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# Probabilistic View-based 3D Curve Skeleton Computation on the GPU

Jacek Kustra<sup>1,3</sup>, Andrei Jalba<sup>2</sup>, Alexandru Telea<sup>1</sup>

<sup>1</sup>*Institute Johann Bernoulli, University of Groningen, Nijenborgh 9, Groningen, the Netherlands*

<sup>2</sup>*Eindhoven University of Technology, Den Dolech 2, Eindhoven, the Netherlands*

<sup>3</sup>*Philips Research, Eindhoven, the Netherlands*

*jacek.kustra@philips.com, a.c.jalba@tue.nl, a.c.telea@rug.nl*

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Abstract: Computing curve skeletons of 3D shapes is a challenging task. Recently, a high-potential technique for this task was proposed, based on integrating medial information obtained from several 2D projections of a 3D shape (Livesu et al., 2012). However effective, this technique is strongly influenced in terms of complexity by the quality of a so-called skeleton probability volume, which encodes potential 3D curve-skeleton locations. In this paper, we extend the above method to deliver a highly accurate and discriminative curve-skeleton probability volume. For this, we analyze the error sources of the original technique, and propose improvements in terms of accuracy, culling false positives, and speed. We show that our technique can deliver point-cloud curve-skeletons which are close to the desired locations, even in the absence of complex postprocessing. We demonstrate our technique on several 3D models.

## 1 INTRODUCTION

Curve skeletons are well-known 3D shape descriptors with applications in computer vision, path planning, robotics, shape matching, and computer animation. A 3D object admits two types of skeletons: *Surface* skeletons are 2D manifolds which contain the loci of maximally-inscribed balls in a shape (Siddiqi and Pizer, 2009). Curve skeletons are 1D curves which are locally centered in the shape (Cornea et al., 2005).

Curve skeleton extraction has received increased attention in the last years (Dey and Sun, 2006; Reniers et al., 2008; Jalba et al., 2012; Au et al., 2008; Ma et al., 2012; Tagliasacchi et al., 2009; Cao et al., 2010a; Tagliasacchi et al., 2012). All these methods work in object space, *i.e.* take as input a 3D shape description coming as a voxel model, mesh, or dense point cloud. Recently, Livesu *et al.* have proposed a fundamentally different approach: They extract the curve skeleton from a set of 2D *views* of a 3D shape (Livesu et al., 2012). Key to this computation is the extraction of a volume which encodes, at each 3D point inside the shape, the probability that the curve-skeleton passes through that point. The curve-skeleton is extracted from this volume using a set of postprocessing techniques. This approach has the major advantage that it requires only a set of 2D views of the input shape, so it can be used when one does not have a complete 3D shape model. However, this method

strongly depends on the quality of the skeletal probability volume.

We present here an extension of the view-based approach of Livesu *et al.*, with the following contributions. First, we propose a different way for computing the curve-skeleton probability and representing it as a dense point cloud. On the one hand, this eliminates a major part of the original proposal’s false positives (*i.e.*, locations where a curve-skeleton point is suggested, but no such point actually exists), which makes our probability better suited for further skeleton extraction. On the other hand, our point-cloud model eliminates the need for a costly voxel representation. Secondly, we propose a fast GPU implementation of the point cloud computation which also delivers the skeleton probability with higher accuracy than the original method. We demonstrate our technique on several complex 3D models.

The structure of this paper is as follows. Section 2 overviews related work on 3D curve skeleton extraction. Section 3 details the three steps of our framework: extraction of accurate 2D view-based skeletons (Sec. 3.1), a conservative stereo matching for extracting 3D skeleton points from 2D view pairs, (Sec. 3.2), and a sharpening step that delivers a point cloud narrowly condensed along the curve skeleton (Sec. 3.3). Section 4 presents several results and discusses our framework. Section 5 concludes the paper.

## 2 RELATED WORK

### 2.1 Preliminaries

Given a three-dimensional binary shape  $\Omega \subset \mathbb{R}^3$  with boundary  $\partial\Omega$ , we first define its *distance transform*  $DT_{\partial\Omega} : \Omega \rightarrow \mathbf{R}_+$  as

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

The surface skeleton of  $\Omega$  is next defined as

$$S_{\partial\Omega} = \{ \mathbf{x} \in \Omega \mid \exists \mathbf{f}_1, \mathbf{f}_2 \in \partial\Omega, \mathbf{f}_1 \neq \mathbf{f}_2, \|\mathbf{x} - \mathbf{f}_1\| = \|\mathbf{x} - \mathbf{f}_2\| = DT_{\partial\Omega}(\mathbf{x}) \}, \quad (2)$$

where  $\mathbf{f}_1$  and  $\mathbf{f}_2$  are two contact points with  $\partial\Omega$  of the maximally inscribed disc in  $\Omega$  centered at  $\mathbf{x}$ , also called *feature transform* (FT) points (Strzodka and Telea, 2004) or *spoke vectors* (Stolpner et al., 2009). Here, the feature transform is defined as

$$FT_{\partial\Omega}(\mathbf{x} \in \Omega) = \arg \min_{\mathbf{y} \in \partial\Omega} \|\mathbf{x} - \mathbf{y}\|. \quad (3)$$

Note that  $FT_{\partial\Omega}$  is multi-valued, as an inscribed ball can have two, or more, contact points  $\mathbf{f}$ . Note, also, that the above definitions for the distance transform, feature transform, and skeleton are also valid in the case of a 2D shape  $\Omega \in \mathbb{R}^2$ .

### 2.2 Object-space curve skeletonization

In contrast to the formal definition of surface skeletons (Eqn. 2), *curve* skeletons know several definitions in the literature. Earlier methods computed the curve skeleton by thinning, or eroding, the input voxel shape in the order of its distance transform, until a connected voxel curve is left (Bai et al., 2007). Thinning can also be used to compute so-called meso-skeletons, *i.e.* a mix of surface skeletons and curve skeletons (Liu et al., 2010). For mesh-based models, a related technique collapses the input mesh along its surface normals under various constraints required to maintain its quality (Au et al., 2008). Hassouna *et al.* present a variational technique which extracts the skeleton by tracking salient nodes on the input shape in a volumetric cost field that encodes centrality (Hassouna and Farag, 2009). Tagliasacchi *et al.* compute curve skeletons as centers of point cloud projections on a cut plane found by optimizing for circularity (Tagliasacchi et al., 2009). A good review of curve skeletonization is given in (Cornea et al., 2007).

One of the first formal definitions of curve skeletons is the locus of points  $\mathbf{x} \in \Omega$  which admit at least two shortest paths, or geodesics, between their feature points (Dey and Sun, 2006; Prohaska and Hege,

2002). This definition has been used for mesh models (Dey and Sun, 2006) and voxel-based models (Reniers et al., 2008). Curve skeletons can also be extracted by collapsing a previously computed surface skeleton towards its center using different variants of mean curvature flow (Tagliasacchi et al., 2012; Cao et al., 2010a; Telea and Jalba, 2012). Alternatively, surface skeletons can be computed using a ball shrinking method (Ma et al., 2012) and then selecting points which match the geodesic criterion (Jalba et al., 2012). However, such approaches require one to first compute the more expensive surface skeleton.

### 2.3 View-based curve skeletonization

A quite different approach was recently proposed by Livesu *et al.*: Noting that the 2D projection of a 3D curve skeleton is close to the 2D skeleton of the projection, or view, of an input 3D shape, they extract curve skeletons by merging 2D skeletal information obtained from several views of the input shape  $\Omega$ . Given two such views  $C_i$  and  $C_i^\perp$ , whose up-vectors are parallel and lines of sight are orthogonal, the silhouettes  $B_i$  and  $B_i^\perp$  of  $\Omega$  are first computed by orthographic projection of the input shape. Secondly, the 2D skeletons  $S_{\partial B_i}$  and  $S_{\partial B_i^\perp}$  of these silhouettes are computed. Next, stereo vision is used to reconstruct the 3D skeleton: Point pairs  $p \in S_{B_i}$  and  $p^\perp \in S_{B_i^\perp}$  are found by scanning each epipolar line, and then backprojected into 3D to yield a potential curve-skeleton point  $\mathbf{x}^1$ . The points  $\mathbf{x}$  found in this way are accumulated into a so-called *probability volume*  $\mathcal{V} \subset \mathbb{R}^3$ , which gives, at each spatial point, the likelihood to have a curve-skeleton passing through that point.

The above method has several advantages compared to earlier techniques. First, it can be used directly on shape *views*, rather than 3D shape models, which makes it suitable for any model which can be rendered in a 2D view, regardless of its representation (*e.g.* polygons, splats, points, lines, or textures). Secondly, the method can be easily parallelized, as view pairs are treated independently. However, this method fundamentally relies on the fast computation of a *good* probability volume which contains a correct estimation of the curve skeleton location. This poses the following requirements:

1. a **reliable** and **accurate** stereo vision correspondence matching, *i.e.* finding the correct pairs of points ( $p \in S_{\partial B_i}, p^\perp \in S_{\partial B_i^\perp}$ ) which represent the projection of the same curve skeleton point in the view-pair  $(C_i, C_i^\perp)$ ;
2. an **accurate** and **efficient** representation of the probability volume  $\mathcal{V}$  for further processing.

<sup>1</sup>Here and next, we denote by italics (*e.g.*,  $p$ ) the 2D projection of a 3D point  $\mathbf{p}$  in a camera  $C$

Requirement (1) is not considered by Livesu *et al.*, where *all* possible point-pairs along an epipolar line are backprojected. This generates, as we shall see in Sec. 3.2.2, a large amount of noise in the probability volume  $\mathcal{V}$ . Removing this noise requires four relatively complex postprocessing steps in the original proposal. Secondly, the probability volume  $\mathcal{V}$  is represented as a voxel grid. This makes the method unnecessarily inaccurate, relatively slow and hard to parallelize, and requires large amounts of memory, thus contradicts requirement (2).

In the following, we present several enhancements that make view-based skeleton extraction compatible with requirements (1) and (2). This allows us to extract a high-accuracy probability volume for further usage in curve skeleton computation or direct visualization.

### 3 ACCURATE PROBABILITY VOLUME COMPUTATION

Our proposal has three steps (see also Fig. 1). First, we extract regularized and subpixel-accuracy 2D skeletons from several views of the input shape (Sec. 3.1). Next, we use additional view-based information to infer a conservative set of correspondences between points in such 2D skeleton pairs, backproject these in 3D, and record the obtained points as a point cloud (Sec. 3.2). Finally, we apply an additional sharpening step on the 3D point cloud, which directly delivers a highly accurate curve-skeleton probability (Sec. 3.3).

#### 3.1 Robust 2D Skeletonization

Given a shape  $\Omega$  and camera specification  $C = (\mathbf{o}, \mathbf{v}, \mathbf{u})$  described by its origin  $\mathbf{o}$ , view direction  $\mathbf{v}$ , and up-vector  $\mathbf{u}$ , we start by computing the silhouette  $B$  of  $\Omega$  by rendering the shape on the camera’s view plane  $(\mathbf{u}, \mathbf{v} \times \mathbf{u})$ . Next, we compute the so-called salience 2D skeleton of  $B$  using the technique presented in (Telea, 2012a). The *salience* of a point  $\mathbf{p} \in B$  is defined as

$$\sigma(\mathbf{p}) = \frac{\rho(\mathbf{p})}{DT_{\partial B}(\mathbf{p})} \quad (4)$$

Here,  $DT_{\partial B}(\mathbf{p})$  is the 2D distance transform of the silhouette boundary  $\partial B$  and  $\rho(\mathbf{p})$  is the so-called skeleton *importance*

$$\rho(\mathbf{p}) = \max_{\mathbf{f}_1, \mathbf{f}_2 \in FT_{\partial B}(\mathbf{p})} \|\gamma_{\mathbf{f}_1, \mathbf{f}_2}\| \quad (5)$$

where  $FT_{\partial B}$  is the feature transform of the boundary  $\partial B$ , and  $\gamma_{\mathbf{ab}}$  is the compact boundary fragment between two points  $\mathbf{a}$  and  $\mathbf{b}$  on  $\partial B$ . The importance  $\rho$  increases

monotonically from the endpoints (tips) of the skeleton towards its center. Intuitively,  $\rho(\mathbf{p})$  associates, to each skeleton point  $\mathbf{p}$ , the length of the longest boundary arc (in pixels) subtended by its feature points. Upper thresholding  $\rho$  with a value  $\rho_0$  will thus remove both skeleton branches created by small boundary wiggles *and* end-parts of important skeleton branches caused *e.g.* by boundary corners. Figure 2 (top row) shows this effect for a silhouette  $B$  of a horse model. For  $\rho_0 = 1$ , we get the full 2D skeleton, which contains many spurious branches. For  $\rho_0 = 5$ , we get the desired skeleton detail at the legs and head, but still have several spurious branches around the rump and neck. For  $\rho_0 = 30$ , we eliminate all spurious branches, but also loose relevant portions of branches corresponding to the legs. This is undesired, since, as we shall later see, we need the important branches at their full length to reconstruct a curve-skeleton reaching into all shape protrusions.

In contrast, the salience metric  $\sigma$  (Eqn. 4) delivers a better result. As shown in (Telea, 2012a),  $\sigma$  is high along the most important, or salient, skeleton branches, and low elsewhere. Hence, we can threshold  $\sigma$  to obtain the skeleton

$$S_{\partial B} = \{\mathbf{p} \in B \mid \sigma(\mathbf{p}) > \sigma_0\} \quad (6)$$

Equation 6 delivers a clean, regularized, skeleton whose spurious branches are eliminated, and whose important branches extend all the way into the shape’s protrusions. Figure 2 (bottom row) shows the saliency-based regularization. For  $\sigma_0 = 0$ , we obtain the same full skeleton as for  $\rho_0 = 1$ . Increasing  $\sigma_0$  over a value of 0.05 practically removes all spurious branches, but keeps the important ones un-pruned (see zoom-ins). As such, we use the value  $\sigma_0 = 0.05$  further in our pipeline.

We further enhance the precision of the computed skeleton by using the subpixel technique presented in (Strzodka and Telea, 2004). As such, skeleton points are stored as 2D floating-point coordinates rather than integers. This will be important when performing the 3D stereo reconstruction (Sec. 3.2.2).

#### 3.2 Accurate Correspondence Matching

We find potential 3D curve-skeleton points along the same key idea of (Livesu *et al.*, 2012): Given a camera  $C = (\mathbf{o}, \mathbf{v}, \mathbf{u})$ , where  $\mathbf{v}$  points towards the object’s origin, we construct a pair-camera  $C^\perp = (\mathbf{o}^\perp, \mathbf{v}^\perp, \mathbf{u}^\perp)$  which also points at the origin and so that the two up-vectors  $\mathbf{u}$  and  $\mathbf{u}^\perp$  are parallel. In this case, projected points  $p$  in  $C$  correspond to projected points  $p^\perp$  in  $C^\perp$  located on the same horizontal scanline. Given such a point-pair  $(p, p^\perp)$ , the generated 3D point  $\mathbf{x}$  is computed by triangulation, *i.e.* by solving

$$\mathbf{x} = \mathbf{p} + k\mathbf{v} = \mathbf{p}^\perp + k^\perp\mathbf{v}^\perp \quad (7)$$

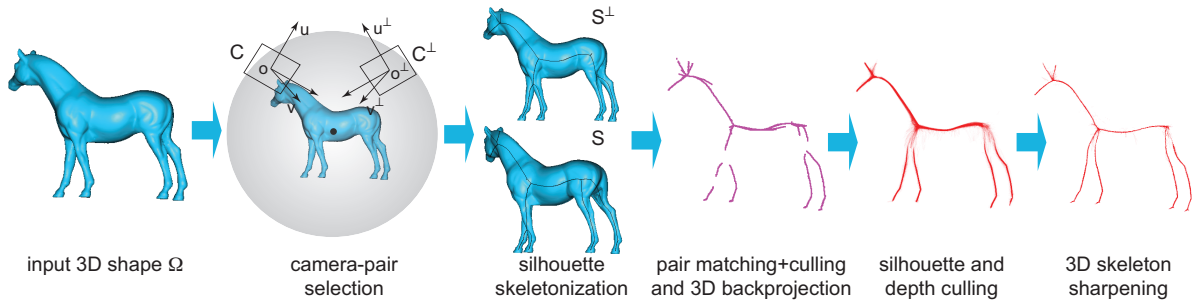


Figure 1: Curve-skeleton probability computational pipeline

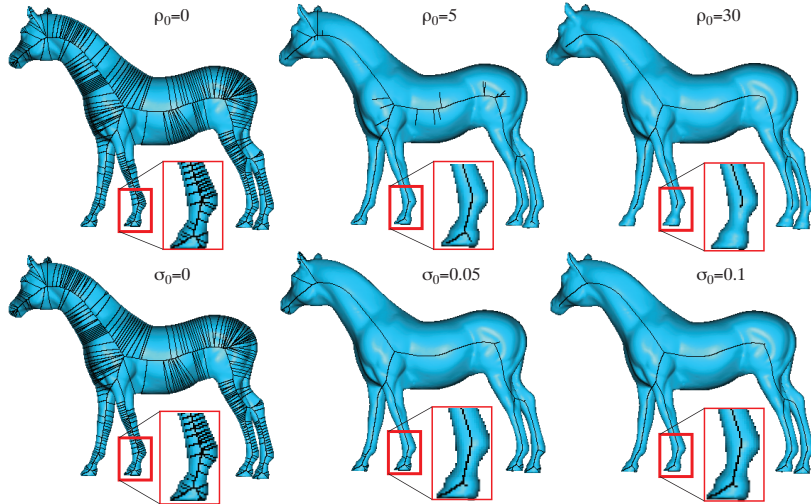


Figure 2: Skeleton regularization. Top row: Importance-based method (Telea and van Wijk, 2002) for three different threshold values  $\rho_0$ . Bottom row: Saliency-based method (Telea, 2012a) for three different threshold values  $\sigma_0$ .

where  $\mathbf{p}$  and  $\mathbf{p}^\perp$  are the 3D locations, in their respective view planes, corresponding to  $p$  and  $p^\perp$  respectively, and  $k$  and  $k^\perp$  are the distances between  $\mathbf{x}$  and the view planes of  $C$  and  $C^\perp$ . Note that  $\mathbf{p}$  and  $\mathbf{p}^\perp$  can be immediately computed as we know the positions of  $p$  and  $p^\perp$  and the cameras' positions, orientations, and near plane locations.

### 3.2.1 Correspondence Problem

However, as well known in stereo vision, the success of applying Eqn. 7 is fundamentally conditioned by having the *correct* 2D points  $p$  and  $p^\perp$  paired in the two cameras. Let us analyze this issue in our context: Consider that a scanline  $y$  intersects a 2D skeleton shape in  $m$  points on the average. Hence, we have  $m^2$  possible point-pairs. These will generate  $m^2$  points in the 3D reconstruction, whereas in reality there are only at most  $m$  such points – that is, if no occlusion is present. The excess of  $m^2 - m$  points are false positives. Given  $N$  such camera-pairs placed uniformly around the object in order to reconstruct its 3D curve skeleton, and consid-

ering a camera viewplane of  $P \times P$  pixels, we have in the worst case  $O(N(m^2 - m)P)$  false-positive points in the curve skeleton. The ratio of false-to-true positives is thus  $\Pi = O(N(m^2 - m)P)/(NmP) = O(m)$ . In our measurements for a wide set of shapes, we noticed that  $m = 5$  on the average. Concretely, at an image resolution of  $P^2 = 1024^2$  pixels, and using the setting  $N = 21$  from Livesu *et al.*, we thus get over 400K false-positive points generated in excess of the  $NmP \simeq 100K$  true-positive skeleton points.

The above false-to-true-positive ratio  $\Pi$  is a conservative estimate: Given a rigid shape  $\Omega$ , the 2D skeleton of its silhouette can change considerably as the silhouette changes, even when no self-occlusions occur. This, and additional self-occlusion effects, reduce the true-positive count and thus increases  $\Pi$ . This ultimately creates substantial noise in the curve-skeleton probability estimation, and thus makes an accurate curve skeleton extraction more complex.

### 3.2.2 Pair-culling heuristic

We reduce the false-to-true-positive ratio  $\Pi$  by using additional information present in our cameras, as follows. Consider a point  $p$  on a scanline  $L$  in  $C$  and all points  $L^\perp = \{p_i^\perp\}$  on the same scanline in  $C^\perp$  (see Fig. 3 e). The 3D reconstructions of all pairs  $(p, p_i^\perp)$  lie along the line  $\mathbf{p} + k\mathbf{v}$  (Eqn. 7). Hence, if we had an estimate of the depth  $k_{est}$  between the correct reconstruction and the viewplane of  $C$ , we could select the best pair  $p_{est}^\perp$  for  $p$  as

$$p_{est}^\perp = \arg \min_{p_i^\perp \in L^\perp} |k_{est} - k| \quad (8)$$

*i.e.* the point in  $C^\perp$  which yields, together with  $p$ , a depth closest to our estimate. We estimate  $k_{est}$  as follows: When we draw the shape in  $C$ , we also compute its nearest and furthest depth buffers  $z_n$  and  $z_f$ , by rendering the shape twice using the OpenGL *GL\_LESS* and *GL\_GREATER* depth-comparison functions respectively. Next, for each point  $p$  in the viewplane of  $C$ , we set  $k_{est} = \frac{1}{2} [z_n(p) + z_f(p)]$  (see Fig. 3 e).

It is essential to note that our heuristic for  $k_{est}$  is *not* an attempt to find the exact value of the depth  $k$ . Indeed, if we could do this, we would not need to apply Eqn. 8, as we could perform the 3D backprojection using a single view. We use  $k_{est}$  only as a way to select the most likely point-pair for 3D reconstruction. This is argued as follows: First, we note that the value  $k$  for the correct point-pair must reside between  $z_n(p)$  and  $z_f(p)$  - indeed, the reconstructed 3D point  $\mathbf{x}$  must be inside the object's hull. Secondly, the curve skeleton is roughly situated in the (local) middle of the object, thus its depth is close to  $k_{est}$ . Thirdly, we note that, when the angle between the cameras' vectors  $\alpha = \angle(\mathbf{v}, \mathbf{v}^\perp)$  decreases, then the depths  $k_i$  yielded by Eqn. 8 for a set of scanline-points  $p_i^\perp \in L^\perp$  get further apart. In detail, if the distance between two neighbor pixels in the scanline  $L^\perp$  is  $\delta$ , the distance between their reconstructions using the same point  $p$  in the other scanline  $L$  is  $\epsilon = \delta / \sin(\alpha)$ , see Fig. 3 c. Hence, if we use a small  $\alpha$  (under  $90^{\text{deg}}$ ), we get fewer depths  $k_i$  close to  $k_{est}$ , so we decrease the probability that selecting the point whose depth is closest to  $k_{est}$  (Eqn. 8) will yield an incorrect point-pair for the 3D reconstruction. In contrast, Livesu *et al.* use  $\alpha = 90^\circ$ , as this slightly simplifies Eqn. 7. Given that low  $\alpha$  values reduce the likelihood to obtain false pairs using our depth heuristic, we prefer this, and set  $\alpha = 20^\circ$ .

Figure 3 shows the results of using our depth-based pairing heuristic. Images (a) and (b) show the two skeletons  $S_{\partial B}$  and  $S_{\partial B^\perp}$  corresponding to the two cameras  $C$  and  $C^\perp$  respectively. The brute-force many-to-many correspondence pairing yields 6046 three-dimensional points. As visible in Fig. 3 c, these points are spread uniformly in depth along the view directions of the two

cameras. This is expected, since 2D skeleton pixels are equally spaced in the image plane. For clarity, we displayed here only those points which pass the silhouette and depth-culling, *i.e.* which are inside the object from *any* considered view (see further Sec. 3.3). The displayed points in Fig. 3 c are thus *final* points in the curve-skeleton probability volume delivered by many-to-many matching.

Figure 3 d shows the reconstructed 3D points when we use our depth-based pair-culling. Since we now only have one-to-one pairs, we obtain much less points (721 vs 6046, see the explanations in Sec. 3.2.1). Moreover, these points are located very close to the actual curve skeleton, as shown by the top view of the model.

Given our conservative point-pair selection, as shown in Fig. 4, we generate much fewer curve-skeleton points than if using many-to-many pairing. Although this is highly desirable for obtaining an accurate (false-positive-free) curve skeleton, it also means that the curve skeleton will be sparser than when using all possible pairs. To counteract this, we simply use more view pairs  $N$ . In practice, setting  $N \simeq 500$  yields sufficiently dense curve skeletons (see results in Sec. 4). An additional advantage of using more views is that we do not need to carefully select the optimal views for stereo reconstruction, in contrast to the original method, where such views are obtained by performing a principal component analysis (PCA) on both the 3D shape and its 2D projections.

We further reduce the number of tested point-pairs (Eqn. 8) by scanning  $L$  from left to right (for  $p$ ) and  $L^\perp$  from right to left (for  $p^\perp$ ). As such, 3D points are generated in increasing order of their depth  $k$ , so  $|k_{est} - k|$  first decreases, then increases. Hence, we stop the scan as soon as  $|k_{est} - k|$  increases, which gives an additional speed improvement.

### 3.3 Probability Sharpening

We collect the 3D points  $\mathbf{x}$  (Eqn. 7) found by the depth-based correspondence matching for a given camera-pair  $(C, C^\perp)$  in an unstructured point cloud  $\mathcal{CS}$ . As  $C$  rotates around the input shape, we keep testing that the projections  $x$  of the accumulated points  $\mathbf{x} \in \mathcal{CS}$  fall inside the silhouette  $B$  in  $C$ , as well as within  $C$ 's depth range  $[z_n(x), z_f(x)]$ . Points which do not pass these tests are eliminated from  $\mathcal{CS}$ . The depth test explained above comes atop the silhouette test which was already proposed by Livesu *et al.* Whereas the silhouette test constrains  $\mathcal{CS}$  to fall within the visual hull of our input shape, we constrain  $\mathcal{CS}$  even further, namely to fall within the exact shape. The difference is relevant for objects with cavities, whose visual hull is larger than the object itself.

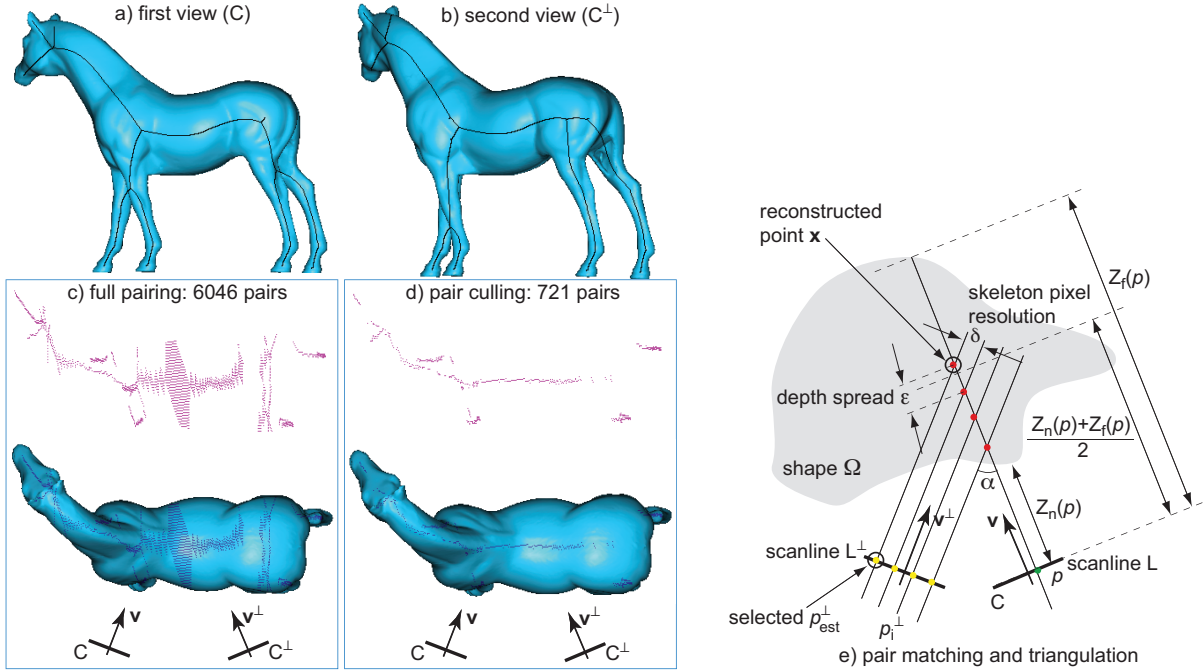


Figure 3: Correspondence matching for curve-skeleton reconstruction. A camera pair (a,b). Reconstructed 3D points when using full pairing (c) and when using our depth-based pairing (d). Depth-based pairing and triangulation (e)

However closer to the true curve-skeleton than the results presented by Livesu *et al.*, our  $CS$  still shows some spread around the location of the true curve skeleton. This is due to two factors. First, consider the inherent variability of 2D skeletons in views of a 3D object: The 2D skeleton is locally centered with respect to the silhouette (or projection) of a 3D object  $\Omega$ . In areas where  $\Omega$  has circular symmetry, the 2D skeleton of the projected 3D object is indeed identical to the 2D projection of the true 3D curve skeleton, *i.e.*, skeletonization and projection are commutative. However, this is not true in general for shapes with other cross-sections. Moreover, self-occlusions, in the case of concave objects, will generate 2D skeletons which have little in common with the projection of the curve skeleton. It is important to stress that this is *not* a problem caused by wrong correspondence matching. Secondly, using a small  $\alpha$  angle between the camera pairs (Sec. 3.2.2), coupled with the inherent resolution limitations of the image-based skeletons, introduces some depth estimation errors which show up as spatial noise in the curve skeleton.

We further improve the sharpness of  $CS$  as follows. For each camera  $C$  which generates a silhouette  $B$ , we move the points  $\mathbf{x} \in CS$  parallel to the view plane of  $C$  with a step equal to  $\nabla DT_{\partial B}$ . Since  $\nabla DT_{\partial B}$  points towards  $S_{\partial B}$ , this moves the curve-skeleton points towards the 2D skeleton  $S_{\partial B}$ . Note that, in general,  $\nabla DT_{\partial B}$  is not zero along  $S_{\partial B}$  (Telea and van Wijk, 2002). Hence, to

prevent points to drift along  $S_{\partial B}$ , and thus create gaps in the curve skeleton, we disallow moving points  $\mathbf{x}$  which already project on  $S_{\partial B}$ . Note also that this advection never moves points outside  $\Omega$ , since  $S_{\partial B}$  is always inside any silhouette  $B$  of  $\Omega$ . Since the above process is done for all the viewpoints  $C$ , the curve skeleton gets influenced by *all* the considered views. This is an important difference with respect to Livesu *et al.*, where a 3D skeleton point is determined only by *two* views.

Figure 4 shows the effect of our three improvement steps: depth-based pairing (Sec. 3.2.2), depth culling, and sharpening. All images show 3D point clouds rendered with alpha blending. Figure 4 a shows the cloud  $CS$  computed following the original method of Livesu *et al.*, that is, with many-to-many correspondence matching along scanlines (Sec. 3.2.1). Clearly, this cloud contains a huge amount of points not even close to the actual curve skeleton. If we decrease the alpha value in the visualization, we see that this cloud, indeed, has a higher density along the curve-skeleton (Fig. 4 b). We see here also that naive thresholding of the density, which is quite similar to decreasing the alpha value to obtain Fig. 4 b from Fig. 4 a, creates problems: If a too low threshold is used, the skeleton still stays thick; if a too high threshold is used, the skeleton risks disconnections (see white gaps in the neck region in Fig. 4 b).

Eliminating the large amount of false positives from the cloud shown in Fig. 4 a is very challenging. To do

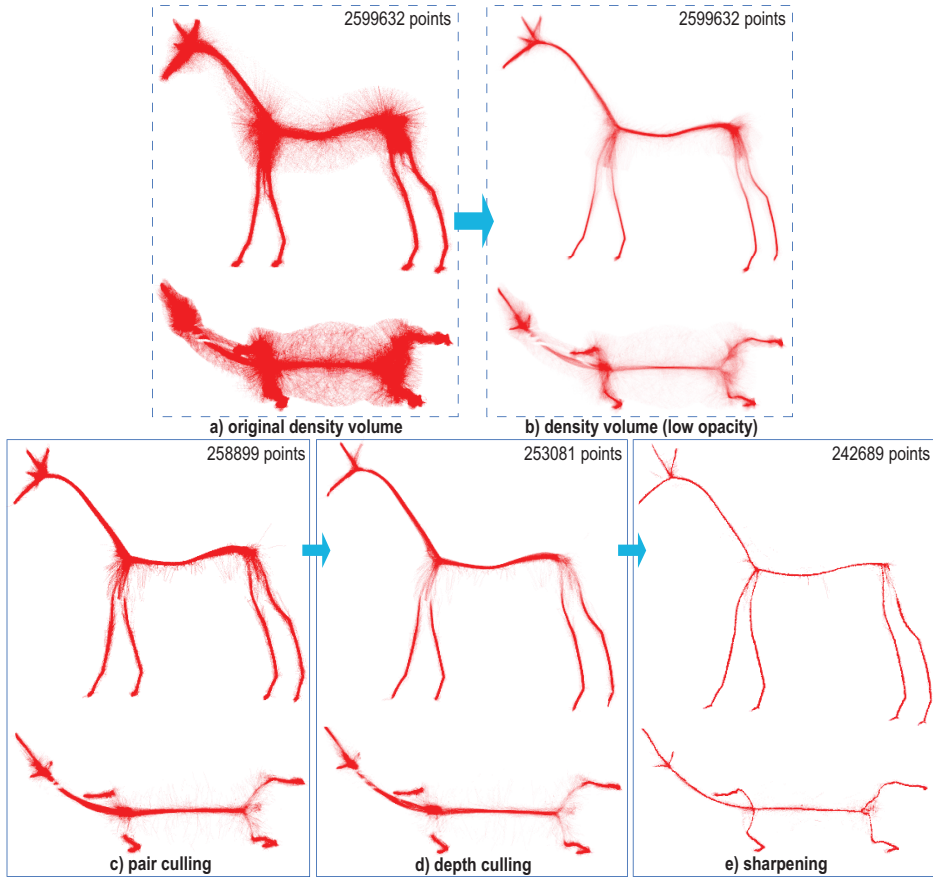


Figure 4: Curve-skeleton probability point-cloud. (a) original method (Livesu et al., 2012). (b) Cloud in (a) displayed with lower opacity. (c) Effect of depth-based pairing. (d) Effect of depth culling. (e) Effect of sharpening (see Sec. 3.3).

this, Livesu *et al.* apply an involved post-processing pipeline: (1) voxelize the cloud into a voting grid; (2) extract a maximized spanning tree (MST) from the grid; (3) detect and prune perceptually salient tree branches; (4) collapse short branches; (5) recover curve-skeleton loops lost by the MST; and (6) smooth the resulting skeleton; for details we refer to (Livesu et al., 2012). Although this is possible, as demonstrated by the results of Livesu *et al.*, this post-processing is highly complex, delicate, and time-consuming.

Fig. 4 c shows our curve-skeleton probability, obtained with the center-based correspondence pair culling (Sec. 3.2.2). The point cloud contains now around ten times less points. Also, note that points close to the true curve skeleton have been well detected, *i.e.*, we also have few false negatives. Applying the depth-based culling further removes a small amount of false positives (Fig. 4 d vs Fig. 4 c). Finally, the density sharpening step effectively attracts the curve-skeleton points towards the local skeleton in each view, so the overall result is a sharpening of the point cloud  $\mathcal{CS}$ , *i.e.* a point density increase along the true curve skeleton and a density de-

crease further from the skeleton (Fig. 4 e).

## 4 DISCUSSION

**Performance:** We have implemented our method in C++ with OpenGL and CUDA and tested it on a 2.8 GHz MacBook Pro with an Nvidia GT 330M graphics card. The main effort is spent in computing the regularized 2D salience skeletons (Sec. 3.1). We efficiently implemented the computation of  $DT_{\partial B}$ ,  $FT_{\partial B}$ ,  $\rho$ , and  $\sigma$  (Eqns. 1-4) using the method in (Cao et al., 2010b), one of the fastest exact Euclidean distance-and-feature transform techniques in existence (see our publicly available code at (Telea, 2012b)). The remaining steps of our pipeline are trivial to parallelize, as points and camera views are treated independently. Overall, our entire pipeline runs roughly at 500 frames/second. Given that we use more views than Livesu *et al.*, *i.e.* roughly 500 vs 21, our CUDA-based parallelization is essential, as it allows us to achieve roughly the same timings as the method of Livesu *et al.*



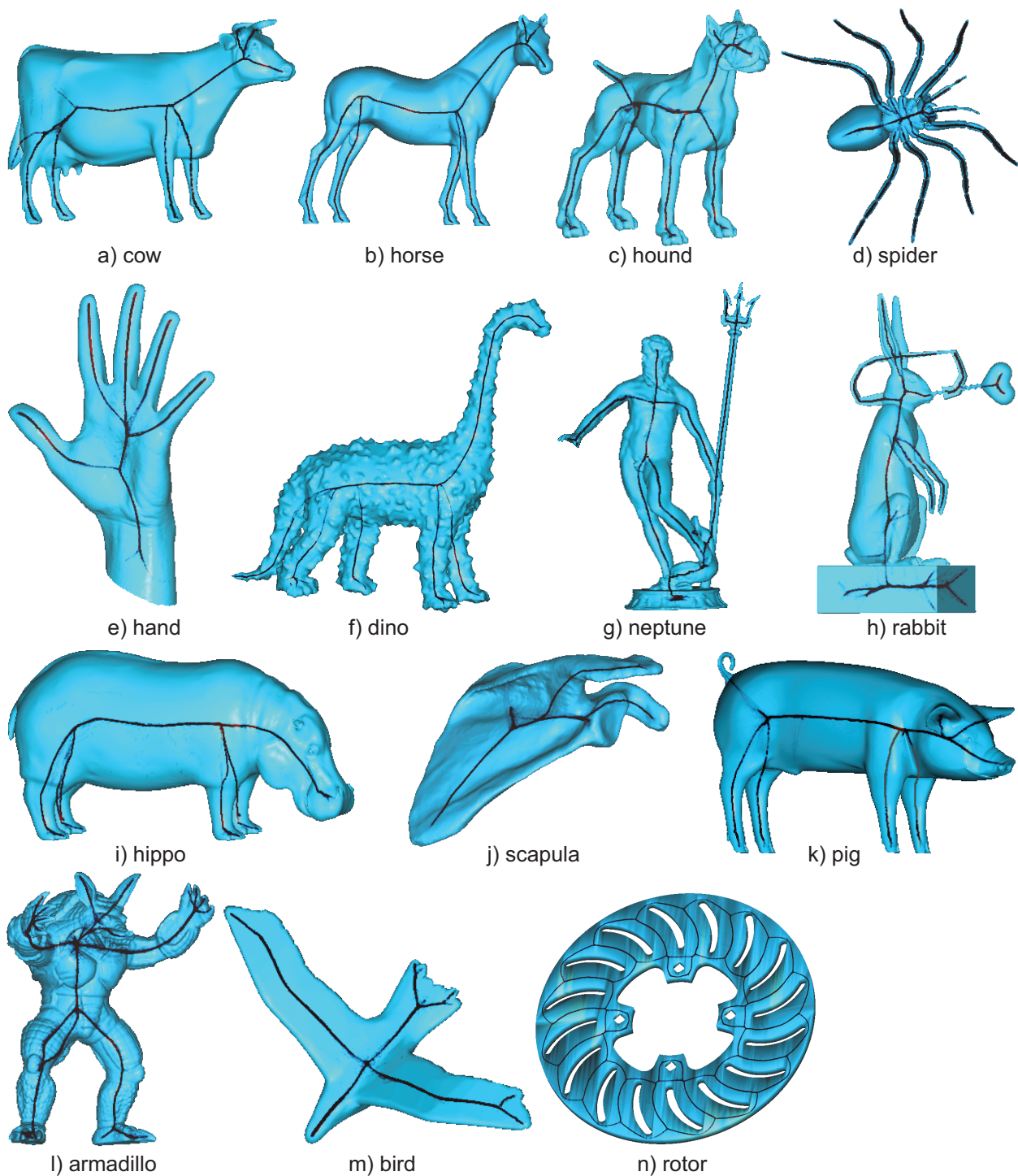


Figure 5: Curve-skeleton probability point-clouds for several models (see Sec. 4).

**Parameters:** All parameters of the method are fixed and independent on the input shape, *i.e.* skeleton saliency threshold  $\sigma_0 = 0.05$  (Sec. 3.1), number of considered views uniformly distributed around a sphere centered in

the object center  $N = 500$  (Sec. 3.2.2), screen resolution  $P^2 = 1024^2$  pixels (Sec. 3.2.1), and angle between the camera-pair view vectors  $\alpha = 20^\circ$  (Sec. 3.2.2). Less views ( $N < 500$ ) will generate sparser-sampled curve

skeletons, as discussed in Sec. 3.2.2. Decreasing the pixel resolution generates slightly thicker point distributions in the curve-skeleton cloud. This is expected, since we have less *and* coarser-spaced 2D skeleton pixels, which also implies higher depth estimation errors (Eqn. 7). Decreasing  $\alpha$  under roughly 5 degrees generates too large inaccuracies in the depth estimation; increasing it over roughly 30 degrees reduces the likelihood of good correspondence pairing; hence, our setting of  $\alpha = 20^\circ$ .

**Results:** Figure 5 shows several results computed with our method. The produced curve-skeleton clouds contain between 100K and 300K points. We render these clouds using small point splats of 2 by 2 pixels, to make them more visible. The key observation is that our skeleton point clouds are *already* very close to the desired 3D location, even in the absence of any cloud postprocessing. In contrast, the equivalent point clouds delivered by the method of Livesu *et al.* are much noisier (see example in Fig. 3 a and related discussion in Sec. 3.2.2), and thus require significant postprocessing to select the true-positives. Since our point clouds are much sharper, we can directly use them for curve-skeleton visualization, as shown in Fig. 5. If an explicit line representation of such skeletons is desired, this can be easily obtained by using *e.g.* the curve-skeleton reconstruction algorithm described in (Jalba *et al.*, 2012), Sec. VIII-C. Thin tubular skeleton representations can be obtained by isosurfacing the density field induced by our 3D point cloud. In this paper, we refrained from producing such reconstructions, as we want to let our main contribution stand apart – the computation of noise-free, accurate point-cloud representations of the curve skeleton probability.

**Comparison:** Figure 6 compares our method with several recent curve-skeleton extraction methods. As visible, our curve skeleton has the same overall structure and positioning within the object. However, differences exist. First, our method produces smoother curve skeletons than (Dey and Sun, 2006) and (Jalba *et al.*, 2012). This is due to the density sharpening step, which does not have an equivalent in the latter two methods. Also, (Au *et al.*, 2008) requires a so-called connectivity surgery step to repair the curve skeleton after the main Laplacian advection has completed. This necessary step has the undesired by-product of creating straight-line internal skeleton branches (Fig. 6 d, palm center). Secondly, we correctly find the skeleton’s ligature and internal branches. This is also the case for all other methods except (Livesu *et al.*, 2012), where all skeleton branches are merged in a single junction point (Fig. 6 b). This fact is not surprising, given the branch collapsing postprocessing step in the latter method. It is not clear to us why this step is required (or beneficial), as it actually changes the topology of the skeleton, and thus may impair operations such as

shape analysis or matching.

**Properties:** Our method maintains all of the desirable properties of curve skeletons advocated by related work (Cornea *et al.*, 2007; Au *et al.*, 2008; Tagliasacchi *et al.*, 2012; Livesu *et al.*, 2012; Jalba *et al.*, 2012): Our skeletons are **thin** and locally **centered** within the object. Higher-**genus** objects (with tunnels) are handled well (see rabbit and rotor models, Fig. 5). The method is **robust** against noise, due to the sharpening step (see dino and armadillo models, Fig. 5). Thin, sharp **detail** protrusions of the models generate curve skeleton branches, as long as these parts project to at least 1 pixel in screen space (see neptune, spider, and rabbit models, Fig. 5). This is due to the usage of the 2D skeleton saliency metric, which keeps 2D skeleton branches reaching into such salient shape details (Sec. 3.1). Input model **resolution**, *e.g.* polygon count, is largely irrelevant to the end result, since 2D skeletons are computed in image space.

**Limitations:** Our method cannot recover complete curve skeletons for shape parts which are not visible from any viewpoint, *i.e.*, permanently self-occluded. This is an inherent problem of view-based 3D reconstruction. For such shapes, the object-space skeletonization methods mentioned in Sec. 2 should be used.

## 5 CONCLUSIONS

We have presented a new method for computing curve-skeletons as unstructured point clouds. Our method extends the view-based curve-skeleton extraction of Livesu *et al.* in several directions: (1) Using salience-based skeletons to guarantee preservation of terminal skeleton branches, (2) using depth information to reduce the number of false-positives in the 3D skeleton reconstruction, and (3) sharpening the obtained point-cloud representation to better approximate the 1D singularity locus of the curve skeleton. We trade off speed for accuracy, by generating more conservative skeleton samples and using more viewpoints. However, by using a GPU implementation, we achieve the same speed as the original method, but deliver a much cleaner and sharper 3D skeleton point-cloud approximation. Overall, our method can be used either as a front-end for reconstructing line-based representations of 3D curve skeletons, or for directly rendering such skeletons as unstructured point clouds.

Future work can improve the point matching accuracy, for example by using optical flow models or exploiting geometric variability properties of 2D skeletons. Separately, implementing the 3D geodesic-based curve-skeleton detector of (Dey and Sun, 2006) by using the 2D collapsed boundary metric  $\rho$  (Eqn. 5) is a promis-

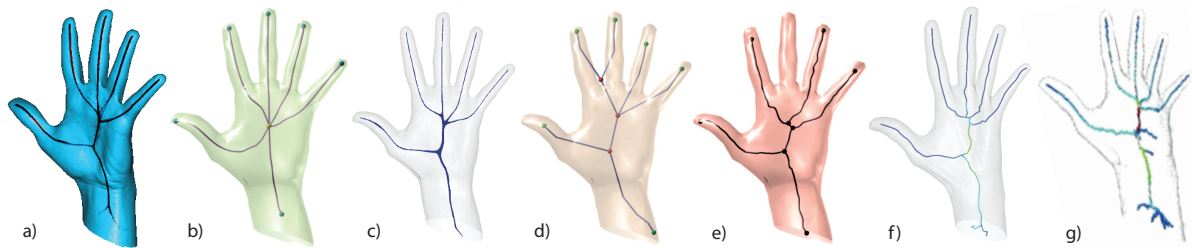


Figure 6: Comparison with related methods: (a) our method; (b) (Livesu et al., 2012); (c) (Telea and Jalba, 2012); (d) (Au et al., 2008); (e) (Dey and Sun, 2006); (f) (Jalba et al., 2012); (g) (Reniers et al., 2008) (see Sec. 4)

ing way for recovering highly accurate curve skeletons in this view-based framework.

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