An Unified Multiscale Framework for Planar, Surface, and Curve Skeletonization

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Abstract—Computing skeletons of 2D shapes, and medial surface and curve skeletons of 3D shapes, is a challenging task. In particular, there is no unified framework that detects all types of skeletons using a single model, and also produces a multiscale representation which allows to progressively simplify, or regularize, all skeleton types. In this paper, we present such a framework. We model skeleton detection and regularization by a conservative mass transport process from a shape’s boundary to its surface skeleton, next to its curve skeleton, and finally to the shape center. The resulting density field can be thresholded to obtain a multiscale representation of progressively simplified surface, or curve, skeletons. Our method is simple to implement, computationally efficient, and easy to use. We show that our results are very similar to the ones produced by several existing 2D and 3D skeletonization methods on a set of complex shapes.

The structure of this paper is as follows. Section II reviews related work. Section III presents our skeletonization method. Section IV details our method’s implementation. Section V compares our results with one 2D, six 3D surface, and 11 curve skeletonization methods. Section VI discusses our results. Section VII concludes the paper.

Index Terms—Medial axes, Skeleton regularization, Physically-based shape processing.

I. INTRODUCTION

Skeletons, or medial axes, are shape descriptors used in virtual navigation, shape matching, shape reconstruction, and shape processing [60]. 3D shapes admit two types of skeletons. Surface skeletons are 2D manifolds which contain the loci of maximally-inscribed balls in a shape [50], [60]. Curve skeletons are 1D curves which are locally centered in the shape [16]. Surface-skeleton points, with their distance to the shape and closest-skeleton points, define the medial surface transform (MST), used for animation, smoothing, and matching [4], [7], [20].

Many methods exist for computing 2D skeletons [20], [47], [69], 3D surface skeletons [27], [53], [59], [64], and 3D curve skeletons [6], [18], [26], [68]. Although recent methods demonstrate high accuracy, insensitivity to noise, and computational efficiency, several challenges remain open. We focus here on two modeling challenges, as follows. First, 2D skeletons, 3D surface skeletons, and 3D curve skeletons are typically extracted, and next simplified, using different methods and metrics. This makes the comparison and the formal reasoning about the properties of the extracted skeletons difficult. Second, few (if any) methods offer a continuous multiscale representation that addresses all skeleton types, i.e., a model which encodes both the geometric importance of any skeleton point (useful for simplifying, or regularizing, noisy skeletons) and the type of skeleton point (non-skeleton, surface skeleton, or curve skeleton).

In this paper, we present a framework for 2D and 3D curve-and-surface skeletonization that addresses the above two goals. We model both the skeleton detection and its importance using an advection principle that collapses mass from a shape boundary to its skeleton and next to the skeleton center (in 2D); and from the boundary to the surface skeleton, next to the curve skeleton, and finally to the latter’s center (in 3D). This allows us to detect all types of mentioned skeletons, and also to regularize them, e.g., to remove detail branches, via a single model and a simple thresholding operation. We propose a single algorithm that unifies skeleton detection and regularization in 2D and 3D, and also establishes a formal connection between surface and curve skeletons. Our method is simple to implement, computationally efficient, and easy to use. We show that our results are very similar to the ones produced by several existing 2D and 3D skeletonization methods on a set of complex shapes.

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II. RELATED WORK

Given a shape $\Omega \subset \mathbb{R}^n$, $n \in \{2, 3\}$ with boundary $\partial \Omega$, we first define its Euclidean distance transform $DT_{\partial \Omega} : \mathbb{R}^n \to \mathbb{R}^+$

$$DT_{\partial \Omega}(x) = \min_{y \in \partial \Omega} \|x - y\|. \quad (1)$$

The skeleton, or medial axis, of $\Omega$ is next defined as

$$S_{\Omega} = \{x \in \Omega \mid \exists f_1, f_2 \in \partial \Omega, f_1 \neq f_2, \|x - f_1\| = \|x - f_2\| = DT_{\partial \Omega}(x)\} \quad (2)$$

where $f_1$ and $f_2$ are the contact points with $\partial \Omega$ of the maximally inscribed balls in $\Omega$ centered at $x$ [23], [53]. The points $f_1$ and $f_2$ are called feature transform (FT) points [65]. The vectors $f-x$ are called spoke vectors [63]. For $n = 2$, $S_{\Omega}$ is a set of curves which meet at the so-called skeleton junction points [20]. For $n = 3$, $S_{\Omega}$ is a set of manifolds with boundaries which meet along a set of so-called Y-intersection curves [14], [17], [36].

In contrast to 2D and surface skeletons (Eqn.2), 3D curve skeletons $C_{S_{\Omega}}$ admit many definitions [16], implemented by a wide variety of methods (discussed further below). As such, a formal relationship between $S_{\Omega}$ and $C_{S_{\Omega}}$ is still not unanimously accepted. For instance, although it is commonly accepted that $C_{S_{\Omega}}$ should be centered within $S_{\Omega}$, only few skeletonization methods use and/or enforce this property [30], [53].

Skeletons can be computed by various methods, as follows.

Thinning: Thinning removes $\partial \Omega$ voxels (or pixels in 2D) while preserving connectivity [8], [48], [52]. Although simple and fast, thinning can be sensitive to Euclidean transformations.

Field methods: These methods find $S_{\Omega}$ along singularities of $DT_{\partial \Omega}$ or related fields [22], [27], [33], [37], [55], [69], [72] and can be efficiently done on GPUs [13], [65], [66]. General-field methods use fields smoother (with fewer singularities) than distance transforms [1], [4], [16], [26], and thus are more
robust for noisy shapes. Siddiqi et al. find the skeleton as the non-zero divergence locus of $\nabla D T_{\partial \Omega}$ [59]. However, $\nabla \cdot (\nabla D T_{\partial \Omega})$, with $\nabla$ the divergence operator, can be non-zero also at non-skeletal points. Torsello and Hancock correct this for a more accurate 2D skeleton detection by a momentum conservation principle $\nabla \cdot (\rho \nabla D T_{\partial \Omega}) = 0$, where $\rho$ is the mass density on the evolving boundary $\partial \Omega$ [5]. Rossi and Torsello extend this idea to compute 3D surface skeletons [54]. However, this method does not compute curve skeletons and does not model the curve-surface skeleton relationship.

**Mesh-based methods:** Field methods volumetrically sample $\Omega$, which can be expensive memory-wise. Mesh-based methods use a surface sampling of $\partial \Omega$, which allows processing higher-resolution shapes. Mesh methods include Voronoi diagrams to compute polygonal skeletons [19]. Amenta et al. compute the Power Crust, an approximation of a surface and its medial axis by a subset of Voronoi points [2]. Other methods use edge collapses [38], starting from a mesh segmentation [32]. Surface skeletons can be extracted from oriented point clouds [29], [41] or polygon meshes [36], [45] by searching for maximally inscribed balls tangent at least two shape points. Curve skeletons can be extracted from point clouds as centers of cloud projections on a cut plane which optimizes for circularity [68]. Contraction techniques are a separate subclass of mesh methods. Like field techniques, they evolve $\partial \Omega$ under various types of normal flows, effectively collapsing it onto the surface-or-curve skeleton. Methods using a (constrained) Laplacian contraction by mean curvature flow deliver high-quality curve skeletons [6], [12], [15], or even ‘meso skeletons’ mixes of surface and curve skeletons [67]. A different approach is taken by Jalba and Telea who contract the surface skeleton to compute its curve skeleton counterpart [30]. A recent review of contraction methods is given in [62].

**Multiscale skeletons:** Clean skeletons are extracted from noisy shapes by thresholding importance measures $\rho: \Omega \to \mathbb{R}^+$. This prunes skeletal branches caused by small details [17], [58]. We distinguish between local and global measures [44], [53]. Local measures cannot separate locally-identical, yet globally-different, contexts (see e.g. [53], Fig.1). Thresholding local measures can disconnect skeletons. Reconnection needs extra work [42], [50], [59], [66], and makes pruning less intuitive [58]. Local measures include the angle between the feature points and distance-to-boundary [2], [21], [66], divergence-based [11], [59], first-order moments [55], and points where $\nabla D T_{\partial \Omega}$ is multi-valued [63], [64]. Leymarie and Kimia topologically simplify point-cloud skeletons to capture Y-intersection curves and skeleton sheet boundaries in medial scaffolds [36]. A good survey of such methods is given in [60].

Global measures monotonically increase from the skeleton boundary $\partial \Omega$ inwards. Thresholding them yields connected skeletons which capture skeleton details at a user-given scale. Miklos et al. approximate shapes by unions of balls (UoB) and use UoB medial properties [24] to simplify skeletons [44]. Dey and Sun introduce the medial geodesic function (MGF), equal to the shortest-geodesic length between feature points [18], [51]. Reniers et al. [53] extend the MGF for surface and curve skeletons using geodesic lengths and surface areas between geodesics, respectively, inspired by the so-called collapse metric used to extract multiscale 2D skeletons [20], [47], [69]. A fast GPU implementation of this extended MGF is given in [29].

The MGF and its 2D collapse metric counterpart have an intuitive geometric meaning: They assign to a skeleton point $p$ the amount of shape boundary that corresponds, or ‘collapses’ to, $p$ by some kind of boundary-to-skeleton mass transport. Skeleton points $p$ with low metric values correspond to small-scale shape details or noise; points $p$ with large metric values correspond to large-scale shape details. This allows an easy simplification of the skeleton: Thresholding by a value $\tau$ eliminates all skeleton points which encode less than $\tau$ boundary length or area units. If the collapse metric monotonically increases from the skeleton boundary to its center, thresholding delivers a set of connected and nested skeleton approximations, also called a multiscale skeleton [18], [20], [53], [69].

### III. Proposed framework

#### A. Preliminaries

Following the above, we aim to create a single model that

1) unifies the representation and detection of 2D skeletons, 3D surface skeletons, and 3D curve skeletons;
2) computes a monotonic, global importance metric for 2D and 3D skeleton regularization and simplification;

Conceptually, we aim to capture the desirable properties of the 2D and 3D boundary collapse metric [20], [47], [53], [69] in a single model, and also connect contraction-based and distance-field based methods in a single framework. Practically, we aim at a single, easy to implement and use, and computationally efficient method that extracts and regularizes all skeleton types.

To achieve this, we first introduce our unified skeleton definition: Given a shape $\Omega \subset \mathbb{R}^{2,3}$, we aim to compute an importance function $\lambda: \Omega \to \mathbb{R}^+$ so that the threshold sets $\lambda_t = \{ x \in \Omega | \lambda(x) \geq t \}$ capture all existing skeleton types and all their simplifications. Specifically, we want $\lambda_0$ to be the full input shape $\Omega$; $\lambda_\varepsilon$ (for a small $\varepsilon > 0$) to be the full (unsmoothed) surface skeleton $S_\Omega$, which implies that $\lambda(x) = 0, \forall x \notin S_\Omega$.

As $\tau$ increases, we want $\lambda_\tau$ to be progressively simplified surface skeletons, and as $\tau$ increases even further, progressively simplified curve skeletons. In the limit, when $\tau = \max_{x \in \Omega} \lambda(x)$, we want $\lambda_\tau$ to be a single point (for genus 0 objects), which we next call the shape center $C_\Omega$. This process can be seen as a ‘recursive’ skeletonization which computes the surface skeleton from the input shape, the curve skeleton from the surface skeleton, and the shape center from the curve skeleton. All skeletons $\lambda_\tau$ should satisfy the well-known desirable properties – centeredness, rotational invariance, homotopy to the input shape $\Omega$, noise robustness, one-pixel (in 2D) and one-voxel (in 3D) thickness, inclusion of the curve skeleton in the surface skeleton, and computational efficiency [16], [61], [62].

#### B. Physically-based skeletonization model

For a shape $\Omega \subset \mathbb{R}^{2,3}$, we model our unified skeletonization as a contraction of $\Omega$ on whose boundary mass is distributed with unit density. Contraction is described by three fields: $\phi(x,t)$, $\rho(x,t)$, and $u(x,t)$, with $x \in \Omega$, and with $t \in \mathbb{R}^+$ being the time parameter, as follows. Similar to phase-field models [18], the field $\phi \to [-1,1]$ is 1 inside $\Omega$ and -1 outside, so that the boundary of the contracting shape is implicitly given by
\[ \Gamma = \{ x \in \Omega \mid \phi(x, t) = 0 \}. \] For now, we assume that \( \phi \) varies abruptly and monotonically over \([-1, 1]\) in a small vicinity around \( \Gamma \). The field \( \rho \to \mathbb{R}^+ \) gives the mass density of \( \Gamma \).

Finally, \( u \to \mathbb{R}^n \) gives the contraction direction of \( \Gamma \).

Our contraction is described by a system of three PDEs:

\[ \frac{\partial \rho}{\partial t} + \nabla \cdot (\rho u) = 0, \quad \frac{\partial \phi}{\partial t} + u \cdot \nabla \phi = 0, \quad \text{with } \Gamma_t \cong \Gamma_0, \]

\[ u = \frac{\nabla \phi}{\| \nabla \phi \|}. \]  

Equation 3 imposes mass conservation on the shrinking boundary. Equation 4 models boundary contraction with the topological homeomorphic constraint \( \Gamma_t \cong \Gamma_0 \). This ensures that the computed skeletons are homotopic to the input shape \( \partial \Omega = \Gamma_0 \). Equation 5 imposes inward contraction of our shape, with unit speed in normal direction to \( \Gamma \).

Eliminating \( u \) from Eqns. 3-5, we obtain

\[ \frac{\partial \rho}{\partial t} = -\nabla \rho \cdot \frac{\nabla \phi}{\| \nabla \phi \|} - \rho \nabla \cdot \frac{\nabla \phi}{\| \nabla \phi \|} = -\nabla \rho \cdot \phi \frac{\nabla \phi}{\| \nabla \phi \|} - \rho \kappa \]

\[ \frac{\partial \phi}{\partial t} + \| \nabla \phi \| = 0, \quad \text{with } \Gamma_t \cong \Gamma_0 \]

where \( \kappa \) is the (mean) curvature of \( \Gamma_t \).

Equations 6-7 are supplemented by the initial conditions

\[ \phi(x, t = 0) = \begin{cases} 1, & \text{if } x \in \Omega \\ -1, & \text{if } x \notin \Omega \end{cases} \]

\[ \rho(x, t = 0) = \begin{cases} 1, & \text{if } x \in \partial \Omega \\ 0, & \text{if } x \notin \partial \Omega \end{cases} \]

Let us define the time-of-arrival function \( T : \Omega \to \mathbb{R}^+ \) so that

\[ \phi(x, t) = T(x) - t. \]

Hence, \( \Gamma_t = \{ x \in \Omega \mid T(x) = t \} \), i.e., \( T(x) \) is the time after which \( \Gamma_t \) passes through \( x \). Using Eqns. 7 and 10, we obtain \( \| \nabla T \| = -\phi(t, x) = 1 \), the well-known Eikonal equation for arrival time \( T \). The Euclidean distance transform \( DT_{\partial \Omega} \) is the weak solution of this equation under Euclidean norm \([57]\). Hence, Eqn. 7 without the constraint \( \Gamma_t \cong \Gamma_0 \) is the PDE generating continuous multi-scale (flat) morphological erosions. Other norms are also possible \([43]\), leading to various distance transforms.

We finally define the skeleton importance \( \lambda \) as the maximum density that has reached a certain location \( x \in \Omega \), i.e.,

\[ \lambda(x) = \max_{t > 0} \rho(x, t). \]

Intuitively, our model describes a conservative advection process where mass, uniformly spread on \( \partial \Omega \), flows on shortest paths from \( \partial \Omega \) to its surface skeleton \( S_{\Omega} \); then, along \( S_{\Omega} \) on shortest paths to the curve skeleton \( C_{\Omega} \); and finally along \( C_{\Omega} \) on shortest paths to the shape center \( C_\Omega \) (Fig. 1). Once all mass has reached \( C_\Omega \), we compute the (simplified) surface and curve skeletons by thresholding \( \lambda \) at increasing values.

**IV. SOLVING THE SYSTEM**

To compute the importance \( \lambda \), we solve the contraction model in Sec. III-B by discretizing \( \Omega \) on a uniform cubic-cell (pixel or voxel) grid embedded in \( \mathbb{R}^2 \) and \( \mathbb{R}^3 \) respectively, as follows.

**A. Topologically-constrained boundary evolution by density-ordered thinning**

As stated in Sec. III-B, Eqn. 7 must be solved with the constraint that \( \Gamma_t \) and \( \Gamma_0 \) are homeomorphic, for all \( t \). Even without this constraint, it is well-known that the evolution of \( \phi \) from Eqn. (7) develops discontinuities of the \( \phi \) derivatives (shocks) within finite time \([43]\), [57]). The skeleton \( S_{\Omega} \) precisely coincides with the locations of these shocks \([59]\).

Since Eqn. 7 can be written as an Eikonal evolution, or boundary-value problem (Sec. III-B), one way to interpret the contraction is as thresholding \( DT_{\partial \Omega} \) inside \( \Omega \) at increasingly higher values, producing multi-scale morphological erosions of \( \Omega \). Additionally, the topological constraint \( \Gamma_t \cong \Gamma_0 \) should also be satisfied by each level set of \( DT_{\partial \Omega} \) corresponding to \( \Gamma_t \). For achieving this, we could consider using topologically-constrained level sets \([25]\). The problem with this approach is that it first performs an un-constrained step to update level-set values, following the motion equation (Eqn. 7). Then, at points \( x \) where the topological constrained is violated, the so-called level-set function \( \psi \) is next ‘fixed’ so that the points \( x \) lie on the corresponding side of the boundary dictated by the constraint. This fix artificially alters the \( \psi \) values, which creates spurious and unwanted discontinuities in \( \psi \), ultimately leading to a not sharply-defined (in terms of our desired sharp transition of the level-set function in \([-1, 1]\)) and/or non-smooth evolution of \( \Gamma_t \).

In turn, this will drastically affect the quality of the extracted skeletons, as we verified in practice. As such, we chose not to use topologically-constrained level-sets for our context.

To handle all above issues, we use topology-preserving morphological thinning to define and steer the evolution (contraction) of \( \Gamma_t \). Our thinning process is ordered both by \( DT_{\partial \Omega} \) and by the density field \( \rho \); As long as \( \Gamma_t \) is far from the skeleton \( S_{\Omega} \), ordering by \( DT_{\partial \Omega} \) ensures a smooth \( \Gamma_t \) while solving Eqn. 7. Additionally, since thinning relies on a binary field, \( \Gamma_t \) is maintained sharp during its evolution.

We next explain why the thinning order is also given by the density field \( \rho \), which is crucial when the evolving \( \Gamma_t \) reaches the (yet unknown) skeleton locations. Recall that such locations correspond to shocks of Eqn. 7. Hence, ordering by \( DT_{\partial \Omega} \) (which is just a viscosity solution of Eqn. 7) becomes meaningless. As sketched in Fig. 1, we want the importance \( \lambda \),...
and thus also the density $\rho$ which determines this importance, to monotonically increase from $\partial \Omega$ to $S_\Omega$, next from $\partial S_\Omega$ to $CS_\Omega$, and finally from $\partial CS_\Omega$ (curve-skeleton endpoints) to $C_\Omega$. Since $\Gamma_t$ shrinks in normal direction (Eqn. 7), this is equivalent to transporting density on shortest paths from $\partial \Omega$ to $S_\Omega$ to $CS_\Omega$ and next to $C_\Omega$. Fig. 1 shows such a path (in white) on which the mass of a point $p \in \partial \Omega$ should flow during its advection to $C_\Omega$. Consider now the set of all such paths from all points on $\partial \Omega$ to $C_\Omega$. For a shape $\Omega$ of genus 0, following a reasoning similar to [53], these paths will form a tree having as leaves all (discrete) points of $\partial \Omega$ and $C_\Omega$ as root. The computation of $\rho$ by means of our contracting $\Gamma_t$ is analogous to traversing this tree from its leaves to the root. To ensure a correct density update, we thus need that, at any junction-point where several subtrees meet, all these trees to have been fully traversed and their roots’ densities to be thus correctly updated. This is why our thinning visits points in $\Gamma_t$ in increasing $\rho$ order.

Figure 2 illustrates our thinning for a 3D shape. When using density-ordered thinning, $\Gamma_t$ (drawn red) is kept smooth during collapsing. In contrast, if not using density ordering, the collapsing $\Gamma_t$ will quickly develop irregularities (Fig. 2 e-h, insets). In turn, these will create irregularities in the signal which will ultimately lead to jagged skeletons $\lambda_t$ after simplification.

### B. Algorithm

Summarizing the observations from Sec. IV-A, our contraction algorithm should:

- $R_1$: provide a sharp definition of the evolving boundary $\Gamma_t$;
- $R_2$: allow interleaved iterative solves of Eqns. 6 and 7;
- $R_3$: ensure a smooth evolution of $\Gamma_t$, steered by $DT_{\partial \Omega}$ and $\rho$;
- $R_4$: allow efficient computation.

Most existing thinning algorithms do not provide a representation of $\Gamma_t$ which satisfies all requirements $R_1-R_3$ above. For example, the divergence-driven thinning algorithm in [50] uses a sorted heap to ensure the correct processing order, thus fails to provide an explicit $\Gamma_t$ representation. In contrast, we use an explicit representation of $\Gamma_t$, modeled as a narrow-band of points (that is, pixels in 2D and voxels in 3D, respectively). Density is transported, according to Eqn. 6, only within this narrow-band, which is computationally efficient ($R_4$).

Let us note that some thinning algorithms combine the detection and removal of a so-called topologically-simple point in a single pass. The thinning result may then depend on the point processing order, as discussed in [28]. In contrast, we use an approach similar to [28], where we first find all simple points (detection phase), and eliminate these next (removal phase).

Our full skeletonization algorithm is now as follows (Alg. 1). During initialization, we compute the Euclidean distance transform $DT_{\partial \Omega}$ on $\Omega$ (line 2). Next, we initialize the full-grid fields $\rho$ (density), $\lambda$ (importance) and $M$ (binary description of the contracting shape) to their default values (line 3). We use $DT_{\partial \Omega}$ to label interior points $x \in \Omega$ with $M(x) = 2$ and initial boundary points $x \in \partial \Omega$ with $M(x) = 1$ respectively (lines 4-8). Finally, we set the density of boundary points to one and gather them in the set $Q_1$. This set will keep, during the algorithm execution, all points processed by the current algorithm iteration.

The main loop (lines 10-29) iteratively solves the system of Eqns. 6 and 7. Here, the field $M$ has two roles. First, $M$ labels points outside ($M = 0$), on the boundary ($M = 1$), and respectively inside ($M = 2$) the shrinking shape, thus efficiently keeps track of this shape. Secondly, we use $M$ to check if a shape point is topologically simple or not: the function $\text{simple}(x, M)$ returns true if removing $x$ from the shape given by $M(\cdot) > 0$ does not change the shape’s topology and false otherwise.

The first inner loop (lines 12-14) fills a set $Q_2$ with unprocessed, 26-connected (8-connected in 2D) neighbors $y \in \mathcal{N}(x)$ of the point $x$ being processed. The set $Q_2$ captures points going to be processed in the next algorithm iteration (detailed further below). Next, we sort the current set $Q_1$ on increasing $\rho$ order (line 15), allowing the second inner loop (lines 17-19) to process these in increasing order of their density values. This second loop performs the detection phase of the thinning algorithm. Additionally, only points which are topologically-simple and within close distance ($\Delta d$) to the current $\Gamma_t$ are added to set $C$ for further processing. The other non-simple points are added to $Q_2$ for processing in the next iterations. The third inner loop (lines 21 to 25) performs the removal phase of the thinning algorithm. Topologically-simple points $x \in C$ are removed (by labeling them with $M(x) = 0$) and collected in the narrow-band set $B$. At this stage, their importance $\lambda$ is also computed (line 24). Non-simple points are added to $Q_2$.

Set $B$ models the current boundary $\Gamma_t$, thus meeting $R_1$. As shown in Alg. 1, $B$ is built from current topologically-simple points (from $Q_1$) in increasing $\rho$ order and by filtering them via the distance-threshold criterion (line 18). Hence, the boundary $\Gamma_t$ is kept relatively smooth (discussed further in Sec. IV-C), thus $R_3$ is met. Once $B$ is available, we can transport density (line 26) from points in $B$ to interior points, thus meeting $R_2$. After density has been conservatively transported away from the current $B$, we set the density to zero at points $x \in B$ (line 27).

### Algorithm 1: Skeletonization algorithm.

```plaintext
1 Skeletonize(Shape $\Omega$, Field $\lambda$)
2 Data: $\Omega$: discretized input shape
3 Result: $\lambda$: importance field
4 $DT_{\partial \Omega}(\cdot)$ ← 3D Euclidean distance transform of $\partial \Omega$;
5 $\rho(\cdot) ← 0$; $\lambda(\cdot) ← 0$; $M(\cdot) ← 0$; $Q_1 ← \emptyset$;
6 foreach $x \in \Omega$ do
7 if $DT_{\partial \Omega}(x) > 0$ then $M(x) ← 2$; // interior points
8 if $DT_{\partial \Omega}(x) < 2$ then $Q_1 ← Q_1 \cup \{x\}$; // boundary points
9 $d ← 0$;
10 repeat
11 $d ← d + \Delta d$; $Q_2 ← \emptyset$;
12 foreach $x \in Q_1$ do
13 if $DT_{\partial \Omega}(x) < d$ then $C ← C \cup \{x\}$;
14 $Q_2 ← Q_2 \cup \{x\}$; $M(x) ← 1$;
15 Sort $Q_1$ in increasing $\rho$ order;
16 $C ← \emptyset$;
17 foreach $x \in Q_1$ // process $Q_1$ in increasing $\rho$ order
18 if $DT_{\partial \Omega}(x) < d \&\& \text{simple}(x, M)$ then $C ← C \cup \{x\}$;
19 else $Q_2 ← Q_2 \cup \{x\}$; $M(x) ← 1$;
20 $C ← C \cup \{x\}$;
21 foreach $x \in C$ do
22 if $\text{simple}(x, M)$ then
23 $M(x) ← 0$; $B ← B \cup \{x\}$;
24 $\lambda(x) ← \max(\lambda(x), \rho(x))$;
25 else $Q_2 ← Q_2 \cup \{x\}$; $M(x) ← 1$;
26 Transport $\rho$ from $x \in B$ to interior points, using Eqn. 6;
27 foreach $x \in B$ do $\rho(x) ← 0$;
28 swap($Q_1$, $Q_2$);
29 until $B = \emptyset$.
```

At the end of the algorithm’s main loop, the sets $Q_1$ and $Q_2$ are swapped (line 28). This is an essential aspect of our algorithm, as it facilitates an explicit and computationally-efficient representation of the boundary $\Gamma_t$ (set $B$ above). Having these sets, we can limit our computations only to a surface-like band of points around the current $\Gamma_t$, thus meeting $R_4$.

The algorithm stops when $B$ becomes empty. For objects of genus 0, this happens when the shrunk shape and $Q_1$ contain only a single point, which is precisely the shape center $C_\Omega$. This point clearly cannot be added to $B$ since the topological constraint would be violated. For objects of higher genus, termination happens when the shrunk shape contains only curve-skeleton loops connected by non-terminal branches, a structure which cannot be shrunk any longer without disconnecting the skeleton (see example further in Sec. IV-D).

Let us now discuss the smoothness of the boundary $\Gamma_t$ captured by the set $B$. Away from singular points of Eqn. 7, $\Gamma_t$ is captured (by the set $B$) as a level-set of $DT_{\partial\Omega}$, which is a Lipschitz-continuous function under the $L_2$ metric. At singular points of Eqn. 7, our thinning still endorses Lipschitz continuity, but under the $L_\infty$ metric. Intuitively, at such points, the order in which new inner points are processed in lines 12-14 reinforces the $L_\infty$ metric when solving Eqn. 6. Note that, although at non-singular points, topologically-constrained level sets [25] provide better smoothness properties of the evolving boundary (due to sub-pixel or sub-voxel precision), such methods have problems regarding the evolution of $\Gamma_t$ along shocks of Eqn. 7, as discussed in Sec. IV-A. In contrast, our approach produces a smooth shrinking $\Gamma_t$, as shown by Fig. 2 a-d.

C. Density transport

We now focus on solving the mass conservation equation (Eqn. 6; Alg. 1, line 26). For 2D and 3D curve skeletons, discretizing Eqn. 6 with the unconditionally-stable, semi-Lagrangian scheme in [35] suffices. However, generating progressively-simpler surface skeletons by simply thresholding the importance field $\lambda$ requires additional work.
at a grid cell $i$ of $\Gamma_{t+1}$ for time-step $t+1$ can be expressed as
\begin{equation}
\rho_i^{t+1} = \rho_i^t + \sum_{j \in \mathcal{N}_i} \frac{1 - \chi_j^t}{\sum_{k \in \mathcal{N}_j} \chi_k^t} \rho_j^t,
\end{equation}
with being $\mathcal{N}_i$ the 26-connected neighborhood centered at $i$ for the 3D case. Let $w_j^t = \frac{1 - \chi_j^t}{\sum_{k \in \mathcal{N}_j} \chi_k^t}$ and assume that cell $i$ receives density contributions from three surrounding cells $j \in \{1,2,3\}$ in $\Gamma_t$ (see Fig. 4), i.e.,
\begin{equation}
\rho_i^{t+1} = \rho_i^t + \rho_i^t w_1^t + \rho_i^t w_2^t + \rho_i^t w_3^t.
\end{equation}
This can be rewritten as
\begin{equation}
\rho_i^{t+1} - \rho_i^t = \sum_{k=1}^3 w_k^t (\rho_k^t - \rho_i^t) + \rho_i^t \sum_{k=1}^3 w_k^t,
\end{equation}
which is a discretization of anisotropic diffusion [49] with an additional reaction term
\begin{equation}
\frac{\partial \rho}{\partial t} = \nabla \cdot (w \nabla \rho) + \rho \mathcal{C}.
\end{equation}

Figure 3. Density transport via advection vs diffusive advection (see Sec. IV-C).

Since our algorithm solves Eqn. 7 under the $L_\infty$ norm along its singular points (see Sec. IV-B), and since noise, small errors and inaccuracies due to the thinning process propagate into the density field evolution (Eqn. 6), simply thresholding $\lambda$ would yield jaggies (indentations) in surface skeletons, for all but trivial shapes (see example in Fig. 3a). To tackle this, we propose a smoothing of the density field $\rho$, which leads to the desired importance field $\lambda$, as follows.

The key idea of [35] for solving conservative-advection PDEs similar to Eqn. 3 is to follow so-called characteristic curves (along which the PDE becomes an ODE) both forwards and backwards in time, while ensuring that interpolation weights are equal to one for all grid cells, i.e., the advected density is conserved. We constrain the density $\rho$ to be zero outside the shrinking shape, so we only need the forward step. Figure 4 left shows a schematic example, assuming that density is transported along surface-skeleton points. For illustration simplicity, and without loss of generality, we next assume that $\rho$ is one at all points in $\Gamma_t$. The density propagation directions, given by $\nabla \phi/\|\nabla \phi\|$ (Eqn. 5), are shown by arrows. Hence, as shown in Fig. 4, the (linear) interpolation weights equal one, and the new density values at grid cells $a$, $b$ and $c$ have the indicated values. Since all weights equal one, mass is conserved, as desired.

One way to tackle the above inaccuracies is by endorsing density advection with a (conservative) diffusive component, yielding a smoother evolution of $\rho$. For this, we propose an anisotropic diffusion process, which we dub diffusive advection (as opposed to the well-known diffusion-advection PDE). That is, instead of transporting density solely in the (potentially-noisy) gradient directions, we also allow density to diffuse to other surrounding nearby cells (Fig. 4 right). As can be seen by following the arrows, each ‘donor cell’ now contributes to multiple nearby cells. The weights along each arrow per donor cell, as well as the new density values of cells $a$, $b$ and $c$, are also shown. More formally, let $\chi'$ be the characteristic function of the shrinking shape, obtained, e.g., by upper-thresholding the field $M(\cdot)$ of Alg. 1 with value 1. Then, the new density value

Figure 4. Conservative advection vs. diffusive advection. Density is transported on the surface skeleton from $\Gamma_t$ to $\Gamma_{t+1}$ by left: conservative advection and right: diffusive advection. Arrows show the directions in which density is transported. The new density values at grid cells $a$, $b$ and $c$ are also shown.

D. Detecting different skeleton types

The importance field $\lambda$ allows us to easily detect both skeleton types and the shape’s center. Specifically, we have that $\mathcal{S}_\Omega = \{ \bm{x} \in \Omega | \lambda(\bm{x}) \geq 2.0 \}$, since surface-skeleton points are, by definition, situated at equal distance from at least two different points on the boundary $\partial \Omega$ (Eqn. 2), and thus have an importance equal to at least that of two (collapsed)
points of $\partial \Omega$, i.e. at least two. For genus 0 objects which admit a center in the sense denoted in Sec. III-B, we have $C_\Omega = \arg \max_{x \in \Omega} \lambda(x)$. Curve-skeleton points could be readily detected by upper thresholding the importance field $\lambda$ with a large threshold $\tau$. However, there are two shortcomings with this approach: First, the resulting curve skeleton may not be always one-voxel thick. Secondly, its extremities may be removed due to the large threshold value used. In other words, for high $\tau$ values, we would obtain a simplified, rather than a full, curve skeleton. To alleviate these issues we detect salient curve-skeleton points, during the shrinking process, using

$$CS_\Omega = \{ x \in \partial \Omega | \rho(x) > c \hat{T}(x) \land \text{endPoint}(x,M) \},$$

where $c > 0$ is a constant (explained next); $\hat{T}$ is a simple estimate for the time-of-arrival (approximating $T$ from Eqn. 10), given by $d$ in Alg. 1; and $\text{endPoint}(x,M)$ returns true if $x$ is a curve-skeleton end point. We justify Eqn. 16 as follows.

First, Eqn. 16 only selects points from the surface skeleton, since only these have a density larger than two. Consider now a $b$ on the skeleton boundary $\partial S_\Omega$, such that $b \in S_\Omega \setminus CS_\Omega$ (Fig. 6). The density $\rho(b)$ equals the length of the circular segment $C(b) \subset \partial \Omega$ (drawn green in Fig. 6), which is $\alpha DT_{\partial \Omega}(b) = \alpha T(b)$ with $\alpha$ its subtended angle. Let $x$ be a neighbour of $b$ such that $T(x) = T(b) + 1$. Using the boundary evolution equation (Eqn. 7) and the arrival-time definition (Eqn. 10), it can be easily shown that $x$ must be in the ‘upstream’ direction from $b$, since $x - b$ and $\nabla \Phi$ are parallel vectors. When the evolving boundary passes through $b$ (i.e., $b$ is removed by density-ordered thinning, see Alg. 1), $\rho(b)$ is ‘pushed’ in the upstream direction through (diffusive) advection transport. Thus, $x$ will directly receive most density of $b$, under advective density transport. Moreover, due to the boundary-propagation order, when the interface is about to pass through $x$, point $x$ must also be found to be part of the surface skeleton, and furthermore, $\rho(x) > \rho(b)$. Indeed, since regular points $x \in S_\Omega \setminus CS_\Omega$ also receive density contributions from its two (or more) feature-points on $\partial \Omega$ (boundary normals are preserved by the Eikonal equation), a rough estimate for the minimum density at $x$ is $\rho(x) > 2 + \alpha T(b)$. By a similar reasoning, for points $y \in S_\Omega \setminus CS_\Omega$ situated in upstream direction from $b$, with $T(y) > T(b) + 1$, we find $\rho(y) > 2T(y) + \alpha T(b)$ as a lower bound on their density. Finally, points $c \in CS_\Omega$ collect density from at least two neighbor-points $y \in S_\Omega \setminus CS_\Omega$; this is so since the curve $CS_\Omega$ locally divides the 2D-manifold surface $S_\Omega$ in two parts. For instance, in Fig. 6, $c$ will receive density from at least two surface-skeleton neighbor points situated at the tips of the red, respective yellow, arrows. Additionally, $c$ receives a (larger) density contribution from (at least) another downstream curve-skeleton neighbor, so $\rho(c) > 4 \sum_i T(y_i) \geq 4T(c)$. Hence, setting $\tau = 4$ in Eqn. 16 performs conservative curve-skeleton detection. We verified empirically that setting $\tau = 4$ cleanly and robustly separates important curve-skeleton points from surface-skeleton points. Additionally, the end-point test in Eqn. 16 ensures that only points which are at the sharp ‘tips’ of the shrinking surface $\Gamma$ are considered as salient curve-skeleton candidates.

Once the points $x \in CS_\Omega$ are found using Eqn. 16, we ‘boost’ their importance during the iterative process by adding a constant fraction $\delta$ of the input shape’s mass $|\partial \Omega|$. As a result, curve-skeleton points will have a significantly higher importance than surface-skeleton points. Also, as $\delta$ is constant for all $x \in CS_\Omega$, the monotonic increase of importance along the curve skeleton branches is preserved. All in all, this allows us to threshold the final importance $\lambda$ at precisely $\delta$ to cleanly and easily detect a full (unsimplified) $CS_\Omega$, and to threshold $\lambda$ at higher values to obtain progressively simplified curve skeletons.

Figure 7 shows the effect of increasing the importance threshold $\delta$ for a shape of genus 3. Skeleton voxels are colored by importance $\lambda$ via a rainbow colormap. The first three $\tau$ values (Figs. 7a-c) yield three increasingly simple surface skeletons. The last of these (Fig. 7c) is a mix of surface and curve skeleton parts, i.e., is an example of the meso skeletons described in [67]. As $\tau$ increases to $\delta$, we find the full curve skeleton (Fig. 7d). Increasing $\tau$ further removes the end branches of the curve skeleton, without disconnecting its loops (Fig. 7e). For visual clarity, we used in Fig. 7e a local normalization of the colormap, based on the range of voxels in the respective simplified skeleton, rather than on the full importance range $[0, \lambda_{\text{max}}]$ used for all other figures. This shows better how the importance increases from the curve skeleton endpoints to its center, and is high over its loops. Finally, for the highest considered $\tau$ value, we get the fully simplified curve-skeleton without disconnections containing only loops connected by internal branches (Fig. 7f).

E. Implementation details and parameter settings

Implementation: Our algorithm is implemented in C++ using OpenGL for skeleton rendering. We compute $DT_{\partial \Omega}$
on the CPU by the method of Meijster et al. [27], or on the GPU (if available) by the method of Cao et al. [13]. Both methods compute the exact Euclidean distance transform and are linear in |Ω|, the number of foreground pixels or voxels in the input shape. We implement simple() using the Euler number for Ω ⊂ R^2 and Malandain’s criterion [9] for Ω ⊂ R^3 respectively. We detect curve-skeleton endpoints (function endPoint(), Eqn. 16) as those voxels x ∈ B with one (26-connected) neighbor y for which M(y) > 0. We use the integral method of Neumann et al. [46] to estimate the gradient directions along which density is transported (Eqn. 5). Sorting Q_1 is implemented by the Standard Template Library (STL) sort function. Algorithm 1 requires K iterations of the loop in line 10, where K is roughly equal to \frac{4}{\pi} \max_{x \in \Omega} DT_\partial \Omega(x), the maximal thickness of Ω. Since at each iteration we sort the set Q_1, which is worst-case equal to the input boundary ∂Ω, the total complexity of our method is O(K |∂Ω| log (|∂Ω|)).

**Parameters:** Throughout the paper we use the following parameter settings. To obtain a good approximation of the motion equation (Eqn. 7), we set the distance step ∆d in Alg. 1 to ∆d = 1. The parameter σ_a, controlling the density spread along gradient directions (Eqn. 15), is set to σ_a = 0.2. The parameter c in Eqn. 16, i.e., the saliency of the detected curve-skeleton points, is set to c = 4.0. The parameter δ, representing the importance difference between curve and surface skeleton points (Sec. IV-D), is set to δ = 0.1 of the total input surface mass |∂Ω|. The above values have been tested on a set of over 60 shapes, voxelized at various resolutions, and have consistently delivered good results like the ones shown in our figures here. As such, the only free parameter is the skeleton simplification threshold τ, whose use is explained in Sec. IV-D.

**V. COMPARATIVE RESULTS**

A. Two-dimensional skeletons

For 2D shapes, we compared our method with the Augmented Fast Marching Method (AFMM) [69]. AFMM is a good example of 2D skeletonization methods that computes centered, accurate, connected, and pixel-thin skeletons regularized by the collapsed boundary-length importance metric (like [20], [47]).

Figure 8 shows skeletons of several shapes from the database in [56] extracted with AFMM and with our method, for several simplification thresholds τ. Our method and the AFMM produce visually identical skeletons, both in terms of position, but also branches kept at a given τ. This is a non-trivial result, given that our method and the AFMM have completely different models behind. Moreover, since \lambda_{AFMM} at a skeleton point x equals the length of boundary that collapses to x when advected in \nabla DT_\partial \Omega, and since \lambda ≈ \lambda_{AFMM} (Fig. 8), this supports the claim that our \lambda is indeed equal to the collapsed boundary length.

B. Surface skeletons

Figure 9 (bottom 6 rows) compares our method with four voxel-based methods: multiscale skeletons (MS) [53], Hamilton-Jacobi (HJ) [59], integer medial axis (IMA) [27], and iterative thinning process (ITP) [31]; and with the multiscale mesh-based skeletonization (MBS) in [29]. Test shapes cover a wide range, including natural and synthetic, smooth and detailed, and objects of various genii (all voxelized at 512^3 resolution by binvox [48]). Our surface skeletons look very similar to those created by other methods, and show similar power in capturing the input shape genus and boundary details.

The comparison with MS is particularly interesting. To our knowledge, MS is the only existing voxel-based technique that computes multiscale surface skeletons whose importance uses a boundary-collapse metric. For MS, this metric is

\[ \lambda_{SS}(x \in S) = \min_{\gamma = (f_1 \leftarrow f_2) \subset \partial \Omega} ||\gamma|| \]

i.e. the length ||\gamma|| of the shortest geodesic path γ on ∂Ω between the two feature points f_1 and f_2 of a skeleton point x.

As for the 2D case (Sec. V-A), we see that our surface skeletons and importance values (color-coded in Fig. 9, row 12, by a rainbow colormap) are very similar to the MS ones (Fig. 9, row 7). Figure 2, two bottom rows, details this insight by showing four surface skeletons obtained by thresholding our importance \lambda, and \lambda_{SS}, at four increasing values. The monotonic increase of both \lambda and \lambda_{SS}, from low values on the surface-skeleton boundary to high values on the curve skeleton, and the resulting skeletons, are similar. This is an even more interesting result than the similarity of our results with the AFMM. Our importance \lambda(x) equals the amount of boundary mass which reaches x subject to Eqs. 6,7. The fact that \lambda ≈ \lambda_{SS} supports the conjectures in [18], [53] that all boundary points on such a geodesic γ, if advected in \nabla DT_\partial \Omega, would reach the skeleton point x. However, a formal proof of these conjectures still lacks. Separately, Fig. 2 (bottom row) shows that our \lambda monotonically increases as we advance inwards on the skeletal structures (Sec. IV), hence that thresholding \lambda yields connected skeletons.

To better qualitatively assess our surface skeletons, Fig. 10 compares these with skeletons computed by the high-resolution MBS method [29], which uses the multiscale-importance in [53], but has a different skeleton detector, and uses a mesh rather than a voxel representation. For more insight, we colored our surface-skeleton border voxels red. Our skeletons are very similar to the MBS ones. Our method captures roughly the same amount of skeleton detail as MBS, even though the latter uses mesh, rather than voxel, skeleton-and-shape representations. Figures 10 (4) shows our reconstruction of the input shape by drawing balls centered at the surface-skeleton voxels x and...
TABLE I

<table>
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<th>Model</th>
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<th>(768 × 768) voxels</th>
<th>Our method</th>
<th>TV</th>
<th>DDS</th>
<th>RT</th>
<th>ITP (curve)</th>
<th>MS (curve)</th>
<th>MS (surface)</th>
<th>HJ</th>
<th>IMA</th>
<th>ITP (surface)</th>
<th>MBS (surface)</th>
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whose radii equal $DT_{Ω}(x)$, using the rendering technique in [29]. As visible, our reconstructions are very close to the input shapes (Figs. 10 (3)). The small bubble-like differences are explained by the fixed resolution of the voxel grid. This verifies the reconstructibility criterion for our method and, implicitly, shows that our skeletons are correctly centered. The last row in Fig. 10 compares our method for the elephant shape from Fig. 7, which has three tunnels, large thin-and-flat areas (ears), near-cylindrical parts (legs), high positive-curvature areas (ear borders), and high negative-curvature areas (ear-head junctions). Our surface skeleton is very close to the one produced by MBS, and also to the skeleton produced by the discrete-scale axis (DSA) mesh-based method of Miklos et al. [44], one of the highest-accuracy existing surface-skeletonization methods.

C. Curve skeletons

Figure 9 (top 6 rows) compares our method with five curve-skeleton methods: Thinvox (TV) [48], distance-driven skeletonization (DDS) [3], robust thinning (RT) [39], iterative thinning process (ITP) [31], and multiscale skeletons (MS) [53]. In contrast to surface skeletons, we see now more variation between the compared methods. Our method delivers consistently thin (unlike MS), smooth (unlike ITP), noise-free (unlike ITP), and genus-preserving (unlike RT and MS) curve skeletons. Figure 11 shows extra insight, by comparing our
Fig. 9. Comparison of our 3D surface and curve skeletons with 10 related methods. Top 6 rows: curve skeletons. Bottom 6 rows: surface skeletons.
method with six additional mesh-based curve-skeletonization methods (Kustra et al. [34]; Livesu et al. [40]; Telea and Jalba [30]; Au et al. [6]; Dey and Sun [18]; and Jalba et al. [29]). As visible, our method yields well-centered curve skeletons which compare favorably, in terms of smoothness and lack of spurious branches, with the highest-quality mesh-based skeletons.

As for surface skeletons (Sec. V-B), let us detail the parallel with MS curve-skeletons. MS detects curve skeletons as those points having at least two equal-length geodesics on \( \partial \Omega \) between their feature points (MGF criterion in [18]). MS extends MGF by assigning a curve-skeleton importance \( \lambda_{CS} \) equal to the area bounded on \( \partial \Omega \) by the above two geodesics. Figures 9 and 11 show that our curve-skeletons and MS are very similar. The importances \( \lambda \) and \( \lambda_{CS} \) are also quite similar (see Fig. 21 vs Fig. 21 and Fig. 9, row 5 vs 6), except for the rockerarm, casting, and frog models, where \( \lambda_{CS} \) is smaller. Upon closer inspection, this shows a defect of MS: Low \( \lambda_{CS} \) points appear on curve-skeleton loops, whose geodesics do not cut \( \partial \Omega \) in two separate parts according to the Jordan theorem [70], so following [53] these points should get a high importance, to prevent loop disconnection when simplifying skeletons. Our method correctly finds such loops and assigns them a high importance.

VI. DISCUSSION

A. Method properties

Unification: Our method extracts 2D skeletons, 3D surface and 3D curve multiscale skeletons. To our knowledge, this is the first time that all these three skeleton types, including multiscale regularization, are computed by a single method which uses a single simplification metric. From both a theoretical and a practical perspective, we believe this to be an important result.

Robustness and accuracy: Our method computes thin, centered, homotopy-preserving, and connected skeletons from potentially noisy 2D and 3D shapes of arbitrary genus. Figure 12 quantitatively compares our skeletons with four other methods, using the technique in [61]. In detail, given two (curve or surface) skeletons \( S_1 \) and \( S_2 \), we first define the distance field

\[
D_{12}(x \in \Omega) = \begin{cases} 
\min_{y \in S_2} \|x-y\| = DT_{S_2}(x) & \text{if } x \in S_1, \\
0 & \text{if } x \notin S_1.
\end{cases}
\]

To compare two same-kind (curve or surface) skeletons, we draw the field \( D_{12} + D_{21} \) over the voxels \( S_1 \cup S_2 \), normalized by
its maximum value, using a rainbow colormap. Close skeleton fragments are blue, while outlier ones are red. Comparing a curve skeleton $CS_1$ with a surface skeleton $SS_2$ shows how well is $CS_1$ contained within $SS_2$. For this, we color voxels in $CS_1$ by $D_{12}$, and voxels in $SS_2 \setminus CS_1$ with gray. Hence, voxels in $CS_1 \cap SS_2$ are blue, and voxels in $CS_1$ far from $SS_2$ become red (Fig. 12 inset). Looking at Fig. 12, we see that our surface skeletons are very similar to those produced by IMA and HJ (Fig. 12 b,d). Warm colors, showing differences, occur mainly on the surface-skeleton boundary, and are due to the different simplification methods (and simplification levels) used by the compared methods. Our curve skeletons are fully contained in our surface skeletons, as expected (Fig. 12 e), but also nearly fully contained in the IMA and HJ skeletons (Fig. 12 g,i). Conversely, the DDS and HJ curve skeletons are well contained in our surface skeletons (Fig. 12 a,c). The largest differences, found between curve skeletons themselves (Fig. 12 f,h), are still quite small in absolute value.

**Scalability:** Table I (column 4) shows the speed of our method, implemented in C++ on a Linux 3.5 GHz PC with 32 GB RAM and an NVidia 690 GTX for the shapes in Fig. 9. Columns 2 and 3 give the shapes’ surface areas $|\partial \Omega|$ and volumes $|\Omega|$, in voxels. We note a high and relatively shape-independent throughput (foreground voxels/second), in line with the complexity stated in Sec. IV-E. Compared to the tested voxel-based methods (columns 5-13), our method is one order of magnitude faster on average. The next-fastest method is TV [48]. However, TV does not compute surface skeletons nor an importance metric. Compared to MS, the only other voxel-based method which computes a multiscale importance metric, we are on average 10 times faster. The last column in Tab.I shows the speed of the MBS mesh-based skeletonization in [29], the second other method we are aware of (apart from MS) which computes multiscale skeletons. Compared to our absolute timings (Tab.I, column 4, figures in brackets), MBS is 2.6 times faster on average. However, MBS is parallelized on the GPU, while our method is sequential and on the CPU.

**Classical properties:** We next summarize the behavior of our method vs several recognized desirable skeleton properties.

1. **Centeredness:** Centeredness is ensured by the unit-speed evolution of $\Gamma_\tau$ (Eqn. 5). We verify centeredness, both for curve and surface skeletons, against several methods known to formally respect this property [3], [27], [53], [69] (see Fig. 10).

2. **Thinness:** Our 2D and 3D skeletons are one-cell (pixel or voxel) thin, by construction. To argue this, suppose that this would not be so. Then, a skeleton $\lambda_\tau$ would (a) be thicker than one cell, and (b) would have, over a cross-section, the same importance $\lambda$. If such a thick cross-section existed, our shrinking algorithm would continue, since the shape can be further shrunk without disconnecting it (see Fig. 2).

3. **Homotopy:** Homotopy of the skeleton with the input shape is guaranteed by construction, by the constraint in Eqn. 4 and its corresponding implementation (Alg. 1).

4. **Reconstructibility:** The ability to reconstruct (smoothed...
versions of) the input shape from (simplified versions of) its skeleton is shown in Fig. 10. As visible, our reconstruction is quite accurate (compare e.g. Fig. 10 with Fig. 4 in [29]), modulo the natural limitation imposed by the fixed voxel grid.

5. Rotational invariance: The discretization of our proposed PDE system (Eqns. 3-5) described in Sec. IV-C is rotationally invariant by construction. We have verified that we indeed obtain nearly voxel-identical multiscale skeletons for the same input shape rotated at random angles with respect to the cell grid (not shown here for sake of brevity).

6. Curve vs surface skeletons: Our curve skeletons are by construction included in the surface skeletons of the same shape, since they are both obtained by thresholding the same single-and-global importance field $\lambda$ (see Sec. IV-D).

7. Multiscale and noise resistance: The field $\lambda$ describes the whole space between the input surface $\partial \Omega$, surface skeleton $S_{\Omega}$, curve skeleton $CS_{\Omega}$, and shape center $C_{\Omega}$. Thresholding $\lambda$ with increasing values yields the $S_0$ from $\Omega$; simplified surface skeletons (without branches due to small-scale shape details); the (simplified) $CS_\lambda$; and the shape center, or zero-dimensional skeleton, of $\Omega$. Reniers et al. get similar results, but they need two separate surface and curve skeleton importances and corresponding algorithms [53]. We compute $\lambda$ making no distinction between the two skeleton types. Telea and Jalba compute multiscale curve skeletons by collapsing surface skeletons inwards, following the idea that the former can be seen as the medial loci of the latter [30]. Yet, as in [53], their surface and curve skeleton algorithms are fundamentally different, and they also do not propose a curve-skeleton importance metric. Thresholding $\lambda$ at values $\tau$ between the average importance of $S_{\Omega}$ and $CS_{\Omega}$ yields a meso-skeleton structure [67] that continuously shrinks from $S_{\Omega}$ towards $CS_{\Omega}$ as $\tau$ increases (see Sec. IV-D).

Limitations: Our method stores four voxel scalar volumes ($\rho, \lambda, M,$ and $DT_{\partial \Omega}$ in Alg. 1), i.e. can handle shapes up to roughly 1000 $^3$ voxels on a 16 GB PC. Mesh-based skeletonization methods [29], [41] need far less memory. For instance, the mesh models for all shapes in this paper, which are up to 1M triangles, need only 24MB with the method in [29]. Separately, we acknowledge that our comparisons highlight differences between our skeletons and those produced by other methods, but do not explicitly show which skeletons are more suitable for a specific application, e.g. shape retrieval, classification, or segmentation. A thorough qualitative comparison with this goal is an important topic for future work.

B. Comparison with Hamiltonian methods

Equation 3 is similar with Torsello and Hancock’s (TH) mass conservation model $\nabla \cdot (\rho u) = 0$ used for 2D skeletonization [5]. Yet, several key differences exist. Numerically, TH transforms the mass conservation $\nabla \cdot (\rho u) = 0$ into a system of two ODEs (Eqns. 7 in [5]) by the substitution $\sigma = \log(\rho)$. They solve these ODEs with a second-order Crank-Nicholson discretization scheme coupled with semi-Lagrangian advection. In contrast, we use the conservative semi-Lagrangian scheme of Fedkiw [35], which only needs linear interpolation and clamping. This has several advantages. First, our scheme is numerically very stable, and conserves density well. Secondly, we do not need the second-order divergence discretization of TH. Thirdly, by computing the density logarithm $\sigma$, TH also needs exponentiation to find the final density $\rho = \exp(\sigma)$. We noticed, in practice, that this creates important numerical problems, such as an infinite value of $\rho$ at several points in $\Omega$.

Torsello and Rossi extend the TH density advection to extract 3D medial surfaces [54]. In their model, density advection stops when reaching the surface skeleton. In contrast, we continue advection by collapsing the surface skeleton to the curve skeleton and the latter to the shape center, yielding all desired skeletal representations within a single process.

VII. CONCLUSIONS

We have presented a unified framework for computing 2D skeletons and 3D surface and curve skeletons. We detect all skeleton types by a single algorithm, and also compute a single importance metric which assigns to each skeletal point the amount of (2D or 3D) input boundary described by that point. Comparing our skeletons and their computed importance with results computed by related methods shows a very good match. We present a simple implementation of our method which achieves good performance results on a range of complex 2D and 3D shapes, i.e., over 3 times faster than the fastest voxel-based skeletonization method we are aware of, and over 10 times faster than comparable multiscale methods.

Future work can target several directions. Porting our method to massively-parallel platforms (e.g., CUDA) using sparse voxel grids will increase scalability. Separately, modulating the input-surface density by e.g. curvature or application-specific metrics would allow different feature-sensitive skeleton simplifications.

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