S$ to $S^{\prime}$ by

$$
\Phi(p)=\Xi^{\prime} \circ \theta \circ \Xi^{-1}(p) \quad \forall p \in S
$$

Any intermediate model can now be generated by linearly interpolating between corresponding points $p$ and $\Phi(p)$ (Figure 14)

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    ${ }^{\dagger}$ Supported by the DFG research center Matheon

[^1]:    ${ }^{1}$ The PSurface library itself provides reading and writing facilities for PSurface objects in the AmiraMesh format 21]. The precise format is described in a text file distributed with the source code.

[^2]:    ${ }^{2}$ For increased robustness this cyclic ordering is not computed by comparing angles. Instead, a graph algorithm is used to determine a longest path in the subgraph of all neighbors of the node.

[^3]:    ${ }^{3}$ At the time of writing, the actual handling of parametrized boundaries in dune-grid has not been standardized yet. Therefore the details are still subject to change. Please consult the dune-grid documentation at www.dune-project. org for up-to-date information.

[^4]:    ${ }^{4}$ www.amira.com

