# OpenCCO: An Implementation of Constrained Constructive Optimization for Generating 2D and 3D Vascular Trees 

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#### Abstract

In this article, we focus on the algorithm called CCO (Constrained Constructive Optimization), initially proposed by Schreiner and Buxbaum [Computer-Optimization of Vascular Trees, IEEE Transactions on Biomedical Engineering, 40, 1993] and further extended by Karch et al. [A Three-Dimensional Model for Arterial Tree Representation, Generated by Constrained Constructive Optimization, Computers in Biology and Medicine, 29, 1999]. This algorithm can be considered as one of the gold standards for vascular tree structure generation. Modeling and/or simulating the morphology of vascular networks is a challenging but crucial task that can have a strong impact on different applications such as fluid simulation or learning processes related to image segmentation. Various implementations of CCO were proposed over the last years. However, to the best of our knowledge, there does not exist any open-source version that faithfully follows the native CCO algorithm. Our purpose is to propose such an implementation both in 2 D and 3 D .


## Source Code

The reviewed source code and documentation associated to the proposed algorithms are available from the web page of this article ${ }^{1}$, as well as the on the GitHub repository (OpenCCOteam $/$ OpenCCO ${ }^{2}$ ). The correspondences between algorithms and source codes are mutually given and compilation with usage instructions are included in the README.md file of the archive. The archives also contains the scripts allowing to reproduce the experiments included in various figures.

Keywords: constrained constructive; optimisation; implementation; simulation; tubular structure; vasculature; bifurcation; 3D images

[^0]
## 1 Introduction

In image processing and analysis, thin structures are among the most difficult to handle. This is especially true for line-like structures in 2-dimensional images or tube-like structures in 3-dimensional images. Such structures are particularly present in the context of (bio)medical imaging. A wide literature has been dedicated to the preprocessing (e.g. filtering), processing (e.g. segmentation) and analysis (e.g. quantification) of vascular structures in so-called angiographic images, which are typical examples of 1-dimensional structures in 2- and 3-dimensional images. Recent surveys on such image processing / analysis approaches [10, 14] summarize the investigation of a wide range of methods for tackling the issues related to blood vessel processing and analysis (from classical image processing to more recent deep learning paradigms).

By contrast with this rich literature, little effort has been dedicated to the generation of tree structures as geometrical objects which could fairly model vascular trees. Being able to build ex nihilo such objects is, however, a crucial research issue. Indeed, the availability of vascular models that present realistic properties with respect to actual vascular trees would open the way to the design of efficient benchmarks for comparing image processing / analysis methods [9], for modeling vascular structures that cannot be easily observed (e.g. at the microvascularization level) [16], for carrying out efficient experiments in computational fluid dynamics [13], or for generating realistic virtual medical images natively endowed with ground-truth [2].

In 1993, Schreiner and Buxbaum [17] proposed an algorithm for building tree structures in $\mathbb{R}^{2}$ with the purpose of endowing them with correct properties related to vascular networks. In 1999, Karch et al. [7] extended this algorithm to the construction of tree structures in $\mathbb{R}^{3}$. In 2010, VascuSynth ${ }^{3}$ [3] was proposed as an open-source software building upon CCO. Although VascuSynth data are frequently used in the angiographic image analysis literature, the underlying software presents various limitations compared to the native version of CCO (e.g. regarding spatial domain definition, optimization strategy for bifurcation geometry computation). In 2019, Jaquet et al. revisited the CCO algorithm [5] with the purpose of using it for cardiovascular modeling. Unfortunately, the associated software was not made available to the scientific community. A couple of years ago, we decided to provide an open-source version of this algorithm. Doing so, we also carefully revisited the algorithm, in order to efficiently deal with some tricky aspects, computationally expensive steps, and to try to simplify its formulation, especially by discriminating its algorithmic and physiology-guided layers.

In this article, we describe our own interpretation of the CCO algorithm, that we aim to make completely clear and as close as possible to the original one, and the details of its implementation. The rest of this article is organized as follows. In Section 2, we propose a synthetic summary of the proposed algorithm. Section 3 explains the principal key points of the algorithm. Section 4 describes one representative iteration of the construction algorithm. Section 5 describes the main parameters of the algorithm. Details related to our implementation of the algorithm are given in Section 6. Experimental results are proposed in Section 7. Section 8 provides concluding remarks and perspectives.

## 2 Overview of the Method

### 2.1 Description of the Constructed Tree

The proposed method builds a tree $\mathfrak{T}$ in $\mathbb{R}^{d}(d=2$ or 3$)$ inside a bounded, closed connected domain $\Omega \subset \mathbb{R}^{d}$.

[^1]From a topological point of view, $\mathfrak{T}$ is composed by a set $\mathcal{V}$ of $n=2 k$ vertices $(k \geq 2)$, including one vertex called the starting vertex, $k \geq 1$ vertices called ending vertices, and $k-1$ vertices called inside vertices; and a set $\mathcal{E}$ of $n-1$ edges linking two distinct vertices. Each ending / starting vertex is incident to exactly one edge. Each inside vertex is incident to exactly three edges. Each edge is incident to at least one inside vertex and at most one ending / starting vertex. The induced graph $(\mathcal{V}, \mathcal{E})$ is connected and acyclic, i.e. it is a tree.

The path between any vertex $v \in \mathcal{V}$ and the starting vertex being unique, we can non-ambiguously define a "parenthood" function $\varphi: \mathcal{V} \rightarrow \mathcal{V}$ such that $\varphi(v) \in \mathcal{V}$ is the successor of the vertex $v$ in the path between $v$ and the root (which is the only vertex where $\varphi$ is not defined). In particular, for any $v \in \mathcal{V}$ except the starting vertex, $(v, \varphi(v))$ is an edge of $\mathcal{E}$, and this specific edge will also be noted $e(v)$. Conversely, we note $\Phi(v)=\{u \in \mathcal{V} \mid \varphi(u)=v\}$.

Let $v_{i} \in \mathcal{V}(i \in \llbracket 0, n-1 \rrbracket)$ be a vertex of the tree $\mathfrak{T}$. From a geometrical point of view, the vertex $v_{i}$ is embedded as a point $p_{i} \in \mathbb{R}^{d}$, whereas the (putative) edge $e\left(v_{i}\right) \in \mathcal{E}$-also noted $e_{i}$ for the sake of concision-is embedded as a straight segment $S_{i} \subset \mathbb{R}^{d}$. In particular, if the edge $e_{i}$ is incident to the two vertices $v_{i}, v_{j} \in \mathcal{V}$, then the segment $S_{i}$ is defined as $S_{i}=\left[p_{i}, p_{j}\right]=\left\{\alpha \cdot p_{i}+(1-\alpha) \cdot p_{j} \in\right.$ $\left.\mathbb{R}^{d} \mid \alpha \in[0,1]\right\}$.

From a morphological point of view, each segment $S_{i}$ has a given thickness defined by a radius parameter $r_{i} \in \mathbb{R}_{+}^{*}$ associated to the edge $e_{i}$. In practice, the 1-dimensional segment $S_{i}$ is then modeled as a $d$-dimensional "tubular" object $C_{i}$ that can be seen as the dilation $C_{i}=S_{i} \oplus B_{r_{i}}$ of $S_{i}$ by the Euclidean ball of radius $r_{i}$, namely $B_{r_{i}}=\left\{x \in \mathbb{R}^{d} \mid\|x\|_{2} \leq r_{i}\right\}$. The whole tree $\mathfrak{T}$ is then defined as the union of these tubular shapes, i.e. $\mathfrak{T}=\bigcup_{i=0}^{n-1} C_{i}$. This tree $\mathfrak{T}$ lies inside $\Omega$, i.e. $\mathfrak{T} \subseteq \Omega$. If two edges $e_{j}$ and $e_{k}$ are not incident to a same vertex $v_{i}$, then we should have $C_{j} \cap C_{k}=\emptyset$, so that the acyclicity of the graph $(\mathcal{V}, \mathcal{E})$ remains valid in the tree $\mathfrak{T} \subset \mathbb{R}^{d}$.

From now on, for the sake of readability, we will sometimes make no distinction between the notions of vertex $\left(v_{i}\right)$ vs. points $\left(p_{i}\right)$ and the notions of edges $\left(e_{j}\right)$ vs. segments $\left(S_{j}\right)$. In other words, we will make no distinction of notation between the tree $\mathfrak{T}$ considered as a graph and as an embedded volume.

### 2.2 Input / Output

The method has the following inputs:

- $\Omega \subset \mathbb{R}^{d}$ : the domain where the tree has to lie;
- $p_{0} \in \Omega$ : the position of the starting point / vertex;
- $k \in \mathbb{N}_{+}$: the number of ending points / vertices;
and provides the following outputs:
- $\mathcal{V}=\left\{p_{i}\right\}_{i=0}^{n-1}$, the set of the $n=2 k$ points / vertices of the tree $\mathfrak{T}$;
- $\mathcal{E}=\left\{S_{i}\right\}_{i=1}^{n-1}$, the set of the $n-1$ edges / segments $S_{i}$ together with their respective radii $r_{i} \in \mathbb{R}_{+}^{*}$.


### 2.3 Sketch of the Algorithm from a Topological Perspective

The tree $\mathfrak{T}$ is incrementally constructed from a trivial tree-composed by two vertices and one edgeby iteratively adding new edges composed of two new vertices. At each iteration, each new edge is connected to the current tree by "breaking" a current edge of the tree into two parts. From the only topological point of view-i.e. if we consider the vertices $v_{i}$ and the edges $e_{i}$, without dealing with

```
Algorithm 1: Tree construction (sketch: topological point of view).
    Input: \(k \in \mathbb{N}_{+}^{*}\)
    Output: \((\mathcal{V}, \mathcal{E})\) with \(\mathcal{V}=\left\{v_{i}\right\}_{i=0}^{n-1}\) and \(\mathcal{E}=\left\{e_{i}\right\}_{i=1}^{n-1}(n=2 k)\)
        Initialization of the tree
    Let \(v_{0}, v_{1}\) be two vertices
    Let \(e_{1}=\left(v_{1}, v_{0}\right)\)
    Let \((\mathcal{V}, \mathcal{E})=\left(\left\{v_{0}, v_{1}\right\},\left\{e_{1}\right\}\right)\)
    // Incremental addition of new vertices and edges
    foreach \(i \in \llbracket 1, k-1 \rrbracket\) do
        Let \(v_{2 i}, v_{2 i+1}\) be two vertices
        Set \(\mathcal{V}=\mathcal{V} \cup\left\{v_{2 i}, v_{2 i+1}\right\}\)
        Choose \(e_{j}=\left(v_{j}, v_{k}\right) \in \mathcal{E}\)
        Set \(e_{j}=\left(v_{j}, v_{2 i}\right)\)
        Let \(e_{2 i}=\left(v_{2 i}, v_{k}\right)\)
        Let \(e_{2 i+1}=\left(v_{2 i+1}, v_{2 i}\right)\)
        Set \(\mathcal{E}=\mathcal{E} \cup\left\{e_{2 i}, e_{2 i+1}\right\}\)
```


(a) Initial

(b) Final

(c) Global tree structure

Figure 1: Addition of a new edge in the tree ( $\mathrm{a}, \mathrm{b}$ ) (see Algorithm 1) and illustration of the structure used in the CCO algorithm (c). The red disk area associated to each terminal point represents the neighboring region receiving $Q_{\text {out }}$ flow at pressure $P_{\text {out }}$.
their spatial embeddings, namely the points $p_{i}$ and the segments $S_{i}$ of radii $r_{i}$-then the algorithmic sketch of CCO is given in Algorithm 1 (with an illustration of edge updating and the global tree structure in Figure 1).

This algorithm is basically the construction of a (nearly) binary tree. The relevance and the main difficulty of CCO lie in the additional handling of the spatial embedding of this tree. It is indeed required to generate this tree by associating to each vertex $v_{i}$ a point $p_{i}$ and to each edge $e_{i}$ a segment $S_{i}$ associated with a radius $r_{i}$, while satisfying geometrical and morphological constraints related to the domain $\Omega$ and various parameters. These constraints are briefly summarized hereafter. Their algorithmic handling will be explained in detail in the next section.

### 2.4 Geometrical and Morphological Constraints

The construction of the tree $\mathfrak{T}$ is enriched / constrained by various factors.
(1) Each edge $e_{i}=\left(v_{i}, v_{j}\right)$ is associated to a segment $S_{i}$ with a radius $r_{i}$ that depends on its spatial embedding (i.e. the distance between the positions of $p_{i}$ and $p_{j}$ in $\Omega$ ) but also on the positioning of the edge into the tree and the radii associated to the other segments. This radius handling is discussed in Section 3.1.
(2) For each edge $e_{i}=\left(v_{i}, v_{j}\right)$ associated to two new vertices $v_{i}, v_{j}$ and associated points $p_{i}, p_{j}$, one of these points, say $p_{i}$, which is an ending point, has a position randomly chosen under constraining criteria. The position of the second point, say $p_{j}$ which is an inside point, is computed by an optimization process. This optimization process is guided by parameters which allow one to model physiological priors. This optimization procedure is discussed in Section 3.2.
(3) During the construction of the tree, it is forbidden for (i) the segment associated to a new edge to be too short and (ii) for the two ending points to be too close. It is also forbidden that (iii) two segments intersect. (Here, the segments are viewed as volumes.) This requires a segment intersection checking at various stages of the process. These checking procedures are discussed in Section 3.3.

## 3 Main Key Points

### 3.1 Radius Computation

In the tree $\mathfrak{T}$ induced by $(\mathcal{V}, \mathcal{E})$, each vertex $v_{i}(i \in \llbracket 0, n-1 \rrbracket)$ is associated to a point $p_{i} \in \mathbb{R}^{d}$, and each edge $e_{i} \in \mathcal{E}(i \in \llbracket 1, n-1 \rrbracket)$ is associated to a segment $S_{i}=\left[p_{i}, p_{j}\right] \subset \mathbb{R}^{d}$ (with $\left.v_{j}=\varphi\left(v_{i}\right)\right)$ of length $\ell_{i}=\left\|S_{i}\right\|_{2}=\left\|p_{i}-p_{j}\right\|_{2}$. In order to complete this spatial embedding, we also aim to determine a radius $r_{i} \in \mathbb{R}_{+}^{\star}(i \in \llbracket 1, n-1 \rrbracket)$ for each segment $S_{i}$.

### 3.1.1 Relative Radii

In a first time, we do not focus on absolute but on relative radii. More precisely, we define for each segment $S_{i}(i \in \llbracket 1, n-1 \rrbracket)$ a value $\rho_{i} \in \mathbb{R}_{+}^{\star}$ which is the ratio between the radius $r_{i}$ of the segment $S_{i}$ and the radius $r_{1}$ of the segment $S_{1}$ which is defined as the only segment containing the starting point $p_{0}$. In particular, for any segment $S_{i}$ we have

$$
\begin{equation*}
r_{i}=\rho_{i} \cdot r_{1} . \tag{1}
\end{equation*}
$$

These values $\rho_{i}$ are defined recursively in a bottom-up fashion by

$$
\rho_{i}= \begin{cases}1 & \text { if } i=1,  \tag{2}\\ \beta_{i} \cdot \rho_{\varphi(i)} & \text { if } i>1,\end{cases}
$$

where, by abuse of notation, $j=\varphi(i)$ means $v_{j}=\varphi\left(v_{i}\right)$. For any vertex $v_{i} \in \mathcal{V}(i \in \llbracket 2, n-1 \rrbracket)$, we set

$$
\begin{align*}
& \beta_{i}=\left(1+\alpha_{i}^{\gamma}\right)^{-\frac{1}{\gamma}},  \tag{3}\\
& \alpha_{i}=\left(\frac{L_{b(i)}}{L_{i}} \cdot \frac{R_{b(i)}}{R_{i}}\right)^{\frac{1}{4}}, \tag{4}
\end{align*}
$$

with $b(i) \neq i$ being the unique index such that $\varphi(b(i))=\varphi(i)$, and

$$
\begin{align*}
L_{i} & =\left(1-\frac{\left|\Phi\left(v_{i}\right)\right|}{2}\right)+\sum_{v_{j} \in \Phi\left(v_{i}\right)} L_{j},  \tag{5}\\
R_{i} & =\kappa \cdot \ell_{i}+\left(\sum_{v_{j} \in \Phi\left(v_{i}\right)} \frac{\beta_{j}^{4}}{R_{j}}\right)^{-1} . \tag{6}
\end{align*}
$$

In particular, these equations involve only two (hyper)parameters, namely $\gamma$ and $\kappa$, that are discussed in Section 5.

It is important to notice that despite the mutually recursive links that exist between some of the values $(R, \beta, \alpha)$, there is no deadlock due to a putative circular definition. More precisely, in Equations (3-4), the values of $\beta$ and $\alpha$ depend on the value of $R$ at the same level, whereas in Equation (6), the value of $R$ at the current level depends on the values of $R$ and $\beta$ (and thus $\alpha$ ) at the lower level of the children nodes ( $\Phi$ ). The computation of these values is then non-ambiguous and can be carried out in a bottom-up fashion, from the leaves of the tree to its root.

The meaning of the computed values is the following:

- $\rho_{i}$ is the radius ratio between segments $S_{i}$ and $S_{1}$;
- $\beta_{i}$ is the radius ratio between the segments $S_{i}$ and the "parent" $S_{\varphi(i)}$;
- $\alpha_{i}$ is a parametric ratio between the "brother" segments $S_{i}$ and $S_{b(i)}$;
- $L_{i}$ is the number of ending points in the subtree starting at segment $S_{i}$;
- $R_{i}$ is a value that models the hydraulic resistance in the segment $S_{i}$, viewed as a vascular structure.


### 3.1.2 Absolute Radii

The radius of the segment $S_{1}$ evolves during the iterative part of Algorithm 1. In particular at iteration $i \in \llbracket 1, k-1 \rrbracket$ (i.e. at a stage where $i+1$ terminal vertices have been set), the radius $r_{1}$ is defined as

$$
\begin{equation*}
r_{1}=\left(\xi \cdot R_{1} \cdot(i+1)\right)^{\frac{1}{4}}, \tag{7}
\end{equation*}
$$

where $\xi$ is a parameter that will be discussed in Section 5. The radii of the other edges are then obtained recursively in a top-down fashion from Equations (1-6).

### 3.1.3 Iterative Computation and Computational Cost

Let us suppose that an intermediate tree $\mathfrak{T}$ has been built at iteration $i-1 \in \llbracket 1, k-1 \rrbracket$ of Algorithm 1 and that the values $R_{\star}, L_{\star}, \alpha_{\star}, \beta_{\star}$ and $\rho_{\star}$ have been computed for this tree $\mathfrak{T}$.

Now, let us suppose that at the iteration $i$, we incrementally update this intermediate tree $\mathfrak{T}$. Since $R_{\star}, L_{\star}, \alpha_{\star}$ and $\beta_{\star}$ are defined in a bottom-up fashion, they only need to be updated for the new two vertices $v_{2 i}$ and $v_{2 i+1}$ added at iteration $i$, the two vertices $v_{j}$ and $v_{k}$ incident to the edge $e_{j}$ that is split at this iteration, and all the vertices located in the unique path between $v_{2 i}$ and the starting vertex $v_{0}$. For a tree $\mathfrak{T}$ correctly balanced, these updates present a time complexity $\Theta(\log i)$. However, all the values $\rho_{\star}$, defined in a top-down fashion, have to be updated, which leads to a time complexity $\Theta(i)$.

As a consequence, the computation of the radii of the final tree $\mathfrak{T}$ ex nihilo presents a time complexity $\Theta\left(k^{2}\right)$.

### 3.2 Segment Construction

Let us now discuss the way to define a new segment in the tree being constructed. At step $i$ of Algorithm 1, an intermediate tree $(\mathcal{V}, \mathcal{E})$ has been built and we now want to define and add two new points $p_{2 i}$ and $p_{2 i+1}$ in $\mathbb{R}^{d}$, and the induced segment $S_{2 i+1}$. One of these points, say $p_{2 i+1}$, is chosen randomly (but under certain constraints to be fulfilled, see Section 3.3). The other point, say $p_{2 i}$,
is defined from the knowledge of $p_{2 i+1}$ and from the position of two points $p_{j}$ and $p_{k}$ that form the segment $S_{j}$ in the current tree.

In practice, we have as input three points, namely $p_{2 i+1}, p_{j}, p_{k}$ such that the last two points already form a segment of the current tree. We aim to compute a fourth point $p_{2 i}$ that will allow us to define three segments: one segment between $p_{2 i}$ and $p_{2 i+1}$ (i.e. the "new" segment) and two segments between $p_{2 i}$ and $p_{j}$ (resp. $p_{k}$ ). These segments will derive from the splitting of the existing segment $S_{j}$ between $p_{j}$ and $p_{k}$ and its "bending" to allow the connection with the "new" segment, see Figure 1 for an illustration.

The point $p_{2 i}$ must lie in the triangle formed by $p_{2 i+1}, p_{j}$ and $p_{k}$. Its position is defined as the result of an optimization process based on the Kamyia algorithm [6], summarized as follows.

For the sake of concision, in the sequel, we note $x_{0}=p_{k}, x_{1}=p_{j}, x_{2}=p_{2 i+1}$ and $x=p_{2 i}$. The expected solution $x$ lies in the plane defined by the three points $x_{i}(i \in \llbracket 0,2 \rrbracket)$, assumed non-colinear. We can then consider this optimization problem as a 2-dimensional problem expressed in this plane isomorphic to $\mathbb{R}^{2}$. From now on, we then assume without loss of generality that we have $x_{i} \in \mathbb{R}^{2}$.

Next, we describe one step of the (iterative) Kamyia optimization algorithm.
At the beginning of the step, we know the current position of $x$, obtained from the previous step, or initialized at the first step as

$$
\begin{equation*}
x=\frac{x_{0}+x_{1}}{2} . \tag{8}
\end{equation*}
$$

For each $i \in \llbracket 1,2 \rrbracket$, we define the length $l_{i}$ of the segment between $x_{i}$ and $x$ as $l_{i}=\left\|x_{i}-x\right\|_{2}$ and we set

$$
\begin{equation*}
\Delta_{i}=\frac{f_{0} \cdot l_{0}}{r_{0}^{4}}+\frac{f_{i} \cdot l_{i}}{r_{i}^{4}} \tag{9}
\end{equation*}
$$

where $f_{i}$ corresponds to the flow ${ }^{4}$ in the segment between $x_{i}$ and $x$, and $r_{i}$ is the radius of this segment (defined in the previous sections, see Equation (1)). This formula derives from Hagen-Poiseuille's law; in particular, $\Delta_{i}$ represents a drop of pressure. When computing the $\Delta_{i}$, both the flows $f_{i}$ and the lengths $l_{i}$ are known (the $l_{i}$ are derived from the current position of the points $x_{i}$ ). The radii $r_{i}$ are obtained from the previous step (see below), except at the initial step of the process, where $r_{0}=r_{1}=r_{2}$ are set from the current tree.

We assume [15] that

$$
\begin{equation*}
f_{i} \propto r_{i}^{3} \tag{10}
\end{equation*}
$$

and [6]

$$
\begin{equation*}
r_{0}^{6}=f_{0} \cdot\left(\frac{r_{1}^{6}}{f_{1}}+\frac{r_{2}^{6}}{f_{2}}\right) . \tag{11}
\end{equation*}
$$

Equations (9) and (11) lead to a two-equation non-linear system

$$
\left\{\begin{array}{l}
\Delta_{1} r_{1}^{4}\left(f_{0}\left(\frac{r_{1}^{6}}{f_{1}}+\frac{r_{2}^{6}}{f_{2}}\right)\right)^{\frac{2}{3}}-f_{0} l_{0} r_{1}^{4}-f_{1} l_{1}\left(f_{0}\left(\frac{r_{1}^{6}}{f_{1}}+\frac{r_{2}^{6}}{f_{2}}\right)\right)^{\frac{2}{3}}=0,  \tag{12}\\
\Delta_{2} r_{2}^{4}\left(f_{0}\left(\frac{r_{1}^{6}}{f_{1}}+\frac{r_{2}^{6}}{f_{2}}\right)\right)^{\frac{2}{3}}-f_{0} l_{0} r_{2}^{4}-f_{1} l_{1}\left(f_{0}\left(\frac{r_{1}^{6}}{f_{1}}+\frac{r_{2}^{6}}{f_{2}}\right)\right)^{\frac{2}{3}}=0,
\end{array}\right.
$$

where the two unknowns are $r_{1}^{2}$ and $r_{2}^{2}$ (the system is then of degree 4). To solve this system, we can use a non-linear solver such as the C++ library Ceres Solver [1], considered in our implementation.

One may then derive the coordinates of $x[6]$ as

$$
\begin{equation*}
x=\frac{\sum_{i=0}^{2} \frac{r_{i}^{2}}{l_{i}} \cdot x_{i}}{\sum_{i=0}^{2} \frac{r_{i}^{2}}{l_{i}}} . \tag{13}
\end{equation*}
$$

[^2]This process allows to compute, at each step, the values $l_{i}$ from the current position of $x$, then $r_{i}$ (Equation (12)) and finally the new position of $x$ (Equation (13)). This iterative procedure terminates when the convergence tolerance of the gradient is reached. (In our implementation this tolerance is set to $10^{-10}$.)

### 3.3 Constraints

### 3.3.1 Preliminary Remark: Evolution of the Size of the Domain

The domain $\Omega \subset \mathbb{R}^{d}$ has a given size $|\Omega|$ (area for $d=2$ and volume for $d=3$ ). If we consider that the tree has $k$ ending points, then the influence area of each ending point is defined as $A=\frac{|\Omega|}{k}$. Since these ending points are added incrementally (one per iteration), we assume that the domain grows at the same rate, i.e. at iteration $i \in \llbracket 0, k-1 \rrbracket$ the domain $\Omega_{i}$ is defined as

$$
\begin{equation*}
\Omega_{i}=\left\{\left.\frac{i+1}{k} \cdot q \right\rvert\, q \in \Omega\right\} \tag{14}
\end{equation*}
$$

where $\frac{i+1}{k}$ is a scale factor between $\Omega_{i}$ and the targeted final domain $\Omega$. Note that the iteration 0 corresponds to the initialization step whereas the last iteration $i=k-1$ corresponds to the end of the process, with $\Omega_{k-1}=\Omega$. In this context, the influence area of an ending point $p$ can be seen as a Euclidean ball of size $A$. If $d=2$ (resp. $d=3$ ), this ball has a radius equal to $\left(\frac{A}{\pi}\right)^{\frac{1}{2}}$ (resp. $\left.\left(\frac{3 A}{4 \pi}\right)^{\frac{1}{3}}\right)$. It is important to note that when going from iteration $i$ to iteration $i+1$, this radius remains constant; however, the positions of the points evolve from $p$ at iteration $i$ to $\left(1+\frac{1}{i+1}\right) \cdot p$ at iteration $i+1$, at the same rate as the evolution of $\Omega_{i}$ to $\Omega_{i+1}$. This growing rate impacts the length $\ell_{i}$ of the segments $S_{i}$, but also their radius $r_{i}$.

### 3.3.2 Distance Constraints

At each iteration $i$ of the process, two new points $p_{2 i}, p_{2 i+1}$ are defined. One of them, say $p_{2 i+1}$, is an ending point and is defined quasi randomly. More precisely, it is chosen randomly but under various constraints:

- $p_{2 i+1}$ must be chosen within the domain $\Omega$;
- $p_{2 i+1}$ must be sufficiently far from the current tree (and a fortiori not in the tree).

In order to satisfy the first constraint, it is sufficient to check that $p_{2 i+1} \in \Omega$. If $\Omega$ is a convex domain defined by $m$ hyperplanes, this can be checked in $\Theta(m)$. If $\Omega$ is an arbitrary (convex or non-convex) domain, then a sufficient condition is to check that $p_{2 i+1}$ belongs to a cover of convex sets that lies inside $\Omega$. It is of course relevant to choose a cover that fits as much as possible $\Omega$. These convex sets can be e.g. a cover of spheres or a partition of cubes (e.g. voxels). (This last strategy was chosen in our implementation.)

In order to satisfy the second constraint, we compute the distance $\delta$ of $p_{2 i+1}$ to the tree, i.e. to each segment $S$ of the tree (considering its radius $r(S)$ ). This distance $\delta$ must be greater that $\delta_{\max }$ with

$$
\begin{equation*}
\delta_{\max }=\left(\frac{1}{i+1} \frac{|\Omega|}{k}\right)^{\frac{1}{d}} \tag{15}
\end{equation*}
$$

Note that, at step $i$, this test requires to compute $i+1$ point-to-segment distances (to this end, we will use the same constant-time algorithm as for segment-to-segment distance [11], see below), leading to a time cost $\Theta(i)$. If no random choice fulfills this constraint after a certain number of attempts (set to 100 in our implementation), then the distance $\delta_{\max }$ is decreased as $\alpha \cdot \delta_{\max }$ with $0<\alpha<1$ ( $\alpha=0.9$ in our implementation), and this process is then iterated as many times as required.

### 3.3.3 Non-intersection Constraints

Once a new ending point $p_{2 i+1}$ has been defined in $\Omega$, three segments have to be computed: one in order to connect $p_{2 i+1}$ to the remainder of the tree, and two by splitting a segment already present in the tree (see Section 3.2). The optimization process leading to the determination of these three segments does not check their validity with respect to the domain $\Omega$ and to the current tree $\mathfrak{T}$.

To tackle the issue of the domain, a solution is to build a distance map of the border $\partial \Omega$ into the domain $\Omega$. Such a distance map can be built in a discrete way on the Cartesian grid $\mathbb{Z}^{d}$ associated to the part of the Euclidean space $\mathbb{R}^{d}$ where $\Omega$ lies. Once computed, it is then possible to determine the (finite) set of pixels / voxels where the considered segment lies. If the distance map value at each of these pixels / voxels is greater than the radius of the segment plus $\sqrt{d}$, then one can guarantee that the segment does not intersect the border $\partial \Omega$ of the domain, i.e. it is actually inside $\Omega$.

It is also mandatory that a new (or updated) segment $S$ does not intersect the other segments, i.e. that there is no self-intersection within the tree. This can be done by computing, for each segment $S^{\prime}$ in the tree (except the segments incident to $S$ ) the distance $D$ between $S$ and $S^{\prime}$, namely

$$
\begin{equation*}
D\left(S, S^{\prime}\right)=\min \left\{\left\|x-x^{\prime}\right\|_{2} \mid\left(x, x^{\prime}\right) \in S \times S^{\prime}\right\} \tag{16}
\end{equation*}
$$

Such a distance can be computed in constant time $\Theta(1)$ [11], leading to a checking in linear time cost $\Theta(2 i)$ with respect to the number of segments within the tree. In particular, if for all the segments $S^{\prime}$, the distance $D\left(S, S^{\prime}\right)$ is greater that the sum of the radii of $S$ and $S^{\prime}$, then we can guarantee that the new (or updated) segment $S$ does not intersect the remainder of the tree.

Figure 2 shows the whole process of vascular tree construction.

## 4 Global View of One Iteration of the Algorithm

In this section, we describe the whole running of one iteration (say iteration $i$ ) of the tree construction algorithm. At this stage, we assume that an intermediate version of the tree $\mathfrak{T}$ has been built, and we aim to add a new segment to this tree with all the induced side effects.

### 4.1 Evolution of the Domain Size

Before adding a new branch and a new ending point to the tree, the size of the domain is increased, as discussed in Section 3.3.1. This step has a time complexity $\Theta(i)$ if the coordinates of all the points are updated (i.e. the size of the domain is increased), or $\Theta(1)$ if the "size of the space" is decreased, which means that the physical positions of the points remain unchanged, but a scale factor is applied, that impacts the computation of the length. This last choice has been chosen in our implementation.

### 4.2 New Ending Point

A new ending point $p_{2 i+1}$ is defined. As discussed in Section 3.3.2, this is a random choice under various constraints. Many attempts may then be carried out, and the time complexity of this step is then $\Theta\left(i \cdot \nu_{p}\right)$ where $\nu_{p}$ is equal to this number of attempts.

### 4.3 Choice of the Candidate Segments for Connection

The creation of a new segment is carried out by connecting $p_{2 i+1}$ to an existing segment $S$ of the tree (that will be split into two segments). In theory, all the segments of the current tree may be considered. However, such an exhaustive strategy would lead to a prohibitive computational cost. In


Figure 2: Workflow of the proposed method of vascular tree construction.
practice, only the nearest segments are investigated. The definition of these $\nu_{S}$ segments is carried out in linear time $\mathcal{O}\left(i \cdot \log \nu_{S}\right)$ via point-to-segment distance computation.

### 4.4 Segment Generation and Connection

For each segment $S$ within the $\nu_{S}$ candidate segments, a connection between the point $p_{2 i+1}$ and the segment $S$ is carried out. This process, documented in Section 3.2, leads to the proposal of a new segment and the update of $S$ as two segments. This proposal is valid if and only if:

- these three segments satisfy the non-intersection requirements discussed in Section 3.3.3. This test of intersection is performed once for a new terminal point (before the optimization process of bifurcation), and during the optimization process of bifurcation;
- a ratio constraint that imposes that for each segment, the ratio length / radius of the segment be greater than 2 (i.e. the segment is "long enough"): $2 r_{i} \leq l_{i}$.

Remark: If the optimization process does not converge or if the obtained bifurcation is not valid, then the candidate segment is discarded. If all the $\nu_{S}$ candidate segments are discarded, then the number $\nu_{S}$ is doubled as many times as necessary. Note that the overall complexity of this step also depends on the complexity of the non-linear solver computation.

### 4.5 Radius Update, Volume Computation and Final Choice

For each successful connection, an updated tree is obtained. The radii of the segments of this tree are updated, based on the procedure documented in Section 3.1. It is then possible to compute the whole volume of the tree as the sum of the volumes of each segment. The tree, which minimizes this volume, is chosen as the new current tree at the end of the iteration.

## 5 Parameters

The proposed algorithm involves various (hyper)parameters introduced in the previous sections. In practice, the purpose is to generate trees that present realistic geometry and morphology with regard to physiological properties. In this context, the tree is seen as a vascular network where blood flows from one unique input (the starting point) towards the $k$ outputs (the ending points).

### 5.1 Hyperparameters

The proposed algorithm relies on the following hyperparameters:

- $\kappa=\frac{8 \mu}{\pi}$, where $\mu$ is a physiological parameter defined in the next subsection;
- $\xi=\frac{Q_{\text {out }}}{P_{\text {in }}-P_{\text {out }}}$, where $Q_{\text {out }}$ and $P_{\text {in }}, P_{\text {out }}$ are flow and pressure values defined in the next subsection.


### 5.2 Physiological Parameters

The algorithm also relies on the following physiological parameters:

- $\mu$ is the viscosity of the fluid in the structure (in the case of the blood, $\mu$ is set to 3.6 cP );
- $\gamma$ is the Murray bifurcation exponent that controls the geometry of the bifurcations (in general, $\gamma$ is set to 3 );
- $Q_{\text {out }}$ is the flow (assumed the same) at each of the $k$ ending points (in our experiments, we set $Q_{\text {out }}$ as $0.125 \mathrm{~mL} / \mathrm{min}$ );
- $P_{\text {out }}$ is the pressure (assumed the same) at each of the $k$ ending points (in our experiments, we set $P_{\text {out }}$ as 60 mmHg );
- $P_{\text {in }}$ is the pressure at the starting point (in our experiments, we set $P_{\text {in }}$ as 100 mmHg ).

Note that, without loss of generality, the pressure, flow and viscosity parameters are expressed in $\mathrm{Pa}, \mathrm{mm}^{3} / \mathrm{s}$ and $\mathrm{Pa} \cdot \mathrm{s}$ in our implementation, respectively.

## 6 Code and Implementation Details

In this section, we provide information about the principal data structure considered in our implementation of the algorithm (Subsection 6.1), some pseudo-codes related to the most important parts of the algorithm (Subsection 6.2) and a brief source-code description (Subsection 6.3).

### 6.1 Data Structure of a Vascular Tree

The data structure of a vascular tree is given in the class CoronaryArteryTree. Each segment of the vascular tree is defined by the structure Segment. It contains the following attributes:

- Distal point of the segment $\left(x_{j}\right):$ myCoordinate;
- Index of the segment ( $j$ ): myIndex;
- Radius of the tubular section $\left(r_{j}\right)$ : myRadius;
- Total number of terminal segments in its children segment $\left(L_{j}\right)$ : myKTerm;
- Hydrodynamic resistance $\left(R_{j}\right)$ : myResistance;
- Flow $\left(f_{j}\right)$ : myFlow;
- Relative radii (radius ratio between the segment and its parent, $\rho_{j}$ ): myBeta.

A vascular tree is then composed of:

- A vector of terminal segments: myVectTerminals;
- A vector of all the segments of the tree $(\mathcal{E})$ : myVectSegments;
- A vector of the left (first) and right (second) segments of each indexed segment ( $\Phi$ ): myVectChildren;
- A vector of the parent of each indexed segment $(\varphi)$ : myVectParent.

Global biological attributes are defined by the structure BioAlgoParameter:

- Number of terminal segments ( $k$ ): my_NTerm;
- Perfusion area / volume $(|\Omega|)$ : my_aPerf.
- Final radius of circular area $(|\Omega| / k)$ : my_rPerf;
- Flow $\left(Q_{\text {out }}\right)$ : my_qPerf;
- Perfusion pressure $\left(P_{\text {in }}\right)$ : my_pPerf;
- Pressure of terminal segment $\left(P_{\text {out }}\right)$ : my_pTerm;
- Bifurcation exponent $(\gamma)$ : my_gamma;
- Viscosity ( $\mu$ ): my_mu;
(Note that 7 of these 8 attributes are parameters, whereas the 8 th one, namely my_rPerf, is calculated from the others.)

Internal algorithm parameters are defined by the structure InternAlgoParameter:

```
Algorithm 2: Generate randomly a point with distance constraints
\(\rightarrow\) see C ++ code: generateALocation() of class CoronaryArteryTree in package src
```

```
Input: A distance threshold myDThreshold
```

Input: A distance threshold myDThreshold
Input: A vector of terminal points myVectTerminals of the current tree
Input: A vector of terminal points myVectTerminals of the current tree
Input: A vector of segments myVectSegments of the current tree
Input: A vector of segments myVectSegments of the current tree
Output: A generated point $p$
Output: A generated point $p$
Output: A Boolean $i s C o m p$ indicating whether the generation has succeeded
Output: A Boolean $i s C o m p$ indicating whether the generation has succeeded
$p \leftarrow$ generate a random point
$p \leftarrow$ generate a random point
// Generated point must have a certain distance to all tree terminals
// Generated point must have a certain distance to all tree terminals
$i s C o m p \leftarrow$ true
$i s C o m p \leftarrow$ true
$i d \leftarrow 1$
$i d \leftarrow 1$
while isComp and id $<$ myVectTerminals.size do
while isComp and id $<$ myVectTerminals.size do
isComp $\leftarrow$ distance $($ myVectTerminals $[i d]$. myCoordinate,$p)>m y D T h r e s h o l d$
isComp $\leftarrow$ distance $($ myVectTerminals $[i d]$. myCoordinate,$p)>m y D T h r e s h o l d$
$i d \leftarrow i d+1$
$i d \leftarrow i d+1$
// Generated point must have a certain distance to all tree segments radii
// Generated point must have a certain distance to all tree segments radii
$i d \leftarrow 1$
$i d \leftarrow 1$
while isComp and id < my VectSegments.size do
while isComp and id < my VectSegments.size do
$i s C o m p \leftarrow$ distance $($ myVectSegments $[i d], p)>$ myVectSegments $[i d]$. myRadius
$i s C o m p \leftarrow$ distance $($ myVectSegments $[i d], p)>$ myVectSegments $[i d]$. myRadius
$i d \leftarrow i d+1$;
$i d \leftarrow i d+1$;
return $(p, i s C o m p)$

```
    return \((p, i s C o m p)\)
```

- Current number of terminal segments: myKTerm;
- Number of nearest neighbors to be tested: myNumNeighbor;
- Average radius of blackboxes: myRsupp;
- Scale factor: myLengthFactor.


### 6.2 Pseudo-codes

In this subsection, we provide some pseudo-codes related to the following tasks:

- Generation of the new points in the tree (Algorithms 2 and 3);
- Intersection checking of the branches in the tree (Algorithms 4 and 5);
- Whole process for creating a new bifurcation (Algorithm 6).


### 6.3 Source Codes and Dependencies

The code of OpenCCO is written in $\mathrm{C}++$. It is composed of the following classes in the src directory:

- CoronaryArteryTree contains the structure of a vascular tree and functions to create a bifurcation;
- DomainController contains the construction of the domain of a vascular tree;
- helpers/GeomHelpers contains functions to generate randomly terminal points, check intersections, Kamyia optimization, ...;
- helpers/ExpandTreeHelpers contains functions to generate a vascular tree;

```
Algorithm 3: Generate a terminal point
\(\rightarrow\) see C++ code: generateNewLocation() of class CoronaryArteryTree in package src
    Input: Number of maximum trials nbTrials ( \(=1000\) by default)
    Output: A generated point \(p\)
    myDThreshold \(\leftarrow\) compute the distance constraint // See Section 3.3.2
    found \(\leftarrow\) false
    \(n \leftarrow\) nbTrials
    while \(\neg\) found and \(n \geq 0\) do
        \(n \leftarrow n-1\)
        (pt, found \() \leftarrow\) generateALocation \((m y D T h r e s h o l d) / /\) See Algorithm 2
        if found then
            \(\llcorner p \leftarrow p t\)
        if \(n=0\) then
            \(n \leftarrow\) nbTrials
            myDThreshold \(\leftarrow\) myDThreshold \(* 0.9\)
    return \(p\)
```

```
Algorithm 4: Check if there is an intersection between two thick segments
\(\rightarrow\) see C ++ code: isIntersecting () of class GeomHelpers in package src/helpers
    Input: Two thick segments given by their endpoints and thickness \((\operatorname{seg} A, \operatorname{seg} B, r A B)\) and \((\operatorname{seg} C, \operatorname{seg} D, r C D)\)
    Output: A Boolean indicating whether they are intersecting
    \(c A B \leftarrow(\operatorname{seg} A+\operatorname{seg} B) / 2\)
    \(c C D \leftarrow(\operatorname{seg} C+\operatorname{seg} D) / 2\)
    \(l A B \leftarrow(\operatorname{seg} A-\operatorname{seg} B) . n o r m\)
    \(l C D \leftarrow(\operatorname{seg} C-\operatorname{seg} D)\).norm
    \(d C \leftarrow(c A B-c C D)\).norm
    \(d \leftarrow \frac{(l A B+l C D)}{2}+r A B+r C D\)
    if \(d C>d\) then
    return false
    distanceSeg \(\leftarrow\) distance between the two segments \([\operatorname{seg} A, \operatorname{seg} B]\) and \([\operatorname{seg} C, \operatorname{seg} D]\)
    if distanceSeg \(>(r A B+r C D)\) then
        return false
    return true
```

- helpers/XMLHelpers contains functions to export a vascular tree into XLM file.

The program (and the online demo) takes into account the following parameters:

- Options for both 2D and 3D cases:
- number of terminal segments/ending points (-n | -nbTerm INT, default=1000)
- perfusion area / volume (-a | -aPerf FLOAT, default=20000)
- value of the gamma parameter (-g | -gamma FLOAT, default=3)
- minimal distance to border (-m|-minDistanceToBorder FLOAT, default=5)
- organ domain using a mask image (-d $\mid$-organDomain TEXT)
- output the resulting graph as xml file (-x|-exportXML TEXT)
- use a squared implicit domain instead of a sphere (-s |-squaredDom, defaut=sphere)
- Specific options in 2D:

```
Algorithm 5: Check if there is an intersection between a thick segment and all segments of
the current tree, except certain segments
\(\rightarrow\) see C ++ code: isIntersectingTree () of class CoronaryArteryTree in package src
```

```
Input: A thick segment given by two points and a radius \((p t A, p t B, r)\)
```

Input: A thick segment given by two points and a radius $(p t A, p t B, r)$
Input: The vector of segments myVectSegments of the current tree
Input: The vector of segments myVectSegments of the current tree
Input: A triplet idExcept containing the 3 segments indices not tested for intersection
Input: A triplet idExcept containing the 3 segments indices not tested for intersection
Output: A Boolean indicating whether there is an intersection
Output: A Boolean indicating whether there is an intersection
// If they are root segment, then ignore the test
// If they are root segment, then ignore the test
if (idExcept $[0]=0$ or $\operatorname{idExcept}[1]=0$ or $\operatorname{idExcept}[2]=0)$ then
if (idExcept $[0]=0$ or $\operatorname{idExcept}[1]=0$ or $\operatorname{idExcept}[2]=0)$ then
return false
return false
foreach $s:$ myVectSegments do
foreach $s:$ myVectSegments do
$i d \leftarrow s$.myIndex
$i d \leftarrow s$.myIndex
if $i d \neq 0$ and $i d \neq i d E x c e p t[0]$ and $i d \neq i d E x c e p t[1]$ and $i d \neq i d E x c e p t[2]$ then
if $i d \neq 0$ and $i d \neq i d E x c e p t[0]$ and $i d \neq i d E x c e p t[1]$ and $i d \neq i d E x c e p t[2]$ then
$p t C \leftarrow$ s.myCoordinate
$p t C \leftarrow$ s.myCoordinate
$p t D \leftarrow$ myVectSegments[myVectParent[id]].myCoordinate
$p t D \leftarrow$ myVectSegments[myVectParent[id]].myCoordinate
if isIntersecting(ptA, ptB,r,ptC, ptD, s.myRadius) then //see Algorithm 4
if isIntersecting(ptA, ptB,r,ptC, ptD, s.myRadius) then //see Algorithm 4
return true
return true
return false

```
    return false
```

- initial position of root (-p |-posInit INT INT, default=image center)
- output the result into EPS format (-o| -outputEPS TEXT, default=result.eps)
- output the result into SVG format (-e | -exportSVG TEXT, default=result.svg)
- Specific options in 3D:
- initial position of root (-p|-posInit INT INT INT, default=image center)
- output the 3D mesh into OFF file (-o| -outputName TEXT, default=result.off)
- output the 3D mesh into text file ( $-\mathrm{e} \mid$-export TEXT)
- display 3D view using QGLViewer (-view)

The program requires the DGtal library (1.3 or later): https://github.com/DGtal-team/DGtal.

## 7 Experimental Results and Comparison

In this section, we propose some experiments and results dedicated to compare the behaviour of our implementation with the state-of-the-art version of the 2 -dimensional and 3 -dimensional algorithm, and variants proposed in the literature.

From a qualitative point of view, we first illustrate, in Figures 3 and 4, some 2- and 3-dimensional vascular trees generated in a disk / sphere domain, by the initial algorithms and our implementation. We also provide, in Figure 5, a 3-dimensional vascular tree generated in a complex arbitrary domain.

We also illustrate, in Figure 6 the difference between a 3-dimensional vascular tree generated by VascuSynth [3] and with our proposed implementation of CCO with the same input parameters, plus a plot of the diameters of the segments depending on the bifurcation level. A quantitative assessment of the diameters of the segments depending on the bifurcation level in a non-open implementation proposed in [4], and our implementation is given in Figure 7. Finally, an analysis of the computation cost of our 2-dimensional and 3-dimensional implementation of CCO is illustrated w.r.t. implicit (i.e. spherical) / masked (i.e. arbitrary) domains and vs. VascuSynth [3], in Figure 8. These measurements were performed on a processor Apple M1 Max.

```
Algorithm 6: Create a new bifurcation from a terminal point to a near segment of the tree
\(\rightarrow\) see \(\mathrm{C}++\) code: isAddable () of class CoronaryArteryTree in package src
    Input: A new terminal point \(p\)
    Input: The index segIndex of a segment close enough to \(p\)
    Input: The maximal number of iterations, nbIter
    Input: The tolerance to assess the convergence of tree volume, tolerance
    Output: A Boolean indicating whether \(p\) can be added to the tree
    \(p\) Current \(\leftarrow\) calculate a first bifurcation from \(p\), the segment segIndex and the image domain
    Let \(s\) Parent be the segment parent of segIndex
    pParent \(\leftarrow\) sParent.myCoordinate
    Let \(s N e w L e f t\), sNewRight, sCurrent be the new left, right and middle segments // see Section 2.4
    \(r 0, r 1, r 2 \leftarrow\) sCurrent.myRadius
    vol, volCurr \(\leftarrow-1\)
    isDone \(\leftarrow\) false
    \(i \leftarrow 0\)
    while \(i<\) nbIter and \(\neg i s D o n e\) do
        res \(1 \leftarrow\) kamyiaOptimization \((p\) Current, pParent, sCurrent.myRadius, sNewLeft, sNewRight,
            1, \(p O p t, r 0, r 1, r 2) / /\) see Section 3.2 (Equations (9)-(13))
        if \(\neg\) res 1 then // Kamyia does not lead to a solution
            return false
        // Check intersection with the current tree for the middle segment
        idSegPar \(\leftarrow\) myVectParent[segIndex]
        \((i d L, i d R) \leftarrow(\) myVectChildren[idSegPar].first, myVectChildren[idSegPar].second \()\)
        res \(2 \leftarrow\) isIntersectingTree( \(p\) Opt,pParent,r0,[idL,idR,idSegPar])// see Algorithm 5
        if res2 then // There is an intersection with the middle segment
            return false
        if \(p\) Current \(\notin\) the image domain or the middle segment \(\notin\) the image domain then
            return false
        Do similar test of intersection for the left and right segments
        // Iterate for the optimization process of the new bifurcation
        Let \(c\) TreeCurr and cTree 1 be a copy of the current tree
        (sNewLeft.myRadius, sNewRight.myRadius \() \leftarrow(r 1, r 2) / /\) Update values for the segments
        cTree1.add(sNewLeft)
        cTree1.add(sNewRight) // Add the left and right segments to cTree1
        cTree1.update() // Update parameters of cTree1: new segments, physiological parameters...
        vol \(\leftarrow c\) Tree1.computeTotalVolume ()\(/ /\) Compute volume of the current tree
        if \(i=0\) then // The first computation of volume
            volCurr \(\leftarrow\) vol
            \(c\) TreeCurr \(\leftarrow c\) Tree 1
        else
            if \(\mid\) volCurr - vol \(\mid<\) tolerance then
            // Check that the resulting segments are long enough (see Section 4.4)
                Let \(l 0, l 1, l 2\) be the length of the segments [ \(p O p t, p\) Parent \(]\), \([p O p t, s N e w L e f t . m y C o o r d i n a t e]\),
                    [pOpt, sNewRight.myCoordinate]
            \(r e s 3 \leftarrow(2 * r 0) \leq l 0\) and \((2 * r 1) \leq l 1\) and \((2 * r 2) \leq l 2\)
            if res3 then // If there is a solution, then save the result
                    isDone \(\leftarrow\) true
                    save cTree 1
            else
            volCurr \(\leftarrow\) vol
            \(c\) TreeCurr \(\leftarrow c\) Tree 1
        Update tree parameters: new position and physiological parameters (see Equations (1-6))
        \(i \leftarrow i+1\)
    return isDone
```



Figure 3: 2-dimensional network generated in a disk domain. The number of terminal points is set to 4000 . This result can be compared to the one depicted in [17, Figure 3, p. 486].


Figure 4: 3-dimensional networks generated in a sphere domain. The number of terminal points is set to 1000 (a) and 4000 (b). These results can be compared to those depicted in [7, Figure 5, p. 31]


Figure 5: 3-dimensional network generated in an arbitrary shape domain. Here, the domain (a) is composed by the nonconvex thick boundary of the Stanford bunny (a topological hollow sphere with holes). The domain (c) is defined by the same construction but by cutting the front mesh to increase visibility. The number of terminal points in the network is set to 20000 in (b) and 3000 in (d).


Figure 6: 3-dimensional networks generated in a cubic domain: VascuSynth (a) and our implementation of CCO (b). (c) Comparison of the segment diameters variation according to the bifurcation level between the VascuSynth reconstruction and our implementation of CCO.


Figure 7: Averaged segment diameters at respective bifurcation levels. (a) Results from [4]: "Radius study along bifurcation levels: mean and standard deviation at each bifurcation level on 10 simulated trees" (text from [4]). (b-c) Results from our implementation in 3-dimensions (b) and 2-dimensions (c). These results can also be compared to those depicted in [17, Figure 4, p. 487]. The behaviours of the curves are comparable, up to the different parameters which may induce distinct quantitative values.


Figure 8: Comparison of computation time: (a) CCO in 2D and (b) 3D with implicit and masked domains, (c) CCO vs. VascuSynth in 3D with implicit domain.

All these experimental results argue in favour of the homogeneity and the correctness of the behaviour of our implementation with the native algorithms proposed in [7] and [17].

## 8 Conclusion and Perspectives

In this article we proposed an implementation of the CCO algorithm, initially designed in [17] for the 2-dimensional case and in [7] for the 3-dimensional case. We did our best to provide an implementation that is as close as possible to these original algorithms.

We believe that our implementation could be of precious use for any researcher interested in obtaining realistic vascular networks (or similar tree-based objects). In particular, it may constitute a useful alternative to existing softwares, such as VascuSynth [3].

It could be possible to build upon this framework in order to propose complementary functionalities that could tackle some recurrent issues that occur in vascular modeling. First, the approach could be extended to generate complementary networks (e.g. arterial and venous trees), which would require the handling of distinct, independent trees in a forest structure, whereas considering the constraints that each tree imposes to the other(s). Another important perspective would concern the ability to design a partial network composed of many branches that constitute the extension of ending points of the vascular structures at a given scale. Such approach would allow to build some vascular models obtained from segmentation at the lowest scales (e.g. from patient-specific analysis of medical images), and from modeling at the highest scales, where the images do no longer provide useful information, but where physiological behaviour from realistic models could be statistically inferred. Such optimizations could find inspiration in recent works dedicated to extend / build upon CCO or similar algorithms, e.g. [12, 8].

In the current version of the algorithm, the output flow is the same for each terminal point of the tree, whereas a uniform probability is considered to determine the putative position of these ending points. More versatile strategies could be developed to allow non-homogeneous output flows and biases in the positioning of the ending points, e.g. for better modeling phenomena such as angiogenesis.

From a more technical point of view, some octree-like hierarchical representations of the tree and of the vascular domain may also be considered for optimizing the computational burden induced by the assessment of the geometrical hypotheses related to the tree-tree and tree-domain constraints.

## Image Credits

Stanford bunny ${ }^{5}$ from Turk and Levoy at Stanford University under two-clause BSD license.
Thickened surface from Stanford bunny by Kerautret, Ngo, Passat, Talbot, Jaquet, license CC BY-SA.

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[^0]:    ${ }^{1}$ https://doi.org/10.5201/ipol.2023.477
    ${ }^{2}$ https://github.com/OpenCCO-team/OpenCCO

[^1]:    ${ }^{3}$ https://vascusynth.cs.sfu.ca

[^2]:    ${ }^{4}$ In this algorithm, we assume that the flow at each ending points is the same. Then, the values of $f_{i}$ considered here are proportional to the number of ending points in the subtrees starting at $x_{i}$. In particular, we have $f_{0}=f_{1}+f_{2}$, and $\frac{f_{1}}{f_{2}}=L_{j}$.

[^3]:    ${ }^{5}$ https://faculty.cc.gatech.edu/~turk/bunny/bunny.html

