Visualizing Diffusion Tensor Images of the Mouse Spinal Cord

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Abstract—
Within biological systems water molecules undergo continuous stochastic Brownian motion. The rate of this diffusion can give clues to the structure of underlying tissues. In some tissues the rate is anisotropic – faster in some directions than others. Diffusion-rate images are second-order tensor fields and can be calculated from diffusion-weighted magnetic resonance images. A 2D diffusion tensor image (DTI) and an associated anatomical scalar field, created during the tensor calculation, define seven dependent values at each spatial location. Understanding the interrelationships among these values is necessary to understand the data. We present two new methods for visually representing DTIs. The first method displays an array of ellipsoids where the shape of each ellipsoid represents one tensor value. The novel aspect of this representation is that the ellipsoids are all normalized to approximately the same size so that they can be displayed in context. The second method uses concepts from oil painting to represent the seven-valued data with multiple layers of varying brush strokes. Both methods successfully display most or all of the information and are easier to read quantitatively. We demonstrate the methods on images of the mouse spinal cord. The visualizations show significant differences between spinal cords from mice suffering from Experimental Allergic Encephalomyelitis (EAE) and spinal cords from wild-type mice. The differences are consistent with pathology differences shown histologically and suggest that our new non-invasive imaging methodology and visualization of the results could have early diagnostic value for neurodegenerative diseases.

I. INTRODUCTION
Within biological systems water molecules undergo continuous stochastic Brownian motion. In different tissues the rate of this diffusive motion can vary by several orders of magnitude – faster in liquids like cerebro-spinal fluid, slower in tissues like muscle. In some tissues the rate is anisotropic, or faster in some directions compared to others. Magnetic resonance imaging (MRI) can acquire images with intensity values sensitive to the diffusion rate of water. A quantitative image of the diffusion rate can be calculated from a set of such MR images [1], [2], [3], [4]. From a 2D slice or 3D volume image of this directionally dependent diffusion rate we can infer underlying tissue structure and better understand the anatomy of the nervous system, neuro-degenerative diseases, and neural development [1].

Diffusion-rate images calculated from MRI measurements are second-order tensor fields. A diffusion tensor image (DTI) and the associated anatomical scalar field, calculated as part of the tensor calculation, define seven dependent values at each spatial location. Often it is useful to simultaneously display more than a few of these values in order to visualize interrelationships among them and to understand the data. We present two new methods for visually representing these diffusion tensor images. The first method, shown in Fig. 1, displays an array of ellipsoids where the shape of each ellipsoid represents one tensor value. The novel aspect of this representation is that the ellipsoids are all normalized to approximately the same size so that they can be displayed in context. The second method, shown in Fig. 2, uses concepts from oil painting to represent the seven-valued data with layers of varying brush strokes. The methods are exploratory – they allow a viewer to explore all of the values comprising a tensor-valued dataset rather than distilling out some simpler derived quantity. Both are implemented as new modules within AVS[5].

In the remainder of the paper we first discuss related work in visualizing second-order tensor fields. We then describe the anatomy of the mouse spinal cord and the pathology of the mouse neurological disease that we use in our example images. Section IV describes diffusion tensor images ac-
Fig. 1. Normalized diffusion ellipsoid visualization. Each ellipsoid represents the tensor value at one pixel geometrically to capture most of the information in the tensor-valued image.

Fig. 2. Diffusion tensor image (DTI) visualization using concepts from painting. This stroke image is composites from layers as shown in Figs. 9 and 10. Component data values are represented in a blue-purple underpainting; in the direction, shape, color, and transparency of a layer of elliptical strokes over that; and in the frequency of a texture applied to the elliptical strokes. The resulting image displays all seven dependent values that comprise a DTI.
quired using MRI. Section V describes the ellipsoid-based method and Section VI describes the painting-motivated method. We summarize and conclude in Section VII.

II. RELATED WORK

References [6], [7], [8], [9], [10] review specific methods for tensor-field visualization and develop several tensor-field visualization methods. Portions of references [11], [12] also give informative reviews of methods. However, these methods are all designed for displaying tensor fields with a different physical structure and interpretation than the diffusion tensor. The methods do not take advantage of the fact that a diffusion tensor has orthogonal eigenvectors and non-negative eigenvalues, and thus devote critical visual bandwidth to tensor attributes not present in diffusion tensors. They also produce visual representations that intuitively represent the physical phenomena that they are representing, but which do not represent diffusion intuitively.

Diffusion tensor imaging data have primarily been displayed by extracting scalar- or vector-valued components of the tensor at each point and visualizing the resulting scalar fields and vector fields [13], [14]. The primary limitation with this approach is that it is difficult to understand the tensor field from individual component images because relationships among the different components are spread across several images. For example, imagine trying to understand a 2D image of 3D vectors by viewing three grayscale images of the $x$, $y$, and $z$ components of the vector. It would be difficult to understand the 3D direction of the vectors without a representation that combines the three values. Another limitation in viewing the components individually is that noise in the acquired images can bias some of the extracted values when they are viewed out of the context of the other values [14].

Arrays of ellipsoids have also been used to represent diffusion tensor data [13], [14]. They have been limited to small regions of interest ($16 \times 16$ pixels) each image provides an iconic representation of 256 tensor values. Fortuitously, the regions displayed have diffusion rates that differ by less than a factor of 10. However, the smaller tensor values are represented by sparsely spaced small ellipsoids, and the connection between the different values is lost. For regions where the diffusion rate is more widely varying this effect would be compounded and the ellipsoids for low-rate tensor values would be all but invisible.

Reference [15] was the first to experiment with painterly effects in computer graphics. More recently reference [16] extended the approach for animation and further refined the use of layers and brush stroke characteristics for creating effective imagery. Both of these efforts were aimed toward creating art, however, and not toward visualizing scientific data. Along similar lines, references [17], [18], [19] used software to create pen and ink illustrations for artistic purposes.

Layering has been used in scientific visualization to show multiple items. In reference [20], [21], transparent stroked textures show surfaces without completely obscuring what is behind them. These results are related to ours, but our application is 2D, and so our layering is not as spatial as it is in the 3D case. Our layering is more in the spirit of oil painting, where layers are used more broadly, often as an organizing principle for a painting.

Much of the inspiration for our approach comes from studying the painting styles of a number of impressionist painters and from consulting books on basic artistic principles and problem solving in painting. We found Van Gogh’s style particularly applicable because he used expressive, discrete strokes that are combined to represent a continuous scene. They can be read differently from different distances and can encode different information from each of those viewpoints.

The goal of our work is to visualize scientific information by building on many of these concepts. From the sources above we discovered many useful concepts and rules-of-thumb for creating effective images. Just as when an artist creates a painting, our goal was to communicate complexity in a direct, expressive, and visually pleasing manner. We used these principles as guidelines for creating our visualization methods.

There is a large body of literature on the perceptual effects of texture, color, icons, and other visual representations of scientific data [22], [23], [24]. In our painting-based work, visual choices were based not on this visualization perception literature, but instead on the artistic guidelines. The effects of many of the components of our visualizations undoubtedly are explained in the visualization perception litera-
III. Anatomy and Pathology of Examples

Our visualization examples are taken from images of the spinal cords of healthy and diseased mice. The diseased mice suffer from Experimental Allergic Encephalomyelitis (EAE), a disease where the mouse immune system attacks the central nervous system [25]. Many aspects of the pathology of EAE mimic multiple sclerosis in humans. The disease causes a stripping away of the insulating myelin sheath (demyelination) of axons which comprise the white matter of the spinal cord. A large amount of inflammation is also observed. EAE causes behavioral symptoms that progress in well-characterized stages: 1) limp tail, 2) uneven gait, 3) hindlimb paralysis, 4) fore and hindquarter paralysis, and 5) moribundity and death.

Fig. 3 shows the anatomy of a spinal cord section in a healthy wild-type mouse. Fig. 4 shows a histological section that has been stained with toluidine blue which labels myelin with a dark blue color. The light-blue butterfly-shaped region in the center of the spinal cord consists of gray matter containing mostly neuronal cell bodies. The darker-blue cardiod-shaped region surrounding the gray matter is white matter. The myelinated axons in this region connect the brain and other levels of the spinal cord. The cardiod shape is the boundary of the spinal cord. Outside of the cord is a layer of spinal nerves that connect the central nervous system to other parts of the body.

IV. Diffusion Tensor Images

The diffusion tensor images used in our visualizations were obtained using novel MRI and numerical methods described in [4]. The data are in the form of a 2D slice image with a second-order tensor value at each pixel. Each second-order tensor value can be represented by a symmetric matrix with non-negative eigenvalues. It can also be represented geometrically by an ellipsoid. The matrix and ellipsoid representations are equivalent [26]. The principal axes of an ellipsoid are the eigenvectors \((n_1, n_2, n_3)\) of the matrix and the radii of an ellipsoid are the eigenvalues \((\lambda_1, \lambda_2, \lambda_3)\) of the matrix. The shape and size of the ellipsoid encodes key aspects of the diffusion tensor values we wish to visualize. It can be interpreted as the shape that a small dot of ink would take on after some fixed diffusion time. Larger ellipsoids represent faster diffusion, spheres represent isotropic diffusion, and eccentric (prolate or oblate) ellipsoids represent anisotropic diffusion.

In the spinal cord, gray matter has nearly isotropic diffusion whereas white matter has very anisotropic diffusion as observed in MRI measurements [1]. In white matter water diffuses more readily along the axon than across it. The pathology of EAE leads to changes in the diffusion characteristics of the spinal cord, especially in certain regions of white matter. We intuitively show these pathologic changes with the help of our visualization methods.

Each sub-image in Fig. 5 shows one element of the matrix representation of a diffusion tensor image. The matrix elements are difficult to interpret intuitively as a tensor-valued image. One reason for this difficulty is that they are not rotationally invariant; they change if the sample is oriented differently. These images are analogous images of the scalar
Fig. 5. Matrix representation of a diffusion tensor image. All of the scalar elements are shown. The upper left image represents the upper left matrix element for each pixel. Understanding the tensor field from this collection of scalar-valued images is not intuitive.

components of a vector. Fig. 6 shows images of the eigenvalues of the same diffusion tensor image. The scalar value in these images are rotationally invariant but display no directional information. These images are analogous to an image of the magnitude of a vector. The directional information that is missing is a significant part of the tensor value that we wish to display.

In addition to the tensor-valued diffusion-rate image, the numerical calculation of the tensor image from MR images also produces a scalar-valued anatomical image as shown in Fig. 7. We name it $I_0$. It shows contrast between anatomical regions using a non-diffusion mechanism, and thus captures additional information.

V. NORMALIZED ELLIPSOID REPRESENTATION

Our first new visualization method for diffusion tensor images captures most of the tensor information into a single image. It takes advantage of the fact that a tensor value with positive eigenvalues can be represented with an ellipsoid. Our first attempts made a direct mapping from tensor values to ellipsoids, but, because the diffusion rates vary by factors of almost a thousand, so did the sizes of the ellipsoids. The ellipsoids could not all be seen in their appropriate locations with such a large variation in size.

We avoid this extreme variance in tensor values by normalizing each tensor value so that its maximum eigenvalue is 1.0. The resulting images, shown in Figs. 1 and 8, contain ellipsoids with their largest radii all equal. The planar grid of ellipsoids is lit from top center with a single directional light source and viewed with perspective projection at a small tilt.

Each ellipsoid has a clear spatial location, and their shapes can be interpreted. Isotropic tensor values, e.g., in the gray-matter regions of Fig. 1, are represented as spheres that completely obscure the background color. Anisotropic tensor values, e.g., in white-matter regions, are skinny and smaller because their largest radius is the same as the largest radius of the spheres but the other radii are much smaller. In these regions the background color shows through more.

Fig. 8 shows an image of a spinal cord from a diseased mouse. The pathology of the disease is apparent in the ventral white matter. The presence of EAE lesions appears to have reduced the diffusion anisotropy throughout much of the white matter and caused the ellipsoids to become much more spherical.
Fig. 8. Ellipsoid representation of diseased spinal cord. Note right-left asymmetries in this image. These and other differences between this image and Fig. 1 of a healthy cord show changes due to the disease. Significant changes are particularly noticeable in the ellipsoid shapes in the myelinated regions in the lower right part of the image.
than in comparable regions of the healthy cord shown in Fig. 1.

This visualization method shows different information at different distances. From farther away, neighboring ellipsoids visually blend together to create a texture that gives an almost-continuous qualitative impression of the underlying anatomical regions. Close up, the ellipsoids are distinct enough to see individually, and show the relative anisotropy and directional information more quantitatively.

These ellipsoid images encode most of the information from the tensor-valued diffusion rate images. They capture the spatial correlation between the tensor values and the underlying space, the orientation of the tensors, and the relative diffusion rates in each direction. They are missing the absolute diffusion rate, which was normalized out of the tensor values so that the ellipsoids would be a consistent size, and they are missing the scalar $I_0$ value.

VI. PAINTING-MOTIVATED REPRESENTATIONS

Our second method applies concepts from oil painting to display diffusion tensor images. We used multiple layers of brush strokes to represent the tensor image and the associated anatomical scalar image, $I_0$. The brush strokes reflect the geometric nature of values derived from the tensors and of the relationships among the values. Also, the use of an underpainting and saturated complementary colors evokes a sense of depth. Together, these painting concepts helped us create a visual representation for the data that encodes all of the data in a manner that allows us to explore the data for a more holistic understanding.

Fig. 2 shows a completed stroke image. We built the visualization in four layers which are illustrated in Figs 9 and 10. Table I summarizes the mapping of data values to image contents. The layers are discussed individually in the following paragraphs.

- **Underpainting** First, we wanted a layer to show the overall form or structure of the anatomy. This is analogous to the function of an underpainting in oil painting. The $I_0$ image provides this information and has the best signal-to-noise ratio (SNR) of any of our scalar variables. It is encoded in the lightness of the purple layer shown in the left image of Fig. 9.

- **Checkerboard layer** Second, we wanted to incorporate the underlying voxel size into the image. Without it, the image can give a false sense of resolution. The voxel-size representation needed to be sub-
Fig. 9. Underlayers of the visualization in Fig. 10. The images show an underpainting layer, a checkerboard layer composited on the underpainting, and a stroke layer composited over that. Fig. 10 shows a fourth and final layer composited over these that captures the diffusion rate in a texture along each stroke.

Fig. 10. Diffusion tensor image (DTI) visualization using concepts from painting of a spinal cord from a mouse with EAE. This stroke image is composited from layers shown in Fig. 9. It displays all seven dependent values that comprise a diffusion tensor image.
Stroke placement is random, with strokes arranged so that they do not overlap in the direction of their widths but do slightly in the direction of their lengths. Thus adjacent strokes create the impression of flow along their direction, similar to Van Gogh’s style in some cases. The flow gives a sense of the directional fibers within the axon tracts. Strokes are not placed where the SNR is insufficient to calculate an accurate tensor. The right image of Fig. 9 shows this stroke layer composited over the two underlying layers.

- **Stroke texture layer** Finally, because the absolute magnitude of the diffusion rate is not correlated with the anisotropy ratio, \( \frac{\lambda_1}{\lambda_3} \), we wanted to encode it in the fourth layer. It is less important than the other values, however, so we encoded it in a form that is interpreted less quickly. We represented the rate with the frequency of the texture on the strokes. Faster diffusion is represented with more stripes along a stroke, and slower diffusion with fewer. The higher-frequency textures create smaller features that are often used artistically to represent higher speeds. This intuitively represents the corresponding higher rate of diffusion.

The complete visualization is shown for a healthy cord in Fig. 2 and for a diseased cord in Fig. 10. In the healthy cord the anatomical regions are clearly distinguished. The anisotropy of the white matter and spinal nerves creates regions of narrow, opaque ellipse shapes. The regions of gray matter are shown with rounder, semi-transparent shapes. Unlike the ellipsoid representation, the stroke images distinguish the diffusion rate difference between white matter and spinal nerves. In the spinal nerves, the diffusion rate is faster, and shows in the higher-frequency texture on the strokes. This difference is consistent with the larger axon diameter of the spinal nerves as shown in histological sections. The differences in the anatomical scalar image also show through between the stroke shapes, particularly when the images are viewed from a distance, incorporating its non-diffusion information to distinguish different regions.

The pathology of the disease is apparent in the ventral white matter. The strokes near EAE lesions are much rounder and more transparent than in other white-matter regions, indicating less anisotropy. The stroke images show different information at different distances. From farther away, the underpainting, strokes, and texture all combine to give a qualitative impression of different anatomical regions. From a closer perspective, the texture becomes more apparent, and more subtle anatomical differences appear. At very close perspectives, more quantitative measures of the tensor can be seen – the anisotropy ratio (ellipse aspect ratio), or the absolute diffusion rate (texture frequency). The quantitative information is easier to read quantitatively from the stroke images than from the ellipsoid images because the stroke images are strictly 2D. There is no distortion from the perspective rendering, the projection from 3D to 2D, or the lighting calculation – all of the geometric information lies directly in the plane of the stroke image. Only the portions of the tensor mapped to less quantitative color and transparency are less easy to quantify.

A stroke image represents all of the information in a diffusion tensor image and in the associated scalar-valued anatomical image. It captures the spatial correlation between the tensor values and the underlying space, the orientation of the tensors, the relative diffusion rates in different directions, and the absolute diffusion rate.

**VII. Summary and Conclusions**

We have presented two new methods for visualizing 2D images of second-order tensor fields representing diffusion rates. Our methods create visually rich images that can represent many values at each spatial location in an image. With this richness we have captured most or all of the information present in the data. Through applying the visualization methods we hope to better understand biological systems imaged with diffusion tensor imaging.

The first method represents the tensor values with ellipsoids. It modifies the direct tensor-to-ellipsoid mapping by normalizing the ellipsoid sizes. As a result, the ellipsoids are all approximately the same size and show the tensor structure effectively everywhere within the image.

The second method we applied concepts borrowed from artists. Varied brush strokes and layering helped to simultaneously display many components of the data. Underpaintings showed form. We used brush strokes both individually, to encode specific values, and collectively, to show spatial connections and to generate texture and a sense of speed corresponding to the speed of diffusion. We used layering and contrast to create depth. Stroke size, texture, and contrast helped to define a focus within each im-
age and also suggested the viewing order for different parts of an image.

The images created with both methods are effective because they display many data values simultaneously. We know of no other methods that simultaneously display as many components of second-order tensor-valued data. Our methods also qualitatively represent the underlying phenomena intuitively and geometrically. From different perspectives, they show the data at different levels of abstraction – more qualitatively from a distance, more quantitatively up close. Finally, the images emphasize different data values to different degrees leading a viewer through the temporal process of understanding the relationship among them.

We have only begun to explore the space of visualization methods that borrow conceptually from painting. We believe that artistic methods, including those we have applied, will continue to lead to many new, effective, and powerful visualization algorithms for multi-valued scientific data from many disciplines.

REFERENCES


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Fig. 9. Ellipsoid representation of diseased spinal cord. Note right-left asymmetries in this image. These and other differences between this image and Fig. 1 of a healthy cord show changes due to the disease. Significant changes are particularly noticeable in the ellipsoid shapes in the myelinated regions in the lower right part of the image.

Fig. 11. Diffusion tensor image visualization using concepts from painting of a spinal cord from a mouse with EAE. This image is composited from layers shown in Fig. 9. It displays all seven dependent values that comprise a diffusion tensor image.
Fig. 4. Histology showing anatomy of a healthy wild-type mouse spinal cord section. The section has been stained with toluidine blue; dark blue regions contain myelin and correlate with regions of white matter.

Fig. 10. Underlayers of the visualization in Fig. 10. The images show an underpainting layer, a checkerboard layer composited on the underpainting, and a stroke layer composited over that. Fig. 10 shows a fourth and final layer composited over these that captures the diffusion rate in a texture along each stroke.