SimpleForest - a comprehensive tool for 3d reconstruction of trees from forest plot point clouds

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Abstract

The here-on presented SimpleForest is written in C++ and published under GPL v3. As input data SimpleForest utilizes forestry scenes recorded as terrestrial laser scan clouds. SimpleForest provides a fully automated pipeline to model the ground as a digital terrain model, then segment the vegetation and finally build quantitative structure models of trees (QSMs) consisting of up to thousands of topologically ordered cylinders. These QSMs allow us to calculate traditional forestry metrics such as diameter at breast height, but also volume and other structural metrics that are hard to measure in the field. Our volume evaluation on three data sets with destructive volumes show high prediction qualities with concordance correlation coefficient \( \text{CCC} \left(r^2_{adj.}\right) \) of 0.91 (0.87), 0.94 (0.92) and 0.97 (0.93) for each data set respectively.

We combine two common assumptions in plant modeling "The sum of cross sectional areas after a branch junction equals the one before the branch junction" (Pipe Model Theory) and "Twigs are self-similar" (West, Brown and Enquist model). As even sized twigs correspond to even sized cross sectional areas for twigs we define the Reverse Pipe Radius Branchorder (RPRB) as the square root of the number of supported twigs. The prediction model \( \text{radius} = B_0 \ast \text{RPRB} \) relies only on correct topological information and can be used to detect and correct overestimated cylinders. In QSM building the necessity to handle overestimated cylinders is well known. The RPRB correction performs better with a \( \text{CCC} \left(r^2_{adj.}\right) \) of 0.97 (0.93) than former published ones 0.80 (0.88) and 0.86 (0.85) in our validation.

We encourage forest ecologists to analyze output parameters such as the GrowthVolume published in earlier works, but also other parameters such as the GrowthLength, VesselVolume and RPRB which we define in this manuscript.

Author Contributions

Hackenberg Jan: Conceptualization, Data Curation, Formal Analysis, Investigation, Methodology, Resources, Software, Validation, Visualization, Writing – Original Draft Preparation, Writing – Review & Editing
Calder Kim: Data Curation, Writing – Review & Editing
Miro Demo: Data Curation, Validation, Writing – Review & Editing
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Alexandre Piboule: Software, Resources, Writing – Review & Editing
Disney Mathias: Conceptualization, Data Curation, Writing – Review & Editing

1 Introduction

Forests are key part of the global carbon cycle [1]. The quantification of global carbon bound in forests generally relies on forest inventory data. In traditional forestry often only two parameters per sampled tree are derived, the diameter at breast height (DBH) and the height (h). Above-ground biomass (AGB) is then derived either by a volume table look-up of DBH and h or by an allometric model utilizing the two parameters as input variables.

Such tables and formulas can be found published [2,3] but this is not always the case. Especially when species provenance and the variation of regional growth patterns are taken into account, one has to rely on destructive data collection [4]. In the last few decades utilizing the point cloud data of terrestrial laser scans to derive directly the volume of trees has been shown as a fast, less costly and non destructive alternative [5].

The complete trees above-ground volume (AGV) can be directly assessed from the topological ordered cylinder structure of so called quantitative structural models (QSMs) [6,7]. QSMs are not only important to collect AGV data which can be multiplied with density predictions [8] for AGB estimates, they allow forest ecology relevant insight into tree architecture itself [9,10]. From a single TLS forest plot scan multiple thousands or even up to millions of geometrical measurements can be derived. Nevertheless, by purely using limited destructive measured data theories such as the West Brown and Enquist (WBE) model [11,12] or the pipe model theory [14,15] have been developed under named limitations.
With QSMs new advances such as the rework of the WBE\cite{16} describing a generalization for asymmetric branched trees are possible, to help understanding the tree architecture\cite{9}.

As an official successor of SimpleTree\cite{7} we provide here the SimpleForest plugin for the Computree platform\cite{17} to retrieve QSMs from plot level raw data with a fully automated pipeline consisting of various filters, a digital terrain model (DTM) algorithm, vegetation segmentation and finally two different QSM modelling approaches. Firstly, the so called SphereFollowing method\cite{18}\cite{18} and additionally the Dijkstra\cite{19} based method such as proposed by\cite{20}\cite{21}. Our implementation of the latter one is known to perform worse than the first and is here-on undiscussed.

SimpleForest’s fully automatic processing chain can be inspected step by step with the powerful visualizer of Computree. SimpleForest’s QSMs can be conveniently extracted in csv format to be postprocessed with AMAPstudio\cite{22} or R software\cite{23}. Ply format allows visual inspection with tools like CloudCompare, Meshlab or Blender. The latter approach is useful for exemplary quality assessment after computation when a larger number of plots is processed in the batch-mode of Computree. The SphereFollowing parameters are automatically searched for in higher dimensional space. This automatic search is especially user friendly for users without informatic knowledge but also did outperform us on expert level.

In case wood vs. leaf separation is desired, separated tree clouds can be processed in a supervised machine learning manner. Remaining leaf fragments in the cloud lead to errors in QSMs which are afterwards automatically handled by filters relying on the WBE\cite{24} theory. Here we use new invented parameters which show a better Radius correlation than traditional ones for the important diameter relations in scaling theories.

Supporting information

S1 Text. The SimpleForest UserGuide.

- Over 100 pages of documentation are contained in the User Guide:
  https://gitlab.com/SimpleForest/computree/-/blob/master/bin/SimpleForestUserGuide.pdf
  https://doi.org/10.5281/zenodo.5138255

All algorithms are described in detail with a special focus on the parameterization of the steps.

1.1 Data

We will evaluate on 5 different data sets. Each data set is online available, in one repository:
https://doi.org/10.5281/zenodo.5131717.

S1 Dataset Hackenberg et al. 2015a data\cite{35} 12 Quercus petraea (leaf-off), 12 Erythrophleum fordii (leaf-on) and 12 Pinus massoniana (needle-on) tree scans are available with harvested ground truth volume data\cite{35}. The Quercus petraea trees have been overgrown with moss leading to larger overestimation of the Radius.

- Erythrophleum fordii clouds raw:
  https://zenodo.org/record/5131717/files/hackenbergErythrophleumRawClouds.zip

- Erythrophleum fordii clouds denoised:
  https://zenodo.org/record/5131717/files/hackenbergErythrophleumDenoisedCloudsCC.zip

- Pinus massioniana clouds raw:
  https://zenodo.org/record/5131717/files/hackenbergPinusRawClouds.zip

- Pinus massioniana clouds denoised:
  https://zenodo.org/record/5131717/files/hackenbergPinusDenoisedClouds.zip

- Quercus petraea clouds denoised:
  https://zenodo.org/record/5131717/files/hackenbergQuercusDenoisedClouds.zip
S2 Dataset  De Tanago et al. 2017 data [36] 29 tropical buttresses trees have been scanned across sites in Peru, Indonesia and Guyana. Harvested reference data is included [36]. All trees have been scanned in leaf-on condition.

- Peru, Indonesia and Guyana clouds raw:
  https://zenodo.org/record/5131717/files/deTanagoRawClouds.zip

- Peru, Indonesia and Guyana clouds denoised:
  https://zenodo.org/record/5131717/files/deTanagoDenoisingScriptsDenoisedClouds.zip

S3 Dataset  Demol et al. 2021 data [37] 65 trees in total of the species Frazinus excelsior, Fagus sylvatica, Pinus sylvestris and Larix decidua have been scanned and published with reference data [37]. The Pinus sylvestris trees have been scanned in needle-on condition, the other species were scanned without foliage.

- Clouds:
  https://zenodo.org/record/5131717/files/demolClouds.zip

S4 Dataset  Disney et al. 2018 data [38, 39] An Acer pseudoplatanus [39] tree cloud of a large and complex individual with DBH larger 1 m is used to show statistical analysis potential. The tree is located in Wytham Woods, UK.

- Clouds:
  https://zenodo.org/record/5131717/files/wythamCloud.zip

S5 Dataset  High resoluted leaf-less plot data. A full plot in leaf off conditions. The plot does not have ground truth data in attached, but the quality of tree segmentation as well as DTM and QSM modeling can be visually assessed.

- Clouds:
  https://zenodo.org/record/5131717/files/leibzigCloud.zip

1.1.1 Processing scripts

S1 Processing scripts  SimpleForest scripts to process S1 Dataset.

- Erythrophleum fordii denoising scripts:
  https://zenodo.org/record/5131717/files/hackenbergErythrophleumDenoisingScripts.zip

- Pinus massoniana denoising scripts:
  https://zenodo.org/record/5131717/files/hackenbergPinusDenoisingScripts.zip

- QSM modeling script:
  https://zenodo.org/record/5131717/files/hackenbergQsm.xsct2

S2 Processing scripts  SimpleForest scripts to process S2 Dataset.

- Denoising scripts:
  https://zenodo.org/record/5131717/files/deTanagoDenoisingScriptsDenoisedClouds.zip

- Poisson reconstruction buttress script:
  https://zenodo.org/record/5131717/files/deTanagoButtressPoisson.xsct2

- QSM modeling script:
  https://zenodo.org/record/5131717/files/deTanagoQsm.xsct2

S3 Processing scripts  SimpleForest scripts to process S3 Dataset.

- QSM modeling script:
  https://zenodo.org/record/5131717/files/demolQsm.xsct2
S4 Processing scripts  SimpleForest scripts to process S4 Dataset.

- QSM modeling script:
  https://zenodo.org/record/5131717/files/wythamQsm.xsct2

S5 Processing scripts  SimpleForest scripts to process S5 Dataset.

- QSM modeling script:
  https://zenodo.org/record/5131717/files/leibzigQsm.xsct2

1.1.2 Validation scripts

S1 Validation scripts  SimpleForest scripts to validate results of S1 Processing scripts.

- Volume validation script:
  https://zenodo.org/record/5131717/files/hackenbergValidation.R

S2 Validation scripts  SimpleForest scripts to validate results of S2 Processing scripts.

- Volume validation script:
  https://zenodo.org/record/5131717/files/deTanagoValidation.R

S3 Validation scripts  SimpleForest scripts to validate results of S3 Processing scripts.

- Volume validation script:
  https://zenodo.org/record/5131717/files/demolValidation.R

S4 Validation scripts  SimpleForest scripts to validate results of S4 Processing scripts.

- Statistical plotting script:
  https://zenodo.org/record/5131717/files/wythamAnalysis.R

S5 Validation scripts  SimpleForest scripts to validate results of S1 Processing scripts, S2 Processing scripts, S3 Processing scripts.

- Combined results data table:
  https://zenodo.org/record/5131717/files/tableAll.csv
- Volume validation:
  https://zenodo.org/record/5131717/files/ValidationScriptAll.R

1.2 Software  Software code repository.

- Under the GPL version 3 license:
  https://gitlab.com/SimpleForest/computree/-/blob/master/pluginSimpleForest/GPL_v3_template
- Inside a subfolder this repository contains a Win10 compiled executable:
  https://gitlab.com/SimpleForest/computree/-/tree/master/bin
- Persistent 5.1.3:
  https://doi.org/10.5281/zenodo.5138255
S2 Software  Online resources for S1 Software

- An homepage: https://www.simpleforest.org/
- A video tutorial channel: https://www.youtube.com/channel/UCq2dqxUF3l0musX0ptk7xhr
- Support forum Computree: http://rdinnovation.onf.fr/projects/computree/boards
- SimpleForest users help users list: https://lists.posteo.de/listinfo/simpleforest

2 Design and Implementation

Computree and SimpleForest rely on the following third party libraries: the Point Cloud Library (PCL) [25], the Boost library, the Gnu Scientific Library (GSL) [26], the OpenCV library, the QT library and the Eigen library.

All steps are implemented multi-threaded with an optimized asymptotic runtime to execute as fast as possible. In Text we describe more than 20 steps regarding the IO data, with the underlying algorithms explained in detail and the parameterization is discussed as well. Here only a short overview is given.

2.1 Computree

Computree is a collaborative platform published under LGPL for 3d data processing for forestry applications. Computree can visualize multiple results of intermediate computation steps at once, Fig 2. If the user wants, he can display one result into a new visualizer windows each. As all visualizers can have a shared camera view the visual inspection is user-friendly implemented.
Fig 2. The user interface of Computree. Four visualizers with a shared camera view are opened. UpperLeft shows the leaf-on condition input cloud, UpperRight the de-noised cloud. On the LowerLeft we see the resulting QSM, with error colors. Brown means good fitted cylinders and green bad fitted ones. On the LowerRight the cylinders are depicted which are considered good enough for an allometric fit.

With the QT user interface steps parameters can be conveniently adjusted. Parameterized step pipelines can be saved and reloaded.

2.2 Point Cloud Filters

SimpleForest implements various point cloud filters for generating slices in a desired height above the DTM, filters to retrieve either ground or stem points by the normal deviation from the z-axis as well as frontends to few PCL \[25\] filters and the Belton filter \[28\].

2.2.1 Gaussian Mixture Models with Fast Point Feature Histograms (FPFH) \[27\]

A semi automatic supervised machine learning technique utilizing gaussian mixture models \[28\], using the FPFH feature space and relying on a single range parameter. The filter was implemented for de-leaving. Up to ten automatic generated clusters have to be labeled by the user into LEAF, WOOD and UNCLASSIFIED. Then on each original point KNN clustering is performed, if more WOOD than LEAF points are in the neighborhood, the final classification is WOOD, otherwise LEAF.

2.3 DTM

A DTM can be computed via a pyramidal approach where raster cell size is halved iteratively from full plot size to a user desired threshold. Into points within one cell MLSAC planes are fitted \[29\]. To be accepted the plane has to surpass a closeness test to the according lower resoluted cell’s plane from the previous iteration.
2.4 Tree Segmentation

The segmentation consists of three main procedures, each implemented in at least a single step to be able to show intermediate results.

2.4.1 Generating the seeds

We use as input a slice of points above the DTM at DBH. The slice points have been filtered with the stem point filter to remove undergrowth branch connections between neighboring trees. By euclidean clustering retrieved clusters serve now as seeds and each cluster receives an unique ID.

2.4.2 Growing seeds to trees with Dijkstra \[19\]

On all points of the full vegetation Dijkstra’s \[19\]’s algorithm is executed in a competitive manner with the seed clusters of the previous step. A point receives the cluster ID of the closest seed point. The distance measure here is the length of the Dijkstra path to a connected seed point. To be able to jump over occlusion gaps the user can scale the cloud by z-coordinate. If the z-coordinate is multiplied with a factor smaller than one, vertical gaps have less impact compared to horizontal gaps. We want to achieve this, because natural non occlusion gaps are large in z-dimension for multi layered forests, but can be rather small in the x,y plane for neighboring trees.

2.4.3 Classifying remaining points with Voronoi regions

Clusters too far away from the main vegetation cluster and not reached by the Dijkstra routine receive the ID of their closest Dijkstra cluster. Here a user given distance threshold has to be passed to reject outlier points with a too large distance to the plot.

2.5 QSM reconstruction

2.5.1 QSM basic reconstruction

The SphereFollowing method \[4,7,18\] utilizes search spheres and onto their surfaces circles are fit in a recursive manner to detect topological ordered cylinders. For details please refer to S1 Text where a multi paged description of the algorithm exists. One improvement to the previously published SimpleTree implementation is the multitude of available cylinder fitting methods: Center of Mass (COM) with median distance radius \[4\], Gauss Newton Least Squares (GNLS) \[4\], Random Sample Consensus (RANSAC) \[30\], M-estimator SAmple Consensus (MSAC), Randomized M-estimator SAmple Consensus (RMSAC), Maximum Likelihood Estimator SAmple Consensus (MLESAC) \[29\], Progressive Sample Consensus (PROSAC) \[31\], Randomized RAndom SAmple Consensus (RRANSAC) \[32\] and Least MEDian of Squares (LMEDS).

We use the COM method by default as it performs very strong in earlier works \[4\]. Nevertheless we provide for interested users the GNLS as well as all sample consensus fitting methods provided in the PCL.

Three SphereFollowing main parameters, see user guide for additional information, are predicted and then internally optimized with an automated parameter search \[7\] utilizing the the downhill simplex method \[33,34\]. This parameter search enables researchers to perform the modeling without algorithmic knowledge.

For the parameter search the computer has to decide for two different models which one is better. The best model is chosen via the cloud to model distance. We provide five different distance metrics. The root mean squared distance (RMSD) of each point to the model and the more robust \[33\] mean distance (MD) of each point to the model. Both RMSD and MD can be used in their more robust form where the distances are cropped to a max value, following the MSAC principle. Lastly the most robust metric out of all is to count the inlier points with less than a threshold distance to the QSM. Good quality clouds should be evaluated with RMSD, those of worse quality with MD. MSAC and the fifth method can be useful if leaf noise exists.

2.5.2 QSM based clustering

We define recursively similar to the GrowthVolume from previous works \[7\]:

The GrowthLength (GL) of a cylinder is the cylinder’s length plus its children’s GL.
We can use the derived QSM after computation to transfer the $GL$ of a cylinder to its closest points. Then points can be clustered by their $GL$ which takes small values for twig regions and large ones for the root area, Fig 3.

Fig 3. A $GL$ clustered point cloud of an *Acer pseudoplatanus*, S4 Dataset. Color codes are ordered from closest to the tips up to close the root: green, orange, red, purple, blue.

2.5.3 QSM advanced reconstruction

In a second modeling run we can use the clusters described in 2.5.2. The three parameters mentioned in 2.5.1 can be estimated and optimized per cluster to receive more accurate cylinders. For example with five clusters we do optimization in a $3 \times 5 = 15$ dimensional parameter space.

2.5.4 Detect outlier cylinders

A step which classifies cylinders into WELLFITTED and BADFITTED cylinders exists. Cylinders close to the tips are classified as BADFITTED cylinders. Additionally for the remaining cylinders we compute the average distance of allocated points to the cylinder. Only cylinders with a small average distance are finally classified as WELLFITTED cylinders.

If we look at the average point distance of QSM cylinders we can see that most errors occur either at the twigs or at branch junctions, Fig 4 and Fig 5.
Fig 4. An unfiltered QSM of an *Acer pseudoplatanus*, S4 Dataset. The fitting error of points to their closest cylinder is colorized. Cylinders with low error are colorized orange and cylinders with large errors purple.
Fig 5. Error plot for an unfiltered QSM of an *Acer pseudoplatanus*, S4 Dataset. The log of fitting error in dependency of the distance to twig.

Some overfitted cylinders can be *Radius* corrected with a QSM median filter, as proposed in earlier works [7]. But even with a good fit quality twig cylinders tend to overestimate the *Radius* [35]. Footprint size of the laser and co-registration errors lead to the fact the cloud regions of small diameter components tend to overestimate.

2.6 QSM filters

2.6.1 Tapering filter

We implemented a taper correction based on the one TreeQSM [42, 43] provides.

2.6.2 GrowthVolume (GV) filter

We applied a GV filter in [7] which handled the overestimation to some extend. There we predicted the *Radius* from the GV with the Eq. (1):

\[
*Radius = B_0 * GV^\theta
\]  

(1)

2.6.3 VesselVolume (VV) filter

The GV of a cylinder is always effected by the error of twigs as the supported twig *Volume* is included in its value. By it’s definition the GV is therefore not the best predictor. We can be more robust against twig *Radius* overestimation by ignoring the *Radius* with the following definitions:

We define the Reverse Pipe Area Branchorder (RPAB): The WBE model considers that the branching architecture is self-similar [16]. In a self-similar branching architecture the *Radius* as well as the cross sectional area of a twig is constant. As we do not know this area in \( cm^2 \) we define that *RPAB* is a tree specific area unit which equals one for a tip cylinder. By following the pipe model theory [14] the *RPAB* of a cylinder at any location inside the QSM equals the number of supported tips. Supported tip cylinders always have a recursively defined parent relation to the query cylinder.

The *RPAB* serves as a proxy for the cross sectional area but is by the definition independent from any *Radius* measurement.

To retrieve a *Radius* independent proxy for the *Volume*, we define:

The VV of a cylinder is the cylinder’s *RPAB* multiplied with the *Length* plus its children’s VV.
We can see in Fig 6 that the VV is invariant during the correction and GV is not. Also notable is that outliers are less strong visible for VV.

Fig 6. Allometric plots of an *Acer pseudoplatanus*, S4 Dataset. (a) The GV extracted from an uncorrected QSM. (b) The GV from the same QSM which has been corrected. The magnitude of the predictor was effected by the correction. (c) The VV of the uncorrected QSM. (d) The VV of the corrected QSM covers the same value range.

SimpleForest provides such a VV predicted *Radius* correction utilizing Eq 2, with intercept $B_1$ estimation being optional.

$$Radius = B_0 \times VV^\theta + B_1$$

(2)

The additive component $B_1$ is relevant in case twig regions have been removed during denoising.
2.6.4 **Reverse Pipe Radius Branchorder (RPRB) filter**

Doing the non linear underlying power function fit unsupervised can lead to the termination in a non desired local minimum if the initial values for $B_0$ and $\theta$ are not good enough. We get an easier mathematical model by using a *Radius* independent proxy for the *Radius* itself:

**The RPRB is the square root of the RPAB.**

This relation is a linear model of a simple form as also the intercept is forced to the origin here:

$$\text{Radius} = B_0 \text{RPRB}$$  \hspace{1cm} (3)

Exemplary data into which equation 3 is fitted can be seen in Fig 7.

![RPRB Scatter plot for an Acer pseudoplatanus, S4 Dataset](image)

Fig 7. RPRB Scatter plot for an *Acer pseudoplatanus*, **S4 Dataset**. Plot of the RPRB on the x-axis and *Radius* as a predicted value.

2.6.5 **Refitting cylinders**

As we only have fitted circles with the SphereFollowing method, we can align points to their closest cylinder and do a cylindrical refit on those points [4], Fig 8.
3 Results

The *Quercus petraea* and denoised *Pinus massoniana* of S1 Dataset were corrected with the Tapering and *Erythrophleum fordii* were corrected with VV approach. In a second modeling run all QSMs were corrected with the RPRB filter. For S2 Dataset we used the VV filter in one run and the RPRB filter in a second one. For S3 Dataset we applied Tapering firstly and then in a second run the RPRB filter, Fig 9.
Fig 9. Volume validation for S1 Dataset, S2 Dataset and S3 Dataset. Modeling results of (a) S1 Dataset with Tapering and VV correction, (b) S1 Dataset with RPRB correction, (c) S2 Dataset with Tapering correction, (d) S2 Dataset with RPRB correction, (e) S3 Dataset with VV correction, (f) S3 Dataset with RPRB correction.
We show for S3 Dataset within one plot SimpleForest results correct one time with \textit{Tapering} and one time with RPRB compared to TreeQsm \cite{42} and in a second plot SimpleForest \textit{Tapering} results which have been filtered by using both tools. SimpleForest \textit{Tapering} corrected results have to be in a 5\% range to TreeQSM \textit{Tapering} corrected results, otherwise the QSM is filtered out, Fig 10.

\begin{figure}[h]
\centering
\begin{minipage}{0.45\textwidth}
\includegraphics[width=\textwidth]{fig10a.png}
\caption{(a) Comparison of SimpleForest \textit{Tapering}, TreeQSM \textit{Tapering} and SimpleForest RPRB results for S3 Dataset.}
\end{minipage}\hfill
\begin{minipage}{0.45\textwidth}
\includegraphics[width=\textwidth]{fig10b.png}
\caption{(b) We only see SimpleForest \textit{Tapering} volume predictions which are close to TreeQSM \textit{Tapering} volume prediction, other QSMs are filtered out.}
\end{minipage}
\end{figure}

Combining SimpleForest results for all data sets with the VV approach for S2 Dataset and RPRB filter otherwise leads to Fig 11.
Fig 11. Combined S1 Dataset, S2 Dataset and S3 Dataset validation. (a) Combined evaluation of all three data sets showing a zoom area of interest used for the next figure. (b) The zoom view showing trees with a maximum volume of 5 m$^3$.

We provide a summary of the results in Table 1.

| Data                | Software | RMSE (m$^3$) | RMSE$_{rel.}$ | CCC  | $r_{adj}^2$ |
|---------------------|----------|--------------|---------------|------|-------------|
| S1 Dataset          | SimpleForest $^a$ | 0.12         | 0.10          | 0.91 | 0.87        |
| S1 Dataset          | SimpleForest $^b$ | 0.24         | 0.16          | 0.79 | 0.83        |
| S2 Dataset          | SimpleForest $^c$ | 3.25         | 0.08          | 0.94 | 0.92        |
| S2 Dataset          | SimpleForest $^d$ | 4.67         | 0.14          | 0.87 | 0.90        |
| S3 Dataset          | SimpleForest $^e$ | 0.14         | 0.06          | 0.97 | 0.93        |
| S3 Dataset          | TreeQSM $^c$     | 0.32         | 0.11          | 0.86 | 0.85        |
| S3 Dataset          | SimpleForest $^f$ | 0.15         | 0.08          | 0.94 | 0.98        |
| S1 Dataset          | SimpleForest $^g$ | 1.85         | 0.05          | 0.97 | 0.96        |

Table 1. Numerical validation of QSM volume measures compared to harvested reference volume measures.

$^a$ = RPRB filtering

$^b$ = VV filter for *Erythrophleum fordii*, Tapering filter otherwise

$^c$ = only VV filter.

$^d$ = only Tapering filter.

$^e$ = only Tapering filter, rejected QSMs not in a 5% range to TreeQSMs.

$^f$ = VV filter for S2 Dataset data, otherwise RPRB filter.
Our plot data did not have ground truth attached, but we use it for visually inspecting the quality of the DTM, Fig 12.

(a) We see the height of ground points visualized increasingly from green to red. A forest road is visible. (b) The DTM height values are colored increasingly from dark to bright. The DTM captures the road without artifacts.

The modeled QSMs also do not show segmentation errors in Fig 13.
Fig 13. Final QSMs of S5 Dataset. Each QSM is colored differently.

The *Acer pseudoplatanus* QSM, S4 Dataset, contains more than 35k cylinders. We use this QSM to show some statistical analysis potential in Fig 14.
Fig 14. Exemplary statistic plots for an *Acer pseudoplatanus* S4 Dataset. (a) We show a traditional stem taper. (b) The branch volume distribution binned along the height of the branch growing out of the stem. The maximum can define the crown base. (c) The branch volume accumulated for each branch order. (d) An analysis in tradition of the Pipe Model Theory [7].
4 Discussion

SimpleForest utilizes an advanced parameter search for optimizing the SphereFollowing method compared to the brute force search published in earlier works leading to better models. Nevertheless, as input data already overestimates the twig regions a radius correction has to be applied. The choice of this correction influences the results significantly. On the S3 Dataset data set we can perform a direct comparison versus an earlier published approach and our RPRB filter performs very strong as can be seen in Fig 10 (a).

We can also see, that even with the less well performing Tapering radius correction we can achieve better results, if we use more than one QSM tool. Outlier QSMs can be simply filtered out if the QSMs of different tools do not have similar volume predictions, Fig 10 (b). In this work we only use TreeQSM, but also 3D Forest can build QSMs nowadays. By using all three tools a more sense of uncertainty can be gained and filtering out QSMs from tree clouds which are hard to model can be automated.

The S1 Dataset was the second data set revealing the strength of the RPRB filter. Only for S2 Dataset this filter performed worse than the comparison filter. The reason behind this performance is simply, that we lost the thin branching structure during the de-noising routine, but forced intermediate sized branches to become thin, Fig 15.

As non TLS derived data to evaluate DTM and segmentation quality was not available, we have to rely on visual inspection. Our example plot reveals no errors and we do not have to improve with manual interaction our segmentation results.

Our QSM validation results are in line with other available tools and sometimes even stronger than results which have
been published by those. At least for leaf-off conditions the complete data processing can be fully automatized on a plot
level. If leaf-on conditions are met, we provide two functional semi automatic leaf filtering procedures.

Regarding that the embedding platform Computree has other plugins for forest modeling in addition and that we
provide as much interfaces as possible to external software tools, we recommend using this software suite for estimating
tree’s above volume as well as gaining insight into forest ecology relevant questions.

5 Availability and Future Directions

5.1 Future work

The \textit{RPRB} filter should not rely on an origin forced intercept, as twigs can be removed during leaf separation. Another
downside of this filter is the fact, that the \textit{RPRB} is same for all cylinders inside a segment \cite{7}. Notice, that we defined a
segment in previous works as the list of all cylinders between two neighboring branch junctions. The \textit{VV} filter does not
have this downside and if the non linear fit can be performed in a more robust way even better filtering might be doable.

Lastly we consider it beneficial to gather with our semi automatic leaf separation approaches training data for a full
automatic leaf separation procedure. Tiny dnn \cite{44,45} is a deep learning header only library which allows easy integration
into SimpleForest code base. A network architecture using FFPHs as input can be implemented fastly.

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