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Even Entem, Loïc Barthe, Marie-Paule Cani, Frederic Cordier, Michiel van de Panne. Modeling 3D animals from a side-view sketch. Computers and Graphics, Elsevier, 2015, 46, pp.221-230. 10.1016/j.cag.2014.09.037 . hal-01073059

HAL Id: hal-01073059

<https://hal.inria.fr/hal-01073059>

Submitted on 8 Oct 2014

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Modeling 3D animals from a side-view sketch

Even Entem^{a,b}, Loic Barthe^a, Marie-Paule Cani^b, Frederic Cordier^c, Michiel van de Panne^d

^aIRIT - University of Toulouse

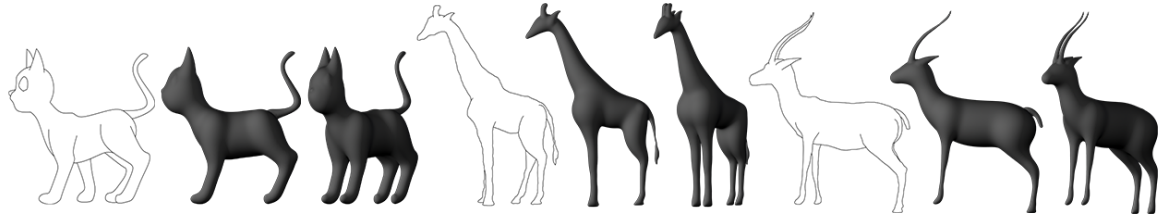
^bUniversity of Grenoble-Alpes, CNRS (Laboratoire Jean Kuntzmann) and Inria

^cUniversity of Haute Alsace

^dUniversity of British Columbia

Abstract

Using 2D contour sketches as input is an attractive solution for easing the creation of 3D models. This paper tackles the problem of creating 3D models of animals from a single, side-view sketch. We use the *a priori* assumptions of smoothness and structural symmetry of the animal about the sagittal plane to inform the 3D reconstruction. Our contributions include methods for identifying and inferring the contours of shape parts from the input sketch, a method for identifying the hierarchy of these structural parts including the detection of approximate symmetric pairs, and a hierarchical algorithm for positioning and blending these parts into a consistent 3D implicit-surface-based model. We validate this pipeline by showing that a number of plausible animal shapes can be automatically constructed from a single sketch.



1. Introduction

With the spread of 3D virtual environments and of 3D printing technologies, many practitioners would like to author their own 3D shapes. Among them, animal models – including imaginary and fantastic ones – are an important category. Being able to easily create and then animate animals would be an important step for generating more lively virtual worlds. Animals are also among the models that the general public, especially children, would typically like to sculpt and print.

There is currently no fast and easy method for creating 3D models of animals. Unfortunately, getting data from 3D scans is much more difficult for animals than for humans, beginning with the obvious challenge of requiring an animal to stand still. In addition, such reconstructions are also limited to existing animals. Standard 3D modeling software, such as Autodesk's Maya or Blender, as well as digital sculpting software such as

Pixologic-Zbrush, can be used for creating animals, but their complexity limits their use to experienced or passionate users. The use of 3D sculpting is possible, but is still difficult: many people are not adept at sculpting animals using real clay and in this case, they are not likely to perform much better in a virtual setting, even with an investment of time in the mastery of digital sculpting interfaces. Sketch-based modeling systems, which only require users to sketch contours in 2D, are probably the most intuitive and accessible class of methods. However, despite these advantages, they either require users to iteratively draw complex shapes part by part, using different viewpoints, or, alternatively, they require an existing data-base of 3D models.

Our work belongs to the category of sketch-based modeling methods and is the first to explore the creation of a 3D animal model from a single, side-view sketch. We are motivated by the belief that many users are capable of drawing a single sketch that depicts the

contour lines and the internal silhouettes of an animal, such as shown in the top-left of Figure 1. If needed, users can use a background drawing or a photo as a guide. The process of inferring 3D geometry from the 2D sketch necessitates the use of relevant assumptions in order to be tractable, and in our context of modeling animal forms, we shall assume smoothness of the resulting shape as well as the presence of structural symmetries. Several further moderate constraints include: (a) restricting oneself to non self-overlapping limbs in the sketch; (b) requiring the user to draw both contours for pairs of symmetric limbs; and (c) ignoring the reconstruction of repetitive details scattered on the surface, such as scales. With these assumptions and constraints in place, the method we develop is capable of automatically converting an input vectorized sketch into a 3D model. The total processing time is less than one second, effectively enabling one to create new animal models in only the time required to sketch them.

Note that in this work, we only tackle the creation of the volumetric shape parts of an animal and that we do not consider the surface components that should be used for ears or scales. The ears we reconstruct are also therefore interpreted as volumes. We are not able to reconstruct large flat parts such as wings. In addition, we reconstruct limbs in a symmetric fashion, even when they were drawn in arbitrary postures in the input sketch. The construction of symmetric 3D models is usually desirable, as it ensures that left and right limbs have identical dimensions. To achieve a desired non-symmetric posture, the 3D model can be deformed, either by using an animation skeleton and the associated skinning weights, or by directly articulating the implicit surface's skeleton.

Our processing pipeline for creating 3D animals from a sketch is summarized in Section 3. It consists of three main steps, which also correspond to our three technical contributions:

1. the identification of the animal's foreground structural parts in the sketch, with completion of the parts that are not explicitly bounded, such as the top of the legs (see the teaser figure)
2. the generation of a hierarchical graph of depths for the structural parts, using the complete results and based on an algorithm for detecting the portions of the sketch that correspond to symmetrical parts of the animal;
3. 3D reconstruction based on a specific choice of implicit surface, scale invariant integral surfaces, which enables us to accurately reconstruct shape parts from their medial axis in the 2D sketch,

and to seamlessly blend them into a single animal model.

2. Related Work

2D sketches only represent the contours, silhouettes and main features of an object. Converting them into a 3D model therefore requires resolving indeterminacies and inferring a large amount of missing data. Four strategies are commonly used to do so, each of which is based on a different level of hypotheses or *a priori* knowledge of the shape being modeled (see [1] and [2] for detailed surveys):

Iterative methods enable the user to build general models part by part, by iteratively adding new shape components from different viewpoints, e.g. the Teddy system [3]. These methods make the hypothesis that the final shape is a combination of parts that all have planar silhouettes from a given viewpoint, and can therefore be inflated from closed planar contours. Animals belong to this category but in practice these methods still require practice and time in order to achieving a convincing result. Other iterative methods have been developed using alternatives to inflation for geometric construction. These include the use of primitives such as generalized cylinders and ellipsoids, as in [4], and methods based on implicit surfaces, as in [5] or [6].

Shape matching approaches match the user sketch with silhouettes, or parts of silhouettes of predefined 3D models, possibly enabling some deformation. This method was successfully applied to organic shapes such as humans or animals [7], as well as for technical models [8]. However, they require a template example of the given general class of animal which imposes a restriction on the family of sketches that can be used. We did not consider this approach in our case, in order to allow the user to imagine any animal shape without further restrictions on the number of limbs, horns, etc.

Multi-view methods, e.g., [9], generate 3D shapes from two or three sketches drawn from orthogonal viewpoints. They require the ability to draw consistent views of the shape to be reconstructed. They are therefore more easily applicable to man-made objects than to organic shapes. Alternatively, a number of man-made objects can be easily built from a network of 3D curves, which can themselves be directly reconstructed from perspective sketches [10, 11]. However, these methods require more user input than a single contour sketch, and are generally difficult to apply to the modeling of free form, organic shapes.

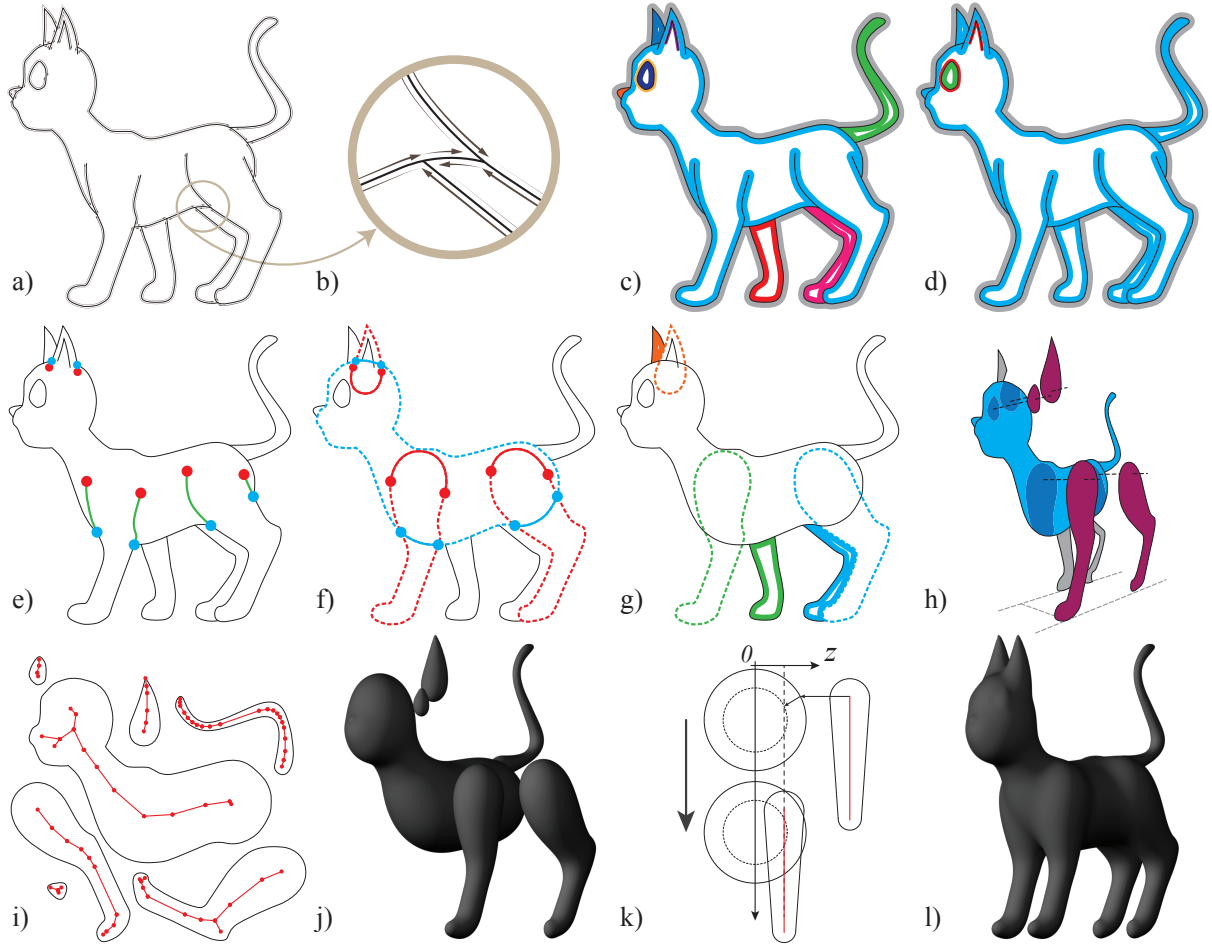


Figure 1: Overview of algorithmic stages: a) Half-edge graph; b) Detailed view of the resulting bounded curves; c) Cycles extracted from the graph, each one depicted in a different color; d) Classification of cycles: the different colors now represent the cycle type; e) Detection of suggestive contours; f) Completion of contours around shape parts; g) Detection of the main body and of pairs of symmetrical parts (displayed using the same color); h) Extracting a depth ordering; i) Skeletons extracted from the medial axes of shape parts; j) Reconstructed shape parts; k) Front view: inferring depth information for shape parts; l) Final result after implicit blending.

Lastly, **direct methods** try to infer 3D shapes from a single complex sketch that depicts complex silhouettes with loops, branches, cusps and T-junctions [12]. This technique is probably the most appealing, since it imposes few restrictions on the sketch being drawn, and mimics the human ability to *see in 3D* when provided a single, 2D-only representation. Different levels of a priori knowledge are used for performing this task, spanning the range from very specific context based hypotheses, as in methods for sketching flowers [13], garments [14], trees [15] or blood vessels [16] to more general methods, such as the one for reconstructing arbitrary shapes under the hypothesis of exact geometrical symmetry [17]. Our research belongs to this last category. In the context of working with animal shapes,

we leverage the hypotheses of symmetrical subshapes with planar contours, smoothly blending with the main body. In contrast to related prior work exploiting symmetry [17], we do not demand exact geometrical symmetry to be present in the input sketch, which would be difficult to draw for most users.

3. Overview

Our method builds a 3D model from a single 2D sketch. This involves the following steps, which are also summarized in Figure 1.

We start from a sagittal-view vectorized sketch of an animal in the (x, y) plane. The hand-drawn input sketch is transformed into a set of parametric curves using a

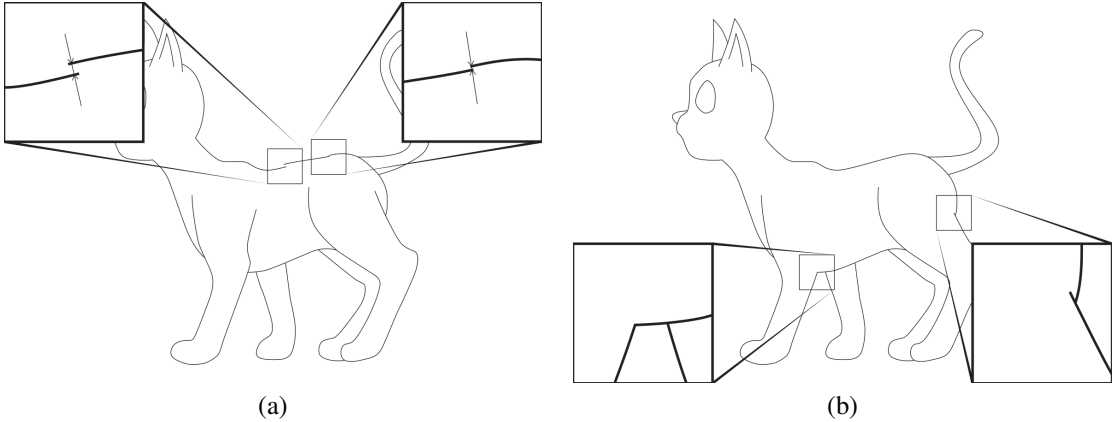


Figure 2: Two cases of failure: (a) Discontinuous sketch: a large discontinuity, as the one on the left, prevents the detection of the main body part in the drawing. In that case our algorithm fails from its first steps. A small gap in the sketch, as on the left, is filtered and considered as a connection during our first step. The reconstruction will perform adequately. (b) Incomplete sketch: on the left, a structural part is missing in the sketch, here, the left inner-body silhouette of the front foreground leg. Our algorithm will not be able to detect the presence of a limb and the reconstruction of the legs will fail. On the right, both inner-body silhouettes of the rear foreground leg are drawn and the rear legs will therefore be adequately reconstructed.

vectorization algorithm such as the one proposed by Noris et al. [18]. These algorithms are relatively robust to noise and provide the smooth curves we seek. What the user must really pay attention to is the correctness of the input hand-drawn sketch topology, i.e., the provision of continuous closed contours that do not have spurious crossings with other contours. Figure 2 illustrates two failure cases, the first, in 2(a), is due to an incorrectly closed contour and the second, in 2(b), is due to an incomplete input.

The goal of the first step, detailed in Section 4, is to identify the strokes that correspond to the same structural part, such as an eye, leg, or body, and to infer any missing curves, i.e., perform contour completion, if the resulting contour is open. This is done in the following way: We build a counter-clock-wise oriented half-edge graph whose edges are in general bounded parametric curves (see Figure 1(a,b)). In this graph, we iteratively follow successive half-edges, and store cycles in a list (Figure 1(c)). Each cycle is classified as being either an *outer-sketch*, *island*, *border* or *other* as explained in Section 4.1 and illustrated in Figure 1(d).

We then define as *inner-edges* those edges where both half-edges belong to the same cycle (the green edges in Figure 1(e)). These lists of inner-edges represent *suggestive curves* where two shape parts merge (such as a leg merging with the body). This implies the presence of hidden silhouettes or cusps [12]. Each suggestive curve has an open endpoint and a closed endpoint, which are depicted in red and blue, respectively, in Figure 1(e). The next step is to pair these suggestive curves and con-

nect their extremities in order to infer new cycles representing the different structural parts of the animal (Figure 1(f)). This is the topic of Section 4.2.

The subsequent step, presented in Section 5, is the computation of a depth hierarchy for structural parts, based on the structural symmetry hypothesis. We first identify the main body part located in the sagittal plane, i.e., the torso, and detect structural symmetries around it such as pairs of ears or pairs of front legs, cf. Section 5.1 and Figure 1(g)). We then use the pieces of information we already gathered, e.g., island cycles, structural parts, chest, and symmetries, to define a depth hierarchy between structural parts, as detailed in Section 5.2 and Figure 1(h).

This leads us to the last step, detailed in Section 6 which consists of the 3D reconstruction of the animal model. For each pair of limbs, only the one in the foreground of the drawing is considered during the reconstruction steps. The 3D reconstruction of the background limbs are added at the end of the process by symmetry. We choose implicit surfaces to represent shape parts, in order to benefit from their blending capabilities. We first compute the medial axis [19] of each structural part (Figure 1(i)). Medial axes are then filtered and specifically simplified in order to be used as skeletons for 3D implicit surface modeling (Section 6.1 and Figure 1(j)). Among the variety of existing implicit models, we use scale invariant integral surfaces (SCALIS) [20] to accurately reconstruct shape parts (Section 6.2). Figure 1(j)) shows the 3D reconstruction of each structural shape part in isolation. Structural parts without sym-

metries are placed in the sagittal plane of the model. The depth of the foreground structural parts is computed using the thickness of their 3D reconstruction and the thickness of the part to which it is connected in the sagittal plane (Section 6.3 and Figure 1(k)). Symmetric background parts are then added to provide the final 3D model (Figure 1(l)). The final shape is obtained using a simple sum to blend the fields from the different implicit primitives.

The next sections give a detailed presentation of our solutions at each stage of this process.

4. Structural parts identification and completion

4.1. Representing the sketch as a set of cycles

The first step is to decompose the curves of the sketch into a set of cycles. This decomposition is a preliminary step towards identifying the different parts of the animal drawn in the sketch. In particular, this step is required for the hidden contour computation and the contour closure, as described in Section 5.

The input data is a hand-drawn sketch composed of a set of non-crossing curves that are connected to each other (see Figure 1(a)). Each of these curves is represented by a pair of directed half-edges¹ of opposite direction (see Figure 1(b)). The curves of the sketch are then represented using a graph whose edges are the half-edges of the curves and whose nodes are the points at which the curves are connected to each other. Next, we decompose this graph into a set of cycles.

Formally, a cycle is defined as a closed sequence of connected half-edges; each half-edge shares its endpoints with at least two other half-edges. We require the half-edges that compose a cycle not to cross each other; we also require the cycles to be topologically equivalent to a disk. Therefore, a cycle divides the sketch into an "interior" region and an "exterior" region. In order to define the interior of a cycle, cycles and their associated half-edges are given a direction which is either clockwise or counter-clockwise; the interior of the region bounded by a cycle lies on the left side of the directed half-edges that compose this cycle.

The first cycle that we identify is the one which is composed of the half-edges located along the outer boundary of the sketch. This cycle is oriented clockwise and is unbounded; it corresponds to the outer region of the sketch. We name this the *outer* sketch cycle. Next, we identify the *border* cycles which are cycles that have

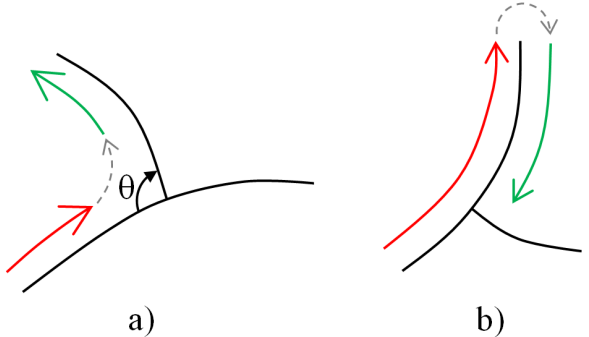


Figure 3: Selecting the next half-edge. In (a), the selection of the next half-edge (shown in green) for T-junctions is chosen such that the angle between the next half-edge and the preceding one (shown in red) is the smallest possible. In (b), the selection of the next half-edge for cusps.

one or several half-edges belonging to the outer boundary of the sketch. All of these cycles are oriented in a counter-clockwise fashion. The process to construct a border cycle is as follows: we first select a half edge which is located along the outer boundary of the sketch and which has not yet been associated to a cycle. We next find its endpoint by following its direction and select the next half-edge. This next half-edge is chosen such that its direction is the same as the previous half-edge and the clock-wise oriented angle between the two half-edges is the smallest possible, as illustrated in Figure 3. The process of finding the next half-edge is iterated until we reach the first half-edge.

Finally, we identify the island cycles; island cycles are single half-edges that correspond to a closed curve with no self-intersection and which are not connected to any other curves of the sketch. These island cycles are also oriented counter-clockwise, meaning that they do not represent a hole but rather a surface feature of the animal (such as the eye of cat model shown in Figure 1). After processing all the cycles, there may remain some cycles that could not be classified as any of the three types (outer sketch cycles, border cycles and island cycles); we mark these cycles as *others*.

4.2. Contour completion

The goal is now to convert the set of cycles into closed contours associated to each structural part of the shape.

Once the graph cycles are classified as described above, we use the hypothesis that structural parts have at least one edge in contact with the outer part of the sketch, and consider only the border cycles. In these cycles, we extract the sets of neighbor embedded edges,

¹Note that each graph edge geometrically correspond to a full curve of the sketch.

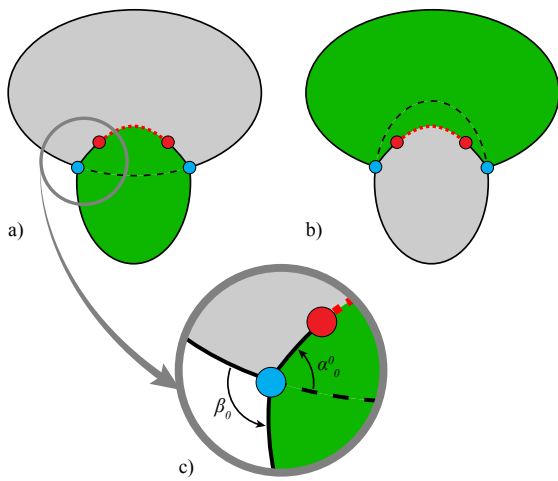


Figure 4: (a) and (b): Illustration of the two possible completions of a pair of closed extremities. (c) Close-up on a closed extremity illustrating the two angles α_0^0 and β_0 used in the computation of the probability p_0^0 .

i.e. edges whose half-edges both belong to the same cycle, and then annotate these as being suggestive curves. Each set is an open path in the graph with one extremity connected to the rest of the graph, i.e., a closed extremity, and one extremity without any connection, i.e., an open extremity (see Figure 1(e)). As we focus on organic models, the suggestive curves come from the drawing of the silhouette of structural parts that stop where the limb smoothly merges with the part of the shape over which they are drawn (for instance the cat legs over the body in Figure 1). We require the user to provide two suggestive curves clearly delimiting each merging limb.

Parts need to be individually closed to be identified and reconstructed. To do so, we compute a set of cubic Bézier curves linking open extremities and closed extremities pairwise with a C^1 continuous connection with the edge curves while minimizing the normalized sum of the curvature variations (“Gestalt principles”):

$$E = \frac{1}{l^2} \sum_i |k_{i+1} - k_i| \quad \text{where } k_i = \frac{\|\dot{c}_i \times \ddot{c}_i\|}{\|\dot{c}_i^3\|}, \quad (1)$$

Here, l is the curve length, k_i is the curvature of the curve at sample point i , \dot{c}_i is the curve first derivative, and \ddot{c}_i is the second derivative at parameter value u_i .

The sum of curvature variations, $l^2 E$, is used as the energy associated with the curve. For curves connecting closed extremities, these energies are denoted E_c and for those connecting open extremities they are denoted

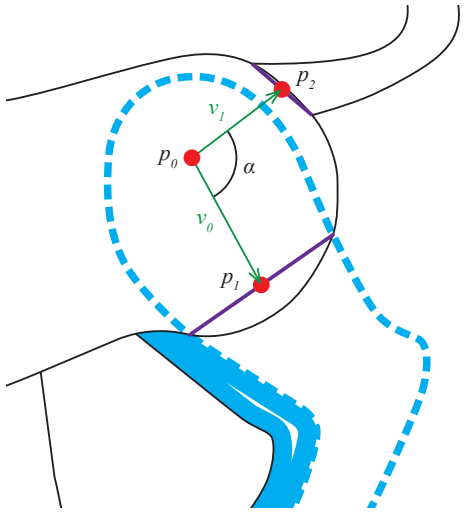


Figure 5: Identification of the cycles belonging to the shape part at the background that is structurally symmetrical to a foreground limb.

E_o . We then compute two energies E_i ($i = 0, 1$) for each pair of suggestive curves, one for each way the closed extremities can be connected (dashed curves in Figure 4(a,b)). These energies are computed as follows:

$$E_i = (p_0^i \cdot p_1^i)(E_o + E_c^i), \quad (2)$$

where E_c^i is the energy associated to the curve connecting closed extremities ($i = 0$ for the closure illustrated in Figure 4(a) and $i = 1$ for the one illustrated in Figure 4(b)), p_0^i and p_1^i are the probabilities that the curve is correctly connected at each extremity respectively. These probabilities are computed as the ratio $p_j^i = \min(\alpha_j^i / \beta_j, 1)$ ($j = 0, 1$) in which angles α_0^0 and β_0 are illustrated in Figure 4(c). All other angles are defined in a symmetric fashion. The best-matching pair of suggestive contours is then chosen as the pair having the minimal sum of energies.

Once the suggestive contours are paired and closed, we define their set of half-edges and derive the new cycles. In these cycles, no edge is embedded in the defined part (i.e. no edge has both half-edges in the same cycle). Together with the remaining border cycles and the island cycles, this gives us the set of the animal structural parts drawn in the sketch.

The minimization of curvature variations is known to provide plausible results when completing organic curves. The selected completion curves linking open and closed extremities does not, in general, intersect other contours. However, if this should occur, it will not be expected as it does not faithfully correspond to the intent of the input sketch, although it does not compro-

mise the rest of the reconstruction. If necessary, the user can correct the completion by editing the extremities.

5. Depth hierarchy of structural parts

5.1. Body and symmetrical parts

In order to assign appropriate depth values, we create a graph that represents the desired depth relations. This begins with the choice of a reference part as the root node of the graph, for which it is natural to use the body or torso. For some animals, this main body part may be smaller than the limbs. As such, we did not find a robust method to automatically label this in the input sketch. Instead, we request the user to select it or to draw the sketch so as to locate it in the center of the drawing canvas. Once labeled, this part defines the node of reference for the graph. The method described below automatically locates the remaining structural parts with respect to the body, in terms of their relative depth. We begin by detecting the sketch strokes that correspond to pairs of symmetric parts, as explained next.

We already know the parts, F_i , that should be located in front of the body and thus that may have a symmetrical part in the background. These are defined by the open contours with suggestive curves we just completed at the previous step. The goal is now to identify the strokes from the sketch corresponding to B_i , the structurally symmetric part with respect to F_i , but located in the background. Indeed, B_i should not be processed any further, since we are going to use symmetric geometry to create the background limbs. In practice, B_i may be composed of several distinct cycles, since it is partly occluded. We use a propagation method for fully selecting it, as we now describe.

We initialize B_i with all cycles not belonging to the shape part under F_i , but that share an edge with F_i . We then compute the medial axis of F_i as described in Section 6. Let us denote as p_0 the medial axis extremity corresponding to the suggestive silhouette we closed (see Figure 5). We then denote as p_1 the middle of the closed extremities of the suggestive curves and then define $v_0 = p_1 - p_0$. Starting from both closed extremities, we march along both sides (one clock-wise and the other counter-clock wise) of the cycle of the structural part which is located under F_i in the sketch, as long as the opposite half-edge belongs to the outside sketch. If we meet an opposite half-edge belonging to a border cycle, we get both extremities of the shared edge and we compute their middle point p_2 . We define $v_1 = p_2 - p_0$ and compute the angle $\alpha = |\widehat{v_0, v_1}|$. If $\alpha < \pi/4$ we consider that this cycle potentially belongs to B_i , since the

angle formed by two symmetrical parts is unlikely to be larger than this value in usual standing poses. Note that this criterion could be relaxed to handle a larger set of poses.

We stop when the half-edge we reach is an occluded edge, i.e., when we meet the next foreground part. If selected on multiple occasions, a potentially symmetric cycle is assigned to the B_i for which the angle α with F_i is minimal. The results of this selection process are depicted in Figure 1(g), where pairs of symmetric parts are depicted using the same colors.

5.2. Graph construction

In order to support the 3D reconstruction of the model, we build a graph whose nodes are the structural parts (and their associated cycle) and whose edges represent the relations *over*, *under*, *adjacent* or *over-in* with respect to the main body. Foreground parts are defined as lying *over* the part they have been separated from. Symmetric counterparts are located *under* the part that their foreground is *over*. All remaining border parts that have not been designated as part of a symmetric pair are defined as being *adjacent* to parts with which they share an edge. Islands are assigned to the graph by first using a simple 2D ray-cast in the sketch plane in order to identify the part within which the island lies. The island cycles are over and inside a structural part, and this part is also then defined as being *under* the island. Each edge is further tagged with additional information, such as the presence of suggestive curves, which help determine if the region being over a particular part is an island or a limb.

6. 3D Reconstruction

6.1. Medial-axis computation & simplification

Animals have the convenient property that their shape is naturally approximated as a set of generalized cylinders. A generalized cylinder is a surface obtained by moving a circle of varying radius along a 3D arbitrary curve. An advantage of generalized cylinders is that their skeleton and associated radii are easily determined from their projected image. Given a 2D simple closed curve C , the skeleton and the associated radii of the generalized cylinder whose orthogonal projection matches C are obtained by computing the medial axis and the associated radius function of the curve C .

We start the 3D reconstruction step by reconstructing each structural part independently. The parts which were detected as being symmetric to foreground parts are not considered. The contour curve of each part is

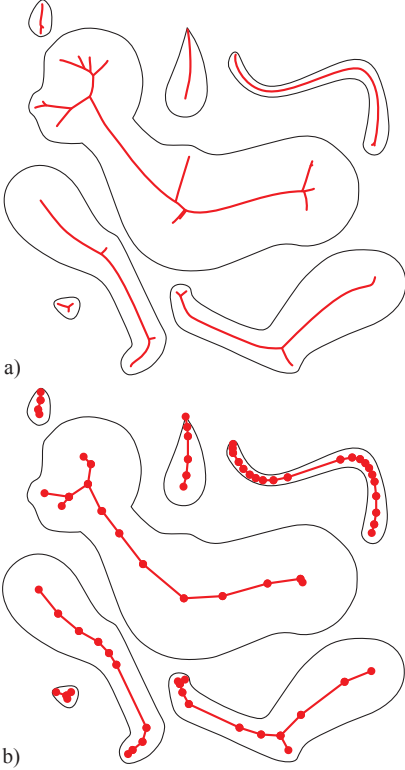


Figure 6: Medial axis computation and simplification. In (a), the medial axes of the sketch components computed using Fortune’s sweep-line algorithm [19]. In (b), the skeletons after removing the branches using the Scale-Axis Transform [21] and after simplifying the medial axes using the Douglas-Peucker algorithm [22].

first sampled so as to obtain a polygonal curve. We then compute the medial axis transform of each of the polygonal curves using the Fortune’s sweep-line algorithm [19]. The medial axes usually contain many branches because of the noise in the polygonal curves. We use the Scale-Axis Transform method by Giesen et al. [21] to remove these branches. In our experiments, the scale factor of the Scale Axis Transform has been set to 1.2.

The last step is to simplify the medial axis; the purpose of this simplification is to lower the computation time for generating the surface of the reconstructed animal. We have implemented a modified version of the Douglas-Peucker algorithm [22] which is a method to reduce the number of points of polygonal curves. A cost function is defined for each point, which is the maximum distance between the original curve and the curve after removing the point. Points are removed iteratively in increasing order of their cost. The algorithm stops when the cost of the point to remove is above a

user threshold. Figure 6(b) illustrate the result of the cat skeletons simplification.

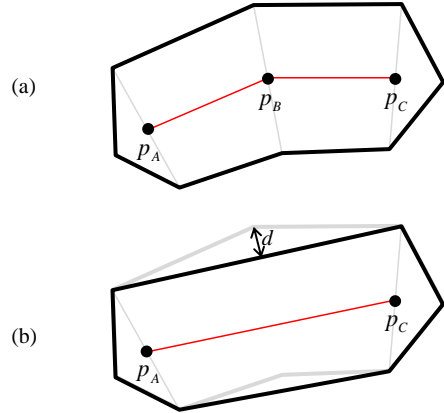


Figure 7: Medial axis simplification. In (a), the polygonal curve (depicted in black) and its medial axis (depicted in red) before the simplification. In (b), the medial axis has been simplified by removing the point p_B . The result is new polygonal curve with fewer segments. The cost value of removing the point p_B is d ; it is the maximum distance between the original curve (depicted in light grey) and the curve after the simplification.

The Douglas-Peucker algorithm has been modified to take into account the radii of the medial axis. This modification consists of a new cost function. Let C be a polygonal curve and p_A, p_B, p_C be three neighboring points of the medial axis of C . If the point p_B is removed, the two segments $\overline{p_A p_B}$ and $\overline{p_B p_C}$ are replaced with the segment $\overline{p_A p_C}$ (see Figure 7). Similarly, the curve corresponding to the medial axis is simplified; two of its segments are removed. The new cost function is defined as the maximum distance d between the original curve and the curve after the simplification of the medial axis.

6.2. Surface reconstruction of the shape components

Once the skeletons of all the shape components have been computed, the next step is to reconstruct the 3D surface of these shape components. We have implemented the SCALe-invariant Integral Surfaces (SCALIS) method proposed by Zanni et al. [20]. This method generates an implicit surface using a skeleton and its radius function. In Zanni’s method, the scalar field is computed with the Homothetic Polynomial kernel of degree 8. In our method, we have implemented the kernel of degree 6 instead of degree 8. Although the quality of the reconstructed surface is slightly lower, the computation time is much smaller, and allows for surface reconstruction at near-interactive rates. The result

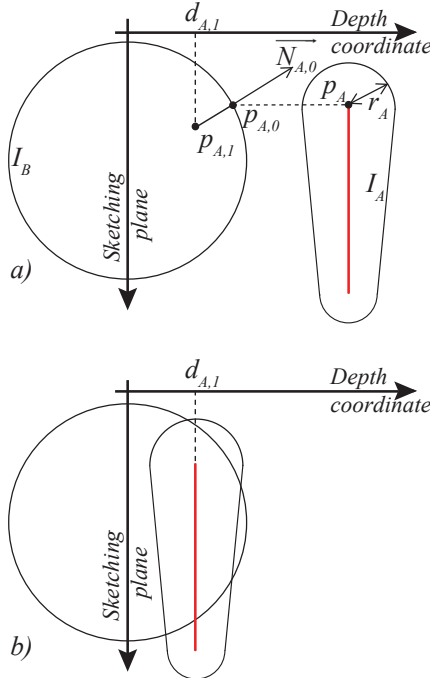


Figure 8: Computation of the depth coordinates of the implicit surface I_A . In (a), the depth coordinate $d_{A,1}$ of the skeleton extremity p_A is computed. In (b), the skeleton (depicted in red) of I_A is translated to the depth coordinate $d_{A,1}$.

is a set of implicit surfaces, one for each shape component (see Figure 1(k)).

6.3. Placement & embedding of the shape components

The last step of the surface reconstruction is to assemble the implicit surfaces to generate the 3D shape of the animal. At this stage, the skeletons of the implicit surfaces are all located in the same plane which is the sketching plane. The goal is to place these implicit surfaces so that their depth order complies with that of the sketch; the shape components that are drawn in the sketch foreground should be placed in front of the sketching plane and those drawn in the sketch background should be placed behind the sketching plane. We make use of the depth graph that has been previously computed (see Section 5.2).

We start by placing the implicit surface corresponding to the main body component in the sketching plane. Other implicit surfaces are sequentially placed as follows. Using the depth graph, we find a shape component that has not yet been placed and which is adjacent to a shape component which has already been placed.

Using the depth order between these two shape components, we compute the depth position of the one to be placed.

Let I_A and I_B be two implicit surfaces, with I_A being in front of I_B . The depth coordinates of I_B have been already computed; those of I_A have to be computed. Let p_A and r_A be the skeleton extremity of the implicit surface I_A and its radius respectively. We first compute $p_{A,0}$ the orthogonal projection of p_A onto I_B . Next, we compute $\overrightarrow{N_{A,0}}$, the surface normal of the implicit surface I_B at the point $p_{A,0}$. Finally, we compute $p_{A,1}$ such that $\|\overrightarrow{p_{A,1}p_{A,0}}\|$ is equal to r_A and such that the two vectors $\overrightarrow{N_{A,0}}$ and $\overrightarrow{p_{A,1}p_{A,0}}$ are parallel and have same direction (see Figure 8(a)). The depth coordinate $d_{A,1}$ of $p_{A,1}$ is then used as the depth coordinate of all the points of the skeleton of I_A (see Figure 8(b)).

Once the depth coordinates of all the implicit surfaces have been computed, the 3D shape of the animal is then generated by blending these implicit surfaces using the well-known Ricci blending operator.

7. Results and discussion

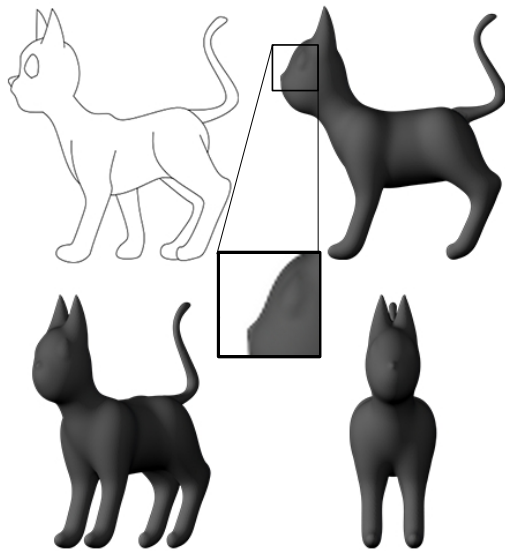


Figure 9: Different views of the 3D reconstruction of the cat sketched in the top-left.

We implement our method as a Maya plugin, enabling us to use the existing sketching interface of Maya to design the vectorized sketches we need as input. Processing the 2D sketches into 3D models is then a fully automatic process.

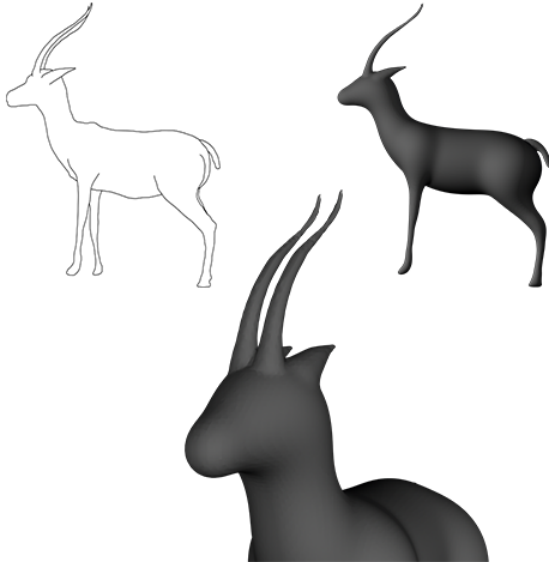


Figure 10: Different views of the 3D reconstruction of the gazelle sketched in the top-left.

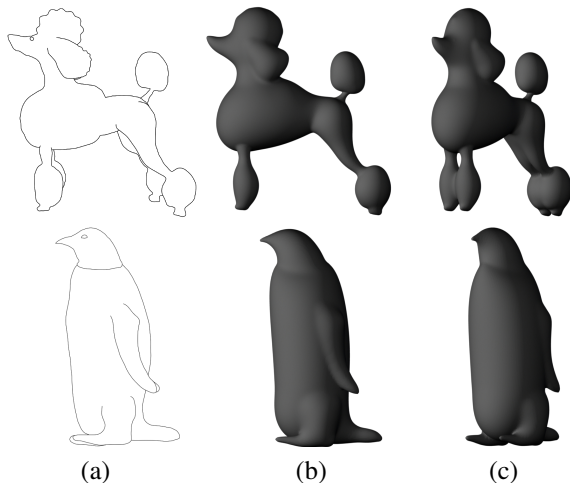


Figure 11: Reconstruction of a dog and a penguin. (a) The input vectorial sketch, (b) the automatically generated result and (c) different view of the reconstructed model.

The teaser image and Figures 9–12 show a set of results from our method. They validate the fact that our method is able to generate plausible rough shapes for various animals from a single sketch. Note that since we only used a structural symmetry hypothesis around the sagittal plane, our animal models may have an arbitrary number of protruding subparts such as limbs, ears (interpreted as volumes) or horns. Moreover, our choice of scale invariant implicit surfaces (SCALIS) as the 3D representation enables us to capture both the smooth-

ness of organic shapes and the singularities that may come from the sketch, such as at the tip of the cat ears.

Pairs of limbs are reconstructed symmetrically, but the user can manually adjust the pose in 1 to 3 minutes by deforming the geometric skeleton (see Figure 1(i) the geometric skeleton of the cat model). Background limbs can be positioned in sketch pose as in Figure 12(c), or the model can be globally deformed as in Figure 12(mantis-d). In Figure 12(elephant), the ears are not reconstructed as they do not represent any of the different cycles considered in the sketch for our reconstruction (as explained in Section 4.1).

The results have been produced on an Intel Core i7 3770K computer, and the code is compiled with ICC 15.0 in release mode. For all our examples, the full sketch-processing and 3D modeling process run in less than one second. This makes the method applicable within an interactive sketching software tool. More precisely, the computational time for the giraffe is 0.35 seconds, for the gazelle is 0.30 seconds and for the cat 0.37 seconds.

To further validate our system we performed a preliminary user study with 4 different users ranging from beginners to a professional computer artist. Their sketches and resulting models are shown in Figure 13. They all found the concept of suggestive curves to delimit the limbs quite easy to understand. When asked how well the 3D model met their expectations, on a scale from 1–9, they gave a score of 7. The time spent to create the input drawings using the Pencil Curve Tool in Autodesk Maya was less than 2 minutes for all the drawings, with the exception of the rabbit, which took 10 minutes to draw.

Limitations:

Our method has a number of remaining limitations.

First, we are limited in the range of sketches we are able to handle:

- The sketch cannot include adjacent internal parts (*islands* in our terminology) such as if the cat had two adjacent eyes. Otherwise, the classification algorithm will fail;
- A shape part cannot self-overlap in the sketch (such as an animal tail looping and self-occluding itself), nor overlap any other shape part (such as a tail occluding the animals body). Our completion algorithm does not handle these cases. Note that we partly avoid this occlusion problem by using symmetry when creating the limbs, but developing a more general solution would be desirable.
- Drawing pairs of suggestive contours is mandatory

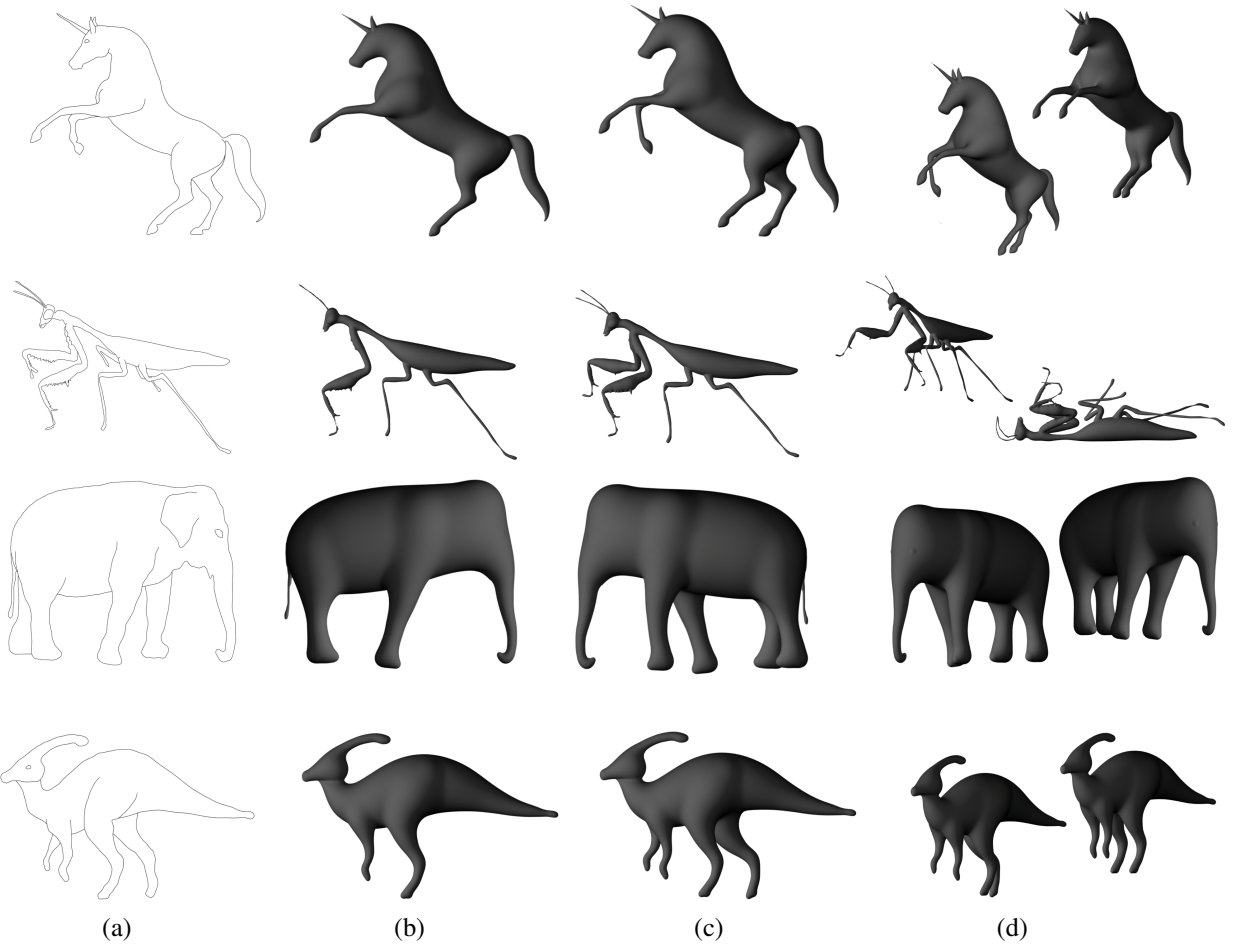


Figure 12: Reconstruction of a mantis, an elephant and a dinosaur. (a) The input vectorial sketch, (b) the automatically generated result, (c) the model with the foreground limbs manually adjusted to the sketch and (d) different views of the reconstructed models. The mantis is shown with different poses in (d).

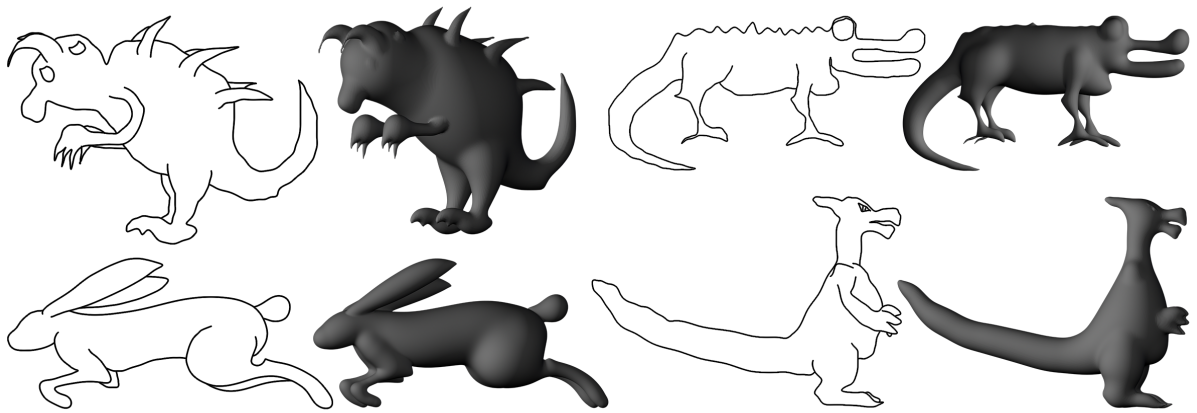


Figure 13: Reconstruction of imaginary creatures and a rabbit made by four different users ranging from beginners to a professional computer graphics artist (the rabbit model).

when two shape parts blend (such as at the top of a front leg, where it blends with the body), and these contours should be such that our completion algorithm keeps the reconstructed part of the contour inside the main body-shape. Although this holds true in most cases, we currently offer no guarantees regarding this point.

A second family of limitations concerns the 3D shape we output:

- All the shape parts we create have a planar medial axis, which we locate in a vertical plane. Although this works quite well for the legs, this method needs to be improved for parts such as the ears or the horns, which should preferably be oriented to lie in a direction that is normal to the surface.
- We only use segment skeletons for generating the implicit shapes, as opposed to surface skeletons, which restricts our construction to generalized cylinders. This prevents us from capturing the flat surfaces that one can readily observe on the flanks of many animals, or flat features such as elephant ears that cannot be reconstructed with our approach (see Figure 12).
- We reconstruct pairs of limbs using symmetric poses. Automatically fitting the reconstructed background parts is non-trivial since we do not reconstruct a rigging skeleton and most of the drawing of the background limbs is missing. Performing this fitting remains a challenging open problem on its own.

8. Conclusions and future work

We have presented the first method, to the best of our knowledge, for creating 3D animal models from a single sagittal-view sketch. Our method handles complex sketches with open curves, closed curves, and T-junctions. Once the main body part is selected, the method is fully automatic for reconstructing a symmetric version of the 3D shape. The reconstructed shape is inferred using only two hypotheses: shape smoothness, except at singular points appearing on the sketch contour; and structural symmetry. The resulting 3D model can then be posed using standard software or by directly manipulating its skeleton.

In future work, we would like to address some of the limitations we listed. In particular, we are in the process of extending our stroke classification and completion method to more general families of sketches. Being

able to convert any sketch to the vector graphic complexes proposed in [23] would be an excellent intermediate step for our application. In addition, exploiting the 2D detection of partial symmetries in the cycles, using techniques such as the one by Mitra et al. [24], may be a promising direction of investigation for improving the pairing stage of our approach. This could also help for handling more complex inputs such as sketches in arbitrary views. A last caveat is that the technique we currently use is not able to reconstruct large flat parts such as wings. A direction is to use more complex skeletons including surface parts as was previously done in the context of reconstruction with convolution surfaces [25].

Acknowledgements

We also thank the anonymous reviewers for their useful comments and suggestions, as well as Cédric ZANNI for sharing his implementation of SCALIS. This work was funded by the advanced grant EXPRES-SIVE from the European Research Council (ERC-2011-ADG_20110209) and by the project IM&M (ANR-11-JS02-007).

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