Texture Mapping via Optimal Mass Transport

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Abstract—In this paper, we present a novel method for texture mapping of closed surfaces. Our method is based on the technique of optimal mass transport (OMT), also known as the "earth-mover’s metric." The original transport problem, proposed by Gaspard Monge in 1781, asks how best to move a pile of soil or rubble to an excavation or fill with the least amount of work. The problem was analyzed and given a modern formulation by the Soviet mathematician and econometrician Kantorovich [43] in the 1940s and so is known now as the Monge–Kantorovich (MK) problem. OMT has found uses in numerous fields including fluid and continuum mechanics, automatic control, statistical physics, shape optimization, expert systems, meteorology, econometrics, data compression, and image processing; see [55], [70], and the references therein.

The texture mapping methodology is based on mapping an image either synthesized or digitized onto a given surface. There is plenty of literature on this topic in the computer graphics community, e.g., see [23] and the references therein. We will not review all the literature on this subject in this present paper. But only some of the more relevant works for the approach taken here. We begin by noting that the mathematical basis of this mapping determines whether the image provides the object with the appearance of surface texture, a specular reflection, or some other effect. In this context, bijectivity is assured if we don’t get triangles some measure of distortion), yet is still bijective. In this paper, we will focus only on mappings such as texture mapping that are a function of surface location and not those, for example, based on surface orientation.

1.1 Connection of Optimal Mass Transport to Texture Mapping

In discrete settings, we often represent surfaces as a collection of samples with their connectivity, typically in the form of a simplicial mesh. Many of the operations we wish to perform on the resulting discrete surface, for processing, analysis, or visualization (such as texture mapping), may be greatly simplified if we perform them in a corresponding canonical domain with the same topology. For example, the surface of the brain is a topological sphere, but is highly convoluted. If one can find a “nice” bijection of the brain surface onto the sphere, this could be very useful for various types of visualizations [69]. Therefore, in general, we wish to find a bijection of the discrete surface lying in 3D space, with a simpler model (e.g., spherical space in the simply connected compact case), such that a certain distortion measure is minimized. This mapping procedure assigns two coordinates to each of the vertices of the original mesh. This procedure is also known as parameterization. The surface to which the mesh is mapped to is typically referred as the parameter domain. Parameterization of polygonal geometric surfaces may be used for image mapping [28], a collection of techniques that includes texture mapping [11], [36], [37], reflection or environment mapping [8], bump mapping [9], and light mapping [51].

The main challenge in mesh parameterization is to produce a corresponding 2D triangulation that best matches the geometry of the 3D mesh (by minimizing some measure of distortion), yet is still bijective. In this context, bijectivity is assured if we don’t get triangles flipping in the mapping process. In this paper, we will show how we can find an optimal mapping, i.e., a mapping which will minimize some predefined measure of distortion, using the OMT formulation.

1.2 Related Work

Surface parameterization was initially introduced to the computer graphics community as a method for mapping textures onto surfaces. Over the last two decades, it has gradually become a common tool, useful for many other mesh processing applications, such as detail mapping and synthesis, morphing and detail transfer between two (or more) objects, creation of aligned objects databases, multiresolution analysis, remeshing, mesh compression, and many more (see [64], [39] for a comprehensive survey). The literature of course is huge, and so in our survey, we decided to only include those works that are most relevant to the approach adopted in the present paper.

Our interest in parameterization arose from our work in medical imaging [5], [54]. A significant portion of the
spheres. Therefore, a good bijective mapping of minimal distortion from such a surface to a sphere may be utilized for registration and visualization. As alluded to above, one of the structures for which such a mapping is particularly useful is the surface of the human brain. After registration, the spherical representation can be analyzed on the sphere using such spherical multiresolution functions such as spherical harmonics or spherical wavelets. Depending upon the application, numerous methods for parameterizing meshes have been developed, targeting diverse parameter domains and focusing on different parameterization properties.

Some key works on parameterization have focused on planar parameterization of meshes with disk-like topology. A number of more recent papers on this subject (e.g., [16], [22], [48], [49], [53], [58], [63]) have proposed certain distortion measures and described methods for their minimization. The parameterization problem is more complicated when the mesh does not have the topology of a disk. One way to circumvent this problem is to cut seams in the surface mesh and then use the methods designed for disks ([29], [62], [65]). A polygonal boundary may be formed by removing an arbitrary triangle from a closed mesh as in [34], [66]. A more elaborate boundary may be formed by cutting along mesh edges. This, however, usually introduces discontinuities, where the edges are cut and may sometimes increase the distortion of the parameterization.

An alternative to cutting the mesh and creating a boundary is a method based on the introduction of cone singularities, first proposed by Kharevych et al. [44]. The main idea here is that instead of introducing artificial boundaries to absorb the undesired curvature, the entire Gaussian curvature of the mesh is redistributed so that it is concentrated at a few designated places, referred to as the “cone singularities.” Once the curvature is redistributed, the edge lengths of a 2D embedding having this target curvature can be determined. The main difference between this method, in which the cut is performed after the new metric is computed, and the techniques presented above, in which the cut is performed before the new metric is computed, is that this approach guarantees that edges on both sides of the cut will be mapped to edges of the same length, so the flattened version of the mesh may be “zippered” back together in the plane at the boundaries. This reduces the discontinuities in the parameter plane. Some recent publications that employ this methodology are those of Ben-Chen et al. [7] and Springborn et al. [66]. The difference between these two methods is the precise algorithm used to manipulate the curvature distribution. These approaches produce parameterizations, which minimize conformal distortion without introducing too much stretch. However, in our work in which we focused on genus zero anatomical surfaces, we preferred to use a mapping to the sphere without introducing seams. This way we could use the sphere as a common domain for registration and analysis of anatomical structures from different patients without any additional constraints on feature points.

Several methods for direct parameterizations on a topological sphere have also been developed. We classify these parameterizations into three groups, based on the type of parametric distortion minimized: methods that do not explicitly consider the problem of distortion, methods that minimize angular distortion, and methods that minimize area distortion. In practice, most existing parameterization techniques belong to the first group, see [46], [1]. For instance, the technique proposed by Alexa [1] is a heuristic iterative procedure, which attempts to converge to a valid parameterization by applying local improvement (relaxation) rules. Here, one starts by computing an initial guess and then moving the vertices one at a time, first computing a 3D position for the vertex using a barycentric formulation, and then projecting the vertex to the unit sphere. This process works well in many cases, but there is no guarantee that it will converge, and, even if it does, the resulting embedding may not be bijective. An interesting alternative has been proposed by Shaprio and Tal [61] using a multiresolution technique. This method begins with a simplification of the mesh until it becomes a tetrahedron (or at least, convex). The simplified model is then embedded in the sphere, and then the vertices are inserted back one by one so that the bijectivity of the mapping is preserved. While this is quite an efficient and stable process, it is difficult to optimize the parameterization.

Methods that attempt to rigorously minimize the angular distortion include those of [32], [33], [34], [35], [41], [63]. For example, in [34], the authors first cut out one triangle, then conformally map the remaining surface onto an infinite plane, and finally, use the inverse stereo projection to map the plane to the sphere. This conformal method was applied to texture mappings in [35]. The presented approach works quite well in practice even for dense meshes and we will use it as a starting point for our algorithm. However, it should be noted that maps that are bijective and conformal for smooth surfaces do not guarantee an embedding when applied to piecewise linear surfaces (meshes) and sometimes produce triangle flips. For instance, the stereographic projection in [34] can flip thin obtuse triangles [39].

Gu et al. [33] introduce a nonlinear optimization method, which is based on minimizing the harmonic energy, to compute global conformal parameterizations for genus zero surfaces. Their optimization is carried out in the tangent spaces of the sphere. Hence, there is no stereographic projection and the method is claimed to be more stable than that of [34]. However, Gu et al. employed “projected” Gauss-Seidel iterations to obtain the parameterization, which were shown by Saba et al. [57] to decrease the residual for only a finite number of iterations. As the result approaches a bijective solution, the scheme ultimately becomes unstable, the residual increases, and the system collapses to a degenerate solution. In order to avoid instability, Gu et al. use a certain trial and error approach to adjust the damping parameter, in order to obtain a value for which the procedure may be terminated sufficiently close to a bijective solution before diverging. Hence, the resulting parameterization is only approximate.

Gotsman et al. [30] showed how to correctly generalize the method of barycentric coordinates, with all its advantages, to the sphere. They provide a quadratic system of equations, which is a spherical equivalent of the barycentric formulation. Using appropriate weights in this scheme can generate a bijective conformal mapping. Saba et al. [57] introduce a method for efficiently solving the system proposed in [30].

Zayer et al. [71] introduce a parameterization method, which cuts the mesh along a line connecting user-prescribed poles. The mesh is then topologically equivalent to a disk and an initial parameterization is found by solving a Laplace equation in curvilinear coordinates. The parameterization distortion is then reduced using a variant of
quasiharmonic maps. The distortion along the seam is also reduced by tangential Laplacian smoothing.

Another method that takes angular distortion into account has been proposed by Sheffer et al. [63]. This is a highly nonlinear optimization procedure utilizing the angles of the spherical triangulation (as opposed to the vertex positions). Here, the authors specify a set of constraints that the angle values have to satisfy to define a planar triangular mesh. They search for angles that are as close as possible to the original 3D mesh angles and satisfy those constraints, and then convert the solution angles into actual vertex coordinates. Using this method, they can define constraints on the angles and even on the areas of the triangles. However, since this method seems to lack an efficient numerical computational procedure, it may not be very practical for large meshes. Finally, another procedure for conformal mapping, which uses redistribution of Gaussian curvature, is the discrete Ricci flow [41]. This method also requires a nonlinear solver.

A major concern with conformal mappings is the distortion in scale. A method for minimizing the area distortion was introduced by Degener et al. [14]. This approach is an extension of a previous method “MIPS” [40], which attempts to minimize angle distortion by optimizing a nonlinear functional that measures mesh conformality. Here, the authors add a term that measures area distortion to their energy functional. They mediate between angle and area deformations by changing the powers of the components in the proposed functional.

1.3 Our Contribution

In general, there may be several natural measures for the “goodness” of the mapping. From one point of view, we wish to obtain a mapping that preserves the local geometry of the original surface and this can be obtained using a conformal (angle preserving) mapping as in [34], [35], [31]. On the other hand, it is reasonable to require the mapping to be area preserving. However, in general, it is not possible to map a simply connected compact surface with non-constant Gaussian curvature to the sphere in a way that preserves both angles and areas. Such a mapping would have to be an isometry and, thus, curvature-preserving [17]. Therefore, as a compromise, in this paper, we propose a method for calculating an area preserving mapping that minimizes the geometrical distortion. This mapping is optimal in the sense that it is both area preserving and attempts to minimize the angle distortion. In general, for a surface which is a topological sphere, there is a unique (up to Moebius transformation) mapping that is angle preserving, but there are many area preserving mappings. Therefore, we will try to find an area preserving map, which gives the smallest distortion of the mesh in the sense of optimal mass transport.

This will be performed via a three-step approach. We assume that all our surfaces are compact simply connected, that is, topological spheres. This assumption will be in force unless stated otherwise throughout this paper. We first start with conformally mapping the given surface $\Omega$ onto the unit sphere $S^2$ using a conformal mapping method, such as the one proposed in [34], [4]. The area distortion of this mapping will be interpreted as a density function $\mu$ on $S^2$. The second step is to find a mapping from $S^2$ with density $\mu$ to $S^2$ with a constant density. We will use the method of Moser ([52], [13]) in this step. This mapping will transport the density $\mu$ to area, and so, a composition of this mapping with the mapping from the first step will give us an area preserving mapping from $\Omega$ to $S^2$. (See Section 2 for details.) This composite mapping will be used as an initial MP mapping for the third step. In this step, we rearrange the area preserving mapping of the second step to make it optimal in the mass transport sense. The composition of the conformal mapping and the optimal mass preserving one will be the required area preserving diffeomorphism to be used for texture mapping. In order to get fast and stable convergence to the optimal map, we use a multiresolution scheme.

The outline of the remainder of this paper is as follows: In Section 2, we sketch the analytical procedure to find the optimal mapping. In Section 3, we discuss some relevant implementation issues. In Section 4, we describe the steps we perform to obtain the texture mapping, in Section 5, we give some illustrative examples of our scheme, and in Section 6, we present an evaluation of the running time and distortion measures of our method. Finally, in Section 7, we summarize our work and give some possible future research directions.

2 Sketch of Relevant Mathematical Theory

In this section, we outline the mathematical justification of our mapping procedure. This is based on the idea of using a minimizing flow to compute the optimal $L^2$ mass transport map as in [34], [4], and its extension to the case of a compact surface [18], [19]. It should be noted that optimal mass transport theory may be carried out on a general Riemannian manifold [21], [70]. For the purpose of texture mapping in the present work, we only consider the genus zero compact surface (topological sphere), since it allows us to use a computationally efficient implementation via spherical wavelets and is an important case in the texture mapping literature.

2.1 Mass Preserving Maps and the Monge-Kantorovich Problem

We first give a precise mathematical formulation of the Monge-Kantorovich problem [21], [70]. Accordingly, consider two oriented compact Riemannian manifolds $\Omega_0$ and $\Omega_1$, each with a corresponding positive density function defined over it, denoted by $\mu_0$ and $\mu_1$, respectively. We assume that the total mass associated with each of the manifolds is equal, i.e.,

$$\int_{\Omega_0} \mu_0(x)dx = \int_{\Omega_1} \mu_1(y)dy,$$

(1)

where $dx$ and $dy$ are the standard area forms induced by the Riemannian metric on $\Omega_0$ and $\Omega_1$, respectively. If this assumption is not satisfied, we can always scale one of the density distributions to make the total amount of mass equal.

We wish to find a smooth mass preserving transport map that takes the first distribution into the second one $u : (\Omega_0, \mu_0) \rightarrow (\Omega_1, \mu_1)$, i.e., a diffeomorphism which satisfies

$$\mu_0 = |\nabla u| \mu_1 \circ u.$$

(2)

Here, $|\nabla u|$ denotes the determinant of the Jacobian map $\nabla u$ and $\circ$ represents composition of functions. This equation is often referred to as the Jacobian equation, which constrains the mapping to be mass preserving (MP) with respect to the given density functions. A mapping $u$ that satisfies this property may, thus, be thought of as defining a redistribution of a mass
of material from one distribution \((\Omega_0, \mu_0)\) to another 
\((\Omega_1, \mu_1)\). We assume that the total surface areas of 
\(\Omega_0\) and \(\Omega_1\) are equal. Then, we say that a diffeomorphism is 
area preserving if it maps the area form of \(\Omega_0\) to the area form of 
\(\Omega_1\) (the area forms are defined with respect to the corresponding 
Riemannian metrics \([17]\)).

There may be many such mappings and we want to 
choose an optimal one in some sense. This can be obtained 
by incorporating into the problem a penalty function of the 
following form:

\[
M(u) = \int_{\Omega} \Phi(x, u(x)) \mu_0(x) dx,
\]

where \(\Phi(x, u(x))\) is a suitable positive convex function, 
typically taken to be the geodesic distance (or its square) on 
the given manifold between \(x\) and \(u(x)\). (Here, \(dx\) denotes 
the area form defined with respect to the Riemannian metric 
on the manifold.) The contribution of this functional to the 
problem is to place a penalty on the distance the map \(u\) 
moves each bit of material, weighted by the material’s mass.

An optimal MP mapping (when it exists) is one that 
minimizes this functional over all while satisfying the mass 
and volume preserving mappings.

A fundamental theoretical result \([21]\) shows that under 
certain mild conditions, a unique optimal MP map 
\(u : (\Omega_0, \mu_0) \rightarrow (\Omega_1, \mu_1)\) exists. In this paper, we will present a 
simple algorithm for the construction of the optimal MP map 
in the case of the sphere.

### 2.2 Transforming a Diffeomorphism to \(S^2\) into an 
Area Preserving Mapping

Let us assume that we have some initial area distorting 
(e.g., conformal) diffeomorphism \(f\) from some compact 
simply connected surface \(\Omega\) to the unit sphere \(S^2\). We can 
quantify the change of area as a density function \(\mu\) at each 
point of the surface so that the integral on the flattened 
surface (the unit sphere) will give us the area measure of 
the original surface:

\[
\int_{\Omega} dx = \int_{S^2} \mu(x) dx.
\]

By change of variables, it is easy to see that this density 
function is the determinant of the Jacobian of \(f^{-1}\), i.e., 
\(\mu = |\nabla f^{-1}|\).

Without loss of generality, we can assume that the total 
surface area of \(\Omega\) equals that of \(S^2\) \((4\pi)\). Consider the problem of 
finding a smooth mass preserving map that takes the 
density \(\mu\) over \(S^2\) into one of constant density \(1\) on \(S^2\):

\[
g : (S^2, \mu) \rightarrow (S^2, 1).
\]

This means that that \(g\) will take \(\mu dx\) to \(dx\), where \(dx\) is the 
standard area form on \(S^2\). One can see then that \(g \circ f\) will 
give us an area preserving mapping from the original 
surface \(\Omega\) to the sphere. Note that we have

\[
|\nabla g| = \mu.
\]

Such a mapping can be constructed using a method 
formulated in \([52], [13]\), which will be further explained in 
Section 4.

### 2.3 Optimal Area Preserving Mappings

We now specialize the Monge-Kantorovich to the case of the 
sphere \(S^2\). We will use the gradient flow ideas described in 
\([4], [18], [19]\) in our solution. So we want to minimize a 
functional of the form

\[
M(g) = \int_{S^2} \Phi(x, g(x)) \mu(x) dx
\]

over MP mappings \(g : (S^2, \mu) \rightarrow (S^2, 1)\). (Again, we will take 
\(\Phi\) to be the geodesic distance or the square thereof) As 
above, \(dx\) is the area form of the standard metric on the 
sphere. Suppose that we have constructed an initial MP 
mapping as above (e.g., using \([52], [13]\)), which will be 
denoted, henceforth, by \(g^0\). Using this mapping, we will 
rearrange the density in the domain of the map in order to 
minimize the cost functional \(M(g)\), while constraining \(g\) so 
that it continues to satisfy (2).

In our approach, we introduce a smooth, \(\mu\) preserving, 
family of diffeomorphisms \(s^t : (S^2, \mu) \rightarrow (S^2, \mu)\) and define 
the family of maps \(g^t\) via

\[
g^0 = g^t \circ s^t.
\]

See Fig. 1. For \(t = 0\), we define \(s^0\) to be the identity map. The 
idea given in \([34], [4]\) is to use the family of maps \(s^t\) to 
rearrange the initial MP map \(g^0\) in such a manner to 
converge to the optimal solution. The family will be defined 
via a certain gradient flow defined for the minimization of 
the functional (6). We now give the details.

If the maps \(g^t : S^2 \rightarrow S^2\) evolve so that

\[
g^0(x) = g^t(s^t(x)),
\]

where \(s^t : S^2 \rightarrow S^2\) preserves the measure \(\mu(x)dx\), and where

\[
\frac{\partial s^t(x)}{\partial t} = v^t(s^t(x)),
\]

then the \(g^t\) satisfy the transport equation

\[
\partial_t g + v \cdot \nabla g = 0.
\]

We can accordingly write the evolution of the cost 
functional as follows:

\[
\frac{\partial M(g^t)}{\partial t} = \int_{S^2} \mu(x) v^t(x) \cdot \Phi(x, g^t(x)) dx.
\]

The maps \(s^t\) will preserve the measure \(\mu(x)dx\) if and only 
if the velocity field \(v^t\) is divergence free, i.e., satisfies

\[
\nabla \cdot (\mu v^t) = 0.
\]
Since $S^2$ is two-dimensional, the general divergence-free vector field is of the form

$$\mu t = -J \nabla H(x),$$  \hfill (10)

where $J$ is the rotation by 90 degree and $H$ is some “Hamiltonian function.” See [18], [19] for all the details.

In general, using (10), one can compute that

$$\frac{\partial}{\partial t} M(g') = -\int_{S^2} (J \nabla H) \cdot \Phi_x(x, g') \, dx$$

$$= \int_{S^2} (\nabla H) \cdot J \Phi_x(x, g') \, dx$$

$$= -\int_{S^2} H \nabla \cdot (J \Phi_x(x, g')) \, dx.$$

Then, in order to obtain a gradient flow, one sets

$$H = \nabla \cdot (J \Phi_x(x, g')) \quad (11)$$

so that the evolution of the cost function will be decreasing. In this setting, the maps should evolve by the following initial value problem:

$$\begin{aligned}
\frac{\partial}{\partial t} g' + v' \cdot \nabla g' &= 0 \\
v' &= -\frac{1}{\mu(x)} J \nabla \{ \nabla \cdot (J \Phi_x(x, g')) \}. 
\end{aligned} \quad (12)$$

### 2.4 Incorporating a Multiresolution Scheme to Our Flow

We now employ a wavelet representation to modify the gradient flow as proposed in the previous section. Accordingly, we can utilize a spherical wavelets basis $\psi_{j,m}$, such as the second generation biorthogonal wavelets introduced by Sweldens [67], [68], to represent $H$:

$$H = \sum_{j,m} \gamma_{j,m}(t) \psi_{j,m}(x). \quad (13)$$

These wavelet functions can be defined intrinsically on the manifold and do not depend on its parameterization. Wavelet functions $\psi_{j,m}$ represent functions on a regular grid at multiple levels of detail. Here, $j$ indexes the scale (resolution of analysis), while higher $j$ corresponds to higher resolution analysis, and $m$ indexes the spatial location of analysis.

Any function $q$ can be decomposed as a superposition of the biorthogonal basis $\psi_{j,m}$ using a set of dual basis functions $\tilde{\psi}_{j,m}$:

$$q = \sum_{j,m} \langle \tilde{\psi}_{j,m}, q \rangle \psi_{j,m} = \sum_{j,m} \gamma_{j,m} \psi_{j,m}, \quad (14)$$

where $\gamma_{j,m}$ denote the expansion coefficients. Therefore, using this biorthogonal wavelet basis to represent $H$ as $\sum_{j,m} \gamma_{j,m} \psi_{j,m}$ leads to the following evolution of the cost:

$$\frac{\partial}{\partial t} M(g') = -\sum_{j,m} \gamma_{j,m}(t) \int_{S^2} \psi_{j,m}(x) \nabla \cdot (J \Phi_x(x, g')) \, dx. \quad (15)$$

This, in turn, leads one to choose $H$ as in (13) with

$$\gamma_{j,m}(t) = \int_{S^2} \tilde{\psi}_{j,m}(x) \nabla \cdot (J \Phi_x(x, g')) \, dx$$

$$= \langle \tilde{\psi}_{j,m}(x), \nabla \cdot (J \Phi_x(x, g')) \rangle \quad (16)$$

so that

$$\frac{\partial}{\partial t} M(g') = -\sum_{j,m} \gamma_{j,m}(t)^2.$$  

Thus, in this case, the maps should evolve by the following initial value problem:

$$\begin{aligned}
\frac{\partial}{\partial t} g' + v' \cdot \nabla g' &= 0 \\
v' &= -\frac{1}{\mu(x)} J \nabla \{ \nabla \cdot (J \Phi_x(x, g')) \}.
\end{aligned} \quad (17)$$

where $\gamma_{j,m}(t)$ is as defined in (16).

In our gradient descent formulation, we can group the wavelet functions $\psi_{j,m}$ according to their resolution level ($j$). Each of these groups can span any function defined over the manifold at a different resolution. We can then run our gradient flow at each resolution level separately, going from coarser to higher resolution, until it converges. This gradient flow will be obtained by iterating the following steps (starting with $J = 1$) until the MK cost functional ceases to decrease (or until $\int H^2 \, dx < \varepsilon$ for some tolerance $\varepsilon > 0$):

1. Using the basis vectors $\{\psi_{j,m}| 1 \leq j \leq J, \forall m\}$, follow the steepest descent.
2. Increase $J$ by 1, so as to include basis functions corresponding to the next level of resolution.

Moreover, the wavelet decomposition creates a natural setting for using multigrid methods, which we use to increase the efficiency of our method. In this case, we will first perform a few iterations (“relaxations”) using the expansion coefficients at all resolution levels. Then we will carry out several iterations, but every few iterations, we will pull out the coefficients of the highest resolution levels from the new representation, leaving a smoothed representation of our functions (this is equivalent to performing low-pass filtering). After reaching the lowest resolution level and performing a few iterations, we start operating in the opposite direction, by adding the coefficients of a higher resolution level every few iterations, until we reach the finest representation level. This will be referred to as the V-cycle. We perform a few such cycles until we obtain convergence to the solution.

### 2.5 Flow on $S^2$

In the case of the sphere, with $\Phi(x,y)$ as the geodesic distance from $x$ to $y$, we have

$$\Phi(x,y) = \arccos(x \cdot y).$$  

$\Phi_x$ is the projection of the derivative of $\Phi(x,y)$ as function from $\mathbb{R}^3 \times \mathbb{R}^3 \rightarrow \mathbb{R}$ onto the tangent plane to $S^2$ at $x$, therefore,

$$\Phi_x(x,y) = -\frac{y - (x \cdot y)x}{\sqrt{1 - (x \cdot y)^2}}.$$  

Since $Jy = x \times y$ and

$$(x \cdot y)^2 + |x \times y|^2 = |x|^2 |y|^2 = 1,$$

we have

$$J \Phi_x(x,g) = -\frac{x \times g}{|x \times g|}.$$
Therefore, in this setting, the maps should evolve by the initial value problem described in (17), where for $\gamma_{j,m}(t)$ defined as in (16), we have

$$\gamma_{j,m}(t) = \int_{S^2} \nabla \cdot \frac{x \times g}{|x \times g|} \bar{\psi}_{j,m}(x) \, dx$$

$$= \left\langle \bar{\psi}_{j,m}(x), \nabla \cdot \frac{x \times g}{|x \times g|} \right\rangle.$$ (18)

In the case of the sphere, we can use for the expansion the spherical wavelet basis that was introduced by Schröder and Sweldens [59], [60], which will explicitly be described in Section 3. Our flow stops when $\nabla \cdot (J\Phi_x(x,g'))$ is orthogonal to all the basis components. These points will now be described in the following section.

3 IMPLEMENTATION ISSUES

In this section, we will focus on how we maintain the correctness of the aforementioned mathematical derivation when we proceed to discrete settings.

3.1 Multiresolution Basis

As alluded to above, in our implementation, we used the construction of the spherical wavelet basis introduced in [59], [60]. These are second generation wavelets, adapted to manifolds with nonregular grids. In the construction of these wavelets, a lifting scheme is used to obtain locally supported, biorthogonal spherical wavelets and their associated fast transforms. The construction of spherical wavelets relies on a recursive partitioning of the sphere into (spherical) triangles. This can be done starting from a Platonic solid of triangles (with 4 triangles forming a tetrahedron, 8 triangles forming an octahedron, or 20 triangles forming an icosahedron), and recursively subdividing the triangles into four child triangles at each stage of the recursion (quadtrees), as shown in Fig. 2.

On this mesh, we denote the set of all vertices obtained after $j \in J$ subdivisions by $K(j)$. The $(j+1)$th resolution mesh is obtained by introducing new nodes, identified by $M(j)$, which subdivide existing edges (typically at their midpoint). The complete set of nodes in the $(j+1)$th resolution mesh is given by $K(j+1) = K(j) \cup M(j)$.

At resolution level $j$, a scaling function $\varphi_{j,k}$ is defined for every vertex $k \in K(j)$. One commonly used scaling function is the hat scaling function that varies linearly from the value 1 at vertex $k$ to 0 at its immediate neighboring vertices. The scaling functions satisfy the refinement relation, therefore, each scaling function $\varphi_{j,k}$ can be written as a linear combination of scaling functions of the next finer level:

$$\varphi_{j,k} = \sum_{l \in K(j+1)} h_{j,k,l} \varphi_{j+1,l}$$

with $\{h_{j,k,l} | j \in J, k \in K(j), l \in K(j+1)\}$ defining a finite filter.

If $q : S^2 \to \mathbb{R}$ is a function defined on $S^2$, we can approximate the function $q$ at a resolution $j$ by an expansion of the form

$$q = \sum_k \lambda_{j,k} \varphi_{j,k}.$$ The coefficients $\lambda_{j,k}$ are called scaling coefficients and formally given by the inner products $\lambda_{j,k} = \langle \varphi_{j,k}, q \rangle$, where $\varphi$ is the dual scaling function which satisfies

$$\langle \varphi_{j,k}, \varphi_{j,k} \rangle = \delta_{k,k}.$$ The scaling coefficients represent the low-pass information of the original signal. Scaling coefficients of lower resolution levels represent smoother versions of the data.

If we wish to perform a multiresolution expansion, we can do so by writing the approximation using wavelet functions that are located at the mesh subdivision points $\psi_{j,m}$. Instead of the scaling functions. For the hat scaling function, the corresponding wavelet functions may be written as

$$\psi_{j,m} = \varphi_{j+1,m} - \sum_{k \in K(j,m)} s_{j,m,k} \varphi_{j,k}.$$ The weights $s_{j,m,k}$ are chosen so that the resulting wavelet has a vanishing integral (for details, see [67], [68]). For $q \in L^2$, we can write the expansion

$$q = \sum_{j,m} \gamma_{j,m} \psi_{j,m}.$$ The coefficients $\gamma_{j,m}$ are called wavelet coefficients and formally given by the inner products $\gamma_{j,m} = \langle q, \psi_{j,m} \rangle$, where $\psi_{j,m}$ is a dual wavelet function.

The intermediate wavelet coefficients represent bandpass information and the highest resolution wavelet coefficient represents the high-pass information. The coarsest level scaling function and all wavelet scaling functions construct a basis for the function space $L^2$. See Fig. 3 for examples of spherical wavelets and a scaling function.

3.2 Discretization

Our approach for discretizing the gradient flow (17) has its mathematical foundation in Discrete Exterior Calculus (DEC) theory that was proposed in [38], [15]. The power of this method is the careful definition of discrete differential quantities, designed to respect structural relationships such as vector calculus identities. This is quite different from previous methods, which focused on satisfying the continuous equations at a discrete set of spatial and temporal samples, but failed to preserve important global structures and invariants. Both fluid mechanics and electromagnetism make heavy use of line integrals, as well as surface and volume integrals. Pointwise evaluations or approximations for such quantities may not be the appropriate discrete analogs, since the defining geometric properties underlying their physical meaning cannot be enforced naturally. Instead, one should consider values on vertices, edges, and faces as proper discrete versions of pointwise functions, line integrals, and surface integrals, respectively.

The main idea behind this approach is the representation of fields through measurements on cells: a 0-form represents a scalar function through its values at vertices
vertices. This is done in Appendices A and B.

Once we find the optimal mapping from a general simply connected compact surface to the sphere, we can then synthesize an image on the sphere, attributing a color to every vertex or triangle, and use the inverse mapping \( g^{-1} \) to map this image onto the original surface.

In the following sections, we will elaborate on the implementation of each of these steps.

### 4.1 Finding the Conformal Mapping

We employ the method for conformal mapping proposed in [5]. We briefly outline the relevant steps. The idea is to remove a triangle, solve the Dirichlet problem, and thus, conformally map the remaining surface onto the complex plane, and finally, use the inverse stereo projection to map the plane onto \( S^2 \). The boundary is mapped to a triangle around the “North Pole” of \( S^2 \).

More precisely, we start with a manifold represented by a triangular mesh with \( N \) vertices from which we remove one triangle \( \triangle ABC \) and then perform the following steps:

- Calculate the matrix \( D \).
- \( D \) is a sparse and symmetric \( N \times N \) matrix whose nonzero elements \( D_{PQ}(P \neq Q) \) are
  \[
  D_{PQ} = -\frac{1}{2} \left( \cot \angle R + \cot \angle S \right),
  \]
  where \( \angle R \) is the angle at the vertex \( R \) in the triangle \( \triangle PQR \) and \( \angle S \) is the angle at the vertex \( S \) in the triangle \( \triangle PQS \) (see Fig. 4a).

  Note that \( D_{PQ} \neq 0 \) only if \( P \) and \( Q \) are connected by some edge in the triangulation. The diagonal elements \( D_{PP} \) satisfy
  \[
  \sum_P D_{PQ} = 0.
  \]

- Calculate the vectors \( a \) and \( b \).
The variables $a$ and $b$ are sparse $N$ vectors with an entry for each vertex. Their nonzero elements are found at the vertices $A, B, C$ of the triangle $\triangle ABC$ that we removed from the mesh:

$$a = \begin{cases} 0 & Q \notin \triangle ABC \\ \|B - A\| & Q = A \\ \frac{1}{\|B - A\|} & Q = B \\ 0 & Q = C, \end{cases}$$

$$b = \begin{cases} 0 & Q \notin \triangle ABC \\ \frac{1 - \theta}{\|C - E\|} & Q = A \\ \frac{\theta}{\|C - E\|} & Q = B \\ \frac{1}{\|C - E\|} & Q = C, \end{cases}$$

where $E$ is the orthogonal projection of $C$ on $AB$ (see Fig. 4a) and

$$\theta = \frac{\langle B - A, C - A \rangle}{\|B - A\|^2}.$$  

- Solve the linear system of equations:

$$\begin{cases} Dx = a , \\
Dy = b. \end{cases}$$

- Map the $x$ and $y$ coordinates from the plane to the unit sphere using inverse stereographic projection:

$$x + iy \rightarrow \left( \frac{2x}{1 + r^2}, \frac{2y}{1 + r^2}, \frac{2r^2}{1 + r^2} - 1 \right).$$

As noted above, the conformal mapping distorts the area of the surface (see Fig. 5, where the nose of the “Max Planck” image is mapped to a smaller area with proportion to his eyes). We quantify the change of area as the density function preserving mapping, but rather just give the final algorithm that is applicable to all of our examples. Details may be found in [52], [13]. As mentioned above, the area preserving mapping from the original surface to the sphere will be the composition of the mapping $g$ and the conformal mapping $f$. The mapping $g$ is calculated by solving two differential equations on the sphere. We first solve

$$\Delta \Theta = 1 - \mu(x)$$

for $\Theta$, with

$$\mu = |\nabla f^{-1}|$$

and then set

$$u_t = -\frac{\nabla \Theta}{(1 - t)\mu + t}$$

for $t \in [0, 1]$. Finally given the family $u_t$, we solve for $g_t$ from the ODE

$$\frac{d}{dt} g_t = u_t \circ g_t$$

with $g_0$ taken to be the identity mapping. Then, $g = g_1$ is our required mapping.

After performing Step 2, we have an area preserving mapping with respect to the original domain $g \circ f$. However, this area preserving mapping is still not optimal in the transport sense defined above. In the next step, we will evolve this mapping to the optimal one.

### 4.3 Finding the Optimal MP Mapping

We now use the MP mapping obtained in Step 2, from now on, denoted by $g'$, as the initial mapping for the optimal transport algorithm. We will evolve $g'$ over time to obtain $g$ using the gradient descent flow as defined by the initial value problem (17). As $t$ goes to infinity, $g'$ will converge to the optimal mapping.

We start by calculating $\mu$ as the area distortion of every face after the conformal mapping. $\mu$ is calculated as the ratio of the...
original area of each triangle face to its area on the sphere after
conformal mapping. It should be emphasized that this
calculation is performed after we scale the original surface
so that its surface area is $4\pi$ (same as the area of the unit
sphere). We then find the vertices of the dual mesh (denoted by $x_d$) as the circumcenters of the triangles. Henceforth, all
quantities referring to the dual mesh will be denoted by a
subscript $d$ and those referring to the primal mesh will be
denoted by a subscript $p$. The vectors of the initial mapping $g^0$
are interpolated at the vertices of the dual mesh and denoted
by $g_d$. The mapping $g^0$ at the original vertices will be denoted
by $g_p$. The following steps are performed iteratively until we
converge to the optimal mapping.

We calculate $g_{p,n}$ at the dual mesh vertices and then
calculate the value of the function $H_p$ at the vertices of the
original mesh as the divergence of $\frac{dx_d}{dx_p}$ \( g_p \). Then, we find
the coefficients of its representation after the spherical
wavelet transform. We will next use only some of these
coefficients—at each iteration, we choose the coefficients
according to our stage in the multigrid algorithm (this
concept will be clarified in the next paragraph). We
perform the inverse transform using these coefficients
only. We then calculate our velocity field (at the vertices of
the dual mesh) as $v_d = -\frac{1}{\mu_d} J\nabla H_p = -\frac{1}{\mu_d} x_d \times \nabla H_p$. Note
that the result of the computation of the gradient is given at
the vertices of the dual mesh.

We next update $g_d$ according to this velocity field as
$g_d = dt \nabla g_p \cdot v_d$. This update is found by first interpolating
$g_d$ at the vertices of the primal mesh $x_p$ and then calculating
the Jacobian of $g_p$ using our formulation for the gradient
calculation. Here, $dt$ is the time step. The process is now
repeated with the updated $g_d$. We use smaller time steps
when we update our mapping using higher resolution
coefficients. The algorithm is sketched in pseudocode (1).

Algorithm 1. Perform One Step Toward Optimal MP Mapping

1: function OMT($g_d$, res)
2: for all triangles do
3: \( \text{temp}_d = \frac{dx_d}{dx_p} \)
4: end for
5: for all vertices do
6: \( H_p \leftarrow \nabla \cdot (\text{temp}_d) \) \{for Divergence
7: \( g_p \leftarrow \text{Interpolate}(g_d) \) \{linear interpolation\}
8: \( g_p \leftarrow \text{Normalize}(g_p) \)
9: end for
10: \( a \leftarrow \text{SWfwd}(H_p) \) \{Spherical wavelet transform,
11: \} see [59]\}
12: for all \( j > \text{res} \) do
13: \( a_{j,m} \leftarrow 0 \)
14: end for
15: \( H_p \leftarrow \text{SWbak}(a) \) \{Spherical wavelet transform,
16: \} see [59]\}
17: for all triangles do
18: \( v_d \leftarrow -\frac{1}{\mu_d} x_d \times \nabla H_p \) \{for Gradient
19: \} see Appendix VIII\}
20: \( g_d \leftarrow g_d - dt \cdot \nabla g_p \cdot v_d \) \{Jacobian using Gradient
21: \} operation\}
22: end for
23: end function

5 Examples

In this section, we give some examples illustrating our use
of optimal mass transport for texture mapping. We should
note that in some previous work [35], conformal mapping
theory was applied to the problem of texture mapping, and
so in the examples below, we explicitly compare the
optimal mass transport approach to this one. In Fig. 13,
we also compare texture mapping using our parameterization
to texture mapping and using the parameterization
proposed in [57] and [30].

Some of the textures we used were created directly on the
sphere, such as the case of the circles pattern, or images taken
with an omnidirectional camera, as can be seen in Fig. 7.
However, most of the images were created on a 2D rectangle
and then mapped to the sphere using the (inverse) equirectangular projection. The equirectangular projection maps the longitudes and latitudes directly into the \( x \) and \( y \) coordinates on the plane, respectively. The poles (zenith, nadir) are mapped to the top and bottom edges and are stretched to the entire width of the image. Areas near the poles get stretched horizontally. This projection is easy to use because of the simple connection between the pixel coordinates on the plane and its azimuth and zenith angles on the sphere. However, it is neither area preserving nor conformal. Therefore, a preliminary step of correcting a given texture image has to take place in order to account for this distortion. This correction can be performed by transforming the azimuth angles. Assume that the image values are mapped vertically onto lines of latitude \(-\frac{\pi}{2} < \theta < \frac{\pi}{2}\) and mapped horizontally onto lines of longitude \(-\pi < \phi \leq \pi\). In order to correct the distortion, there is no need to modify \( \theta \), but \( \phi \) is scaled as we approach the two poles by \( \cos(\theta) \). In Fig. 8, one can see the lines on the 2D image that correspond to the lines of longitude after the transformation. An example of this mapping can be seen in Fig. 9.

Once we have the synthesized texture over the sphere, we use the inverse of our mapping (derived via optimal mass transport) to map this texture from the sphere to the 3D object as described above. In the mapping results in Figs. 10 and 11, one can see how our mapping corrects the area distortion with respect to the conformal but still manages to minimize the geometrical distortion. Note that in Fig. 10, the conformal mapping maintains the shape of the circles but may scale them differently, so that in some places, the circles are very large, while in others, they are quite small. In our mapping, all the circles are practically the same size, whereas the geometrical distortions are local and small. In Fig. 11, one can see the same effect for the puzzle texture. In Fig. 12, this improvement in quality is even more striking, since in this case because of the high curvature changes, the conformal mapping causes major area distortions, and almost all the land of the earth is mapped to a small portion of the bucket. Our mapping places the north pole on the bottom inside the bucket and the south pole on the bottom of the bucket from outside, so the rest of the world is evenly spread on the whole
In Fig. 13, texture mapping using our method is compared to the texture mapping results obtained using the parameterization method proposed in [57] and [30] for the Stanford bunny. In this case, while the extremities of the bunny's ears impose on most other parameterization methods a very dense parameterization, in our method, this is avoided. However, in this case, the geometrical distortion is fairly noticeable, despite being local.

6 Evaluation

We experimented with our code using a Matlab and mixed C++ implementation on a standard 2 GHz CPU with 2 GB RAM processor. Most of the code was implemented in Matlab, except for some spherical wavelet transform functions.

In order to quantify the distortion of the parameterization, we compute both the angle and area distortion metrics defined using the signed singular values of the Jacobian of the transformation for each triangle, as presented in [40], [14], and [49]. Small angular and area distortions are indicated by a distortion value approaching 2. We ran our algorithm, referred to as MK, on a variety of inputs, in order to evaluate the computation time and convergence rate. We compared the results of the MK algorithm with the results obtained after running the algorithm of Haker et al. [34] and that of Moser’s [52]. We also compared our results to those obtained using the method of Gotsman et al. [30] with Tutte weights and Mean-Value weights (using the implementation of Saba et al. [57]). The values of the distortion measures obtained by the various algorithms are summarized in Tables 1 and 2. One can see that the proposed algorithm produces maps with very small area distortion that also minimizes the angular distortion.

Running time of our algorithm for meshes of varying sizes is summarized in Table 3. The experiments we performed show that the number of iterations required for convergence is not determined solely according to the mesh size, and is in fact, even more affected by the extremities of the excrescences of the meshes. It should be noted that in our research within the medical imaging field, the meshes we use are those of anatomical structures, which do not have such extremities, and thus, we did not have to deal with this issue. However, we note that in case of such large extremities, besides longer computation time, triangle flipping may also occur in the Moser algorithm, as discussed below.

The calculation that dominates the computation time is that of finding the coefficients of the spherical wavelets representation. Otherwise, the computations in each iteration of the algorithm can be performed in parallel, therefore, an implementation on a GPU would probably eliminate most of the increase in running time versus mesh size that is evident in Table 3.

The convergence behavior of our algorithm for the Squirrel example is shown in Fig. 14. Obviously, one can see that the convergence rate is not linear and it takes a few hundred iterations to converge to the optimal solution. However, after several hundred iterations, the algorithm reaches a solution that is fairly close to the optimal solution.

It should be noted that our method is immune to triangle flipping. In the continuous setting, this is imposed by the bijectivity of our mapping (that can easily be shown using the maximum principle). In the discrete setting, we enforce bijectivity by using an implementation of the differential equations according to the DEC theory. As shown in Appendix B, this implementation allows us to maintain the mass preserving property of our velocity field even in the discrete settings, and thus, prevent triangle flipping. However, since the methods described in [34] and [52] are not bijective in the discrete settings, triangle flipping may occur in the preparation stages for our algorithm. Our algorithm

<table>
<thead>
<tr>
<th>Model</th>
<th># of faces</th>
<th># of iterations</th>
<th>time (sec)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Squirrel</td>
<td>5k</td>
<td>814</td>
<td>971</td>
</tr>
<tr>
<td>Gargoyle</td>
<td>20k</td>
<td>1687</td>
<td>1451</td>
</tr>
<tr>
<td>Max-Planck</td>
<td>25k</td>
<td>2120</td>
<td>2460</td>
</tr>
<tr>
<td>Skull</td>
<td>40k</td>
<td>1371</td>
<td>2132</td>
</tr>
<tr>
<td>Bunny</td>
<td>70k</td>
<td>2530</td>
<td>3093</td>
</tr>
</tbody>
</table>
cannot recover from triangle flipping in these stages. Therefore, for meshes with large extremities in which triangle flipping may occur (rarely in the method of [34] and more often in the method of [52]), an additional step of fixing these flips has to be applied to the mesh. Of course, this step may come at the cost of locally increasing the distortions.

7 CONCLUSION AND FUTURE WORK

In this paper, we introduced a novel method for the parameterization of 3D objects. In the proposed method, we find the optimal mapping in the sense that it is the area preserving mapping produced via optimal mass transport. The optimal mass transport map is found via a gradient flow directly computed on the sphere, implemented using a multiresolution scheme for fast convergence. We have also shown how we can use our method to map textures onto closed compact simply connected surfaces and compared our results to texture mappings derived from conformal mapping theory.

There are several directions that we plan to pursue in future work. First of all, we believe that we can improve the rate of convergence of our algorithm by refining the multigrid scheme. Further, we want to extend our methodology to higher genus surfaces. This is not completely straightforward since in this case, the general divergence-free vector field (on a two-dimensional surface) involves a harmonic term. In the simply connected case, this does not appear. In general, for a surface of genus $g$, the space of such harmonic forms has dimension $2g$. However, we believe that this is tractable and this extension will make our method applicable for texture mapping on more complicated surfaces. It should also be noted that an algorithm for computing harmonic 1-form group basis of a high genus surface has been introduced by Gu and Yau based on Hodge theory in [32].

Finally, we plan to apply our algorithm to real medical data including brain structures such as the caudate nucleus and hippocampus (whose surfaces are topological spheres) to support the work in statistical shape analysis for our research in schizophrenia.

APPENDIX A

DEC OPERATORS

In our implementation, we use the formulation of the DEC theory in [38]. Here, we repeat some of the key points in the derivation of the gradient and divergence operators.

A.1 Gradient

For a function $f$ (a primal 0-form), the gradient can be computed in the interior of the triangles (the primal simplices) by first interpolating the function from the primal vertices to the interior using affine, barycentric interpolation functions, and then taking the gradient. Since the interpolation is affine, the gradient is a constant vector, and we can associate it with the dual of the simplex.

The set of interpolating functions we use is defined as follows. The interpolation function $\varphi_i$ has the value 0 at all vertices other than vertex $i$ and the value 1 at vertex $i$. The piecewise linear interpolation of the discrete function $f$ at a point $x$ in any triangle $\triangle ABC$ can be written as

$$\hat{f}|_{\triangle ABC}(x) = \sum_{m=A,B,C} f(m) \varphi_m(x).$$

Taking the usual gradient of this smooth function in $\triangle ABC$, we have

$$\nabla \hat{f}|_{\triangle ABC}(x) = \sum_{m=A,B,C} f(m) \nabla \varphi_m(x). \quad (19)$$

In general, we have

$$\sum_{m=A,B,C} \nabla \varphi_m(x) = 0.$$

By substituting this into (19), we get

$$\nabla \hat{f}|_{\triangle ABC}(x) = \sum_{m=A,B} (f(m) - f(C)) \nabla \varphi_m(x).$$

The gradient value is independent of which vertex is chosen as the distinguished vertex $C$. The gradient of the interpolating function in triangle $\triangle ABC$ is perpendicular to the edge opposite the vertex from which the interpolating function stems, and its magnitude is $\frac{h}{r}$, where $h$ is the length of the perpendicular line from the vertex to the edge, as shown in Fig. 15.

A.2 Divergence

Let us denote the directed edges stemming from a vertex $i$ by $e_{ij}$ for $j = 1, 2, \ldots, n$, where $n$ is the number of mesh triangles that contain the vertex $i$. We denote by $e_{ij}^*$ an edge perpendicular to $e_{ij}$ that connects the circumcenters (the dual vertices) of the triangles adjacent to the edge $e_{ij}$. (Note that this is not a directed edge.) Each of these edges has two triangles adjacent to it. We denote the two triangles adjacent to an edge $e_{ij}$ by $\triangle ijk$ for $k = 1, 2$. The divergence of a vector field $V$ at vertex $x_i$ can then be calculated as...
calculate the gradient of the scalar function

\[ \text{PROOF OF MP PROPERTY} \]

Appendix A for the gradient and divergence, we maintain here, we show that, if we use the operators defined in formulations of these operators and their justification, we can show that

\[ \nabla \cdot (J \nabla H) = 0 \]

if \( H \) is a scalar function that takes its values on the vertices. The notation used in the following proof is elaborated in Fig. 16. Without loss of generality, we need only consider the contribution to the divergence at vertex \( x_A \) from triangle \( \triangle ABC \).

Our velocity field \( v \) is calculated as follows. First, we calculate the gradient of the scalar function \( H \):

\[ \nabla H|_{\triangle ABC} = (H(C) - H(A)) \nabla \varphi_C(x) + (H(B) - H(A)) \nabla \varphi_B(x). \quad (20) \]

We calculate the gradients of the interpolating functions as

\[ \nabla \varphi_B(x) = \frac{\hat{c}^i}{a \sin(\beta)}, \]

\[ \nabla \varphi_C(x) = \frac{\hat{b}^i}{a \sin(\gamma)}. \]

Substitution of these two expressions into (20) gives us

\[ \nabla H|_{\triangle ABC} = \frac{H(C) - H(A)}{a \sin(\beta)} \hat{c}^i + \frac{H(B) - H(A)}{a \sin(\gamma)} \hat{b}^i. \quad (21) \]

Therefore, the velocity field, obtained by 90 degree rotation of the above, is

\[ \mu v = -J \nabla H = \frac{H(C) - H(A)}{a \sin(\beta)} \hat{c} - \frac{H(B) - H(A)}{a \sin(\gamma)} \hat{b}. \quad (22) \]

We now take the divergence of our velocity field:

\[ \nabla \cdot (\mu v) = \mu v \cdot (l_e \cdot \hat{c} + l_b \cdot \hat{b}). \quad (23) \]

Using the properties of the circumcenter, we get

\[ l_e = \frac{c}{2} \cot(\gamma) \]

and

\[ l_b = \frac{b}{2} \cot(\beta). \]

Substitution of these two equations into (23) results in

\[ \nabla \cdot (\mu v) = \mu v \cdot (l_e \cdot \hat{c} + l_b \cdot \hat{b}) = \frac{H(C) - H(A)}{2a\sin(\beta)} \left( c \cdot \cot(\gamma) + b \cdot \frac{1}{2} \cos(\alpha)\cot(\beta) \right) - \frac{H(B) - H(A)}{2a\sin(\gamma)} \left( c \cdot \cot(\gamma) \cos(\alpha) + b \cdot \frac{1}{2} \cot(\beta) \right) \]

\[ = \frac{h(C) - h(A)}{2a\sin(\beta)} \left( c \cdot \cot(\gamma) + b \cdot \frac{1}{2} \left[ \sin(\alpha) \sin(\beta) - \cos(\gamma) \right] \right) - \frac{H(B) - h(A)}{2a\sin(\gamma)} \left( c \cdot \sin(\gamma) \sin(\alpha) - \cos(\beta) + b \cot(\beta) \right). \quad (24) \]

Now, using the sine formula \( \frac{c}{\sin(\gamma)} = \frac{b}{\sin(\alpha)} \), we get

\[ \nabla \cdot (\mu v) = \frac{H(C) - H(A)}{2a\sin(\beta)} \left( c \cdot \frac{\cos(\gamma)}{\sin(\gamma)} - b \cdot \frac{1}{\sin(\beta)} \cos(\gamma) + b \cdot \frac{1}{\sin(\beta)} \right) \]

\[ - \frac{H(B) - H(A)}{2a\sin(\gamma)} \left( c \cdot \frac{\cos(\beta)}{\sin(\gamma)} + b \cdot \frac{1}{\sin(\beta)} \right) \]

\[ = \left( H(C) - H(A) \right) \frac{b \cdot \sin(\beta)}{2a\sin(\beta)} \]

\[ - \left( H(B) - H(A) \right) \frac{c \cdot \sin(\alpha)}{2a\sin(\gamma)} \]

\[ = \frac{1}{2} \left( H(C) - H(A) \right) - \frac{1}{2} \left( H(B) - H(A) \right) \]

\[ = \frac{1}{2} \left( H(C) - H(B) \right). \quad (25) \]

Recall that \( \frac{1}{2} \left( H(C) - H(B) \right) \) is the contribution to the divergence at vertex \( x_A \) from one of its adjacent triangles. If we sum over the contributions from all surrounding triangles, we will get that the value of the scalar function
H at each of the surrounding vertices will show up once with a positive sign and once with a negative sign—and will, therefore, cancel out, adding to a total divergence of zero.

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