Measurement and Analysis of Brain Deformation During Neurosurgery


Abstract—Recent studies have shown that the surface of the brain is deformed by up to 20 mm after the skull is opened during neurosurgery, which could lead to substantial error in commercial image-guided surgery systems. We quantitatively analyze the intraoperative brain deformation of 24 subjects to investigate whether simple rules can describe or predict the deformation. Interventional magnetic resonance images acquired at the start and end of the procedure are registered nonrigidly to obtain deformation values throughout the brain. Deformation patterns are investigated quantitatively with respect to the location and magnitude of deformation, and to the distribution and principal direction of the displacements. We also measure the volume change of the lateral ventricles by manual segmentation.

Our study indicates that brain shift occurs predominantly in the hemisphere ipsi-lateral to the craniotomy, and that there is more brain deformation during resection procedures than during biopsy or functional procedures. However, the brain deformation patterns are extremely complex in this group of subjects. This paper quantitatively demonstrates that brain deformation occurs not only at the surface, but also in deeper brain structure, and that the principal direction of displacement does not always correspond with the direction of gravity.

Therefore, simple computational algorithms that utilize limited intraoperative information (e.g., brain surface shift) will not always accurately predict brain deformation at the lesion.

Index Terms—Interventional MRI, intraoperative brain deformation, nonrigid image registration.

I. INTRODUCTION

IMAGE-GUIDED surgery (IGS) systems register preoperative tomographic images to the intraoperative coordinate system of the patient. The resulting transformation or mapping is used to display the position and orientation of tracked surgical instruments on reformatted image slices and renderings of the brain and anatomical structures of interest. In neurosurgery, commercial IGS systems make the assumption that a patient’s head and brain is a rigid body. Recent studies have shown that this assumption is not valid and report on significant brain deformation (brain shift) after the skull is opened and before the interventional procedure is started, e.g., [1]–[6]. If the tissue deformation is large relative to the amount of surgical accuracy required, then the overall accuracy of the IGS system will be substantially reduced.

Several approaches have been developed to address the brain deformation problem. A magnetic resonance (MR) scanner that has been modified for intraoperative surgery can be used to scan the patient multiple times during the intervention and monitor the brain deformation (e.g., [7]). Alternatively, real-time ultrasound systems have been proposed to correct for the brain deformation [8], [9]. Although these devices can provide surgeons with updated images during the intervention, the use of preoperative data is still of clinical interest. For instance, it may be desirable to have images from other modalities (e.g., positron emission tomography, functional MR imaging) and preoperatively prepared data (e.g., segmentations of clinically important anatomical structures) displayed in an IGS system. In particular, the fusion of these data with intraoperatively acquired images provides surgeons with additional information. Furthermore, since interventional MR scanners are expensive, complicate access to the patient and prevent the use of standard metallic surgical instruments because of their high magnetic fields, their usage is restricted and for some cases it may be preferable to use preoperatively acquired images.

Recent studies develop biomechanical models which estimate displacements in order to update the preoperative images [10]–[15]. These models are based on physical brain deformation and require intraoperative measurements to constrain their model. In general, these studies assume, that the brain deformation can be estimated by introducing simple physical models for the cause of deformation, e.g., direction of gravity or size of the resection [14], [15].

In this paper, we qualitatively and quantitatively investigate brain deformation based on MR images from 24 patients that underwent cranial neurosurgical procedures at the University of Minnesota. The images were acquired at the start and end of each procedure using an interventional MR scanner in the operating room. The aim of this paper is to investigate the pattern of brain deformation with respect to location, magnitude, and direction and to consider the implications of this pattern on models for correcting brain deformation or on the errors in IGS systems due to brain deformation. We focus on three questions: 1) Where...
is the deformation, e.g., does it occur only at the brain surface or also in deeper brain structures? 2) What is the magnitude of the deformation? 3) Can simple rules describe or predict the deformation, e.g., loss of cerebrospinal fluid (CSF) causes the brain to sink in direction of gravity?

The volume change in each lateral ventricle is calculated after manually segmenting the lateral ventricles in the start- and end-procedure images. The start-procedure image is automatically registered to the end-procedure image using a nonrigid registration algorithm. The resulting transformation is a deformation vector field, which is used to quantify deformation between the images. Since the nonrigid transformation defines a displacement vector for each point in the image, in contrast to the manually determined ventricular volume changes, this approach provides deformation measurements throughout the entire brain. The deformation field is investigated quantitatively with respect to the location of the deformation, the magnitude of deformation in relation to the distance to the brain surface, the principal direction of the deformation, and the distribution of the displacement direction.

Studies prior to the mid-1990s mentioned brain shift (e.g., [16]), but did not quantify the deformation. More recent studies determined either the deformation at the brain surface, [2]–[4], [17], or the volume change of manually segmented anatomical structures [1], [5], [6], [17]. While these studies reported manual and/or local measurements of the deformation at certain surfaces or structures during neurosurgery, we are interested in an approach that provides deformation values throughout the entire brain. A similar approach has already been used to quantify the volume change of anatomical structures in the developing brain of children [18] and of the temporal lobe of Alzheimer’s disease patients [19], [20]. Hata et al. [21] (see, also, [22] and [23]) applied a similar method to measure brain deformation during neurosurgery, but did not apply any complex operators to the deformation field, did not consider the location of deformation, and did not analyze volume changes of specific anatomical structures.

The paper is organized as follows: Section II describes the image acquisition and the methods we used to quantify and localize the brain deformation. Section III presents the results of the quantitative measurements, presents a visual assessment of the deformation, and provides illustrations of the deformation fields.

II. MATERIALS AND METHODS

This section describes the image acquisition and the methods we used to quantify and localize the brain deformation. First, the volume change of the lateral ventricles is determined by manual segmentation in the start-procedure and end-procedure image. Then, in order to investigate the deformation throughout the whole brain the start- and end-procedure images are registered nonrigidly and the resulting deformation vector field is analyzed with respect to the magnitude, direction, and location of displacement. At the end of this section the deformation localization methods are described.

Fig. 1. Example interventional start-procedure (left) and end-procedure (right) image (case resection8), (top) Sagittal plane (as acquired) and (bottom) reformatted transaxial plane.

A. Image Acquisition

This paper is based on MR images from 24 patients that underwent cranial neurosurgical procedures in the interventional MR suite at the University of Minnesota. During surgery, magnetization-prepared rapid gradient echo (MP-RAGE) volume images were acquired with a 1.5-T interventional MR scanner (Philips ACS-NT, Philips Medical Systems). In addition to the normally acquired images, which generally contain only a small number of slices, additional whole brain images volumes were obtained at the start and at the end of each intervention (see Fig. 1). The voxel dimensions are typically $0.9 \times 0.9 \times 1.5$ mm, and the readout gradient is oriented in the cranial–caudal direction with a magnitude of 4.7 mT/m. Since the images were acquired using an open-style radio-frequency coil (a “synergy” phased array coil), some images have substantial nonuniform signal intensity. Table I summarizes details of the 24 cases studied in this investigation.

Gradient echo images have high resolution (small pixel size), and the relatively high readout-gradient magnitude (4.7 mT/m) used in this study produces images with relatively little geometrical distortion in the read-out direction due to static field inhomogeneity [24]. However, gradient echo images are sensitive to magnetic susceptibility differences in the object being imaged and, consequently, there can be signal loss at the boundary between air and soft tissue. Spin echo images do not have this potential problem of signal loss. For patients undergoing resections, the brain is exposed to the air and air can enter the head. The magnitude of any brain deformation could, therefore, be exaggerated by signal loss due to susceptibility artifacts. To establish the extent of this problem, we also acquired T2-weighted turbo spin echo images from some patients.
TABLE I
DETAILS OF 24 CASES STUDIED USING INTERVENTIONAL MR. IN THE CASES MARKED WITH *1*, THE SECOND IMAGE WAS ACQUIRED IN THE MIDDLE OF THE INTERVENTION AND NOT AFTER THE INTERVENTION

<table>
<thead>
<tr>
<th>Case</th>
<th>Age</th>
<th>Gender</th>
<th>Lesion position</th>
<th>Lesion type</th>
</tr>
</thead>
<tbody>
<tr>
<td>resection1</td>
<td>53</td>
<td>male</td>
<td>left occipital</td>
<td>lung metastasis</td>
</tr>
<tr>
<td>resection2</td>
<td>5</td>
<td>male</td>
<td>left occipital</td>
<td>astrocytoma with necrosis</td>
</tr>
<tr>
<td>resection3</td>
<td>69</td>
<td>male</td>
<td>left frontal</td>
<td>glioblastoma multiforme</td>
</tr>
<tr>
<td>resection4</td>
<td>3</td>
<td>female</td>
<td>right medio temporal</td>
<td>low grade glioma</td>
</tr>
<tr>
<td>resection5</td>
<td>39</td>
<td>female</td>
<td>right occipito parietal</td>
<td>meningioma</td>
</tr>
<tr>
<td>resection6</td>
<td>41</td>
<td>female</td>
<td>right frontal</td>
<td>glioma</td>
</tr>
<tr>
<td>resection7</td>
<td>36</td>
<td>female</td>
<td>superficial left temporal</td>
<td>glioma</td>
</tr>
<tr>
<td>resection8</td>
<td>57</td>
<td>male</td>
<td>left frontal</td>
<td>glioblastoma multiforme</td>
</tr>
<tr>
<td>resection9</td>
<td>57</td>
<td>male</td>
<td>left temporal</td>
<td>glioblastoma multiforme</td>
</tr>
<tr>
<td>resection10</td>
<td>26</td>
<td>female</td>
<td>left temporal</td>
<td>astrocytoma</td>
</tr>
<tr>
<td>resection11</td>
<td>34</td>
<td>male</td>
<td>right medio frontal lesion</td>
<td>low grade glioma</td>
</tr>
<tr>
<td>resection12</td>
<td>60</td>
<td>male</td>
<td>left temporal</td>
<td>glioblastoma multiforme</td>
</tr>
<tr>
<td>resection13</td>
<td>54</td>
<td>male</td>
<td>occipital</td>
<td>glioblastoma multiforme</td>
</tr>
<tr>
<td>biopsy1</td>
<td>38</td>
<td>female</td>
<td>left frontal lesion</td>
<td>oligodendroglioma</td>
</tr>
<tr>
<td>biopsy2</td>
<td>72</td>
<td>female</td>
<td>left parietal</td>
<td>glioblastoma multiforme</td>
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<tr>
<td>biopsy3</td>
<td>47</td>
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<td>left temporal parietal</td>
<td>glioblastoma multiforme</td>
</tr>
<tr>
<td>biopsy4</td>
<td>25</td>
<td>female</td>
<td>left parietal</td>
<td>astrocytoma</td>
</tr>
<tr>
<td>biopsy5</td>
<td>73</td>
<td>male</td>
<td>left frontal</td>
<td>glioma</td>
</tr>
<tr>
<td>functional1</td>
<td>50</td>
<td>female</td>
<td>bilateral</td>
<td>obsessive compulsive disorder</td>
</tr>
<tr>
<td>functional2</td>
<td>43</td>
<td>male</td>
<td>right temporal</td>
<td>brain damage and tremor</td>
</tr>
<tr>
<td>functional3</td>
<td>58</td>
<td>male</td>
<td>left</td>
<td>tremor</td>
</tr>
<tr>
<td>functional4</td>
<td>61</td>
<td>male</td>
<td>left</td>
<td>tremor</td>
</tr>
<tr>
<td>functional5</td>
<td>84</td>
<td>female</td>
<td>left</td>
<td>tremor</td>
</tr>
<tr>
<td>functional6</td>
<td>61</td>
<td>male</td>
<td>right</td>
<td>tremor</td>
</tr>
</tbody>
</table>

B. Segmentation of the Lateral Ventrices

In order to estimate brain deformation in deeper brain structures during the surgical procedure, we manually segment the lateral ventricles in the start- and end-procedure images using Analyze (Mayo Clinic). Based on these segmentations, volume changes were calculated separately for the lesion’s ipsi- and contra-lateral ventricles, and for both ventricles together.

C. Calculation of the Deformation Vector Field

While manual segmentation provides the volume change of certain structures, nonrigid registration of the start- and end-procedure images produces displacement values throughout the brain for each voxel in the image. The resulting displacement map can be used to investigate deformation in the entire brain and to detect typical deformation patterns.

First, we rigidly register images to correct for any rigid-body motion between the start- and end-procedure images by maximizing the normalized mutual information (NMI) in the joint probability distribution between the two images [25]. Then we nonrigidly register images by deforming a regular grid of control points in the start-procedure image [26]. The nonrigid registration algorithm moves the control points; tissue motion is described by free-form deformation using B-spline approximation between the control points. The control point spacing determines the flexibility of the grid and was set in our investigation by 15 mm. Again, NMI is used as a measure of the similarity of the images. This algorithm was previously evaluated for the registration of three-dimensional (3-D) breast MR images [27] and of 3-D brain MR images [28]. In the latter study, the displacement vectors determined by the nonrigid algorithm were compared with the displacements of manually determined point landmarks in deformed brain MR image. It was shown that 68% of the values of the displacement map are within 0.5 mm of the interactively measured values and 90% are within 1.0 mm. Since the reproducibility error of the interactive measurements was up to 0.9 mm, this suggests that the results of the automatic nonrigid registration algorithm agree with the interactive measurements to the precision of these.

The registration algorithm results in a 3-D deformation field \( D \) which is mathematically defined as a vector field \( \mathbb{R}^3 \rightarrow \mathbb{R}^3 \) and maps a point \( \mathbf{p}_{\text{source}}(x, y, z) \) from the source image to the target image \( \mathbf{p}_{\text{target}} = D(\mathbf{p}_{\text{source}}) \).

**Displacement Vectors:** Usually the deformation field is visualized as a displacement field \( d \) (see, e.g., Fig. 6). This field represents the displacement of each point and is defined on the basis of the deformation field: \( D(\mathbf{p}) = \mathbf{p} + d(\mathbf{p}) \).

We are not only interested in the magnitude of the displacement vectors, but also in the distribution of their directions and in the main direction of the whole displacement field. In order to analyze the displacement directions, principal component analysis (PCA) is applied on the displacement field in the following way: the displacement vectors are considered as points in a 3-D coordinate system, whereby the axes of the coordinate system represent the components of the displacement vectors (see Fig. 2). Then, a vector is adjusted through the point cloud in such way that the variance of the points is maximal along the vector. In contrast to the classical PCA, the vector is determined with respect to the origin of the diagram and not with respect to the mean of the points. This vector can be considered as the principal direction of the displacement field and the variance in its direction as a measure for the over-all magnitude of the deformation. Another vector perpendicular to the first is determined that maximizes the variance of the points in its direction again. A third vector is perpendicular to the first and second axis, and the variance in this direction is calculated. Usually, the variances in the direction of these three vectors...
Fig. 3. Distance map for resection. (left) End-procedure image and (right) corresponding distance map. The gray values inside the brain represent the distance to the brain surface (darker values correspond to larger distance).

Fig. 4. Example sagittal and coronal slices from resection. The brain surface contour of the spin-echo image is overlaid on the gradient-echo image. The spin-echo image were taken 3 min before the gradient-echo image. The good agreement between contours suggests that signal loss due to susceptibility difference at the air/tissue interface is not a cause of large errors.

The Jacobian operator can be used to measure local relative tissue volume change throughout the brain. It is defined at the point \( p \) as the determinant of the Jacobian matrix of the deformation field

\[
J_{\text{ac}}(D) = \det(\nabla_{\text{p}} D) = \begin{vmatrix}
\frac{\partial D_x}{\partial x} & \frac{\partial D_x}{\partial y} & \frac{\partial D_x}{\partial z} \\
\frac{\partial D_y}{\partial x} & \frac{\partial D_y}{\partial y} & \frac{\partial D_y}{\partial z} \\
\frac{\partial D_z}{\partial x} & \frac{\partial D_z}{\partial y} & \frac{\partial D_z}{\partial z}
\end{vmatrix}.
\]

The Jacobian operator relates an elementary volume \( \delta V_{\text{source}} \) in the source image to the corresponding deformed volume \( \delta V_{\text{target}} \) in the target image

\[
\delta V_{\text{target}} = J_{\text{ac}}(D) \cdot \delta V_{\text{source}}
\]

and can be considered as a local expansion of the volume and responses less than one as a local shrinking of the volume at point \( p \). The response \( J_{\text{ac}}(D) = 1 \) means that the volume is preserved locally at this point. We apply the Jacobian operator to the deformation field determined by the nonrigid algorithm to analyze the local volume change for each point in the image.

D. Localization of Brain Deformation

In addition to the magnitude of deformation, we are also interested in the location of the deformation and whether a typical pattern can be found in their spatial distribution. First, we separately analyze the values in the ipsi-lateral and in the contra-lateral hemisphere to the lesion. Second, a distance map is used, which specifies for each voxel inside the brain its distance to the brain surface.

Separating Ipsi- and Contra-Lateral Hemisphere: In order to investigate the deformation measurements for each hemisphere separately, the brain is segmented into the left and right hemispheres. We use the segmentation of the brain atlas Internet Brain Segmentation Repository [31] and perform an affine registration of the reference MR image of the atlas to the end-procedure images. The resulting affine transformation is used to transform the atlas segmentation to the coordinate system of the end-procedure image. Since we are not interested in the exact boundary of the brain, but only in the deformation values inside each hemisphere, the approximate segmentation obtained with this method is sufficiently accurate for our investigation.

Distance to the Brain Surface: In order to quantify the deformation with respect to the distance to the brain surface, a distance map is calculated for each patient [32]. The brain is segmented in the end-procedure images, and, starting with the voxels at the brain surface, the voxels are marked with a value that represents the distance to the surface. The result is an image in which the value of a voxel inside brain represents the distance to the brain surface (see Fig. 3). This image is used to analyze the deformation measurements with respect to distance by considering for each distance value only those deformation measurements corresponding to voxels in the distance map with that distance value.
III. RESULTS

First, we compare the spin-echo images with the gradient-echo image to assess the signal loss at the boundary between air and soft tissue in the gradient-echo image which could exaggerate the magnitude of brain deformation. Fig. 4 shows the contour of the spin-echo image overlaid on the gradient-echo image. In three patients, we carried out visual assessment of the gradient echo MR volume with the brain boundary from spin echo overlaid. For these three cases, the boundaries were judged on average $2.1(\pm 1.8)$ mm, $0.4(\pm 0.5)$ mm, and $1.4(\pm 0.7)$ mm of one another, respectively. Given the difference in read-out gradient strength between gradient-echo and spin-echo imaging, the time delay between acquisitions (between 3 and 6 min), and a partial volume effect due to different slice thicknesses, these are consistent with the gradient echo and spin echo images providing equivalent information about the deformation. Thus, signal loss in the gradient echo images at the air–tissue boundary is not likely to be a major cause of error in this study.

A. Visual Assessment of Brain Deformation

The size and location of the lesion varies significantly for the cases investigated and, therefore, the comparison of the start-
and end-procedure image shows different types of deformation. In Fig. 5, four example resection cases and their corresponding subtraction images after rigid and nonrigid registration are depicted. Case resection7 (first row in Fig. 5) underwent a small resection in the left temporal lobe and the subtraction image after rigid registration (second column) mainly shows deformation at the surface of the brain while the contra lateral hemisphere is hardly deformed. In contrast, a deep lesion was resected in case resection10 which caused a considerable deformation of the ventricular system. Even though the deformations at the contra lateral ventricle are much less in comparison to the ipsi lateral ventricle, the brain also deforms substantially in the contra-lat-

eral hemisphere. The subtraction image of this case suggests that there is less deformation at the brain surface than at the ventricu-

lar system. The resection of the lesion at the frontal lobe in the case resection11 caused both a brain shift at the brain surface and a deformation of deeper brain structures (see lateral ventricle). The lesion of case resection12 was located in the left temporal lobe and the image shows similar deformation pattern to resection7. In all of these cases except in case resec-
tion10, the subtraction image suggests that the main direction of the brain shift is in the direction of gravity (indicated by an arrow in the end-procedure images in the first column). We will see in the next section, where we analyze the distribution of the displacement vectors, that this result may not apply to all cases.

The deformation map overlaid on the end-procedure image (see Fig. 6) indicates that the displacement is greatest close to the lesion or craniotomy, but can vary considerably. For instance, while the deformation in case functional6 is greatest immediately beneath the craniotomy, and near the mid-line has diminished to virtually zero, case resection10 also shows significant displacements in the contra-lateral hemisphere.

The Jacobian operator is applied in Fig. 7 on the displace-

ment field to visualize the relative volume change throughout the brain. Dark regions in these images indicate a shrinking of the brain and bright regions indicate an expansion. Comparing the displacement field with the Jacobian operator responses, demonstrates that a large displacement at a given position does not yield a significant volume change at that position. For ex-

ample, in case resection7 in Fig. 7 the largest displacement is at the surface while a significant volume change appears in deeper brain structures. In case functional1, significant displacement vectors are spread over the frontal lobe while the larger volume change is concentrated in a smaller region.

B. Deformation in the Ipsilateral and Contra-Lateral Hemisphere to the Procedure

The deformation values in the ipsi- and contra-lateral hemisphere are considered separately for each case using the segmen-
tation yielded by registering the brain atlas to the end-procedure
Fig. 8. Analysis of the displacement vectors in the ipsi- and contra-lateral hemisphere. (top) In order to ignore statistical outliers, we consider in the upper diagram the 95th percentile of the displacement magnitudes. (bottom) Degree of distribution of the displacement vector directions measured by the volume of the covariance ellipsoid based on the PCA of the deformation field (see Section II-C).

image (see Section II-D). The displacement vectors of the non-rigid algorithm are analyzed in two ways: First, the magnitude of the displacement field and the distribution of the displacement vectors determined by the PCA (see Section II-C) are investigated and illustrated in Fig. 8. Second, the volume change is determined for each voxel on the basis of the deformation field and the fifth percentile of the volume change—representing the maximal volume change—is depicted separately for each hemisphere in Fig. 9.

Comparing the magnitude of deformation for each hemispheres in Fig. 8 (top), in general, larger deformation occurs in the ipsi-lateral hemisphere than in the contra-lateral hemisphere. But the case resection6 also yields significant deformation values in the contra-lateral hemisphere. If you compare the resection cases with the biopsy and functional cases, it turns out, that the magnitude of displacement for the resection cases is, in general, larger. However, for instance, case functional6 shows a deformation with magnitude comparable to the magnitude of some resection cases. Fig. 8 (bottom) demonstrates that the direction of the displacement vectors in the resection cases can be much more distributed than in the biopsy and functional cases. In particular, whereas the deformation magnitude in case functional6 is substantially larger than, for instance, in case resection13, the direction of the displacement vectors is much less distributed. Since the magnitude of deformation is smaller in the contra-lateral hemisphere than in the ipsi-lateral hemisphere, the displacement directions are also less distributed in this hemisphere. However, in some cases with significant brain shift in the contra-lateral hemisphere the displacements occur in nearly the same direction, i.e., the distribution is nearly zero (for example, case resection10 and resection13).

Along with a larger displacement magnitude in the ipsi-lateral hemisphere, we also, in general, observe larger volume change in this hemisphere as illustrated in Fig. 9. However, the cases resection1 and functional5 show a larger shrinking in the contra-lateral hemisphere. Comparing the mean of the maximal volume change values separately for the resection, biopsy, and functional cases (compare the diamonds with each other), shows that, in general, the volume change in the resection cases is larger than in the biopsy cases, and the volume change in the biopsy cases is larger than in the functional cases. However, the standard deviation is relatively large in each group and, thus, the groups cannot be clearly separated.

C. Volume Change of the Lateral Ventricle

Fig. 10 summarizes, for each case, the volume change of the lateral ventricle determined by manual segmentation. The change is depicted for both ventricles together and for the ipsi- and contra-lateral hemisphere separately. The variability in
These diagrams make clear that by the end of the procedure the location of the curve maximum, i.e., the maximal distance from the brain surface to the center of the ventricle, shows a shrinking close to the brain surface, but also in deeper brain structures.

Comparing the displacement vector diagrams with the volume change diagrams, demonstrates that the position of the maximal volume change does not correspond with the position of the maximal displacement. For example, the maximal shrinking in resection12 and resection10 occurs in deeper brain structures than the maximal displacement. While resection12 shows an expansion close to the brain surface, resection10 shows no peak in the expansion curve. The large displacement in resection7 close to the surface does not yield to a significant volume change, either. The case biopsy4 has smaller displacement vectors but shows considerable shrinking.

Fig. 10. Volume change of the lateral ventricle determined by manual segmentation. The y axis represents the volume ratio of the lateral ventricle in the start- and end-procedure images, i.e., a value “1" represents no volume change; values larger than “1" represent an expansion, and values less than “1" represent a shrinking of the ventricle. The circles depict the volume change in the ventricle ipsi-lateral to the lesion, while the squares depict the volume change in the contra-lateral ventricle. The diamonds represent the volume change of both lateral ventricles together.

volume change measure assessed by repeated segmentation of a start- and end-procedure ventricle pair was 5.4%.

Significant volume change is measured mainly in the resection cases, while both a shrinking (e.g., cases resection4 and resection6) and an expansion (e.g., cases resection8 and resection10) of the structure occurs. In general, the biopsy and functional cases show less volume change, but the ventricle also deforms significantly in some of these cases (e.g., case functional1). Comparing the volume change of the ipsi-lateral ventricle with the contra-lateral ventricle, shows that either the values are similar or the ipsi-lateral ventricle changes to a greater extent. The volume change in the ipsi-lateral ventricle varies between half (case resection4) and twice the original volume (case resection10).

D. Deformation Values With Respect to Brain Surface Distance

The distance map for each case described in Section II-D is used to analyze the displacement with respect to the distance to the brain surface in order to determine whether the shift is greatest at the brain surface. The displacement vector with the largest magnitude is selected for each distance and displayed in a diagram which produces scan lines shown in Fig. 11 (left). Analogously, the maximum and minimum volume changes—which represents the maximal expansion and maximal shrinking—are analyzed for each distance and visualized in Fig. 11 (right).

Comparison of the displacement diagrams in Fig. 11 shows that the location of the curve maximum, i.e., the maximal displacement in the whole brain, varies from case to case. Separating the distance from the brain surface to the center of the brain into five equal sized parts, reveals that for five cases the maximum is in the first fifth of this distance (i.e., close to the brain surface), for 13 cases it is in the second fifth, for four cases it is in the third fifth, and for two cases it is in the fourth fifth. In Fig. 11, one representative diagram for each group is depicted. These diagrams make clear that by the end of the procedure the main displacement does not only occur at or close to the brain surface, but also in deeper brain structures.

Comparing the displacement vector diagrams with the volume change diagrams, demonstrates that the position of the maximal volume change does not correspond with the position of the maximal displacement. For example, the maximal shrinking in resection12 and resection10 occurs in deeper brain structures than the maximal displacement. While resection12 shows an expansion close to the brain surface, resection10 shows no peak in the expansion curve. The large displacement in resection7 close to the surface does not yield to a significant volume change, either. The case biopsy4 has smaller displacement vectors but shows considerable shrinking.

E. Relationship Between Displacement in the Hemisphere and Volume Change of the Lateral Ventricle

Both the investigation of the displacement magnitudes and the volume change of the lateral ventricle show that brain structures in the ipsi-lateral hemisphere deform more than in the other hemisphere. In order to determine whether a significant brain shift is always associated with a volume change of the lateral ventricles, the largest displacement vector in one hemisphere is depicted in comparison to the manually determined volume change of the lateral ventricles (see Fig. 12).

The measurements of the contra-lateral hemisphere in the diagram are much more clustered than the measurements of the ipsi-lateral hemisphere. Since this cluster is close to “small displacement vectors” and “no volume change,” this again indicates that the contra-lateral hemisphere deforms less, in general. In comparison with this the ipsi-lateral measurements are less clustered and in some cases, e.g., resection10 and resection6, a large displacement is associated with a large volume change of the ventricle. However, in some other cases, e.g., resection1 and biopsy3 large displacements were measured whereas almost no volume change occurred.

Altogether, these results suggest that a large displacement is not always associated with a volume change of the lateral ventricle.

F. Direction of the Displacement Vectors

The subtraction images in Fig. 5 suggest that the brain shift occurs mainly in the direction of gravity. In order to verify this observation, we compare the direction of gravity which is provided for each slice by the MR scanner with the direction of the first principal component of the displacement vectors (see Fig. 8), which can be considered as the main direction of the displacement field. The angle between these two vectors is depicted in Fig. 13 for 16 selected cases. In this diagram, cases are left out where magnitudes of deformation are so low, that the direction cannot be determined reliably (we omit cases which showed in Fig. 8 a maximal displacement less than 2 mm).

Comparing the angle between these vectors shows that in some cases the main direction of the displacement field aligns well with the direction of gravity (e.g., case resection9, biopsy4, and functional1). In other cases, however, the brain shift occurs in a direction significantly different to that of gravity. For instance, in case resection13 the main direction of the displacement field is nearly perpendicular to gravity. In Fig. 14, the ex-
ample slice of resection13 suggests that because of the resected lesion a major part of the displacement vectors point in a different direction to the gravity vector and some even point in the opposite direction.

IV. DISCUSSION

Twenty-four patients undergoing neurosurgery with intraoperative MR imaging were studied in order to quantify the interventional brain shift. The magnitude and location of deformation, and the distribution of the displacement directions were investigated using measurements determined by a nonrigid registration algorithm which provides displacement measurements throughout the brain. In addition, the volume change of the lateral ventricle was measured by manual segmentation. Careful study of brain deformation throughout the brain is important in ascertaining how easily computational algorithms could correct for brain deformation in IGS systems.

Our results are consistent with the results of previous studies (e.g., [17], [21], and [28]) and can be summarized as follows:

- **Location of Deformation:**
  The manual segmentation of the lateral ventricles as well as the displacements determined by the nonrigid algorithm indicate that larger deformation occurs in the hemisphere ipsi-lateral to the lesion than in the contra-lateral hemisphere. These results agree well with previous studies [28]. However, some cases (both resection and no-resection cases) also show a significant brain shift in the contra-lateral hemisphere. Furthermore, considering the displacement scan-lines with respect to the brain surface distance, our study demonstrates that brain deformation does not only occur at the brain surface, but also in deeper brain structure. Even if the main displacement appears close to the surface, the resulting local volume change of the brain can occur further below the surface.

- **Magnitude of Deformation:**
We measured a shrinking of the lateral ventricles to up to half of the original volume and an expansion up to double the volume. In general, the magnitude of the displacement is larger in the resection cases in comparison with the functional or biopsy cases, but this varies significantly in each group.

- **Predicting Deformation:**

  A simple view of the brain shift problem might be that the shift is caused by the brain collapsing under the force of gravity to fill space previously occupied by CSF and resected tissue. While gravity may be a major driving force in brain deformation, our results show that the direction of gravity is a poor predictor of the direction of brain shift. In many subjects, the direction of the displacement field was not aligned with the direction of gravity (see, e.g., Fig. 14). The direction of shift is likely to be a consequence of a complex interplay between the force of gravity, boundary conditions (e.g., resected regions), fluid pressure, and other forces. Furthermore, substantial deformation is not always associated with substantial CSF loss from the lateral ventricles (Fig. 12), which suggests that measurement of CSF loss is not a good predictor of the magnitude of brain deformation. Thus, this investigation quantitatively confirms the comment in [17], that measuring only the volume change of the lateral ventricle is not reliable enough to quantify brain deformation.

Overall, the investigation demonstrates that though, in general, the deformation is less in the biopsy and functional than in the resection cases, significant brain shift can also occur in these cases (see, e.g., case functional6).

The pattern of brain deformation, both in terms of magnitude and direction of displacement, and of volume change, is extremely complex in this group of subjects. Therefore, simple computational algorithms that make use of very limited intraoperative information (e.g., brain surface shift) are unlikely to accurately predict brain deformation at the lesion for all cases. Simple biomechanical models might be able to predict brain deformation if they had sufficiently good boundary conditions. Our results suggest that it will be difficult to obtain these boundary conditions without using intraoperative imaging. For example, in some subjects, there is negligible deformation contra-lateral to the side of resection, but in other cases there is substantial deformation on both sides of the brain. An algorithm cannot, therefore, assume that the mid-line of the brain is fixed in all cases. Further research is needed in order to ascertain how well a biomechanical model needs to know mechanical properties of tissue and the boundary conditions in order to model the complex types of deformation we have observed in these subjects. Furthermore it is a mistake when using a traditional IGS system to assume that the brain shift at the surface is always a useful predictor of the amount of deformation further deep in the brain, or that the main direction of brain shift is in the direction of gravity.
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REFERENCES


