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Assessing the Feasibility of Estimating Axon Diameter using Diffusion Models and Machine Learning

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ABSTRACT

Axon diameter estimation has been a focus of the diffusion MRI community for the past decade. The main argument has been that while diffusion models always overestimate the true axon diameter, their estimation still correlates with changes in true value. Until now, this remains more as a discussion point. The aim of this paper is to clarify this hypothesis using a recently acquired cat spinal cord data set, where the diffusion MRI signal of both a multi-shell and AxCaliber acquisition have been registered with the underlying histology values. We find that the axon diameter as estimated by signal models and AxCaliber does not correlate with their true sizes for axon diameters smaller than 3 µm. On the other hand, we also train a random forest machine learning algorithm to map signal-based features to histology values of axon diameter and volume fraction. The results show that, in this dataset, this approach leads to a more reliable estimation of physically relevant axon diameters than using sophisticated diffusion models.

Index Terms— Diffusion MRI, Axon Diameter, Validation, Histology, MAP-MRI, 1D-SHORE, AxCaliber, Machine Learning, Random Forest

1. INTRODUCTION

For over twelve years has axon diameter estimation been a subject of interest in the diffusion MRI community [1]. However, recent works suggest that the diffusion signal is not sensitive enough to variations in axon diameters within the physiological range [2, 3]. We attempt to investigate this claim by using a recent public data set of a cat spinal cord [4], where both high-quality diffusion MRI acquisitions and underlying histology values are registered to the same template.

Our aim is to verify whether histological variations in mean axon diameter correlate with estimates produced by various state-of-the-art diffusion MRI models. We evaluate the AxCaliber multi-compartment model [5], as well as “apparent axon diameter” estimates by signal models such as Mean Apparent Propagator (MAP)-MRI [6] and 1D-Simple Harmonic Oscillator Reconstruction and Estimation (1D-SHORE) [7]. We produce these correlations for both the dataset as a whole and for separate segmentation of the data based on different axon diameter ranges. In this way, we can evaluate for which axon diameter ranges these methods are actually sensitive.

We also evaluate a machine learning approach to axon diameter estimation [8], where we use a random forest algorithm to learn a direct mapping between signal-based metrics and the underlying histology. As we will show, this algorithm picks up on correlations in the data that allows us to predict not only axon diameter, but also intra-axonal and even myelin volume fractions, using metrics that on their own do not have this interpretation.

2. MATERIALS

We base our experiments on a recent dataset where both AxCaliber and multi-shell diffusion MRI acquisitions have been registered to one axial slice of cat spinal cord [4]. The data is 64 × 64 voxels with resolution 0.16 × 0.16 × 0.16 mm³. Histology: For every voxel, the mean axon diameter, restricted volume fraction, and myelin volume fraction is known, see top Fig. 1. We segment the data into 4 ROIs with diameters [1 − 2; 2 − 3; 3 − 4; > 4] µm, see top-right Fig. 1.

Diffusion MRI: The data was scanned on an Agilent 7T animal scanner equipped with 600 mT/m gradients. A single shot EPI sequence was used: BW=250kHz, TR=2s. One AxCaliber acquisition was acquired (perpendicular to the axon axis) with parameters δ=3/8/8/8 ms, ∆=7/12/25/40 ms, G = [0 .. 849] mT/m (199 increments) and TE minimized (36 - 62ms). The data was TE-normalized by dividing the data for every TE by the G=0 signal. The multi-shell acquisition was acquired with parameters δ=3ms, ∆=30ms, 4 shells with bvalue=[40,189,1680,6720] s/mm², TE=47ms, with a total of 796 diffusion weighted images.
3. AXON DIAMETER ESTIMATION

Diffusion models relate the measured diffusion MRI model to axon diameters by making several assumptions on both the dMRI acquisition and tissue composition. An essential one is modeling axons as an ensemble of parallel cylinders, which is reasonably accurate in our spinal cord data. We compare signal-based and compartment-based approaches, whose particular model assumptions we clarify in this section.

3.1. Axon diameter using signal models

Signal models can relate the diffusion signal to the “apparent axon diameter” (AAD) by calculating the Return-To-Axis probability (RTAP) of the Ensemble Average Propagator (EAP) as $AAD = \frac{2}{\sqrt{\pi}} \frac{1}{RTAP}$ [7, 6]. The EAP $P(r)$ with $r$ the 3D displacement vector is estimated as the Fourier Transform (FT) of the measured signal attenuation $E(q)$ with $q$ the 3D q-space vector. The FT is facilitated by representing $E(q)$ with a Fourier basis expansion

$$\hat{E}(q) = \sum_{i=0}^{N_q} c_i \hat{\Phi}_i(q) \overset{\text{FT}}{\Rightarrow} \hat{P}(r) = \sum_{i=0}^{N_q} c_i \hat{\Psi}_i(r)$$

with $\Phi = FT(\Psi)$ and $c$ the basis coefficients. The AAD is called “apparent” because this interpretation relies on several strong assumptions that the tissue is composed of only intraaxonal space, the gradient pulse length is infinitesimally small ($\delta \approx 0$), and the diffusion time is long enough for diffusing particles to be restricted.

We estimate the AAD using Laplacian-regularized MAP-MRI [9, 10] on the multi-shell acquisition, and 1D-SHORE [7] on the $\Delta = 30$ms segment of the AxCaliber one to have the same $\Delta$ between the two. MAP-MRI is the 3D-generalization of 1D-SHORE, so results are expected to be similar.

3.2. Axon diameter using compartment models

Compartment models separate the signal contributions of the intra- and extra-axonal space by fitting a combination of biophysical models. We consider the AxCaliber model [5] for the AxCaliber acquisition. This model represent the measured signal $E(q, \Delta, \delta)$ as

$$E(q, \Delta, \delta) = f \cdot E_{\text{intra}}(q, \Delta, \delta) + (1 - f) \cdot E_{\text{extra}}(q, \Delta, \delta)$$

where $E_{\text{intra}}$ is given by the Van Gelderen cylinder model [11] and $E_{\text{extra}}$ by a Gaussian. We fit AxCaliber with 3 free parameters; mean axon diameter $\langle D \rangle$, intra-axonal volume fraction $f$ and extra-axonal diffusivity $D_\perp$.

3.3. Axon Diameter using Random Forest Regression

We also include a Random Forest (RF) regression approach to estimate axon diameters from an ensemble of randomly trained decision trees [8]. From the multi-shell acquisition, we use it to learn a mapping between rotationally invariant signal-based features, derived from DTI [12] and MAP-MRI, to the ground truth histology parameters. For every shell separately we compute the DTI metrics Fractional Anisotropy (FA); Mean Diffusivity (MD); parallel and perpendicular diffusivity ($D_\parallel$ and $D_\perp$). Using the multi-shell data at once we also compute the MAP-MRI metrics Return-to-Origin, Return-to-Axis and Return-to-Plane Probability (RTOP, RTAP, RTPP); overall, parallel and perpendicular non-Gaussianity ($NG, NG_\parallel, NG_\perp$); and Propagator Anisotropy (PA). This results in 26 features per voxel.

We train the RF regressor, containing 100 trees of maximum depth 20, on a random selection of 90% of all voxels in the mask and predict the histology parameters on the remaining 10%. We repeat this process 10 times, predicting a different 10% of the data each time, such that every voxel has...
In Table 2 we show that we can use RF regression to not only predict histology-based axon diameter, but also restricted volume fraction and even myelin volume fraction. Note here that the RF algorithm only picks up on correlations between the input signal features and the output histology parameters. This is emphasized by the observation that we use signal models to predict myelination, despite the fact that myelin has no diffusion signal contribution.

Our experiment evaluates which features, from which models and which shells of the multi-shell acquisition, produce the best prediction of the tissue histology. From top to bottom in Table 2, for every histology metric, we predict histology values using DTI features of 1) only the first shell \((b = 41s/mm^2)\); 2) the last shell \((b = 6720s/mm^2)\); 3) all shells combined; 4) only MAP-MRI on all shells; and 5) all DTI and MAP-MRI features combined.

The first thing to notice is that for any feature subset the Pearson correlation is always very high \((\rho_{\text{min}} = 0.84)\) with p-values close to zero. Using only DTI features of shell 1 produces the lowest \(\rho\) for all histology parameters, with FA and \(D_\parallel\) consistently the most relevant features. Using only the DTI features of shell 4 not only raises \(\rho\) in all cases, but also changes the ordering of relevant features for every histology parameter. Using the DTI metrics of all shells together raises the correlation to nearly linear with \(\rho = 0.96\) and up. The relevant features show that in this case, the DTI-metrics derived from the shells with the highest b-values are the most important. Using only MAP-MRI features also produces similar nearly linear correlations. Finally, using all features together does not further increase \(\rho\), but the most relevant features do show that MAP-MRI features are never the most important in predicting histology parameters.

### 5. DISCUSSION AND CONCLUSION

In this work we assessed, using a spinal cord data set with known ground truth histology, whether diffusion model-derived estimates of axon diameter are sensitive to axon diameter changes within the physically relevant range. While the results on this spinal cord data set don’t directly generalize to brain tissue, it does give a good idea about what performance we can expect in similar nervous tissues. The results show that even the most sophisticated dMRI models have trouble in being sensitive to axon diameters smaller than \(3\mu m\). This was not previously pointed out by \([4]\). As we

<table>
<thead>
<tr>
<th>ROI</th>
<th>Method</th>
<th>(\rho, p)</th>
<th>(\mu \pm \sigma, \rho, p)</th>
<th>(\mu \pm \sigma, \rho, p)</th>
<th>(\mu \pm \sigma, \rho, p)</th>
<th>(\mu \pm \sigma, \rho, p)</th>
</tr>
</thead>
<tbody>
<tr>
<td>All</td>
<td>1(\mu m - 2\mu m)</td>
<td>2(\mu m - 3\mu m)</td>
<td>3(\mu m - 4\mu m)</td>
<td>&gt;4(\mu m)</td>
<td>&gt;4(\mu m)</td>
<td></td>
</tr>
<tr>
<td>1D-SHORE</td>
<td>-0.04,3.3e-1</td>
<td>5.7±0.5, 0.58, 1.2e-1</td>
<td>5.5±0.5, -0.60, 1.8e-3</td>
<td>4.7±0.4, -0.08, 2.5e-1</td>
<td>4.8±0.3, 0.40, 1.9e-16</td>
<td></td>
</tr>
<tr>
<td>AxCaliber</td>
<td>0.26,7.4e-11</td>
<td>5.5±0.2, -0.04,9.2e-1</td>
<td>5.3±0.3, -0.79,3.5e-6</td>
<td>5.2±0.2,0.38,6.1e-8</td>
<td>5.2±0.2,0.55, 2.5e-32</td>
<td></td>
</tr>
<tr>
<td>MAPMRI</td>
<td>0.01,7.7e-1</td>
<td>10.6±1.6,0.28,4.9e-1</td>
<td>11.7±1.5, -0.05,8.1e-1</td>
<td>10.0±1.1,-0.25,6e-4</td>
<td>10.4±0.9, 0.23, 5.1e-6</td>
<td></td>
</tr>
</tbody>
</table>

4. RESULTS

4.1. Correlations of Signal and Compartment Models

We show the predicted axon diameters of AxCaliber, the AAD of 1D-SHORE and MAP-MRI and RF prediction in Fig. 1. As expected, all approaches significantly overestimate the axon diameter compared to their actual values top-left, except for RF. Of the modeling approaches, the AxCaliber result appears to be most similar to histology, showing the same general pattern. On the other hand, AAD estimates from both signal-based models appear quite flat compared to histology. The RF approach is closest to the ground truth.

Taking a closer look at the modeling approaches in Table 1, we show the mean and standard deviation of the axon diameter estimates within the axon diameter-based tissue segmentation in Fig. 1-top-right. We also produce the Pearson correlation with the histology diameters \((\rho > 0\)) and the two-tailed p-value \((p < 0.05\)) with histology the numbers are bold-faced.

4.2. Histology prediction through Machine Learning

In Table 2 we show that we can use RF regression to not only predict histology-based axon diameter, but also restricted volume fraction and even myelin volume fraction. Note here that the RF algorithm only picks up on correlations between the input signal features and the output histology parameters. This is emphasized by the observation that we use signal models to predict myelination, despite the fact that myelin has no diffusion signal contribution.

Our experiment evaluates which features, from which models and which shells of the multi-shell acquisition, produce the best prediction of the tissue histology. From top to bottom in Table 2, for every histology metric, we predict histology values using DTI features of 1) only the first shell \((b = 41s/mm^2)\); 2) the last shell \((b = 6720s/mm^2)\); 3) all shells combined; 4) only MAP-MRI on all shells; and 5) all DTI and MAP-MRI features combined.

The first thing to notice is that for any feature subset the Pearson correlation is always very high \((\rho_{\text{min}} = 0.84)\) with p-values close to zero. Using only DTI features of shell 1 produces the lowest \(\rho\) for all histology parameters, with FA and \(D_\parallel\) consistently the most relevant features. Using only the DTI features of shell 4 not only raises \(\rho\) in all cases, but also changes the ordering of relevant features for every histology parameter. Using the DTI metrics of all shells together raises the correlation to nearly linear with \(\rho = 0.96\) and up. The relevant features show that in this case, the DTI-metrics derived from the shells with the highest b-values are the most important. Using only MAP-MRI features also produces similar nearly linear correlations. Finally, using all features together does not further increase \(\rho\), but the most relevant features do show that MAP-MRI features are never the most important in predicting histology parameters.
Table 2. Results of random forest regression algorithm for different subsets of signal features. We show Pearson correlation ($\rho$), p-value and the three most relevant features in the prediction for axon diameter, restricted volume fraction and myelin volume fraction.

<table>
<thead>
<tr>
<th>Axon Diameter</th>
<th>Most Relevant Features</th>
<th>Features</th>
<th>$\rho$</th>
<th>p-value</th>
<th>#1</th>
<th>#2</th>
<th>#3</th>
</tr>
</thead>
<tbody>
<tr>
<td>DTI Shell 1</td>
<td></td>
<td></td>
<td>0.90</td>
<td>2e-215</td>
<td>FA-S1</td>
<td>D$_{\parallel}$-S1</td>
<td>D$_{\perp}$-S1</td>
</tr>
<tr>
<td>DTI Shell 4</td>
<td></td>
<td></td>
<td>0.94</td>
<td>1e-282</td>
<td>MD-S4</td>
<td>D$_{\parallel}$-S4</td>
<td>D$_{\perp}$-S4</td>
</tr>
<tr>
<td>DTI Shell 1-4</td>
<td></td>
<td></td>
<td>0.96</td>
<td>0</td>
<td>MD-S4</td>
<td>D$_{\parallel}$-S4</td>
<td>FA-S3</td>
</tr>
<tr>
<td>MAP</td>
<td></td>
<td></td>
<td>0.96</td>
<td>0</td>
<td>RTPP</td>
<td>NG</td>
<td>MD-S3</td>
</tr>
<tr>
<td>DTI + MAP</td>
<td></td>
<td></td>
<td>0.96</td>
<td>0</td>
<td>MD-S4</td>
<td>NG</td>
<td>MD-S3</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Restricted Volume Fraction</th>
<th>Most Relevant Features</th>
<th>Features</th>
<th>$\rho$</th>
<th>p-value</th>
<th>#1</th>
<th>#2</th>
<th>#3</th>
</tr>
</thead>
<tbody>
<tr>
<td>DTI Shell 1</td>
<td></td>
<td></td>
<td>0.87</td>
<td>9e-191</td>
<td>FA-S1</td>
<td>D$_{\parallel}$-S1</td>
<td>MD-S1</td>
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<tr>
<td>DTI Shell 4</td>
<td></td>
<td></td>
<td>0.94</td>
<td>5e-295</td>
<td>FA-S4</td>
<td>MD-S4</td>
<td>D$_{\parallel}$-S4</td>
</tr>
<tr>
<td>DTI Shell 1-4</td>
<td></td>
<td></td>
<td>0.97</td>
<td>0</td>
<td>FA-S4</td>
<td>D$_{\parallel}$-S3</td>
<td>D$_{\perp}$-S2</td>
</tr>
<tr>
<td>MAP</td>
<td></td>
<td></td>
<td>0.96</td>
<td>0</td>
<td>RTPP</td>
<td>NG$_{\perp}$</td>
<td>RTPP</td>
</tr>
<tr>
<td>DTI + MAP</td>
<td></td>
<td></td>
<td>0.96</td>
<td>0</td>
<td>FA-S4</td>
<td>D$_{\parallel}$-S3</td>
<td>D$_{\perp}$-S2</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Myelin Volume Fraction</th>
<th>Most Relevant Features</th>
<th>Features</th>
<th>$\rho$</th>
<th>p-value</th>
<th>#1</th>
<th>#2</th>
<th>#3</th>
</tr>
</thead>
<tbody>
<tr>
<td>DTI Shell 1</td>
<td></td>
<td></td>
<td>0.84</td>
<td>6e-162</td>
<td>FA-S1</td>
<td>D$_{\parallel}$-S1</td>
<td>MD-S1</td>
</tr>
<tr>
<td>DTI Shell 4</td>
<td></td>
<td></td>
<td>0.95</td>
<td>6e-299</td>
<td>MD-S4</td>
<td>FA-S4</td>
<td>D$_{\parallel}$-S4</td>
</tr>
<tr>
<td>DTI Shell 1-4</td>
<td></td>
<td></td>
<td>0.96</td>
<td>0</td>
<td>D$_{\parallel}$-S3</td>
<td>FA-S4</td>
<td>MD-S4</td>
</tr>
<tr>
<td>MAP</td>
<td></td>
<td></td>
<td>0.96</td>
<td>0</td>
<td>RTPP</td>
<td>NG$_{\perp}$</td>
<td>NG</td>
</tr>
<tr>
<td>DTI + MAP</td>
<td></td>
<td></td>
<td>0.97</td>
<td>0</td>
<td>D$_{\parallel}$-S3</td>
<td>FA-S4</td>
<td>RTPP</td>
</tr>
</tbody>
</table>

showed here, using a machine learning approach like Random Forest regression can overcome this limitation. Though, machine learning prediction is also limited in that it is hard to generalize to data that was not included in the training set. But similarly, diffusion MRI models also cannot produce accurate results in data for which their model assumptions are not met. This is particularly evident in our signal model results using MAP-MRI and 1D-SHORE, who have the strongest tissue assumptions. Their AAD estimates were found to have Pearson correlations close to zero overall in the data, but did correlate positively in voxels with diameters larger than 4$\mu$m. Moreover, AxCaliber showed to correlate better with histology for smaller axon diameters compared to the multi-shell techniques. This might also be a consequence of a lower axon diameter resolution limit [3] due to the higher gradient strength available (849 mT/m vs 600 mT/m), not only the model assumptions. Nonetheless, the results clearly show that, in this particular (limited) data set of one slice, the machine learning approach vastly outperformed the modeling approaches in the estimation of histology parameters, which should be considered in future works.

Acknowledgments

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