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A COMPUTATIONAL TEMPLATE FOR 3D MODELLING OF THE VASCULAR SCAFFOLD OF THE HUMAN THYROID GLAND

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ABSTRACT

We recently designed an innovative scaffold-bioreactor unit for the bioengineering of a three-dimensional (3D) bioartificial human thyroid gland or its miniaturized replica as a part of a microfluidic chip test system. This device is based on the evidence that the 3D geometry of the intraglandular stromal / vascular scaffold or SVS (i.e. the fibrous and vascular matrix) of mammalian viscera plays a key role in guiding growth and differentiation of *in vitro* seeded cells. Therefore, we initiated a research program focused on computer-aided reconstruction of the 2nd to 4th order intralobar arterial network (IAN) of the human thyroid gland as a reliable surrogate for its 3D SVS, to be used as an input for rapid prototyping of a biomaterial replica.

To this end, we developed a computational template that works within the *Mathematica* environment, giving rise to a quasi-fractal growth of the IAN distribution, constrained within an approximation of the thyroid lobe shape as a closed surface. Starting from edge detection of planar images of real human thyroid lobes acquired by *in vivo* real-time ultrasonography, we performed data approximation of the lobar profiles based on splines and Bezier curves, providing 3D lobar shapes as geometric boundaries for vessel growth by a diffusion limited aggregation model. Our numerical procedures allowed for a robust connection between development of lobar arterial trees and thyroid lobe shape, led to a vascular self-similarity consistent with that of a cadaveric lobar arterial cast, and reproduced arterial vessels in a proportion not statistically different from that described for the real human thyroid gland.

We conclude that our algorithmic template offers a reliable reproduction of the extremely complex IAN of the adult human thyroid lobe, potentially useful as a computational guidance for bioprinting of thyroid lobe matrix replicas. In addition, due to the simplicity and limited number of morphometrical parameters required by our system, we predict its application to the design of a number of patient-tailored human bioartificial organs and organs-on-chip, including parenchymal viscera and bones.

IMPACT STATEMENT: The study introduces the computer simulation of the 3D intrinsic vascular matrix of the human thyroid gland, offering a general concept applicable to a number of other human viscera. Indeed, it provides a flexible software tool for reproduction of a 3D surrogate of the organ's 3D stromal matrix, suitable for eventual 3D bioprinting with biomaterials, and recellularization with organ-specific stem cells / progenitors. The final expectation is the design of patient-tailored, 3D organ's matrices upon clinical request.

INTRODUCTION

It is increasingly recognized that the natural organ's matrix or stromal / vascular scaffold (SVS) plays a critical regulatory role in developmental morphogenesis and related functional maturation of mammalian viscera.¹ In epithelial organs three-dimensional (3D) tissue patterning, cellular differentiation, vascular morphology, organ size, and hormonal secretory activity are topobiologically influenced by the interactions occurring between SVS and growing cells,² including the thyroid gland.^{3,4} Accordingly, we proposed that the 3D geometry of the natural SVS may act *per se* as a pivotal mechano-chemical input for growth and differentiation of thyroid stem cells / progenitor cells.^{5–9} Consistently, we showed that a natural SVS obtained by decellularization of the rat thyroid lobe drives *in vitro* self-assembly and functional differentiation of colonizing rat thyrocytes.^{6,9,10–13}

Based on animal models,^{7,14,15} recent investigations tackled the *in vitro* 3D reconstruction of human thyroid tissue. Investigations relied on the capacity of normal and cancerous, human thyroid follicular cells and thyroid-differentiated human embryonic and induced pluripotent stem cells to self-assemble into 3D organoids, either spontaneously or driven by a biocompatible 3D scaffold, including a decellularized xenogenic thyroid lobe SVS or exploiting microgravity.^{16–20} However, none of these approaches allowed for a robust 3D anatomical reconstruction of a patient-tailored, human thyroid lobe.

Therefore, we designed a cell culture bioreactor for self-assembly of a human bioartificial thyroid gland directly on the laboratory bench (i.e. *ex situ*), including its miniature replica as a part of a microfluidic chip test system. This device includes a core that replicates the 3D architecture of the adult human thyroid SVS fabricated with biomaterials. A basic assumption is that the 3D organ-like (i.e. organomorphic) geometry of the SVS may provide vascular conduits able to supply trophic media and O₂ inside the mass of stem cells / progenitor cells eventually seeded, thus favouring their growth and differentiation up to formation of a complete viable human thyroid gland organoid *ex situ*.^{5,6,9}

For this purpose, we initiated a computational and reverse engineering program to identify the best approaches for both replication of the 3D geometry of the adult thyroid SVS, and its physical reconstruction by 3D bioprinting/additive layer manufacturing and biomaterials.^{21–27} Here, we present an original core of numerical and rendering routines aimed at virtually reconstructing the intralobar arterial network (IAN) of the human thyroid, chosen as a reliable replica of its entire SVS, constrained within the limits of a thyroid lobe simulation. Preliminary analysis of few building blocks of this computational structure were presented elsewhere.^{21,22,25,27}

MATERIALS AND METHODS

Human subjects, 2D imaging of the thyroid lobe and its 3D simulation

In vivo ultrasonographic images of the human thyroid lobe were obtained from three euthyroid, control subjects (a male, two females, age range 17 - 43 years, body mass index or BMI range 22.4 - 24.1 kg/m²) without visible and/or palpable thyroid nodules, screened in one outpatient endocrine clinic (OSTEONET Unit, Galliera Medical Center / CMG - DIMEC, UNIPR, Italy) for a routine examination. Data were collected following approval by the Ethical Committee of the clinical Center, and used under patient's Informed Consent. A Toshiba Nemium Premio real-time, ultrasound machine with 9-12 MHz linear probes was used. To obtain an approximate representation of the lobe profile, a single lobe projection was chosen, specifically a bidimensional longitudinal section taken at the longest sagittal axis of the lobe. In this manner, we avoided complex computations based on a set of lobe slices, as currently occurring in 3D reconstruction of many human organs by medical rendering techniques. In addition, our approach allowed for determination of features of approximated symmetry (here coined as pseudosymmetry) of the lobe profile, with the aim of obtaining an adequate sampling of points on the lobe contour.

Finally, an analytical description of the entire simulated lobe surface was reached within the computer algebra environment of *Mathematica*,²⁸ and based on the following computational steps:

- Edge detection and spline interpolation of the thyroid lobe profile;
- Homotopy-like parametrization and 3D reconstruction of the thyroid lobe;
- Estimate of the thyroid lobe volume.

Details on each of these steps are available in Supplementary Materials.

Fractal growth of the intralobar vascular tree via Diffusion Limited Aggregation

To achieve a 3D reconstruction of the lobar SVS, we focused on the simulation of the intraglandular arterial branches, chosen as a reliable geometric replica of the thyroid SVS. To this aim, we developed an original quasi-fractal algorithm based on the mathematical principle of Diffusion Limited Aggregation or DLA, leading to a fractal growth as a binary Brownian tree.²⁹ Details are given in Supplementary Material under:

- Auxiliary DLA subroutine;
- Template of DLA subroutines.

In addition, to ensure growth of the IAN as a reliable replica of its entire SVS, constraints were imposed on the DLA to ensure that the simulated intralobar vascular tree remained within the boundaries of the reconstructed thyroid lobe, and according to specific tree branching rules. Details on this can be found in Supplementary Material under:

- Constraints of DLA growth.

Finally, the 3D fractal dimension (fd) of the simulated intralobar vascular tree was computed using an original implementation of classical box counting within the *Mathematica* scientific environment, as outlined in Supplementary Material under:

- Computation of the fractal dimension of the simulated intralobar vascular tree.

Validation procedures

Validation of the computational robustness of our IAN simulations with respect to the vascular anatomy of the human thyroid gland was performed by evaluating the 3D fd of the DLA structure with respect to an injection-corrosion cadaveric thyroid vascular cast (TVC). Briefly, the gland of a human subject (male, age 59 years, BMI 24 kg/m²) obtained

at autopsy from the Section of Forensic Medicine of the SBIBIT Department at the University of Parma (courtesy of Giovanni Lanzi), Italy, was injected through the superior (STA) and inferior (ITA) thyroid arteries with a solution of 3% polyvinyl chloride-co-vinyl acetate (Pevikon C 870, Sigma- Aldrich) diluted in acetone, to fill the IAN. After resin polymerization, the thyroid tissue was digested by immersion in a solution of 5% HCL / pepsin for 48-72 hours, leading to retrieval of the glandular IAN.^{5,8,22,25} In a second step, the TVC was scanned using the microtomographic system Skyscan 1172 (Bruker mCT, Belgium) at the nominal resolution of 30 μm , via a rotation up to 180° with 0.4° rotation step and a frame averaging of 4, and reconstructed in three orthogonal planes using the Skyscan NRecon software (version 1.6.2.0). Finally, the right lobe of the reconstructed TVC was selected, and a mean 3D *fd* calculated using the Bruker CT-Analyser software (version 1.12.0) as previously applied.²³

An additional validation was achieved by comparing the percentage of lobar arteries belonging to each vascular order between computer reconstructions and literature data on the human thyroid gland.^{30–32} In a first step, we calculated the absolute number of: a) node calibers (including roots) of 2nd order vessels; b) node calibers of 3rd order vessels; c) node calibers of 4th order vessels up to an intermediate caliber of 0.025 mm (instead of the smallest caliber of 0.0125 mm; see the caliber intervals table in Supplementary Materials). This procedure was repeated for each simulation. The need for considering an intermediate caliber was due to excessive computational time; as a consequence, the number of 4th order nodes remained incomplete. Therefore, we estimated all of them (i.e. up to caliber 0.0125 mm) by doubling the number of computed 4th order nodes. This choice was based on the heuristic that a maximum of eight applications of the Roux - Murray Law (with exponent $\delta=2.69$; see formula (5) in Supplementary Material) were needed to span the entire 4th order interval of calibers, as opposed to the four applications expected to reach the caliber 0.025 mm.

Node calculations was achieved by performing an inverse scanning of the simulated tree, whose information were stored in four arrays organized in List form as follows:

- nodeList: containing the spatial coordinates of the nodes in the cluster;
- sizeList: containing the calibers of the nodes in the cluster;
- fatherList and sonList: containing integer indexes relating each node to its sons and father up to the 2nd order root.

Then, an indexing method was developed. Specifically, this is an algorithm with memory of performed steps where each node (or bifurcation) was tagged exploiting the List structure in *Mathematica*. This allowed for identification of familial relationships including ancestor and descendant particles, and their coordinates and calibers as well. In this way, information on nodes (including 2nd order roots) belonging to each vessel order were gathered for all orders.

In a second step, we developed a logical frame to estimate the number of 3rd and 4th order vessels in our simulations as follows:

- i. the number of simulated 2nd order arteries was arbitrarily chosen at the minimum of 16 roots, corresponding to the minimal number of 2nd order arteries in a thyroid; 30–32
- ii. based on the number of computed 2nd order nodes (and roots), and having evidence that their values represented 16 real arteries (i.e. the same number arbitrarily chosen to start our simulations), we estimated that the mean number of algorithm iterations necessary to reconstruct 16 arteries per simulation was the ratio between the mean of 2nd order nodes (and roots) and the starting number of arterial roots (i.e. 16). This resulted in an average of 47 algorithm iterations per tree (equivalent to the number of 2nd order nodes / vessel) that spanned the entire caliber interval (of width equal to 0.34 mm) of 2nd order vessels;
- iii. to obtain an estimate of the unknown number of 3rd order vessels in our simulations, we reasoned that if 47 algorithm iterations spanned the 2nd order interval width (0.34 mm), then the number of algorithm iterations to span the entire caliber interval (of width equal to 0.056 mm) of 3rd order vessels could be obtained as a proportion between these values, resulting in about 8 algorithm iterations per tree on average (specifically, 7.76).

Thus, the number of 3rd order estimated vessels resulted as the ratio between the mean simulated 3rd order nodes and these 8 iterations;

iv. finally, to obtain an estimate of the unknown number of 4th order vessels in our simulations, we reasoned that if an average of 8 algorithm iterations spanned the entire 3rd order interval width (0.056 mm), then the number of algorithm iterations to span the entire caliber interval (of width equal to 0.016 mm) of 4th order vessels could be obtained as a proportion between these values, resulting in about 2 iterations per tree on average (specifically, 2.15). Thus, the number of 4th order estimated vessels resulted as the ratio between the mean estimated 4th order nodes and these 2 iterations.

In a third step, an estimate of the number of 2nd, 3rd and 4th order vessels for an average human thyroid lobe was obtained taking into account the following anatomical data: 30–34

- a) the number of 1st order arteries collectively stemming from STA and ITA (4 to 5 branches);
- b) the number of 2nd order vessels stemming from each 1st order artery (4 to 5 branches);
- c) the total number of follicles (1.2×10^6 - 32×10^6) and the number of follicles in each lobule (20-40 follicles);
- d) the anatomical evidence that, on average, each thyroid lobule is vascularized by a single 3rd order vessel;
- e) the number of 4th order vessels stemming from each 3rd order vessel (2 to 5 branches).

Using these informations: a) minimal, mean, and maximal interval values for the number of vessels in each order were determined; b) each of these intervals was used to provide the percentages of vessels in a thyroid lobe. Finally, we calculated the distance ($\Delta\%$) between mean simulated and real vascular percentage values.

Statistical analysis

Differences in the 3D *fd* (either by order or averaged) of the simulated arterial trees were determined using one-way ANOVA. An *r* Pearson's correlation coefficient was computed

for the overall mean 3D fd distribution of both simulated cases and vascular cast. Finally, differences in the distance ($\Delta\%$) between mean simulated and real vascular percentage values were calculated by initial arcsin transformation, and comparison of the transformed percentages using a z-value in a so-called simplest statistical test.³⁵ All results were considered statistically significant if $p < 0.05$.

RESULTS

Figure 1 summarizes the steps for 3D reconstruction of the thyroid IAN in the three subjects studied. Lobar simulation started from a sagittal ultrasonographic image of the right thyroid lobe. Following edge detection and spline approximation of its outlined contours, the lobe was reconstructed as a polygonal mesh, providing a closed space for growth of the IAN by the constrained DLA algorithm. Vascular simulations started from 16 vascular roots, each representing one of 16 real 2nd order thyroid arteries placed on the lobar surface keeping a reciprocal uniform distance. A progressive increase in the IAN density (from 2nd to 4th order vessels) occurred and, for mere rendering reasons, reconstructed arteries were coloured in red and blue depending on roots located above and below the barycentric plane parallel to the XY-plane, respectively. Similar to normal anatomy, superior and inferior vessel groups tended to overlap at the level of the barycentric plane and, moving from ovoid-like (case 1) to conic-like (cases 2 and 3) lobar morphologies, a diffusion of superior vessels within the upper lobe pole occurred with respect to inferior vessels segregating in the lower lobe pole.

Figure 2 depicts the values of the 3D fd of the IAN in each of the simulated cases in comparison to that of a cadaveric case. A statistically significant difference emerged in the mean value of 3D fd for 2nd order vessels with respect to mean values of 3rd and 4th order vessels, as expected for a multifractal arterial pattern. However, a close degree of self-similarity in arterial geometry was found among the three IAN reconstructions for all order vessels, leading to absence of any statistically significant difference in their mean 3D fd values. These data linearly correlated in a statistically significant manner with the mean 3D fd value of the virtual microtomographic reconstruction of the cadaveric vascular cast.

Figure 3 summarizes results of the indexing method to calculate the number of roots and nodes of 2nd order arteries, and nodes of 3rd and 4th order arteries, in the simulations of the three euthyroid control cases. It also reports the interval of values for the number of 2nd to 4th order arteries in an average thyroid gland as deduced from literature data, and the differences in the proportion (percentage) of arteries at each vascular order between simulated and real cases. Both in simulated and real cases, the number of arteries increased following a similar trend, with a vascular “explosion” (around 80 times) from 2nd to 3rd order, and a more limited ramification (around 12 times) from 3rd to 4th order. Differences in arterial percentages between simulated and real arteries were not statistically significant, confirming robustness of our algorithm in reproducing an IAN consistent with known anatomical data.

Figure 4 shows the flow-chart of the procedure for simulation of the IAN of the human thyroid lobe. Phase 1 was set to describe the surface of the 3D thyroid lobe, giving rise to a 3D numerical model graphically rendered, whose volume could be estimated. The output of phase 1 provided a primary constraint for the 3D growth of the IAN, set in phase 2. The latter represented the core of the template, finalized to the generation of the IAN using a constrained DLA algorithm. The output of phase 2 constituted the set of data whose organization was exploited in phase 3, to compute the 3D fractal dimension (*fd*) of the IAN. This 3D *fd* worked as a control block to either proceed to the final output in STereoLithography or standard to layer (STL) format, or to break the flow and feedback to phase 2 for modification of DLA parameters. When 3D *fd* was considered acceptable, an additional control block could be activated (indexing) to count the number of simulated nodes (and roots) belonging to different vascular orders (optional phase 4), to provide a mean percentage of the IAN branches at different orders as an indicator of IAN model accuracy. This mean vascular percentage could be used to either proceed to the final STL output, or to break the flow and feedback to phase 2 to improve the DLA simulation.

DISCUSSION AND CONCLUSIONS

In the present study, we developed an original computational template of numerical and rendering routines for simulating the 3D intralobar arterial network (IAN) of the human thyroid gland, based on a diffusion limited aggregation (DLA) algorithm. A peculiarity of

our approach was to constrain the vessel growth within the 3D shape of a virtually-reconstructed human thyroid lobe, with boundaries set through an homotopic-like parametrization involving splines and Bezier curves.^{36,37} This choice (detailed in Supplementary Material) stemmed from previously tested building blocks developed to provide a robust and flexible framework yielding a reproducible structure to be eventually used for additive layer manufacturing / bioprinting of the stromal matrix or stromal - vascular scaffold (SVS) of human organs, primarily the human thyroid gland.^{5,7,21–27}

Indeed, the 3D geometry of the SVS acts as a “photographic negative” of the macro - microscopic organ morphology, being *per se* capable of mechano-chemically and topobiologically guiding cells during embryonic development and postnatal growth.^{1,38} Therefore, we raised the possibility that the human thyroid lobe is organized as a topological “phase space” where growing thyrocytes may self-assemble driven by rules inherent in the 3D geometry of the SVS,^{5–7} that acts as a fractal-like chaotic attractor.^{7,9,21–25} Indeed, fractality is a property belonging to numerous human vascular networks.^{39,40}

Hence, we conceived a thymomorphic scaffold-bioreactor unit potentially scalable as a part of a microfluidic chip test system whose basic requirement resides in the strict similarity between the 3D geometry of the *ex situ* engineered / bioprinted SVS and the natural thyroid lobe SVS.^{5,8,9} We recently substantiated this perspective showing that a decellularized organomorphic, rodent and pig SVS is suitable for *in vitro* reconstruction of lung, liver, thymus, adrenal, anterior pituitary, cerebral cortex, pancreas, kidney, skin, and bone tissues by recellularization.^{8,41–43}

Consistently, we simulated a patient-tailored SVS constrained within the geometric boundaries of the virtually-reconstructed lobar “phase space”, and modelled as an hybrid fractal network with a “deterministic boost” provided by average values of some standard vascular parameters (for details see Supplementary Material). Exploiting the evidence that in the normal thyroid gland there is spatial uniformity of the lobe fitting an ovoid-like or conic-like geometry,^{5,6,9} we reproduced the 3D lobar shape from a single ultrasonographic lobar image (for details see Supplementary Materials). The ensuing lobar

volume (data not shown) resulted substantially coincident with those found using more routinely used calculations.^{44–48}

Once the DLA was iterated up to 4th order arteries, we applied an *ad hoc* box counting method to derive 3D *fd* values, resulting statistically coincident in all three reconstructed thyroid lobes and linearly correlated with those of a vascular cast from a cadaveric thyroid, prepared prospectively as an internal control. However, comparison between mean 3D *fd* of 2nd with respect to 3rd and 4th order arteries depicted a statistically significant difference, suggesting multifractality in the branching pattern, as occurs in a number of vascular arborizations of human viscera.⁴⁹

To substantiate the similarity between DLA simulations and current anatomical data on thyroid lobe vessels,^{30–33} we developed an heuristic based on a control routine counting simulated roots and nodes by vessel order (so-called indexing method) coupled to an estimate of the number of algorithmic iterations needed to “utilize” the entire interval of real vessel calibers (as deduced from known literature data) in each simulated SVS. In this manner, an estimate of the number of 2nd to 4th order simulated vessels was reached (for details on this procedure see Supplementary Material). As a result, we obtained a percentage of 2nd to 4th order vessels not statistically different from that predictable by current anatomical data (details on this frame are in Supplementary Material). However, beyond automation of the indexing method, a patient-tailored SVS simulation by DLA might benefit of a machine learning procedure aimed at classifying and recognizing vascular patterns corresponding to those acquired by *in vivo* imaging techniques on each single subject. Similarly, DLA has recently been combined with neural network algorithms to improve diagnosis of brain injury of individual patients.⁵⁰ Whether this optimization should proceed by supervised inputs or reinforcement mechanisms⁵¹ remains to be tested.

Finally, we developed an operational flow-chart of the entire process aimed at the 3D virtual reconstruction of the thyroid SVS, savable in STL format for its prototyping with biomaterials.^{9,21,22,24–26} Use of an STL-based 3D blueprint of ramified vascular structures is a well known requirement for their fabrication by extrusion bioprinting^{52,53} and material jetting technologies.⁵⁴ Consistently, in previous studies based on

preliminary blocks of our computational template, we tested the suitability of a Multi Jet Modelling approach to fabricate with UV-curable resins a single branch prototype of the human thyroid SVS. Results showed that minimal calibers of printed endings varied from 0.11 to 0.41 mm, with an excess dimensional deviation of 0.01 - 0.03 mm with respect to the STL simulation,²² thus depicting acceptable limits of dimensional accuracy for 2nd order vessels. Recent development of material extrusion and jetting technologies, like layer-by-layer ultraviolet assisted extrusion-based and drop-on-demand inkjet bioprinting with hydrogels,^{54,55} promise to reach dimensional limits and accuracy compatible with those requested by 3rd and 4th order SVS branches. Eventually, when *in vivo* tomographic technologies will allow for collection of volumetric images of the intrinsic vascularity of human organs, VAT polymerization will prove ideal for patient-tailored reconstructions^{56–58} including thyroid SVS.

To model fluid dynamics in mammalian vessels, a number of computational approaches relied upon fractal geometry^{59–61} coupled to vascular permeability, pressure drop and difference, Poiseuille's equation, and Reynold's number.⁶² However, our primary focus was on a geometric - structural similarity with the anatomical reality of the native thyroid SVS, at a scale level above that of capillary beds requiring different network laws.^{39,63,64} Since embryonic and fetal development of vascular systems leads to an adult vascular geometry through a hemodynamic-induced remodelling action,^{65,66} it may be assumed that the 3D geometry of an adult SVS reasonably retains structural features compatible with the organ-specific fluid dynamics.⁹ For these reasons, we predicted that the low / medium geometric constraints offered by our 2nd to 4th order IAN would allow for perfusion dynamics of culture media, favouring maintenance and self-aggregation of metastable toroidal patterns of seeded cells into luminous modules (follicle-like structures). Later, co-cultured endothelial precursors would be expected to give rise to a dense interwoven perifollicular capillary bed (so-called thyroid baskets) whose highly geometric constraints could lead to self-limited follicular inflation, and arrest of glandular growth within the limits of the thyroid lobe boundaries.^{5,9}

In conclusion, we believe that the computational template introduced here might prove efficient to engineer an organomorphic scaffold reproducing the 3D macro-microscopic architecture of the adult human thyroid lobe. This would offer a powerful tool for its *ex situ* reconstruction using human adult thyroid stem cells and endothelial precursors, up to fabrication of a thymorphic microfluidic chip test system. In addition, relying on standard morphometric variables like vessel order and size, bifurcation law, number of branches per vessel order, and fractal dimension, it provides a general concept for SVS fabrication, suitable to *ex situ* bioprinting of various human organs upon clinical request.

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Author contributions

GS and MF: conceptualization and analysis of numerical mathematical procedures, ideation and structure of the computational kernel, software resources; FB, GC, SM: selection and anatomical recognition of ultrasonographic thyroid images, collection of thyroid cadaveric anatomical specimens, injection-corrosion procedure for vascular identification, application of routines for statistical calculations; RT: research conceptualization, development and supervision of the entire scientific program, collection of clinical ultrasonographic images, evaluation of appropriateness of all methodologies (imaging, anatomical, mathematical, geometric, statistical), data evaluation and heuristic of estimates, manuscript writing, editing, and supervision including English language.

Disclosure Statement

No competing financial interests exist.

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Supplementary Material

Supplementary Material includes details on mathematical, numerical, and computational steps, and four explanatory images (Supplementary Image Homotopy, Supplementary Image Bounding Box, Supplementary Image Collision Network, Supplementary Image Vessel Branching).

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FIGURE LEGENDS

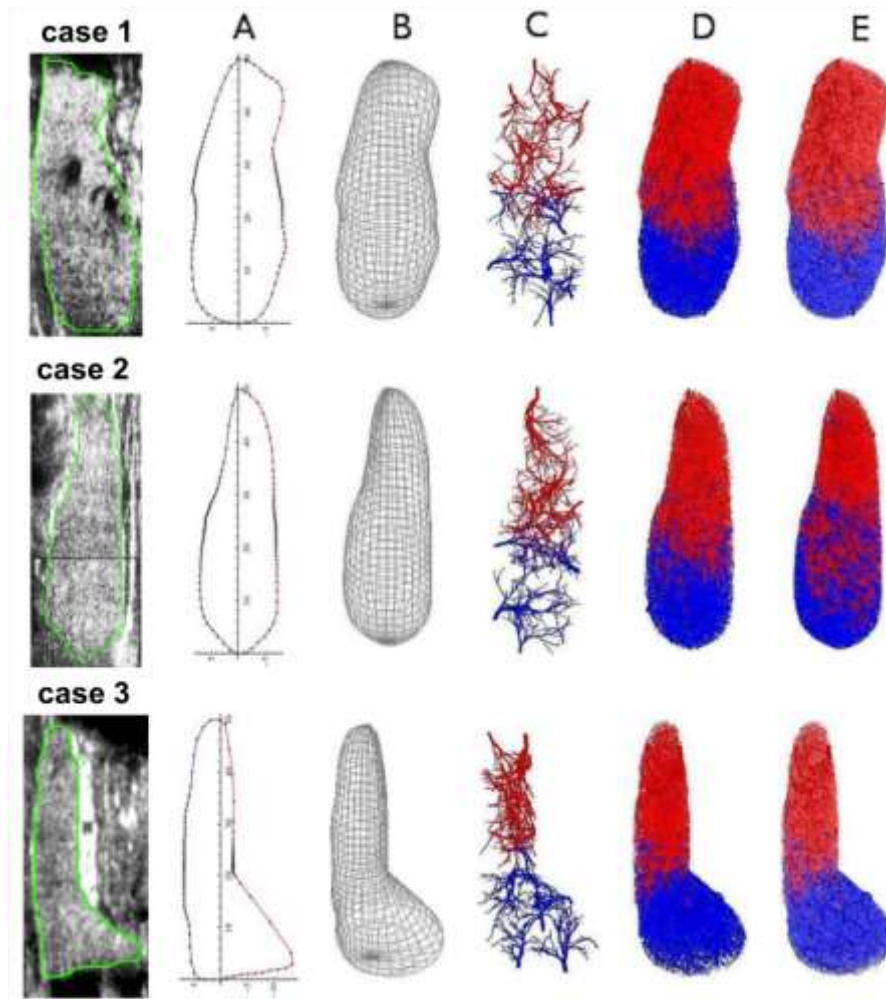


Figure 1. Schematics of the steps for 3D reconstruction of the IAN in the three subjects studied (cases 1, 2, 3). The procedure started from an ultrasonographic longitudinal section of their right thyroid lobe taken at its longest sagittal axis; A) thyroid lobe profiles after edge detection and spline approximation, including their pseudosymmetry axis; B) 3D reconstruction of the right lobe of each patient; C-E) DLA growth of the IAN of the right thyroid lobe of each patient from 2nd to 4th order vessels, respectively. For mere rendering reasons, arteries stemming from roots located roughly above and below the barycentric plane parallel to the XY-plane were coloured in red and blue, respectively.

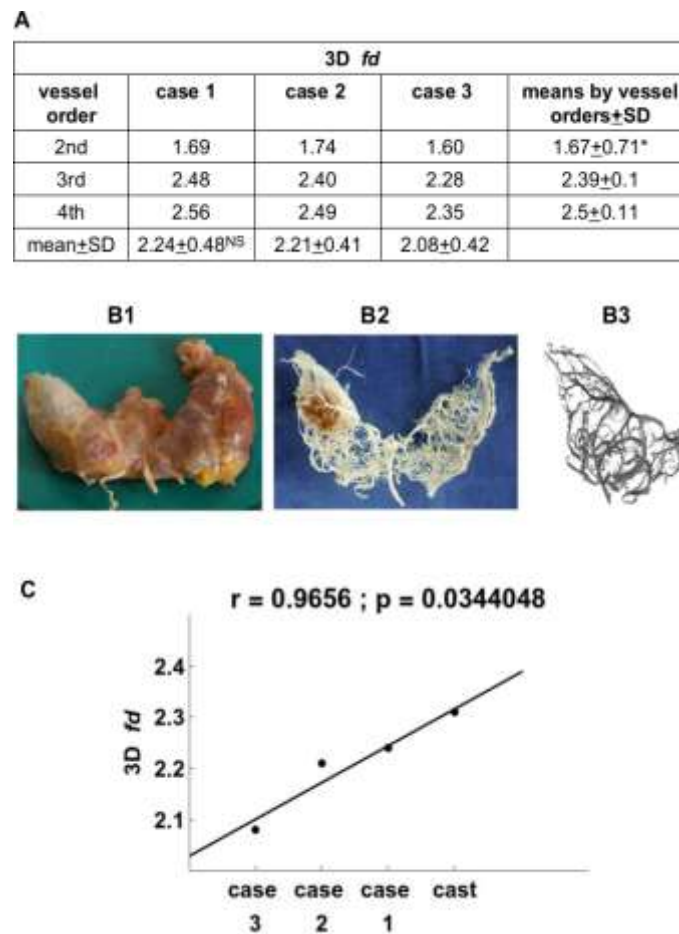


Figure 2. A) Values of the 3D *fd* of the IAN in each of the simulated cases analyzed. Note that a statistically significant difference occurs in the mean value of 3D *fd* for 2nd order vessels with respect to mean values of 3rd and 4th order vessels, suggesting a multifractal vascular pattern. In contrast, no statistically significant differences emerged between mean 3D *fd* based on all vascular orders, confirming consistency of the generative process of our algorithm; B1) human thyroid gland collected at autopsy; B2) corrosion cast of its IAN obtained after injection of synthetic resin. Note the presence of thyroid tissue in the upper pole of the right lobe, due to a still incomplete process of tissue digestion; B3) rendering of the reconstruction by microtomographic analysis of the resin-injected IAN of the right thyroid lobe, after completion of the tissue digestion process; C) linear regression analysis of the mean 3D *fd* of the IAN values of both simulated cases and vascular cast. Note the statistically significant correlation between all data. * = $p < 0.05$ with respect to 3rd and 4th order vessels; NS = not statistically significant with respect to cases 2 and 3.

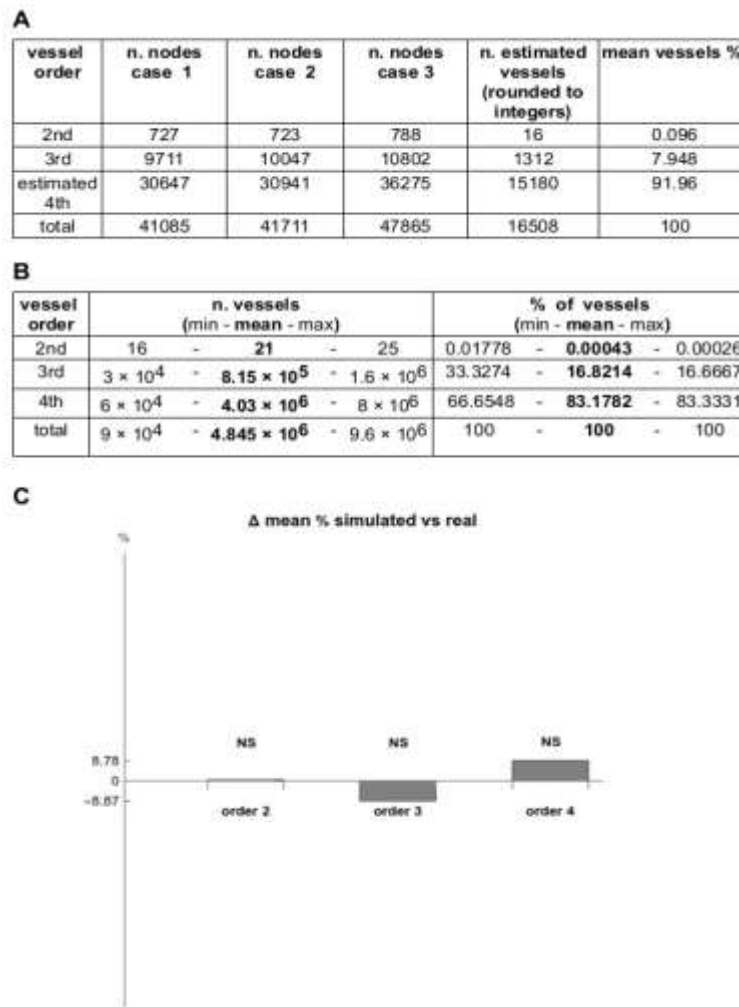


Figure 3. A) Number of roots and nodes of 2nd order arteries, and nodes of 3rd and 4th order arteries, as determined by the indexing method in the simulations of the IAN of our three euthyroid control cases. Values in the last column refer to the mean percentage of vessels at each artery order with respect to the total of the lobe arteries; B) estimated ranges of the number and percentages of arteries in the adult human thyroid lobe, as determined by anatomical literature data. Note the wide intervals, spanning from minimal to maximal through mean values (in bold); C) distribution of the difference in percentage between number of arteries in simulated vs real cases using mean artery values. The simulations did not yield statistically significant oscillations in the amount of simulated vessels with respect to known anatomical data, suggesting that the generative principles of our algorithm robustly overlapped those of the native real vessels; min = inferior extreme of the interval; mean = average level of the interval; max = superior extreme of the interval; NS = not statistically significant.

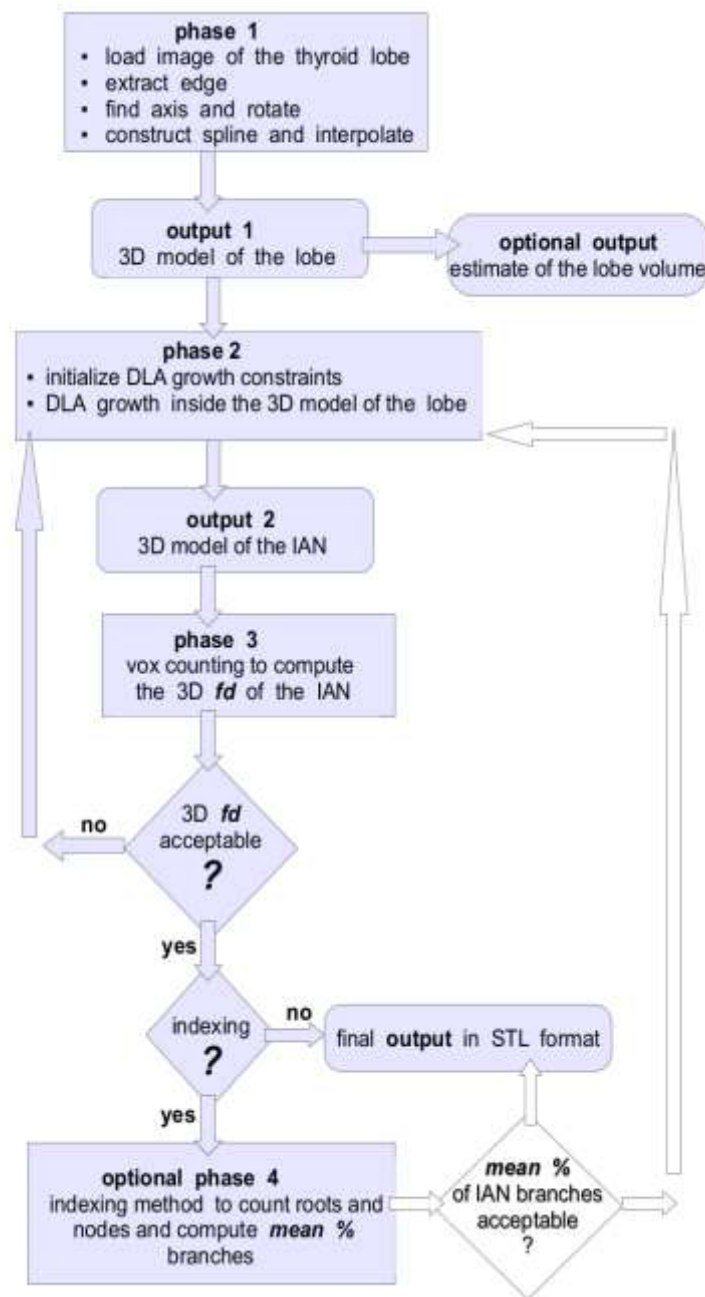


Figure 4. Computational template summarizing the 3D modelling of the IAN of the adult, human thyroid lobe. After phase 3, note the presence of two subsequent control blocks based on the 3D fractal dimension (fd) and indexing method. The latter measures the number of roots and nodes at each vascular order, to allow for estimation of vessel numbers with respect to their anatomical reality. Both are decision switches for either feeding back into the DLA growth process or exiting to a computational format suitable for 3D bioprinting, like standard to layer (STL). White block and arrows signify that this part of the template can be activated at request.