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Progressive and Efficient Multi-Resolution Representations for Brain Tractograms

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Abstract
Current tractography methods generate tractograms composed of millions of 3D polylines, called fibers, making visualization and interpretation extremely challenging, thus complexifying the use of this technique in a clinical environment. We propose to progressively simplify tractograms by grouping similar fibers into generalized cylinders. This produces a fine-grained multi-resolution model that provides a progressive and real-time navigation through different levels of detail. This model preserves the overall structure of the tractogram and can be adapted to different measures of similarity. We also provide an efficient implementation of the method based on a Delaunay tetrahedralization. We illustrate our method using the Human Connectome Project dataset.

1. Introduction
Tractography from diffusion MRI is currently the only technique able to non-invasively explore the white matter architecture of the brain. It results in a tractogram which is a bundle of 3D polylines, usually called fibers, which are estimates of the trajectories of large groups of neural tracts. Tractography has been proven to be an invaluable tool for clinicians and researchers. It is nowadays used on a daily basis by neurosurgeons for pre-operating planning and
during surgical operations [JDML17]. It also offers important information for studying pathological processes in neurological diseases [CCJB*08].

Recent tractography methods produce up to one million of fibers [TML11]. This can complicate the rendering, visualization and interpretation of tractograms, thus limiting the aforementioned clinical applications. Furthermore, the considerable number of fibers can make computationally intractable processes such as non-linear registration or atlas construction [GCMK*16], which are important for research purposes. Many fibers might have a similar trajectory and connectivity, making the tractogram redundant. For this reason, several authors have proposed new geometric representations and visualization techniques to simplify tractograms. One of the most popular approaches consists in grouping similar fibers into clusters [GCMK*16, GBC*12, GPR*11, MWW*07, ZL02] which are then approximated with one representative fiber usually called prototype [GBC*12, GPR*11]. Other authors have also proposed to represent the spatial extent of the clusters using an encompassing geometry [MWW*07]. These methods are usually controlled by one parameter, e.g. a threshold [ZL02], thus presenting only one level of resolution at a time. Furthermore, information such as the number of fibers or the spatial extent (i.e. the volume) of the cluster might be lost in the process. To improve the visualization quality and efficiency, the geometric models used to represent the fibers are often computed directly on the GPU [PPK07, RBE*06, ESM*05]. Other methods, such as in [EBB*15], ease visualization by bringing similar fibers closer to each other. In this paper, we do not focus on single resolution approximations (i.e. clustering), but propose a multi-resolution representation based on progressive mergings, that preserves the overall structure of the tractogram and whose continuous levels of resolution can be traversed in real-time.

**Progressive simplification** methods, which focus on decimating complex 3D objects while preserving both important geometric features and topological relationships, have so far not been explored to approximate large scale tractograms. In [ZL02], Zhang and Laidlaw propose to use a hierarchical clustering algorithm to progressively group together similar fibers. They only applied it on bundles composed of a small number of fibers. Their goal was to divide tractograms into clusters and not to propose a new geometric representation or visualization technique (as it is the case in this paper). Taking inspiration from error-driven surface mesh simplification [GH97], we propose a progressive merging strategy for grouping fibers into generalized cylinders. The proposed method reduces the redundancy of the tractogram, producing a multi-resolution structure, which is organized into a nested hierarchy of levels of detail. Every fusion of fibers (or cylinders) represents a new level of resolution. Once the entire multi-resolution representation is computed, it is possible for the user to navigate through different levels of detail in a continuous fashion and in real-time, while maintaining the overall structure of the original tractogram. Furthermore, we also propose an efficient implementation based on a Delaunay tetrahedralization which makes it possible to use our method on large tractograms containing millions of fibers.

2. Method

**Fiber Decimation** We propose a tractogram simplification method (see Fig.1) based on the progressive mesh methodology [GH97, HDD*93]. Given a tractogram with $N$ fibers, we first look for the two most similar fibers based on a similarity measure. Once detected, the couple is collapsed into a single generalized cylinder – all input fibers can be seen as generalized cylinders with a null radius. The process is then iterated until obtaining a single large cylinder (i.e. $N - 1$ iterations) or using a stopping criterion to prevent over-simplification.

The proposed “progressive brain tractograms” algorithm is general and can be used with any similarity measure. It is inherently multi-resolution, where every level of detail corresponds to the fusion of two cylinders. Once the sequence of fusions is computed, we can visualize the tractogram at any resolution and switch among levels in real-time.

In the following, we will use three dissimilarity measures: the Mean of Closest distances (MC) [GPR*11], the minimum average direct-flip (MDF) [GBC*12] and one based on the computational model of Weighted Currents (WC) [GCMK*16]. Let $X = \{x_i, i = 1...N\}$ and $Y = \{y_j, j = 1...M\}$ be two fibers composed of $N$ and $M$ points respectively the MC distance is defined as:

$$MC(X, Y) = \text{mean}(d_{\text{MC}}(X, Y), d_{\text{MC}}(Y, X))$$

where $d_{\text{MC}}(X, Y) = \frac{1}{N} \sum_{i=1}^{N} \min_{j \in Y} ||x_i - y_j||$. The MDF is defined in a similar way, by assuming point-wise correspondence between fibers. The similarity measure of WC is instead defined as:

$$WC(X, Y) = \left[ K_a(||f^a - t^a||) K_b(||f^b - t^b||) \right] \sum_{i=1}^{N-1} \sum_{j=1}^{M-1} \frac{1}{g(i)} K_c(||c_i - d_j||) B_j$$

where $a_i$ and $a_t$ (respectively $b_i$ and $b_t$) are the centers and tangent vectors of $X$ (respectively $Y$), $f^a$, $f^b$ and $t^b$, $t^b$ are the corresponding endpoints of $X$ and $Y$ respectively, and $K_a$, $K_b$ and $K_c$ are three Gaussian kernels parametrized by $\sigma_a$, $\sigma_b$ and $\sigma_c$ respectively.

**Delaunay Tetrahedralization** Comparing all fibers to each other leads to quadratic complexity and intractable computations. Furthermore, most of the computations would be useless since similar close-by fibers should be merged in priority, thus indicating that during surgical operations [JDML17], it is possible for the user to navigate through different levels of detail in a continuous fashion and in real-time, while maintaining the overall structure of the original tractogram. Furthermore, we also propose an efficient implementation based on a Delaunay tetrahedralization which makes it possible to use our method on large tractograms containing millions of fibers.
Figure 2: Scheme of the merging of two fibers/cylinders $C_1$ and $C_2$ into $C_3$.

Geometric Representation In this paper, we consider that every fiber is described by its geometry and connectivity. The employed geometric representation should preserve these properties. To this end, we propose to use generalized cylinders with an elliptical basis as geometric representations for the merged fibers. We define one ellipse per vertex of the center-curve such that the resulting cylinder incorporates the trajectory and endpoints of the fibers (or cylinders) of the previous level of resolution.

Given two fibers/cylinders $C_1$ and $C_2$, Fig. 2 schematically presents how they are merged into cylinder $C_3$. We use as reference the center-curve (or fiber) with less points ($C_1$ in Fig. 2). For each point of $C_1$, we look for the closest point in $C_2$. The corresponding point in $C_3$ is computed as their weighted mean, where the weights $W_1$ and $W_2$ correspond to the number of original fibers that $C_1$ and $C_2$ represent (and we set $W_1 = W_1 + W_2$ accordingly).

To avoid flickering, the thickness of rendered cylinders is clamped to a minimum. Moreover, it is important to notice that we use the Euclidean distance between the endpoints of $C_1$ and $C_2$ to compute the extremities of $C_3$. In order to make sure that the extremities of $C_3$ would actually lie at the border with the gray matter surfaces, one should use a (computationally expensive) geodesic distance. This is left as future work.

Implementation We implemented our method in C++ using Qt for the graphical user interface and OpenGL for the rendering. To improve efficiency and memory usage, the geometry is computed on the GPU, using the hardware tessellation unit to synthesize on-the-fly our visual approximations. We will make the code publicly available at https://perso.telecom-paristech.fr/comercier/. We provide a video of the proposed interactive system as a complementary material (same url).

3. Results and Discussion

Dataset We conducted our experiments on the HCP dataset (https://db.humanconnectome.org/). Tractograms are obtained using the SDSTREAM deterministic tracking algorithm of MRTrix3 [TCC12]. All tracts employed in the experiments (Uncinate Fasciculus, Ifof and Thalamocortical bundle) are extracted using either WMQL [WMR+ 16] or manual segmentation. Please note that our method is general and could be used with any streamline tractography algorithm.

Comparison with QuickBundles In Fig. 3, we compare our method with QuickBundles (QB) [GBC+12], a well-known approximation algorithm for brain white matter tractograms based on prototypes. We first executed QB on a bundle composed of 19,782 fibers. We used different thresholds in the range 5-10mm as suggested in [GBC+12]. We chose the middle one (7) for comparison. It resulted in 275 prototypes. We then executed our algorithm (once) with the MDF metric, used by QB, and then interactively change the resolution to obtain Fig. 3(c,d and e). Our result, in Fig. 3e, has as many cylinders as prototypes in Fig. 3b. The proposed method, after a single and fast pre-computation (see Tab. 1), creates an encompassing representation that well approximates the original bundle at each level of resolution. On the contrary, QB produces prototypes that, depending on a user-defined threshold, might not preserve the overall structure and volume of the original bundle. Moreover, one might need to execute QB several times before finding an optimal threshold (for a given clinical/research application).

Multi-resolution Figure 4 shows results on three different bundles: the Uncinate Fasciculus (a, b), the Ifof (c), and a whole brain (d). The measures used for the experiments are MC (see (1)) for (a), WC (see (2)) for (b and d), with $\sigma_a = 6\text{mm}$, $\sigma_b = 6\text{mm}$ and $\sigma_e = 8\text{mm}$, and MDF for (c). Computation times for these experiments are given in Tab. 1 and were obtained on an Intel Xeon E5-1650V4. These results suggest that the overall structure of the bundles is preserved across the multi-resolution representation, even at low resolutions, e.g. 15%. The graphs on the right of Fig. 4, illustrate the mean and max valences across the resolutions. The valence of a fiber is defined as the number of its neighbors. The evolution of the valence depends on the order in which the fibers are merged, and therefore on the employed distance/similarity measure. A measure that favors geometrically well distributed cylinders – composed of a similar number of original fibers – preserves a bounded valence. In this case, the size of the priority list remains linear in the number of original fibers $n$, and our strategy exhibits a total time complexity of $O(n \log(n))$. These complexities were observed in our experiments (see Tab. 1). In this table and in Fig. 4, we notice that the mean valence remains relatively stable using all distance measures, even with one million of fibers (Fig. 4d). This suggests that our method, with the proposed dissimilarities, should preserve the overall structure of the brain since fibers are aggregated all around the tractogram and not only in a specific area.

The proposed method results in a multi-resolution representation which approximates the original tractogram with a decreasing precision, and is not intended to produce anatomically reproducible clusters among subjects. Moreover, it is initialized with a Delaunay tetrahedralization of the extremities, which means that we inherently assume that two fibers are similar if their extremities are close to each other. The employed metric, as the ones proposed here, should therefore be consistent with this assumption.

For similarity measures defined on inner product spaces, we also propose an automatic stopping criterion to prevent oversimplification (e.g. a single cylinder). Two cylinders are not merged together if they are orthogonal (or almost) to each other. Furthermore, all fibers not merged at the end of the algorithm (and cylinders representing very few fibers) can be considered as outliers and thus discarded.

4. Conclusion

We introduced a new multi-resolution representation for brain tractograms that reduces the redundancy, easing the visualization and interpretation. It supports any similarity measure between fibers.
Figure 3: a) Thalamocortical bundle with 19,782 fibers, b) reduced to 275 prototypes with QuickBundles (Threshold=7mm), c), d) and e) reduced to 6925, 1422 and 275 cylinders with our method.

Figure 4: Bundles at different resolutions: a) Uncinate Fasciculus, computed with the mean of closest distances; b) Uncinate Fasciculus, computed with the dissimilarity of the weighted currents; c) Ifof bundle, computed with the dissimilarity of the weighted currents; d) whole brain tractogram, with 1 million fibers, computed with the dissimilarity of the weighted currents. The graphs represent the valence of the fibers in function of the resolution. The maximum valence is in red, and the average valence is in blue.
Table 1: Maximum and average valence with computational times for some bundles. The computational time for QuickBundles is presented on the last line as comparison. The threshold used for obtaining a single simplification was 7mm.

<table>
<thead>
<tr>
<th>Bundle</th>
<th>Metric</th>
<th>Number of fibers</th>
<th>Maximum valence</th>
<th>Average valence</th>
<th>Tetrahedralization time (s)</th>
<th>Total time (s)</th>
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<tbody>
<tr>
<td>Uncinate Fasciculus</td>
<td>MC</td>
<td>940</td>
<td>85</td>
<td>25.0 ± 3.2</td>
<td>0.01</td>
<td>1.3</td>
</tr>
<tr>
<td></td>
<td>WC</td>
<td>940</td>
<td>98</td>
<td>23.8 ± 3.2</td>
<td>0.01</td>
<td>1.9</td>
</tr>
<tr>
<td>If of</td>
<td>MC</td>
<td>1,983</td>
<td>217</td>
<td>28.3 ± 4.3</td>
<td>0.02</td>
<td>4.3</td>
</tr>
<tr>
<td></td>
<td>WC</td>
<td>1,983</td>
<td>197</td>
<td>27.7 ± 3.4</td>
<td>0.02</td>
<td>6.9</td>
</tr>
<tr>
<td></td>
<td>MDF</td>
<td>1,983</td>
<td>216</td>
<td>29.0 ± 3.8</td>
<td>0.02</td>
<td>1.7</td>
</tr>
<tr>
<td>Thalamocortical bundle</td>
<td>MC</td>
<td>19,782</td>
<td>760</td>
<td>30.3 ± 3.3</td>
<td>0.28</td>
<td>34.0</td>
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<tr>
<td></td>
<td>WC</td>
<td>19,782</td>
<td>717</td>
<td>29.3 ± 2.5</td>
<td>0.28</td>
<td>55.6</td>
</tr>
<tr>
<td>Whole Brain</td>
<td>MC</td>
<td>1,000,000</td>
<td>9,392</td>
<td>29.6 ± 2.5</td>
<td>15.42</td>
<td>1,893.7</td>
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<tr>
<td>Tractogram</td>
<td>WC</td>
<td>1,000,000</td>
<td>1,331</td>
<td>27.6 ± 1.8</td>
<td>15.42</td>
<td>3,094.2</td>
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<tr>
<td></td>
<td>MDF</td>
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<td>373</td>
<td>29.9 ± 2.7</td>
<td>15.42</td>
<td>616.3</td>
</tr>
<tr>
<td></td>
<td>QB (MDF)</td>
<td>1,000,000</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>4,825.2</td>
</tr>
</tbody>
</table>

We also proposed a method to determine adequate candidates for efficient merging of groups of fibers. Our progressive method makes it possible to display the input tractogram at any resolution in a continuous and real-time way. From a technical point of view, our two main contributions are a multi-resolution representation for tractograms based on a progressive decimation algorithm, and a combinatorial strategy based on a Delaunay tetrahedralization to make it computationally tractable. Visualizing groups of similar fibers as single generalized cylinders and being able to easily change the level of resolution may be very useful for clinicians. For instance, it can help neurosurgeons identify relevant anatomical tracts which should not be severed during the operation, thus reducing post-operative complications and improving the clinical outcome. It is to note that we received very positive feedbacks from our neurosurgeons colleagues of the Ste Anne hospital in Paris.

Future Work: To improve visualization, in particular the 3D perception of the bundles, ambient occlusion could be efficiently implemented, exploiting the hierarchical structure. Furthermore, we also plan to expand the proposed representation by adding functional information defined, for instance, as a scalar field (e.g. Fractional Anisotropy (FA)) which could be visualized using color or texture.

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