A Smoothness Energy without Boundary Distortion for Curved Surfaces

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Fig. 1. Solving an interpolation problem on an airplane. Using the Laplacian energy with zero Neumann boundary conditions (left) distorts the result near the windows and the cockpit of the plane: the isolines bend so they can be perpendicular to the boundary. The planar Hessian energy of Stein et al. [2018] (center) is unaffected by the holes, but does not account for curvature correctly, leading to unnatural spacing of isolines at the front and back of the fuselage. Our Hessian energy (right) produces a natural-looking result with more regularly spread isolines, unaffected by the boundary.

Current quadratic smoothness energies for curved surfaces either exhibit distortions near the boundary due to zero Neumann boundary conditions, or they do not correctly account for intrinsic curvature, which leads to unnatural-looking behavior away from the boundary. This leads to an unfortunate trade-off: one can either have natural behavior in the interior, or a distortion-free result at the boundary, but not both. We introduce a generalized Hessian energy for curved surfaces, expressed in terms of the covariant one-form Dirichlet energy, the Gaussian curvature, and the exterior derivative. Energy minimizers solve the Laplace-Beltrami biharmonic equation, correctly accounting for intrinsic curvature, leading to natural-looking isolines. On the boundary, minimizers are as-linear-as-possible, which reduces the distortion of isolines at the boundary. We discretize the covariant one-form Dirichlet energy using Crouzeix-Raviart finite elements, arriving at a discrete formulation of the Hessian energy for applications on curved surfaces. We observe convergence of the discretization in our experiments.

CCS Concepts: • Mathematics of computing → Discretization; Partial differential equations; Numerical differentiation; • Computing methodologies → Mesh geometry models;

Additional Key Words and Phrases: geometry, biharmonic, laplacian, hessian, curvature, interpolation, smoothing

1 INTRODUCTION

Smoothness energies are used as objective functions for optimization in geometry processing. A wide variety of applications exists: smoothness energies can be used to smooth data on surfaces, to denoise data, for scattered data interpolation, character animation, and much more. We are interested in quadratic smoothness energies formulated on triangle meshes.

It is desirable for a smoothing energy to have minimizers with isolines whose spacing does not vary much across the surface—the gradient of the function is sufficiently constant. When the gradient of the function is sufficiently constant, the function only changes very gradually, resulting in a smooth function. In the same vein, a good smoothing energy should have minimizers whose isolines are not distorted anywhere: their spacing is not influenced (on the interior) by the surface’s curvature, and they are not biased by the boundary of the surface—they behave locally as if the boundary were absent. Such behavior is relevant for applications where the boundary is not directly related to the actual problem that is being solved, e.g., when the boundary is an artificial result of faulty surface reconstruction resulting in a shape with many extraneous holes. One class of energies with the desired behavior in the interior are energies whose minimizers solve the biharmonic equation, the prototypical elliptic equation of order four. [Gazzola et al. 2010, viii]. Such energies are pertinent as smoothness energies in computer graphics applications [Jacobson et al. 2010].

One such energy is the squared Laplacian energy—the squared Laplacian of a function integrated over the surface. We henceforth
refer to the energy as simply the Laplacian energy. Its minimizers solve the biharmonic equation; as a result, they are very smooth, and their isolines behave well on curved surfaces, if the surfaces are closed. The energy’s most popular discretization, however, comes with zero Neumann boundary conditions. Thus, if a surface has boundaries, the minimizers are distorted near the boundary (see Figure 1), since at the boundary they are as-constant-as-possible.

The issue of boundary distortion is addressed by the Hessian energy of Stein et al. [2018]. For planar domains, they provide an energy whose minimizers solve the biharmonic equation and are as-linear-as-possible at the boundary. These boundary conditions lead to decreased distortion. The Hessian energy of Stein et al. [2018], however, is only defined for subsets of the plane $\mathbb{R}^2$. Stein et al. [2018] offer a way to compute an energy for curved surfaces, but, as they point out, their approach does not account for the curvature of the surface correctly. The approach of Stein et al. [2018] does not solve the biharmonic equation on curved surfaces; this leads to global distortions in the isolines of the solution (see Figure 1).

Contributions

1. **Generalized Hessian energy.** We generalize the Hessian energy to accommodate curved surfaces. Our new Hessian energy is

$$E(u) = \frac{1}{2} \int_{\Omega} (\nabla u) : (\nabla u) + \kappa \|\nabla u\|^2 \, dx,$$

(1)

where $\nabla$ is the covariant derivative of differential forms, $d$ is the exterior derivative, $\kappa$ is the Gaussian curvature, and $\nabla u$ denotes the contraction of two operators in all indices that corresponds to $A : B = \text{tr}(A^T B)$ (where the transpose $^T$ takes the metric into account). This energy

- corresponds to the Laplacian energy in the case of a domain without boundaries;
- corresponds to the Hessian energy of Stein et al. [2018] for surfaces in $\mathbb{R}^2$, $\frac{1}{2} \int_{\Omega} \|\mathbf{H}_{uu}\|^2 \, dx$, where $\mathbf{H}_{uu}$ is the $2 \times 2$ Hessian matrix of $u$, and $\|A\|_F$ is the Frobenius norm of $A$;
- has the as-linear-as-possible natural boundary conditions of the Hessian energy of Stein et al. [2018] for flat domains in $\mathbb{R}^2$. These boundary conditions lead to decreased distortion at the boundary.

Figure 1 shows how our Hessian energy manages to achieve the best of both worlds.

2. **Discretization.** We also introduce a discretization of this curved Hessian energy that uses Crouzeix-Raviart finite elements “under the hood”, but, after the energy matrix has been assembled, relies solely on piecewise linear hat functions. We observe convergence of the discretization for a wide variety of numerical experiments, given certain regularity conditions, and apply it to various smoothing and interpolation problems.

2 RELATED WORK

This work extends Stein et al. [2018]. They introduce a smoothness energy with higher-order boundary conditions whose minimizers are biased less by the shape of the boundary than energies using low-order boundary conditions such as zero Neumann. Our goal is to extend their approach to curved surfaces. Section 5.3.1 mentions that their work does not correctly account for curved surfaces, and this shortcoming is addressed in this work.

2.1 Smoothing energies

Smoothing energies are used for many applications in computer graphics, image processing, machine learning, and more. Quadratic smoothing energies are particularly interesting, since they are easy to work with and fast to optimize [Nocedal and Wright 2006]. The Laplacian energy is used for surface fairing and surface editing [Botsch and Kobbelt 2004; Crane et al. 2013a; Desbrun et al. 1999; Sorkine et al. 2004], for geodesic distance computation [Lipman et al. 2010], for creating weight functions used as coordinates in character animation [Jacobson et al. 2011; Weber et al. 2012], data smoothing [Weinkauf et al. 2010], image processing [Georgiev 2004], and other applications [Jacobson et al. 2010; Sýkora et al. 2014].

Geometric energies that share some of the properties of our Hessian energy have been studied in the past: in image processing, Hessian-like energies are popular for their boundary behavior, but their formulations in general do not extend to curved surfaces [Didas et al. 2009; Lefkimmiatis et al. 2011; Lysaker et al. 2003]. Similar energies are also used for data processing and machine learning, but are not discretized for polyhedral meshes there [Donoho and Grimes 2003; Kim et al. 2009]. Wang et al. [2015, 2017] explicitly enforce boundary conditions on a discrete quadratic fourth-order energy in order to make minimizers of the energy less dependent on the boundary shape, but do not discuss any continuous model corresponding to their method or which equations their minimizers satisfy.

Stein et al. [2018] present a Hessian energy for triangle meshes, however, minimizers of their discretization extended to $\mathbb{R}^3$ do not fulfill the biharmonic equation, leading to artifacts that are discussed in detail in Section 7. Liu et al. [2015] explicitly enforce higher-order boundary conditions on a smoothness energy based on a fourth-order PDE. Their energy, however, is in general not quadratic, and the boundary conditions are different than the ones presented in this article, as they are missing the as-linear-as-possible property.

A special case of a quadratic smoothness energy is the Dirichlet energy, which solves the harmonic equation $-\Delta u = 0$, a simpler version of our biharmonic equation $\Delta^2 u = 0$. The Dirichlet energy can be used, for example, to create smooth character deformations [Baran and Popovic 2007; Joshi et al. 2007; Weber et al. 2007], and for image processing [Levin et al. 2004]. While the Dirichlet energy has advantages, such as a discrete maximum principle, which is preserved in some discretizations [Wardetzky et al. 2007], there are disadvantages due to the energy being first-order: because of reduced freedom around constraints, minimizers fail to be smooth, which can lead to artifacts when applied to shape deformation [Jacobson et al. 2011, Fig. 9], or worse results in image processing [Peter et al. 2016]. Higher-order smoothness energies, such as the ones derived from the biharmonic equation, are better at fitting to existing data, and tend to distort results less [Georgiev 2004; Jacobson et al. 2011, 2012; Weber et al. 2012]. Additionally, the Dirichlet energy does not admit higher-order boundary conditions
work of Knöppel et al. [2015], where it is used to compute stripe
∇
we use to derive the natural boundary conditions of our Hessian
An important part of the discretization of our curved Hessian energy
2.3 Discretization of the vector Dirichlet energy
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A main theme in our work is the difficulty of generalizing expres-
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(like biharmonic energies), which makes it more difficult to use
as a smoothing energy without boundary bias.

2.2 Generalizing the Hessian energy to curved surfaces
A main theme in our work is the difficulty of generalizing expres-
sions formulated on flat domains to curved surfaces. The presence
of curvature will result in an additional term in the definition of our
energy, which is absent in the planar Hessian energy of Stein et al.
[2018]. This mirrors many other areas of geometry where, with
the introduction of curvature, properties of flat domains cease to apply.

One such example of curvature making calculations more elabor-
ate is parallel transport. While parallel transport of vectors is
trivial on flat surfaces, this is no longer true for curved surfaces.
In the presence of curvature, the parallel transport of a vector along a
closed curve might result in a different vector than the initial one
[Harder et al. 2006, pp. 156-157]. The difficulties that this phenomenon
introduces to applications are discussed, for example, by Bergou et al.
[2008], Crane et al. [2010]; Polthier and Schmies [1998]; Ryët al.
[2009]. Our discretization method simplifies the treatment of paral-
lel transport by employing linear finite element basis functions that
are only supported on two adjacent triangles, since this necessitates
discontinuous basis functions, this approach is less common.

Another instance of difficulties arising from the curved setting
occurs in the numerical analysis of finite element methods. In order
to apply standard finite element methods to curved surfaces, the
discretization has to account for the curvature of the surface. For the
case of the Poisson equation, for example, this can be either achieved
by inscribing all the vertices on the limit surface while imposing
triangle regularity conditions [Dziuk 1988], or by demanding a
specific kind of convergence of the vertices as well as the normals
of the mesh [Hildebrand 2006; Wardetzky 2006] together with
specific triangle regularity conditions. Similarly, in some of our
own numerical experiments, we require vertex inscription and the
triangle regularity condition to achieve convergence.

2.3 Discretization of the vector Dirichlet energy
A classical smoothness energy for a surface \( \Omega \subseteq \mathbb{R}^3 \) is the Laplacian
energy with zero Neumann boundary conditions. When using this
method, one solves the optimization problem
\[
\min_u \frac{1}{2} \int_{\Omega} |\nabla u|^2 \, dx \quad \text{subject to} \quad \frac{\partial u}{\partial n} |_{\partial \Omega} = 0 \,,
\]
where \( \Delta \) is the Laplace-Beltrami operator, and \( \frac{\partial u}{\partial n} |_{\partial \Omega} \) is the normal derivative at the boundary. For meshes that are non-planar
using methods such as the mixed finite element method (FEM) [Jacobs et al. 2010]. Using this method,
the zero Neumann boundary condition does not need to be imposed

\[\text{argmin}_{u} \quad \frac{1}{2} \int_{\Omega} |\nabla u|^2 \, dx \quad \text{subject to} \quad \frac{\partial u}{\partial n} |_{\partial \Omega} = 0 \,,
\]

1 Of course, simply minimizing (2) results in the zero function. However, when combined
with additional Dirichlet boundary conditions, this gives a nontrivial result for the
biharmonic equation \( \Delta^2 u = 0 \). That is, when combined with the additional energy term
\( \int_{\Omega} |f|^2 \, dx \) it gives a result for the biharmonic Poisson-type equation \( \Delta^2 u = f \).
on top of the discretization, it is simply “baked in” by squaring the classical cotan Laplacian. The cotan Laplacian is also known as the Lagrangian linear FEM for the Poisson equation (it goes back to Duffin [1959] and MacNeal [1949], and its convergence for the Poisson equation was shown by Dziuk [1988]).

The minimizers of \( E_{\varphi} \), however, are biased by the shape of the boundary. Their isolines are significantly distorted near the domain boundary: they are perpendicular to it as they have to fulfill the zero Neumann boundary conditions (as-constant-as-possible). Simply removing the zero Neumann boundary conditions, and minimizing the Laplacian energy without any boundary conditions is not a good alternative. Minimizations without explicit boundary conditions lead to natural boundary conditions. The natural boundary conditions of the Laplacian energy are too permissive [Stein et al. 2018, Fig. 3]. This behavior is one of the motivations for the Hessian energy of Stein et al. [2018]. It is formulated as the following minimization problem. For a surface \( U \subseteq \mathbb{R}^2 \),

\[
\text{argmin}_{u} \frac{1}{2} \int_{U} \nabla u \cdot \nabla u \, dx
\]

where \( \nabla u \) is the \( 2 \times 2 \) Hessian matrix of \( u \), and \( A : B = \text{tr}(A^T B) \). Minimizers of this energy solve the biharmonic equation in \( \mathbb{R}^2 \). Its natural boundary conditions lead to as-linear-as-possible behavior on the boundary. This makes minimizers less biased than the zero Neumann boundary condition.

Stein et al. [2018] demonstrate the benefits of the natural boundary conditions of the Hessian energy with applications for curved surfaces in \( \mathbb{R}^2 \) as well. Their discretization of the planar Hessian energy for curved surfaces is achieved by extending every operator involved in the \( \mathbb{R}^2 \) discretization to three dimensions. This approach (the discretization, as well as the smooth formulation) does not account for the curvature of surfaces correctly, and its minimizers do not solve the biharmonic equation on curved surfaces [Stein et al. 2018, Section 5.3.1]. We refer to this generalization as the planar Hessian energy \( E_{\mathbf{H}} \) when talking about it in the context of curved surfaces. This planar Hessian energy is suitable for some applications, but leads to global deviations from the natural-looking isolines produced by \( E_{\mathbf{A}}(u) \) (see Figure 1) or an implementation of the Hessian energy which does account for curvature (see Figure 2) in others.

4 WARM-UP: THE DIRICHLET ENERGY ON CURVED SURFACES

As a warm-up, we consider the simple and well-known Dirichlet energy: it is easy to generalize to curved surfaces. We will perform the calculation for this generalization here. The calculation is well-known, and this didactic exercise will inform our generalization of the planar Hessian energy to curved surfaces later.

4.1 From the energy to the PDE

Let \( H^k \) denote the Sobolev space of real-valued functions with \( k \) weak derivatives in \( L^2 \). The Dirichlet energy for domains \( U \subseteq \mathbb{R}^2 \) is defined, for \( u \in H^2(U) \), as

\[
E_{\varphi}(u) := \frac{1}{2} \int_{\Omega} \nabla u \cdot \nabla u \, dx ,
\]

where \( \nabla \) is the vector of partial derivatives of \( u \), \( \nabla u = (\partial_x u \quad \partial_y u)^T \), the normal two-dimensional gradient in \( \mathbb{R}^2 \).

Minimizers of the Dirichlet energy solve the Laplace equation [Evans 2010]. Indeed, consider the variation

\[
\begin{align*}
\int_{U} (\partial_i u) (\partial_j u) \, dx &= 0 \quad \forall v \in H^2(U) ,
\end{align*}
\]

where \( \partial_i \) is a partial derivative, and summation over repeated indices is implied. This is a standard technique of variational calculus. Using integration by parts (where \( \mathbf{n} \) is the boundary normal)

\[
\begin{align*}
0 &= \int_{U} (\partial_i u) (\partial_j u) \, dx \\
&= \int_{\partial U} (\partial_i u) v \, \mathbf{n}_i \, dx - \int_{U} (\partial_i \partial_j u) v \, dx .
\end{align*}
\]

Here a boundary term appeared as a result of integration by parts. The second term of the second line corresponds to the standard two-dimensional planar Laplacian \( \Delta \equiv \nabla \cdot \nabla \), and so we conclude

\[\text{We choose to formulate this energy for } u \in H^2(U) \text{, although it is well-defined for } u \in H^1(U) \text{, since we will continue our calculations with the same } u \text{ right away, and we will need additional smoothness.}\]
that minimizers of the energy $E_{\nabla^2}(u)$ fulfill the two-dimensional planar Laplace equation $\Delta u = 0$. The additional boundary term, the first term of the second line in (6), determines the natural boundary conditions of the Dirichlet energy. They are called natural boundary conditions because they naturally emerge from solving the variational problem over the set of all functions, without explicitly enforcing additional boundary conditions. In this case we can see that the natural boundary conditions are zero-Neumann boundary conditions:

$$\partial_u n_i = \nabla u \cdot n = 0 \quad \text{on } \partial U . \quad (7)$$

4.2 From the PDE to a new energy

We now generalize the Dirichlet energy to curved surfaces. This means we are looking for an energy whose minimizers solve a curved version of the Laplace equation, and fulfill a curved version of the natural boundary conditions (7). While we were able to write the calculations in terms of coordinates in the flat setting, this is much harder to do in the curved setting. This is why we perform calculations in the curved setting in a coordinate-free fashion.

The curved analog of the planar Laplace equation is $\Delta u = 0$, where $\Delta$ is the Laplace-Beltrami operator [Jost 2011, Chapter 3]. It holds for a function $u \in H^2(\Omega)$ (where $\Omega$ is a compact surface immersed in $\mathbb{R}^3$) that

$$\Delta u = \delta du , \quad (8)$$

where $d$ is the exterior derivative and $\delta$ is the codifferential, the (formal) dual of the exterior derivative under integration by parts. For planar surfaces, the Laplace-Beltrami operator $\Delta$ corresponds to $-\Delta$.

We start with an integral formulation of the Laplace equation, and then use integration by parts. For all $\nu \in H^2(\ Omega)$ it must hold that

$$0 = \int_{\Omega} (\Delta u) \nu \, dx = \int_{\Omega} (\delta du) \nu \, dx$$

$$= - \int_{\partial \Omega} \langle du, n \rangle \nu \, dx + \int_{\Omega} (du) \cdot (du) \, dx ,$$

where the natural (metric-independent) pairing of one-forms and vectors is indicated using the angle bracket, and $\cdot$ is the dot product of one-forms.

Using the definition of the gradient $\nabla$ on curved surfaces, $\nabla u \cdot w := \langle du, w \rangle$ for a vector $w$ (where $\cdot$ is the dot product of vectors and the angle bracket $\langle \cdot, \cdot \rangle$ denotes the pairing of a one-form with a vector) [Jost 2011, (3.1.16)], we can write

$$0 = - \int_{\partial \Omega} \nabla u \cdot n \, dx + \int_{\Omega} \nabla u \cdot \nabla u \, dx . \quad (9)$$

Walking back through the variation from (5), this now motivates the definition of a curved Dirichlet energy

$$E_{\nabla^2}(u) := \frac{1}{2} \int_{\Omega} \nabla u \cdot \nabla u \, dx . \quad (10)$$

We have shown that minimizers of this energy solve the curved Laplace equation, and by the boundary term in (9) it is also clear that minimizers fulfill a curved zero Neumann boundary condition:

$$\nabla u \cdot n = 0 \quad \text{on } \partial U . \quad (11)$$

Thus we have successfully generalized the Dirichlet energy to curved surfaces. Even though we went through the work of using differential geometric operators, we ended up with something quite similar to what we started with, but with $\nabla$ replaced by $\nabla$. For more complicated energies this will no longer be the case.

5 THE HESSIAN ENERGY ON CURVED SURFACES

We now seek to derive a smooth Hessian energy on surfaces that generalizes the Hessian energy in $\mathbb{R}^2$, while ensuring that minimizers of the energy solve the biharmonic equation. This will follow the approach we used in Section 4 to generalize the planar Dirichlet energy to curved surfaces.

5.1 From the energy to the PDE

For the planar Hessian energy $E_{\nabla^2}$, it is a straightforward calculation to prove that minimizers fulfill the biharmonic equation. This calculation is mentioned, for example, in Stein et al. [2018, Section 4.2], and we will repeat it here for convenience. Our setting is a compact planar domain $U \subseteq \mathbb{R}^2$. The linear equation fulfilled by minimizers of (3) derived with standard variational calculus is: find $u \in H^2(U)$ such that

$$\int_U (\partial_i \partial_j u)(\partial_i \partial_j \nu) \, dx = 0 \quad \forall \nu \in H^2(U) , \quad (12)$$

where, as before, $\partial_k$ is a partial derivative, and summation over repeated indices is implied. Using integration by parts (where $n$ is the boundary normal) we know that

$$0 = \int_U (\partial_i \partial_j u)(\partial_i \partial_j \nu) \, dx$$

$$= \int_{\partial U} (\partial_i \partial_j u)(\partial_i \nu) n_i \, dx - \int_U (\partial_i \partial_j u)(\partial_i \nu) \, dx$$

$$= \int_{\partial U} (\partial_i \partial_j u)(\partial_i \nu) n_i - (\partial_i \partial_j u)n_j \, dx$$

$$+ \int_U (\partial_i \partial_j \partial_i \partial_j u) \nu \, dx . \quad (13)$$

Since all partial derivatives commute in the plane, in the very last term we can write $\partial_i \partial_j \partial_i \partial_j u = \partial_i \partial_i \partial_j \partial_j u = -\Delta u$. As a result, we can conclude that minimizers of the Hessian energy satisfy the biharmonic equation with some additional boundary terms. This commutation will not be that easy for curved surfaces.

After some rearranging, these boundary terms can be seen to imply the natural boundary conditions

$$n^i \nabla u_i \cdot n = 0 \quad \text{on } \partial U$$

$$\nabla \nabla u \cdot n + \nabla (n^i \nabla u_i) \cdot t = 0 \quad \text{on } \partial U , \quad (14)$$

where $n$ is the normal vector at the boundary, and $t$ is the tangential vector of the (oriented) boundary. A derivation of (14) can be found in the work of Stein et al. [2018, Section 4.3].

A naive approach to a Hessian energy for curved surfaces. Since our goal is to generalize the Hessian energy for surfaces, it seems natural to simply replace the planar Hessian $\nabla^2 u$ with an analog for curved surfaces, and minimize this generalization of
we can already see that we will not be able to repeat our approach which degenerates to the Laplace-Beltrami operator

\[ \Delta = \delta^2 + \delta d \]

where \( \delta \) is the (formal) dual covariant derivative. The formal dual is defined via integration by parts on a closed manifold \( M, \int_M X : \nabla \omega \, dx = \int_M \nabla^* X \cdot \omega \, dx \). It holds that

\[ \Delta = \nabla^* \nabla + \text{Ric} \quad (18) \]

where \( \text{Ric} \) is the Ricci curvature tensor [Petersen 2006, Chapter 7]. This formula dates back to Bochner [1946] and Weitzenböck [1885]. It is used, together with the fact that \( d^2 = 0 \), to continue our calculation from (17).

\[
\int_{\Omega} (\delta \delta u) \cdot (\delta v) \, dx = \int_{\Omega} ((\delta + \delta \delta)u) \cdot (\delta v) \, dx \\
= \int_{\Omega} (\nabla^* \nabla u) \cdot (\delta v) + \text{Ric}(du, dv) \, dx \\
= - \int_{\partial \Omega} n^i (\nabla du)_{ij} \cdot (\delta v)_j \, dx \\
+ \int_{\Omega} (\nabla du) : (\nabla dv) + \text{Ric}(du, dv) \, dx ,
\]

where indices have been added to make clear which contraction happens in which index.

The term involving the Ricci curvature tensor \( \text{Ric} \) can be further simplified. For the case of two-dimensional manifolds we know that we can write the Ricci curvature tensor as simply

\[ \text{Ric} = \kappa g \quad (20) \]

where \( \kappa \) is the Gaussian curvature, i.e., half the scalar curvature [Petersen 2006, pp. 38-41].

Putting (17), (19), and (20) together then gives

\[
0 = - \int_{\partial \Omega} \langle \delta \delta u, n \rangle \, v + n^i (\nabla du)_{ij} \cdot (\delta v)_j \, dx \\
+ \int_{\Omega} (\nabla du) : (\nabla dv) + \kappa \, du \cdot dv \, dx .
\]

This is, in the case of a planar surface (for which it holds \( \kappa = 0 \), exactly the term from our earlier calculation with the planar Hessian energy from (13). Here we also see why minimizers of the naive Hessian energy \( E_{\text{HF}} \) do not solve the biharmonic equation on curved surfaces: the energy \( E_{\text{HF}} \) lacks the curvature correction term involving \( \kappa \) (see Figure 3).

The result from (21) motivates the definition of the following curved Hessian energy:

\[ E(u) := \frac{1}{2} \int_{\Omega} (\nabla du) : (\nabla du) + \kappa |du|^2 \, dx \quad (22) \]

Minimizers of the energy \( E \) solve the biharmonic equation on a surface in \( \mathbb{R}^3 \), unlike minimizers of \( E_{\text{HF}} \).

It remains to check what the natural boundary conditions of \( E \) are. We can find them by checking which biharmonic functions \( u \) fulfill the boundary terms

\[ 0 = \int_{\partial \Omega} \langle \delta \delta u, n \rangle \, v + n^i (\nabla du)_{ij} \cdot (\delta v)_j \, dx \quad \forall v \in H^4(\Omega) .
\]
We use the same strategy as Stein et al. [2018, Section 4.3]: testing with specific subsets of all valid test functions. These subsets are purpose-built to expose the natural boundary conditions of the energy. First, we test with all functions \( v \) that vanish on the boundary, and thus only have nonzero differential in the normal direction \( (v = 0, \langle dv, w \rangle = gn \cdot w \) for some smooth \( g \). It follows that
\[
\n^t (\nabla u)_{ij} n = 0 \quad \text{on } \partial \Omega,
\]
(23)
i.e., the (curved) Hessian of the solution is linear across the boundary; the second derivative of the function across the boundary is zero. This mirrors the “as-linear-as-possible” condition of Stein et al. [2018, (17)].

Using the same strategy of testing the expression with a specific subset of functions to expose boundary behavior, if we plug in all functions that have zero differential in the normal direction at the boundary \( (\langle dv, n \rangle = 0) \), we get
\[
(\delta_{ij} + \delta_{j\Omega, j\partial \Omega} (n^t (\nabla u))_{ij}) = 0 \quad \text{on } \partial \Omega,
\]
(24)
where \( 1_{\partial \Omega} \) is the natural projection of one-forms on the surface to one forms on the boundary, and the subscript on the codifferential implies that this is the codifferential of the boundary manifold in the index \( j \). This mirrors the condition from Stein et al. [2018, (18)]. In fact, the two natural boundary conditions (23) and (24) of the Hessian energy are exactly the ones of the planar Hessian energy if the domain is a planar surface.

The Hessian energy natural boundary conditions. Like the natural boundary conditions of \( E_{\Pi} \) from Stein et al. [2018, Section 4.3], the natural boundary conditions (23) and (24) of the Hessian energy \( E \) guarantee that its minimizers

- continue linearly across the boundary in the normal direction \( (n^t (\nabla u))_{ij} n^t = 0 \),

as discussed by Stein et al. [2018, Section 4.3]. Both boundary conditions are fulfilled by minimizers of \( E \) in the absence of explicitly enforced boundary conditions.

On planar surfaces, these boundary conditions mean that the null space of the energy contains all linear functions, in contrast to the Laplacian energy with zero Neumann boundary conditions \( E_{\Pi} \), whose null space only contains constant functions. On closed surfaces, the null space of \( E \) and \( E_{\Pi} \) is the same: all constant functions.

The natural boundary conditions of the Hessian energy have a physical interpretation. Consider a deforming flat thin plate where displacement is modeled by the function \( u \). The plate is not clamped or supported at the boundary in any way; it is a free plate. Then the conditions (24) are the boundary conditions fulfilled by \( u \) [Courant and Hilbert 1924, pp. 206-207]. These boundary conditions go back at least as far as Rayleigh [1894, p. 355].

Its natural boundary conditions make the Hessian energy a good choice for ignoring the boundary as much as possible, while maintaining biharmonic behavior everywhere away from the boundary (see Figure 4 where they are contrasted with zero Neumann boundary conditions).

6 DISCRETIZATION

We offer a discretization for the curved Hessian energy \( E \) derived in Section 5. The approach presented here is a simple method using only linear finite elements, intended to make the Hessian energy easily accessible. There are, however, other conceivable ways to discretize this energy, such as, for example, higher-order conforming finite elements [Braess 2007, II.5].

![Fig. 4. Using the Laplacian energy \( E_{\Pi} \) (top) for scattered data interpolation gives a result that is influenced by the boundary: adding holes makes the isolines near them bend towards the holes. Our Hessian energy \( E \) (bottom) is less distorted at the holes and produces a very similar result without and with holes.](image)

![Fig. 5. A scalar Crouzeix-Raviart basis function for the edge \( e \) (top left). The parallel and perpendicular one-forms for the edge \( e \), represented by their dual vectors (top right). Crouzeix-Raviart functions and their sums are, in general, discontinuous. Continuity is only guaranteed at edge midpoints (bottom).](image)
we extend the surface here
to try to assign curvature

Fig. 6. For the boundary of a continuous, piecewise linear surface (top) there is no way to uniquely assign curvature at the boundary. The surface can be extended in many different ways that yield different curvatures at the boundary, examples leading to positive (bottom left), no (bottom center), and negative (bottom right) curvature are shown.

6.1 Computing the Hessian energy

Discretizing the Hessian energy $E$ (22) as written would require us to discretize functions that can be differentiated twice. To avoid this complication we use the mixed finite element method [Boffi et al. 2013] by introducing an intermediate variable $w = du$ and formulate the problem of minimizing $E$ as

$$\arg\min_u \frac{1}{2} \int_{\Omega} (\nabla w) : (\nabla w) + \kappa |w|^2 \, dx, \quad w = du. \tag{25}$$

Using Lagrange multipliers to enforce the constraint $w = du$, we can write the optimization problem as the saddle problem (where our goal is finding a stationary point)

$$\text{saddle}_{w, w, \lambda} \frac{1}{2} \int_{\Omega} (\nabla w) : (\nabla w) + \kappa |w|^2 \, dx - \int_{\Omega} \lambda \cdot (w - du) \, dx. \tag{26}$$

We discretize the space of scalar functions (containing $u$) using standard continuous, piecewise linear functions which are a very commonly used finite element. Definitions are found, for example, in Braess [2007, II.5]. The basis of this discrete space consists of the $\varphi_i$, $i = 1, \ldots, n$, sometimes called “hat functions” (see inset). We write $u = \sum_i u_i \varphi_i$, and we have the vector $u = (u_1, \ldots, u_n)^T$.

The space of one-forms (containing $w$) is discretized using Crouzeix-Raviart one-forms (CROFs), which are described in Section 6.2. The basis of this discrete space are the functions $\eta_i$, $i = 1, \ldots, m$.

We write $w = \sum_i w_i \eta_i$, and we have the vector $w = (w_1, \ldots, w_m)^T$.

Using these discretizations we can construct the one-form Dirichlet matrix

$$L_{ij} = \int_{\Omega} (\nabla \eta_i) : (\nabla \eta_j) \, dx,$$

the differential matrix

$$D_{ij} = \int_{\Omega} \eta_i \cdot d\phi_j \, dx,$$

the mass matrix

$$M_{ij} = \int_{\Omega} \eta_i : \eta_j \, dx,$$

and the curvature matrix

$$K_{ij} = \int_{\Omega} \kappa \eta_i : \eta_j \, dx.$$

The matrix entries are provided in Appendix A.

Using these matrices, we write the discrete version of (26) as seeking a critical point of the expression

$$\frac{1}{2} w^T (L + K) w - \lambda^T (Mw - Du) ,$$

for $u \in \mathbb{R}^n$, $w, \lambda \in \mathbb{R}^m$. Differentiating with respect to $\lambda$ gives $Mw = Du$. As $M$ is invertible, we get the system

$$\arg\min_u u^T D^T M^{-1} (L + K) M^{-1} Du . \tag{27}$$

This optimization problem can now be solved with a variety of constraints, or mixed with other energy terms, depending on the application.

6.2 Crouzeix-Raviart One-Forms

While there are multiple approaches to discretizing tangent one-forms for triangle meshes, we choose to base our approach on Crouzeix-Raviart finite elements (see Section 2 for a discussion). The advantage of this approach is its simplicity. Crouzeix-Raviart basis functions are only ever nonzero on two adjacent triangles, so every basis function lives on an intrinsically flat domain: the two triangles can be unfolded without distortion. This means that our discretization will account for curvature correctly in the end, without having to explicitly address issues like parallel transport during construction.

Fig. 7. The first nonzero eigenvector of the Laplacian energy $E_{\Delta}$ (left), the Hessian of Stein et al. [2018] (center), and the curved Hessian energy $E$ (right). The eigenvectors of $E_{\Delta}$ and $E$ look similar, since they both discretize the biharmonic energy. The method of Stein et al. [2018] visibly disagrees.
6.2.1 Introduction to Crouzeix-Raviart. The scalar Crouzeix-Raviart basis function for the edge $e_{ij}$ is defined to be 1 on the edge itself, $-1$ on the two vertices $k$, $l$ opposite the edge, and linear on the two triangles $T_{ijk}, T_{ijl}$ [Braess 2007, p. 109]. For boundary edges, only one triangle needs to be considered. As a result, it is 0 on the midpoints of the edges $e_{ik}, e_{kl}, e_{lj}, e_{ij}$ (see Figure 5, left). The scalar Crouzeix-Raviart element is not continuous, except at the midpoints of edges. This makes it a non-conforming element, and if it is used in a Galerkin method, one speaks of the discontinuous Galerkin method. Despite being nonconforming, it is known to converge for certain problems, most notably the Poisson equation in $\mathbb{R}^2$. [Braess 2007, III, Theorem 1.5].

6.2.2 One-Forms. The scalar Crouzeix-Raviart element can be used to define a finite element space for one-forms. At the midpoint of every edge $e$ of a flat triangle pair, the space of one-forms is spanned by the two forms $\omega_e^l, \omega_e^r$, such that
\[
\langle \omega_e^l, \tau_e \rangle = 1,
\langle \omega_e^l, \nu_e \rangle = 0
\]
\[
\langle \omega_e^r, \tau_e \rangle = 0,
\langle \omega_e^r, \nu_e \rangle = 1,
\]
where $\nu_e$ is the (oriented) perpendicular vector of the edge $e$ in each triangle, $\tau_e$ is the (oriented) tangent of the edge $e$, and the angle bracket $\langle \cdot, \cdot \rangle$ denotes the pairing of a form with a vector. See Figure 5 (right) for an illustration. The definition of $\omega_e^l$ depends on which triangle one is in, but only in an extrinsic way: in the intrinsic geometry of the triangle pair, the edge is completely flat, thus the two covectors $\omega_e^l$ defined in each triangle are the same covector intrinsically. Because of this, both $\omega_e^l$ and $\omega_e^r$ can be easily extended to the triangles adjacent to $e$: since the triangle pair (or the one triangle) is intrinsically flat, parallel transport along the triangles is trivial, and we can easily extend the definition of $\omega_e^l$ and $\omega_e^r$ to the interior of the triangles adjacent to $e$.

If $b_e$ is the Crouzeix-Raviart basis function for the edge $e$, then we define its two CROF basis function as
\[
b_e^l \equiv \omega_e^l b_e
\]
\[
b_e^r \equiv \omega_e^r b_e.
\]

Defined this way, CROFs have the correct notion of parallel transport, without having to explicitly account for it. Consider a path $y$ through all edge midpoints of edges emanating from a vertex $v$ in a counterclockwise direction (see inset). We start with a single tangent vector on the midpoint of one edge, corresponding to a combination of two basis functions, and see what angle we pick up when going around the vertex $v$ using our basis functions. We now go along the path $y$, moving from edge to edge by choosing successive basis functions so that the sum of the basis functions from two adjacent edges is constant on the shared triangle. Doing that corresponds exactly to parallel transport on a cone manifold: the tangential part of the vector at each edge does not change extrinsically at all when crossing the edge along $y$. The perpendicular basis function jumps extrinsically: the angle between normal vectors on each side of the edge is $\pi$ minus the dihedral angle of the edge. At the end of our journey along $y$, when we are back at our original edge, our starting vector picked up angle defect corresponding to the discrete curvature of the mesh. The CROF basis functions have accounted for the discrete curvature of the mesh in the sense of curvature on cone manifolds [Wardetzky 2006] without having to explicitly account for parallel transport during the construction of the basis functions.

Since every basis function is only supported on at most two triangles, the matrices $L, M, D, K$ will be sparse. The matrix $M$ is diagonal, which makes it easy to invert. The matrix entries can be found in Appendix A.

6.2.3 The curvature term. Special care needs to be applied when computing the matrix $K$. The Gaussian curvature $\kappa$ of an intrinsically flat pair of triangles would appear, at first, to be 0. But actually, the Gaussian curvature of a polyhedron is entirely concentrated on its vertices (and is zero anywhere else). The integrated Gaussian curvature at a vertex is also known as the angle defect
\[
\kappa_v \equiv 2\pi \sum_{f \in N(v)} \theta_v^f,
\]
where the sum is over all faces $f$ in the set of faces containing the vertex $v$, and $\theta_v^f$ is the angle at vertex $v$ in face $f$ [Grinspun et al. 2006]. The idea of angle defects is very old: it goes back all the way to at least Descartes c. 1630, who showed that the sum of all angle defects of a polyhedron with spherical topology is $4\pi$ [Federico 1982].

We thus interpret the Gaussian curvature of the polyhedron as a collection of delta functions at every vertex, i.e.
\[
\kappa \equiv \sum_v \kappa_v \delta_v,
\]

error in eigenvalue
\[
\begin{array}{c|c|c|c}
\text{average edge length} & 10^{-1} & 10^{-2} & 10^{-3} \\
\hline
\text{mesh gen. every step (w. regularity)} & O(h) & & \\
\text{mesh gen. every step (no regularity)} & & & \\
\end{array}
\]

Fig. 8. Computing the fourth eigenvalue of the Hessian energy $E$ on an ellipse that was distorted in the third dimension (bottom left). Both refinement through Loop subdivision and projection to a given smooth surface, as well as generating a planar mesh of the desired resolution with regular triangles at every step and then projecting to a given smooth surface show convergence to the highest resolution. For simple mesh generation without triangle regularity no convergence is observed.
where \( \delta_v \) is the Dirac delta. This means that the integral of \( \kappa g \), where \( \kappa \) is the Gaussian curvature and \( g \) is any continuous function over the triangle \( T_{ijk} \) with vertices \( i, j, k \), can be written as:

\[
\int_{\Omega} \kappa g \, dx = \sum_v \kappa_v g(v) .
\]  

(32)

If the function \( g \) itself is only continuous in each triangle, then we need to distribute the contribution of each triangle accordingly. Let \( s_{\kappa,f} > 0 \) for each vertex \( v \) and each face \( f \) in the neighborhood of \( v \) be coefficients that average the contribution of each face at a vertex, i.e., the sum of the \( s_{\kappa,f} \) over all faces \( f \) in the neighborhood of \( v \) is one. Then

\[
\int_{\Omega} \kappa g \, dx \approx \sum_v \kappa_v \sum_{f \in N(v)} s_{\kappa,f} g_f(v) ,
\]

where \( g_f \) is the function \( g \) in the triangle \( f \) and \( N(v) \) is the set of all faces in the neighborhood of \( v \). We choose to average by tip angle, which corresponds to an integral along a small circle around the vertex. We did not explore other reasonable choices, such as averaging by face area. This formula is used to compute the entries of \( K \), they are given in Appendix A.

One remaining issue with the angle defect as Gaussian curvature is that the angle defect is not defined at boundary vertices. The problem stems from the fact that the notion of curvature at the boundary of meshes (continuous, piecewise linear surfaces) is not in and of itself meaningful: by choosing to extend the surface in different ways at the boundary we can achieve any arbitrary Gaussian curvature, as can be seen in Figure 6. We choose to set the angle defect to 0 for all boundary vertices, thereby choosing the most developable (intrinsically linear) extension of all possible extensions. This fits in with our as-linear-as-possible boundary conditions, but differs from some conventions of angle defect at the boundary, which define it as the sum of tip angles subtracted from \( \pi \) (which is a discretization of geodesic curvature).

6.3 Observed Numerical Convergence

Using our CROF discretization of the Hessian energy to solve a variety of problems, we observe convergence on the order of the average edge length \( h \) (Figure 9). As can be seen in Figure 8, a successful strategy for obtaining convergence is making sure that the vertices are contained in a smooth surface, and then either refining the mesh through Loop subdivision [Loop 1987] with a fixed smooth boundary, or generating meshes that fulfill the triangle regularity condition: the ratio of circumcircle to incircle of each triangle (the triangle regularity) is bounded from above and below independent of refinement level. This condition is standard for finite elements [Braess 2007, Definition 5.1 (uniform triangulation)]. The order of convergence and the triangle regularity condition correspond to the discretization of the Laplacian energy with zero Neumann boundary conditions, \( E_{\Delta_v} \), with mixed FEM in the flat setting [Jacobson et al. 2010; Scholz 1978]. However, we do not have a proof of convergence for our method to confirm this convergence rate.

Our method correctly reproduces the first eigenvector of the Laplacian energy on closed surfaces in the experiment proposed by Stein et al. [2018, Section 5.3.1] on a refined mesh (Figure 7). As mentioned in Stein et al. [2018, Section 4.5], discretizations can sometimes exhibit spurious modes in the kernel of the energy, which lead to wrong solutions. We have not proved that this does not happen for our CROF discretization of the Hessian energy, but we have not observed it in our experiments (see Figure 10 for the cheeseman example domain mentioned in Stein et al. [2018, p. 7]).

Further experiments can be found in Appendix B: Figures 15 and 16 feature additional convergence experiments confirming the order of convergence.
the 6 smallest eigenvalues

Fig. 10. The six lowest eigenvalues of the Hessian energy discretized with CROF on the cheese-man (top). As expected, there are only three zero eigenvalues. The three lowest eigenvectors (bottom) are the linear functions, which corresponds to the smooth Hessian energy.

7 APPLICATIONS

We implement the optimization of (27) by constructing a sparse matrix in C++ using Eigen [Guennebaud et al. 2010], and then manipulating and optimizing it in MATLAB [MATLAB 2019] with mex. For linear equality constraints, we use the optimizer of Jacobson et al. [2019a, min_quad_with_fixed] via the library of Jacobson [2019]. Using this approach, complicated constraints are also possible, such as linear and quadratic inequality constraints for more complicated applications. Since the Hessian energy is a quadratic energy, optimizers using the interior point method (such as the solver of Andersen and Andersen [2000]) are appropriate.

7.1 Scattered data interpolation

Like any smoothness energy, the Hessian energy can be used for scattered data interpolation. One solves the following minimization problem, for some given interpolation data \( u(x_i) = f_i, \quad i = 1, \ldots, n \)

\[
\arg\min_{u} E(u) \quad u(x_i) = f_i, \quad i = 1, \ldots, n.
\]  

(33)

As long as at least three interpolation points are provided, this problem has a solution. This is because the null space of the Hessian energy can have at most all linear functions in it, which is a three-dimensional space, and the null space of the Laplacian energy with zero Neumann boundary conditions contains only constant functions, which is a one-dimensional space [Stein et al. 2018].

The choice of smoothness energy will greatly influence the quality of the result. The Laplacian energy with zero Neumann boundary conditions, \( E_{\Delta z} \), is a popular method, since it produces smooth, evenly spaced isolines, which results in natural-looking interpolation and extrapolation. This is because the gradient of the solution is relatively uniform across the surface. As can be seen in Figure 11, our curved Hessian energy \( E \) reproduces the desirable behavior of the Laplacian energy for surfaces without boundary. The implementation of the planar Hessian energy \( E_{\Pi} \) for curved surfaces by Stein et al. [2018] fails to do so: the distance between the isolines varies greatly, for example on the legs. The isolines also experience significant bunching at the rump and back of the horse.

On the other hand, the Laplacian energy is known to produce bias near domain boundaries due to its low-order boundary conditions: isolines of solutions bend so they can be perpendicular to the boundary. This was one of the motivations of Stein et al. [2018], and thus their planar Hessian energy minimizes the influence of the boundary by employing natural boundary conditions that make the function as-linear-as-possible. Figure 4 shows that our Hessian energy \( E \) does not show the bias at the boundary that the Laplacian energy does: this is because it also has as-linear-as-possible natural boundary conditions.

For this application, our Hessian energy \( E \) combines the two worlds of Laplacian energy and planar Hessian energy to produce a smoothness energy that is suited for scattered data interpolation on curved surfaces while unbiased by the presence of boundaries (Figure 1, Figure 12). This is helpful if the boundaries of the surface don’t have any physical meaning: perhaps they are the result of a faulty laser scan, or perhaps surface information is simply not available there. The Hessian energy’s natural boundary conditions make a best guess everywhere the data is missing by extrapolating the function linearly across the boundary.

7.2 Data smoothing

Another popular application for smoothness energies is the eponymous data smoothing. This can be used to simply smooth arbitrary data, to denoise noisy data, or to smooth the surface itself via surface fairing. One solves the following Helmholtz-like optimization
problem: given an input function $f$ to be smoothed,
\[ u = \arg \min_u E(u) + \alpha \int_\Omega (f - u)^2 \, dx , \quad (34) \]
where the parameter $\alpha > 0$ is a trade-off between the input data and the smoothness of the output data.

Figure 2 shows our Hessian energy $E$ applied to such a smoothing problem. Correctly accounting for curvature by modeling a curved biharmonic equation using the Laplace-Beltrami operator is important here: the figure shows that the approach of Stein et al. [2018] causes distortion in high-curvature regions when smoothing a step function. In this figure the smoothing parameters are chosen to give visually similar amounts of smoothing, which means a slightly larger parameter $\alpha$ for the method of Stein et al. [2018].

It is natural to ask why the fact that minimizers of $E_{\text{H2}}$ do not solve the biharmonic equation leads to worse results when smoothing the step function of Figure 2, but not for the smoothing problems solved by Stein et al. [2018, Fig. 1, Fig. 11, Fig. 13]. These examples all smooth very noisy functions with a lot of variation everywhere on the surface. The step function is the opposite of that: the variation is much more sparse. This allows the error of not accounting for curvature correctly to manifest. In Figure 13 such a denoising problem is solved using the energies $E_{\text{H2}}$ (with zero Neumann boundary conditions), $E_{\text{H2}}$ (with the implementation of Stein et al. [2018]), and $E$. It can be clearly seen that $E_{\text{H2}}$, the Laplacian energy with zero Neumann boundary conditions, is biased by the boundary, and the isolines near the boundary are distorted so they can be normal to it. The denoised solution using the Hessian energy $E$ does not suffer from this, and the isolines ignore the boundary. In regions far away from the boundary it can be observed that the result of denoising with the Hessian energy $E$ matches the Laplacian energy with zero Neumann boundary conditions $E_{\text{H2}}$, while the planar Hessian energy $E_{\text{H2}}$ differs.

The smoothing problem can also be used to smooth the geometry of the surface itself if the input data $f$ from (34) is the vertex positions in each coordinate, and the output data $u$ is the new vertex positions. If such a smoothing operation is applied repeatedly, one has a smoothing flow. Figure 14 shows our Hessian energy $E$ applied to such a problem. While the method of Stein et al. [2018] can lead to some artifacts due to not accounting for curvature, this does not happen with our curved Hessian energy $E$.

8 CONCLUSION

In this work we have introduced a smoothing energy for curved surfaces, the Hessian energy. Its minimizers solve the biharmonic equation, and it exhibits the as-linear-as-possible natural boundary conditions in the curved setting that the planar Hessian energy of Stein et al. [2018] exhibits in the flat setting. This Hessian energy can be used in many applications where smoothness energies are required, these smoothness energies should be unbiased by the boundary, and it is crucial that the minimizers of the energy solve the biharmonic equation.

8.1 Limitations

We have no numerical analysis proof for the convergence of our discretization method. We also do not provide any theoretical analysis of the spectrum of our discrete operator. Both are needed to make this discretization reliable, and to improve understanding of the method, where it works, and where it does not.

8.2 Future work

One interesting avenue for future work is to explore alternate discretizations. Higher-order versions of Crouzeix-Raviart basis functions, such as cubic or quintic basis functions, would be an interesting potential improvement. Alternatively, instead of choosing the intermediate variable $w = dv$ for the mixed formulation as in (25), a discretization where $w = \nabla \cdot v$ sounds very promising. This would more closely mirror the mixed formulation of Stein et al. [2018]. The CROF approach can be used to define a basis for tensors in the same way as is done for vectors in Section 6.2, based on the parallel and the perpendicular vector at each edge. Using other finite elements to discretize the space of one-forms could also produce new methods. Moreover, future work could explore discretizations of the smooth energy on other surface representations beyond triangle meshes.
A rich source of future work is the numerical analysis of our method. We do not have any proof of convergence, or a solid mathematical analysis of the spectrum of our operator, and while the experiments in Section 6.3 provide some evidence for problems that can be solved with our discretization of $E$, a thorough numerical analysis treatment of our discretization would be valuable to exactly identify the strengths and weaknesses of our method. Our Crouzeix-Raviart discretization is a potential candidate for spurious modes, since the finite element is non-conforming, even though we have not observed them in practice. The method of English and Bridson [2008] is an example of a Crouzeix-Raviart discretization that works for many cases, but where specific triangle configurations exist that lead to spurious modes [Quaglino 2012, Section 4.4.2]. The properties of minimizers of the discrete energies also warrant further investigation: it is unclear which properties of smooth minimizers they actually inherit, and which properties only hold in the limit.

Another interesting direction for future work is to consider additional applications. Smoothness energies have many uses, and if such an application has to be unbiased by the boundary even on heavily curved surfaces, our Hessian energy $E$ is a powerful tool. Applications could include animation [Jacobson et al. 2011], distance computation [Crane et al. 2013b], and more.

Moreover, our simple Crouzeix-Raviart discretization of the one-form Dirichlet energy containing covariant derivatives from Section 6.2 offers an interesting approach to discretize the vector Dirichlet energy in a wide variety of applications. Potential applications include vector field design [Knöppel et al. 2013], parallel transport of vectors [Sharp et al. 2018], and many more [Azencot et al. 2015; Corman and Ovsjanikov 2019; Liu et al. 2016].

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REFERENCES


Fig. 15. Error plot for six different boundary value problems. The minimizer of the Hessian energy $E$ discretized with our discretization is compared to a high-resolution solution with the same discretization. Refinement happens via loop subdivision with various types of fixed boundary. The high-resolution solution as well as the wireframe of the lowest-resolution mesh are displayed for each problem.

Fig. 16. Error plot for six different forward problems. The domains are curved surfaces of the form $(x, y, z(x, y)) \in \mathbb{R}^3$, so the integrand of the Hessian energy can be exactly computed pointwise using the properties of Monge patches [Weisstein 2019]. Quadrature is then used to compute the exact value of $E(f)$. The high-resolution function $f$ as well as the wireframe of the lowest-resolution mesh are displayed for each problem. Refinement happens via loop subdivision, and then projection to the given smooth surface.
The entries of the differential matrix $D$ on the triangle $T_{ijk}$ for each edge $e$ are

$$
\begin{align*}
-D_{i,j,el}^{ijk} &= \frac{A_{ijk}}{6l_{ij}} \\
-D_{j,k,el}^{ijk} &= \frac{A_{ijk}}{6l_{ij}} \\
-D_{k,i,el}^{ijk} &= -\frac{l_{ij}}{6l_{ij}} \cos \theta_{ijk}^{ij} \\
-D_{j,i,el}^{ijk} &= -\frac{l_{ij}}{6l_{ij}} \cos \theta_{ijk}^{ij} \\
-D_{k,j,el}^{ijk} &= \frac{1}{6} \\
-D_{i,k,el}^{ijk} &= 0
\end{align*}
$$

where $i$ is the vertex at the tail of the edge $e$, and $j$ is at its tip. If one of the edges has reversed orientation in the triangle $T_{ijk}$ with respect to its global orientation, its entries get multiplied by $-1$.

The entries of the diagonal CROF mass matrix $M$ on the triangle $T_{ijk}$ are

$$
\begin{align*}
M_{i,el}^{ijk} &= M_{j,el}^{ijk} = \frac{A_{ijk}}{6l_{ij}} \\
M_{k,el}^{ijk} &= 0
\end{align*}
$$

where $A_{ijk}$ is the double area of the triangle $T_{ijk}$, $\theta_{ijk}^{ij}$ is the angle in the triangle $T_{ijk}$ at the vertex $i$, and $l_{ij}$ is the length of the edge from vertex $i$ to $j$. If one of the edges has reversed orientation in the triangle $T_{ijk}$ with respect to its global orientation, its off-diagonal entries get multiplied by $-1$. These are only the terms for the triangle $T_{ijk}$. One must add the terms for all triangles and all pairs of edges in that triangle to compute the full matrix $L$. We suggest looping through all triangles, and adding the terms for each triangle to the respective entries of the matrix corresponding to the edges. This can easily be parallelized with a parallel_for loop.

The entries for each of the matrices defined in Section 6.1 needed to construct the system matrices used in (27) are as follows. Let $e$ be an oriented edge from the vertex $i$ to $j$. The two triangles adjacent to $e$ are $T_{ijk}$ and $T_{jil}$, and $f$ is an oriented edge from the vertex $k$ to $i$. The entries of the symmetric CROF vector Dirichlet matrix $L$ on the triangle $T_{ijk}$ are

$$
\begin{align*}
L_{e^i,el}^{ijk} &= L_{e^j,el}^{ijk} = \frac{2}{A_{ijk}} \\
L_{e^i,el}^{ijk} &= L_{e^j,el}^{ijk} = 0 \\
L_{e^i,el}^{ijk} &= L_{e^j,el}^{ijk} = \frac{2}{A_{ijk}} \cos \theta_{ijk}^{ij} \\
L_{e^i,el}^{ijk} &= L_{e^j,el}^{ijk} = \frac{2}{l_{ij}^{el}} \cos \theta_{ijk}^{ij}
\end{align*}
$$

where $A_{ijk}$ is the double area of the triangle $T_{ijk}$, $\theta_{ijk}^{ij}$ is the angle in the triangle $T_{ijk}$ at the vertex $i$, and $l_{ij}$ is the length of the edge from vertex $i$ to $j$. If one of the edges has reversed orientation in the triangle $T_{ijk}$ with respect to its global orientation, its off-diagonal entries get multiplied by $-1$. These are only the terms for the triangle $T_{ijk}$. One must add the terms for all triangles and all pairs of edges in that triangle to compute the full matrix $L$. We suggest looping through all triangles, and adding the terms for each triangle to the respective entries of the matrix corresponding to the edges. This can easily be parallelized with a parallel_for loop.

The entries of the differential matrix $D$ on the triangle $T_{ijk}$ for each edge $e$ are

$$
\begin{align*}
-D_{i,j,el}^{ijk} &= \frac{A_{ijk}}{6l_{ij}} \\
-D_{j,k,el}^{ijk} &= \frac{A_{ijk}}{6l_{ij}} \\
-D_{k,i,el}^{ijk} &= -\frac{l_{ij}}{6l_{ij}} \cos \theta_{ijk}^{ij} \\
-D_{j,i,el}^{ijk} &= -\frac{l_{ij}}{6l_{ij}} \cos \theta_{ijk}^{ij} \\
-D_{k,j,el}^{ijk} &= \frac{1}{6} \\
-D_{i,k,el}^{ijk} &= 0
\end{align*}
$$

where $i$ is the vertex at the tail of the edge $e$, and $j$ is at its tip. If one of the edges has reversed orientation in the triangle $T_{ijk}$ with respect to its global orientation, its entries get multiplied by $-1$.

The entries of the diagonal CROF mass matrix $M$ on the triangle $T_{ijk}$ are

$$
\begin{align*}
M_{i,el}^{ijk} &= M_{j,el}^{ijk} = \frac{A_{ijk}}{6l_{ij}} \\
M_{k,el}^{ijk} &= 0
\end{align*}
$$

where $A_{ijk}$ is the double area of the triangle $T_{ijk}$, $\theta_{ijk}^{ij}$ is the angle in the triangle $T_{ijk}$ at the vertex $i$, and $l_{ij}$ is the length of the edge from vertex $i$ to $j$. If one of the edges has reversed orientation in the triangle $T_{ijk}$ with respect to its global orientation, its off-diagonal entries get multiplied by $-1$. These are only the terms for the triangle $T_{ijk}$. One must add the terms for all triangles and all pairs of edges in that triangle to compute the full matrix $L$. We suggest looping through all triangles, and adding the terms for each triangle to the respective entries of the matrix corresponding to the edges. This can easily be parallelized with a parallel_for loop.

The entries of the differential matrix $D$ on the triangle $T_{ijk}$ for each edge $e$ are

$$
\begin{align*}
-D_{i,j,el}^{ijk} &= \frac{A_{ijk}}{6l_{ij}} \\
-D_{j,k,el}^{ijk} &= \frac{A_{ijk}}{6l_{ij}} \\
-D_{k,i,el}^{ijk} &= -\frac{l_{ij}}{6l_{ij}} \cos \theta_{ijk}^{ij} \\
-D_{j,i,el}^{ijk} &= -\frac{l_{ij}}{6l_{ij}} \cos \theta_{ijk}^{ij} \\
-D_{k,j,el}^{ijk} &= \frac{1}{6} \\
-D_{i,k,el}^{ijk} &= 0
\end{align*}
$$

where $i$ is the vertex at the tail of the edge $e$, and $j$ is at its tip. If one of the edges has reversed orientation in the triangle $T_{ijk}$ with respect to its global orientation, its entries get multiplied by $-1$.

The entries of the diagonal CROF mass matrix $M$ on the triangle $T_{ijk}$ are

$$
\begin{align*}
M_{i,el}^{ijk} &= M_{j,el}^{ijk} = \frac{A_{ijk}}{6l_{ij}} \\
M_{k,el}^{ijk} &= 0
\end{align*}
$$

where $A_{ijk}$ is the double area of the triangle $T_{ijk}$, $\theta_{ijk}^{ij}$ is the angle in the triangle $T_{ijk}$ at the vertex $i$, and $l_{ij}$ is the length of the edge from vertex $i$ to $j$. If one of the edges has reversed orientation in the triangle $T_{ijk}$ with respect to its global orientation, its off-diagonal entries get multiplied by $-1$. These are only the terms for the triangle $T_{ijk}$. One must add the terms for all triangles and all pairs of edges in that triangle to compute the full matrix $L$. We suggest looping through all triangles, and adding the terms for each triangle to the respective entries of the matrix corresponding to the edges. This can easily be parallelized with a parallel_for loop.

The entries of the differential matrix $D$ on the triangle $T_{ijk}$ for each edge $e$ are

$$
\begin{align*}
-D_{i,j,el}^{ijk} &= \frac{A_{ijk}}{6l_{ij}} \\
-D_{j,k,el}^{ijk} &= \frac{A_{ijk}}{6l_{ij}} \\
-D_{k,i,el}^{ijk} &= -\frac{l_{ij}}{6l_{ij}} \cos \theta_{ijk}^{ij} \\
-D_{j,i,el}^{ijk} &= -\frac{l_{ij}}{6l_{ij}} \cos \theta_{ijk}^{ij} \\
-D_{k,j,el}^{ijk} &= \frac{1}{6} \\
-D_{i,k,el}^{ijk} &= 0
\end{align*}
$$

where $i$ is the vertex at the tail of the edge $e$, and $j$ is at its tip. If one of the edges has reversed orientation in the triangle $T_{ijk}$ with respect to its global orientation, its entries get multiplied by $-1$.
The entries of the curvature correction matrix $K$ on the triangle $T_{ijk}$ are

$$K^{ijk}_{e\parallel,e\parallel} = \frac{1}{l_{ij}} \left( \frac{\theta^{ijk}}{s_i} \kappa_i + \frac{\theta^{ijk}}{s_j} \kappa_j + \frac{\theta^{ijk}}{s_k} \kappa_k \right)$$

$$K^{ijk}_{e\parallel,e\perp} = 0$$

$$K^{ijk}_{e\perp,f\parallel} = K^{ijk}_{e\perp,f\perp} = \frac{\cos \theta^{ijk}}{l_{ijk}} \left( \frac{\theta^{ijk}}{s_j} \kappa_j + \frac{\theta^{ijk}}{s_k} \kappa_k - \frac{\theta^{ijk}}{s_i} \kappa_i \right)$$

where $\kappa_v$ is the angle defect at the vertex $v$ and $s_v$ is the angle sum at the vertex $v$. If one of the edges has reversed orientation in the triangle $T_{ijk}$ with respect to its global orientation, its off-diagonal entries get multiplied by $-1$.

**B ADDITIONAL EXPERIMENTS**

Figure 15 features a series of convergence experiments that shows the convergence of a boundary value problem on a variety of meshes to the highest-resolution solutions. In Figure 16, a series of forward problems is solved, where the Hessian energy of a function is measured on a curved surface, and because both the function and the surface embedding are known, the exact solution is also known. This is used to measure the error. In both these examples, convergence of the order of the average edge length is observed.

Figure 17 shows that for different meshings of the same surface, very similar results are achieved, and the method is thus robust to remeshing. In Figure 18 our CROF implementation of the Hessian energy is compared with various Hessian energies discussed by Stein et al. [2018] in the flat annulus setting, where the exact solution is known.