ABSTRACT
Accurate and robust identification of the gyri and sulci of the human brain is a pre-requisite of high importance for modelling the brain surface and thus to facilitate quantitative measurements and novel classification concepts. In this work we introduce a watershed-inspired image processing strategy for topographical analysis of arbitrary surfaces in 3D. Thereby the object’s topographical structure represented as depth profile is iteratively transformed into cyclic graph representations of both, the lowest and the highest characteristics of the particular shape. For graph analysis, the surface elements are partitioned according to their depth value. Neighbouring regions at different depth levels are iteratively merged. For region merging, the shape defining medial axes of the involved regions have to be connected by the optimum path with respect to a fitness function balancing shortness and minimal depth level changes of the solution.

Keywords: topographical surface analysis, cyclic graph representation, sulcus and gyrus classification

1. INTRODUCTION
The accurate quantification of metabolic processes from functional emission tomography imaging modalities like positron emission tomography (PET) and single photon emission computed tomography (SPECT) for diagnosis of neurodegenerative diseases necessitates a precise and patient-specific segmentation and classification of the brain. For segmentation and classification tasks, morphological image modalities as magnetic resonance imaging (MRI) have to be fused with the data acquired by functional emission imaging. Thus, the segmentations and classifications evaluated based on the anatomically-precise imaging modalities can be applied to the emission data, facilitating quantitative analysis of the metabolic activity with respect to pre-classified anatomical regions. The classification concept addressed in this work is the partitioning of gray and white brain matter according to the gyrus and sulcus characteristics.

Any computer-based functional or anatomical classification requires binary segmentation as pre-processing. Utilizing T1-weighted brain MRI data, segmentation of gray and white matter can be achieved, using k-means clustering (Kanungo et al. 2002; Ibanez et al. 2005) for determination of the tissue types to discriminate and region growing for ensuring connectedness.

Based on a binary segmentation of the brain surface, several strategies for sulcus and gyrus classification have been presented and published in the past. A morphologic closing operation, i.e. dilation followed by erosion, with subsequent subtraction of the original MRI data allows processing of the sulcus volume via 3D skeletonization for extraction of the sulcus and gyrus folds (Lohmann 1998). In contrast to applying Euclidean distances, the use of a geodesic depth profile accounts for complexity and partial occlusion of the sulcus folds (Kao et al. 2006). Besides these morphologic concepts, curvature analysis of a surface mesh calculated from gray and white matter can be utilized for detection of the gyrus and sulcus course (Vivodtzev et al. 2003) with respect to convexity and concavity.

In this paper we present a generic strategy for topographic analysis of arbitrary shapes and transformation of the depth profiles into cyclic graph representations. Thereby we account for imbalances in the local depth profiles due to asymmetries and deformations of the brain. The minimum graph connecting all local maxima and minima respectively is calculated, normalizing the local depth levels similar to the watershed segmentation concept. Our strategy is perfectly applicable for the task of gyrus and sulcus modelling as concave and convex paths can be identified. Based on the graph representations of the sulcus and gyrus courses, modelling of the brain surface can be easily achieved via distance-based classification utilizing morphologic operators as presented and discussed in the following sections.

2. MEDICAL BASICS
Classification of the human brain can be accomplished at different levels of granularity. At a top level, the main anatomical structures, like cerebrum, cerebellum and the brain stem can be identified. The cerebrum is
subdivided into two hemispheres and the main anatomical components, like white matter, gray matter, cerebro-spinal fluid (CSF), ventricle, fat, bones and the arterial and venous vessel systems are demarcated. The brain tissue composed of white and gray matter is subclassified into frontal lobe, parietal lobe, occipital lobe and some more, all specific areas responsible for diverse neurological functions of the body (Pschyrembel 2002). Each lobe comprehends several gyrus and sulcus areas, forming the brain surface. Thereby the gyri refer to the convex bulges on the brain surface that are delimited by convex trenches, the so called sulci. The notable main sulci and gyri are named, listed and charted in anatomical atlases (Ono et al. 1990).

The topography of gyrus and sulcus characteristics is highly applicable for registration tasks in case of multi-modal image processing or follow-up examinations. Furthermore, modelling of the gyrus segments facilitates the quantitative analysis of metabolic activities with respect to defined anatomical structures.

3. DATA
For testing of the gyrus and sulcus modelling concept, \(n=20\) T1-weighted MRI datasets of simulated brainweb database (Cocosco et al. 1997; Kwan, Evans, and Pike 1999) and associated reference segmentations are used. Further test runs and validations will be performed utilizing \(n=12\) anonymous multi-modal patient studies comprising morphologic image acquisitions (T1, T2, PD, …) as well as functional images (SPECT, PET).

4. METHODOLOGY
Prior to performing the analysis process, a binary representation of the targeting object’s surface, not addressed in this work, and a 3D depth profile must be pre-processed.

4.1. Estimation of the Reference Shape
For calculation of the depth profile of an arbitrarily shaped object, the reference shape, i.e. the smoothed shape without the vales and ridges, must be estimated.

Processing a solid body with an approximately spherical shape, like the human brain, calculation of the 3D convex hull (Barber, Dobkin and Huhdanpaa 1996; Sonka, Hlavac and Boyle 2007) as reference shape is highly feasible, see Fig. 1.

For other more complex shapes, where a spherical approximation would be too imprecise, an alternative calculation of the reference shape is feasible. When calculating a winged-edge isosurface of a binary 3D body (Baumgart 1972; Baumgart 1975; Ritter 2007; MeVis 2011), utilizing the quality factor, allows steering of the smoothing effect by polygonal reduction, i.e. up to which level, vales and ridges should influence the depth profile calculation, see Fig. 2 (b). The resulting isosurface is projected back to regular 3D voxel grid for further processing. Furthermore, morphologic closing operations as dilation followed by erosion can be utilized for smoothing the surface and calculation of the reference shape, see Fig. 2 (c).

Figure 1: The stack of 2D MRI slices (a) assembles a 3D volume of the brain (b). After binary segmentation (c), the calculated convex hull (d) is the reference shape for depth profile calculation.

Figure 2: The precisely calculated surface model (a) can be smoothed for use as reference shape via rough isosurface calculation (b). As an alternative reference shape calculation strategy, a morphological closing operation with dilation kernel size \(7x7x7\) followed by erosion \(6x6x6\) can be applied (c).

4.2. Calculation of the Depth Profile
The depth profile is calculated as the minimum Euclidean distance between the surface of the object and the reference shape, see Fig. 3 as illustration of 2D depth profile calculation. A distance map calculation is used to represent the depth profile in 3D with the Euclidean neighbourhood weights for the adjacency constellations \(N_6\), \(N_{12}\) and \(N_8\) in \(3x3x3\) neighbourhood according to the distance from the hot spot as

\[
\begin{align*}
    w(N_6) &= 1.0; \\
    w(N_{12}) &= \sqrt{2}; \\
    w(N_8) &= \sqrt{3}.
\end{align*}
\]

Starting at the convex hull, the distance weights are propagated to the particular neighbours to set or update their values. Whenever the depth value of a neighbour gets adapted, the change is recursively propagated to all adjoined neighbours. Calculation of the depth profile is finished, when the depth value of each voxel refers to the minimal Euclidean distance from the convex hull.
The depth profile is calculated utilizing an Euclidean distance transform (Sonka, Hlavac and Boyle 2007). For calculation of the distance metrics, morphologic propagation of the outer surfaces, similar to the concept of grassfire transform (Blum 1967) is applied for fast approximate calculation of the Euclidean distance map from the object’s borders. Results of the depth profile calculation are presented in Fig. 4. For the depth profile only the outer depth values below a threshold $t$ are of relevance for gyrus and sulcus modelling. Depth profile values in the inner ventricular area are to exclude.

4.3. Watershed-Inspired Topography Analysis

Based on the calculated depth profile, the shortest graph interconnecting all local depth minimums is calculated, as well as a graph for connection of all local depth maxima. In the following delineation of the method, only the graph creation for the local maxima is addressed. The graph modelling for the local minimums can be derived by changing the leading signs and processing order.

The iterative process of graph construction is outlined in pseudo-code listing 1 and explained in detail in the following paragraphs.

For the autonomous regions to be interpreted as local maxima (condition 1), the first part of the graph is calculated via skeletonization. Below a region count of $R=50$ voxels, the region element closest to the centre of mass is taken as starting sub-branch of the graph, whereas for larger regions the medial axis, precisely extracted via skeletonization (Jonker 2002; Zwettler et al. 2009), is applied as starting sub-branch of the graph. Regions with only one adjoined neighbouring region (condition 2), necessitate no discrete skeleton calculation. Instead, the region elements are just merged...
with the neighbouring region, already defining a sub-
branch.

If the current region is surrounded by more than
one neighbour region at higher depth profile values
(condition 3), besides a merge with the first neighbour
region, all involved adjoined regions need to be
cumulated. As all of the involved neighbouring regions
have an already defined sub-part of the graph, these
segments must be iteratively linked together. This link
operation, described in the following sub-chapter, is a
crucial task as it significantly influences the resulting
graph after processing all depth levels from the deepest
to the lowest profile values, see Fig. 5 for illustration of
described iterative topography analysis. Starting at the
deepest values with autonomous regions and the first
skeleton parts Fig. 5 (c), the regions are enlarged
whenever adjoined new regions at lower depth profile
values are reached Fig. 5 (d-e). In case of reaching
regions with already defined skeletons, the optimum
connective path must be found Fig. 5 (f-g) utilizing
detection of the optimal path, see Fig. 6. That way the
topography describing path can be iteratively
constructed until one final region remains Fig. 5 (h).

```
regions;
  for(depth=maxVal; depth>=minVal; depth--)
    currRegions;
    for(xIdx=0; xIdx < sizeX; xIdx++)
      for(yIdx=0; yIdx < sizeY; yIdx++)
        for(zIdx=0; zIdx < sizeZ; zIdx++)
          if((distanceMap[xIdx][yIdx][zIdx] ==
            depth)&&
            (!classified(xIdx,yIdx,zIdx)))
            currRegions.Add(
              getRegionAt(xIdx,yIdx,zIdx));
    for(region : currRegions)
      neighbourRegions = getNeighbours(region);
      if(neighbourRegions.size == 0)
          region.CalculateSkeleton();
          regions.Add(region);
      else if(neighbourRegions.size == 1)
          merge(neighbourRegions.first, region)
      else
          merge(neighbourRegions.first, nRegion)
          for (nRegion : neighbourRegions)
            if(nRegion != neighbourRegions.first)
              link_merge(
                neighbourRegions.first, nRegion)

Code Listing 1: Illustration of the topography analysis
algorithm implemented in pseudo code. Regions at the
same depth level are grouped together via region
growing and stored in currRegions. Then for each
region in currRegions, the neighbouring regions are
identified. In case of neighbourhood condition 1 with
N=0, a seed region has been detected and the first
skeleton is calculated. Seed regions are added to the
global region container regions. If there is exactly
one neighbouring region at higher depth level
(condition 2), a region merging is performed. In case of
additional neighbours (condition 3), the shortest
skeleton path linking the involved regions is calculated
prior to performing the merge operation.

Figure 5: Illustration of the topography analysis on an
cross-shaped 2D terrain.

4.3.1. Assembling Graph from Sub-Segments

When assembling two regions \( r_1 \) and \( r_2 \) with pre-
calculated skeletons, i.e. graph sub-segments, the
optimum path between the two skeletons \( skel_1 \) and \( skel_2 \)
must be found. The search for the optimal connection
path \( p_{\min} \) is defined as minimization problem of a fitness
function \( F \), accounting for the Euclidean distance \( \text{dist}(i) \)
of the path and depth gradients of the path as \( \text{depth}(i) \)
with respect to the maximum profile depth \( \text{depth}_{\text{max}} \), as

\[
F(p) = \sum_{i=1}^{L} \text{dist}(i) \cdot (1 + \text{depth}_{\text{max}} - \text{depth}(i)) \quad (2)
\]

Thereby the target path \( p_{\min} \) is that path of all possible
non-cyclic connections between \( skel_1 \) and \( skel_2 \) with a
minimal value for \( F \).

For calculation of the optimal path, the metric
value \( F \) of the current sub-path is propagated to its
neighbours, starting at the skeleton elements \( skel_1 \) of
region $r_1$. The sub-path fitness value is recursively propagated to the particular neighbours and updated for each added path element. Whenever the first element of $skel_2$ is reached, an upper border for the optimal path fitness value is given. That allows reduction in the calculation complexity as all sub-paths exceeding this upper border fitness value can be aborted.

After convergence of the fitness values is reached, the optimum path can be traced back starting at the element $e_2$ of skeleton $skel_2$ with an adjoined neighbour showing the lowest fitness. Starting at $e_2$ the way to skeleton $skel_1$ is traced back by selecting the element with lowest fitness in each particular neighbourhood. The optimal path is finished, when the first neighbour of $skel_1$ has been reached. The described algorithm is illustrated in Fig. 6.

![Figure 6: Detection of the optimal path between the two regions in Fig. 5 (f). Starting at the skeleton of the first region, all neighbours in $N_8$ are set or updated with the minimal path value according to the metrics defined in Eq. 2. Whenever a path value gets changed, all adjoined neighbours must be checked for propagation of the new value (b). Finally the path connecting the skeletons $skel_1$ and $skel_2$ can be traced back by picking each neighbour with the lowest value until skeleton of $r_1$ is reached (d).](image)

5. RESULTS
In this chapter first results of the discussed method are presented and discussed.

5.1. Results of Sulcus Classification
The graph model resulting from sulcus classification highly correlates with the main anatomical folds, see Fig. 7. Some pruning and smoothing operations on the graph can be utilized in future to remove dispensable bifurcations and redundant node elements.

![Figure 7: Visualization of sulcus analysis results in sagittal view (a-d) and axial view (e-h). The resulting tree model for the sulcus folds (b)(f) is amplified via morphologic dilation operation utilizing a 3x3x3 structuring element for better visibility (c)(g). Correlation of original brain surface (a)(e) and the sulcus models is presented in (d)(h) via overlay.](image)

As our presented algorithm only interconnects adjoined region skeletons bridging shallow sections, foreshortening could arise if the depth profile is strictly monotonic increasing. That problem can only be exploited by processing artificial testing data. As the first results show, even minimal deflection from strict monotonic behaviour prevents any foreshortening of the particular sub-branch, processing real-world medical image data.
6. DISCUSSION AND CONCLUSIONS
In this paper we introduce a novel concept for classification of the gyrus and sulcus folds and representation as graph model.

The presented iterative topography analysis with step-wise assembling and interconnecting the final graph allows handling of local imbalances of the depth profile, e.g. due to deformation or asymmetries of the brain. Besides roughly estimated upper threshold value for the profile depth to process with respect to imaging resolution, no further parameterization is required.

Linking the final graph representations and the depth profile allows later restrictions on the granularity of sulcus and gyrus representation.

Utilizing the presented sulcus and gyrus graph models, classification of the brain areas can be easily achieved by iterative assigning the gray matter and white matter voxels to the closest neighbouring sub-trees, similar to vasculization-based anatomy classification (Zwettler, Backfrieder and Pichler 2011). Thereby the gyrus course can be used for iterative classification of the surrounding tissue and the sulcus lines are incorporated as additional barriers to prevent invalid merging of adjoined brain sections.

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