Survey of the Visual Exploration and Analysis of Perfusion Data

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Abstract—Dynamic contrast-enhanced image data (perfusion data) are used to characterize regional tissue perfusion. Perfusion data consist of a sequence of images, acquired after a contrast agent bolus is applied. Perfusion data are used for diagnostic purposes in oncology, ischemic stroke assessment or myocardial ischemia. The diagnostic evaluation of perfusion data is challenging, since the data is complex and exhibits various artifacts, e.g., motion artifacts. We provide an overview on existing methods to analyze, and visualize CT and MR perfusion data. The integrated visualization of several 2D parameter maps, the 3D visualization of parameter volumes and exploration techniques are discussed. An essential aspect in the diagnosis of perfusion data is the correlation between perfusion data and derived time-intensity curves as well as with other image data, in particular with high resolution morphologic image data. We discuss visualization support with respect to the three major application areas: ischemic stroke diagnosis, breast tumor diagnosis and the diagnosis of coronary heart disease.

Index Terms—Medical visualization, multi-parameter visualization, volume rendering, perfusion data.

1 INTRODUCTION

Compared to static image data, where the morphology of anatomic and pathological structures is represented with high spatial resolution, dynamic image data characterizes functional processes, such as metabolism and blood flow, which is often essential to detect diseases at an early stage or to discriminate pathologies with very similar morphology.

Important examples of dynamic medical image data are functional MRI, where activations of brain areas are imaged, dynamic PET and SPECT, where the temporal distribution of a radioactive tracer is measured to assess metabolic processes and perfusion imaging, where the blood flow is measured. We focus on CT and MR perfusion data which are acquired to support essential diagnostic tasks, e.g., ischemic stroke diagnosis, the assessment of different types and stages of tumors and the detection and diagnosis of coronary heart disease (CHD).

With modern CT and MRI devices, the effects of perfusion can be measured in high spatial and temporal resolution. In perfusion imaging, the distribution of contrast agents (CAs) is registered to assess blood flow and tissue kinetics. Signal intensities after the administration of a CA are recorded. Whether or not a CA is delivered and subsequently absorbed within a particular region, how long it takes until the maximum amount of CA is delivered as well as other perfusion parameters are determined for medical diagnosis. These parameters are substitutes for physiological parameters such as tumor perfusion and vessel permeability, which e.g., characterize the malignancy of a tumor [1].

Visual exploration of perfusion data is particularly challenging. It is primarily based on the derived perfusion parameters, which represent features of time-intensity curves (TICs). These parameters are derived for each voxel of the perfusion data and represent a high-dimensional space, usually of five to eight parameters. The correlation between these parameters as well as the local distribution of single perfusion parameters are essential. Since the time-dependency is not represented in the perfusion parameters, often perfusion maps along with TICs have to be analyzed. The comprehensible and simultaneous display of these curves and perfusion maps poses considerable challenges for the layout. The visual exploration is also challenging due to the character and the quality of the data: they exhibit various artifacts and thus the visualization also serves the assessment of the reliability of the original data and also the assessment of preprocessed data, where artifacts are reduced. In contrast to static CT data, no absolute scale for the intensity values exists. Therefore, simple visualization techniques with predefined (absolute) settings are not applicable. As a consequence of these difficulties, image processing and visualization have to be tightly integrated and a variety of visualization techniques is needed to detect and characterize important features.

This paper is organized as follows: In Section 2, we give a brief overview on the medical background in selected application areas. In Section 3, we briefly describe the image data processing which enhances the expressiveness of simple visualization techniques and is indispensable for advanced visualization support. Basic
visualization techniques, which are widely available in commercial software, are presented in Section 4. More advanced techniques for the visual exploration, data analysis techniques as well as the combination of both are discussed in Section 5. The application of advanced techniques is described in separate case studies on cerebral perfusion (Section 6), tumor perfusion (Section 7) and myocardial perfusion (Section 8). In Section 9, we propose guidelines for the use of the basic and advanced visualization techniques.

2 MEDICAL BACKGROUND

In perfusion imaging, a certain amount of a CA is injected intravenously and its distribution is measured by a repeated acquisition of subsequent images covering the volume of interest [2]. The CA causes signal changes and works as a tracer of perfusion which means that the uptake of the tracer is relative to the blood flow. The time-dependent behavior of the signal depends on the type of CA applied. The CA circulates through the body in several passages until it gets completely excreted. In perfusion imaging, normally only the first-pass, which shows the most significant signal changes, is evaluated. Perfusion imaging differs strongly from static imaging, since greater care must be exercised in injection rate and dose, image timing and image analysis. The diagnostic quality of perfusion data and the derived perfusion parameters depends on the type of CA, the amount of CA and the speed of injection. Currently, such imaging techniques are mainly performed in a research context [1].

Depending on the physiological process, either the short-term blood flow or the long-term (> 1 min) diffusion process of the tracer particles through the membranes of the micro-vessels are represented in the varying signal of the image voxels. Different imaging sequences are used for perfusion imaging: T1-weighted MRI data are typically used for breast cancer diagnosis, whereas T2-weighted MRI data and CT images are employed for the diagnosis of ischemic stroke. In T1-weighted imaging and CT imaging, a signal enhancement is achieved in areas of CA accumulation. In contrast, T2-weighted imaging leads to a decrease of signal intensity where the CA accumulates. To facilitate a consistent processing of the data from both sequencing modalities, the signal intensities in T2-weighted datasets are often inverted prior to any visualization task. This leads to the more intuitive mapping of CA accumulation to signal enhancement (rather than attenuation). The T2-weighted cerebral perfusion datasets used to generate Fig. 1 (left) and Fig. 4 have been inverted. Another MR imaging technique—which is not yet used in clinical routine—is arterial spin labeling. With this non-invasive approach, a CA does not necessarily need to be injected. The CA can be excited endogenous protons [3].

Perfusion datasets from different application areas considerably differ in spatial and temporal resolution. Table 1 lists typical parameters for data from MR perfusion imaging. In contrast to cerebral and breast tumor perfusion data which continuously cover the volume of interest, myocardial perfusion data exhibit large gaps (e.g., slice thickness: 6 mm and 12 mm gaps).

<table>
<thead>
<tr>
<th></th>
<th>matrix</th>
<th>$R_S$</th>
<th>$D_S$</th>
<th># slices</th>
<th>$R_T$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cerebral</td>
<td>128$^2$</td>
<td>2</td>
<td>7</td>
<td>10–15</td>
<td>1–2 (40–80)</td>
</tr>
<tr>
<td>Breast tumor</td>
<td>512$^2$</td>
<td>0.7</td>
<td>2</td>
<td>60–80</td>
<td>60–90 (3–10)</td>
</tr>
<tr>
<td>Myocardial</td>
<td>128$^2$</td>
<td>1.5</td>
<td>18</td>
<td>3–4</td>
<td>0.5–1 (&gt;40)</td>
</tr>
</tbody>
</table>

**Perfusion parameters.** For the diagnosis, regions of interest in healthy and suspicious tissue are defined, and TICs—averaged over all voxels in a selected region—are analyzed. Typical TICs from cerebral and breast tumor perfusion are presented in Fig. 1. The curves observed in myocardial perfusion diagnosis are similar to those of cerebral perfusion. In both application areas, regions exhibiting no significant or a delayed and diminished enhancement (red and green curves in Fig. 1 (left)) are of interest. However, in breast tumor perfusion, regions showing a high early enhancement followed by rapid wash-out, i.e. a decrease of signal intensity afterwards, are especially suspicious (red curve in Fig. 1 (right)). To achieve a more quantitative description of curve shape, perfusion parameters are derived from the TICs. Depending on the application area, different sets of perfusion parameters are relevant. However, some parameters are of general interest for almost all application areas (see Fig. 2). Before we describe these parameters, we introduce three auxiliary variables necessary for a reliable evaluation.

![Fig. 1.](image)

Left: TICs for regions of gray matter in the brain (40 measurements). The blue curve shows normal brain perfusion. The red curve indicates no significant perfusion in the infarction core. The green curve shows decreased and delayed perfusion around the core. Right: TICs of different regions in breast tissue (5 measurements). The enhancement relative to the signal intensity at the first points in time is shown. The red curve is especially suspicious because of its strong wash-out, which is typical for malignant tumors.
Fig. 2. A typical TIC in myocardial perfusion with a significant first pass and an alleviated second pass of CA traversal annotated with the essential parameters to evaluate the first pass.

The CA arrival represents the point in time when the signal enhancement actually starts, whereas TimeEnd refers to the end of the first CA passage. The Baseline represents the average intensity before CA arrival (see Fig. 2). These auxiliary variables are determined to focus the evaluation of the TIC on the relevant portion.

Assessing perfusion considering the actual CA arrival (time), TimeEnd and the Baseline is essential to compare perfusion analysis results from different scanning devices and patients. Major diagnostically relevant perfusion parameters are:

- **Peak Enhancement** (PE). The maximum value normalized by subtracting the baseline.
- **Time To Peak** (TTP). The point in time where PE occurs, normalized by subtracting the CA arrival time. This parameter allows assessing whether blood supply is delayed in a particular region. If the peak is not a significant maximum or the temporal resolution is low, the TTP value is not expressive. The signal change in the interval between CA arrival and TTP is referred to as wash-in, whereas the signal change in the time between TTP and TimeEnd is referred to as wash-out.
- **Integral**. For a certain time interval (often representing one cycle, or pass, of blood flow) the area between the curve and the baseline, the approximated integral, is computed. Together, PE and Integral allow to assess whether the blood supply is reduced in a particular region. Reduced and delayed blood supply is a strong indicator for a damaged region for example in ischemic stroke or CHD diagnosis.
- **Mean Transit Time** (MTT). In the time interval used for the integral calculation, MTT specifies the point in time where the integral is bisected. It is normalized by subtracting CA arrival.
- **Slope** characterizes the steepness of the curve during wash-in. Depending on the temporal resolution, different regression methods, such as the Gamma-Variate and a linear fit are used to characterize the curve progression. The term *Up-slope* in CHD diagnosis relates to the maximum slope between two or three subsequent timesteps between CA arrival and TTP.
- The *DownSlope* characterizes the steepness of the descending curve during wash-out and is computed similar to the **Slope**.

**Application areas.** Throughout this paper, we focus on ischemic stroke, breast tumor and CHD diagnosis. However, perfusion analysis bears a great potential in other diagnostic tasks as well. As an example, it has been shown that renal perfusion [4] and lung perfusion [5] enhance selected diagnostic processes, such as detecting disorders of pulmonary vessels and acute pulmonary embolism.

**Commercial perfusion software.** The evaluation of perfusion data is supported by a variety of specific tools dedicated to a particular diagnostic question and to a particular modality. Examples are the PERFUSION 3 SOFTWARE, as package for the General Electric, Advantage Windows workstation, the SIEMENS Syngo, Neuro Perfusion software and Philips CT perfusion software. The basic features, the ability to analyze TICs for selected pixels and regions as well as the display of parameter maps are common to all these systems. In nuclear medicine, the EMORY CARDIAC TOOLBOX and Cedars-Sinai’s QUANTITATIVE GATED SPECT SOFTWARE are the two most widely used packages for PET and SPECT data analysis.

## 3 Data Processing

Data processing techniques are experimental, often subject of active research and therefore not widely available. Depending on the specifics of an application area, e.g. the temporal resolution and the amount of motion, data processing techniques have to be applied.

**Motion correction.** The analysis and visualization of perfusion data relies on comparable image data. Otherwise, subtraction images and TICs are misleading. Comparability means that a voxel with coordinates \((x, y, z)\) at time \(t_1\) corresponds to a voxel with the same coordinates at time \(t_2\). Often, a motion correction has to be carried out to achieve comparability. This is essential in assessing tumor and myocardial perfusion where the inter-voxel correspondence is hampered due to breathing, patient movement, muscle relaxation or heartbeat (Fig. 3). Without motion correction, the subtraction volume is filled with bright artifacts. Motion artifacts might hide relevant signal changes, but also pretend signal changes that are actually not present.

In Dynamic Contrast-Enhanced (DCE)-MRI mammography, breathing and muscle relaxation result in considerable soft tissue deformations. Here, rigid registration approaches are not appropriate. Elastic registration that considers local transformations enables a better registration quality. The registration algorithm described by
Rueckert et al. [6] is a good basis, which employs normalized mutual information [7] as similarity measure. Originally developed for motion correction in DCE-MRI mammography, it is now also used for motion correction in other application areas.

**Calibration of signal intensities.** While CT imaging provides calibrated signal intensities in “Hounsfield” units, MRI signals are dependent on the scanning sequence used. Therefore, it is very important to calibrate the raw signal intensities versus CA concentration. For gradient-echo MR sequences, the calibration can be based on the assumption of a linear correspondence between CA concentration and signal intensity [1]. When user-individual vascular input functions are used for the quantitative analysis instead of a standardized assumption about the inflow of CA into the region of interest, this calibration can be neglected [8]. In this case, input functions correspond to the intensity curves of reference tissue with high vasculature.

**Temporal denoising.** Since TICs exhibit high-frequency noise, smoothing in the temporal dimension is essential for a reliable analysis. Lysaker et al. [9] introduced an appropriate filter based on partial differential equations, which simulate a diffusion process and applied it to DCE-MRI mammography data. For the generation of parameter maps, such as MTT and Integral, the “right” points in time must be chosen (recall Sect. 2, auxiliary variables). The smoothed visualization supports this selection.

### 4 Basic Visualization Techniques

Some straightforward techniques to visualize and analyze perfusion data are:

- **cine-movies**, which step through all points in time for a selected slice,
- subtraction images, depicting the intensity difference between two selected time points, and
- color-coded parameter maps for a selected slice. A parameter map depicts the value of a perfusion parameter in a pixelwise manner.

**Cine-movies.** The cinematic depiction of gray scale images in a movie loop is helpful to assess image noise and artifacts [1], but especially for the assessment of enhancement patterns. In CHD diagnosis, cine-movies are not only applied for assessing myocardial perfusion but for evaluating the left ventricular wall motion in functional MR data. This special type of data is acquired to image the contraction of the myocardium.

**Subtraction images.** Subtraction images may also be used for quality control; the injection of a CA leads to an increase of signal intensity. If the subtraction for two early points in time, $t_2$ and $t_1$ with $t_2 > t_1$, leads to a negative value it is likely that the pixels do not correspond to each other due to motion artifacts. If this occurs, motion correction is indispensable for a meaningful analysis. In T2-weighted imaging, where the intensity decreases after CA arrival, the quality control must be adapted (recall Sec. 2).

In Figure 4, two subtraction images are shown, which are used for the diagnosis of an ischemic stroke. Both reveal a dark area in the right hemisphere (left part of the images). This is suspicious, since it does not occur in the corresponding region of the left hemisphere. The region, which is dark in both images, depicts the core of an ischemic stroke. Around this region, a larger area appears dark in the early subtraction image (left), but bright in the subtraction image which refers to a later time (right). This region shows the “tissue at risk” around a stroke core.

![Fig. 4. Subtraction images to analyze cerebral perfusion.](image)

Left: difference between $t_6$ and $t_2$. The low perfusion in a larger portion of the right hemisphere (left part in the images!) characterizes the infarction zone. Right: difference between $t_{17}$ and $t_2$. The late enhancement in a part of the right hemisphere represents the “tissue at risk”. It is characterized by a high signal intensity. (Data is courtesy of Jonathan Wiener, Boca Raton Community Hospital).
Several parameter maps in one image, information derived from perfusion data with morphologic information from another dataset, and advanced visualization techniques is presented in section 5.4. The combination of data analysis and advanced visualization techniques is presented in section 5.5.

Subtraction images provide valuable information for the diagnosis. However, there is no assistance in choosing the “right” points in time for subtraction images. Moreover, the 2D data are only used to visually detect abnormalities. Subtraction images do not provide quantitative temporal and spatial information, which could make the diagnostic results more reproducible.

Parameter maps. Parameter maps are displayed as color-coded images (see Fig. 5). Besides parameter images, it is common to compute TICs for user-selected regions of interest. Often, a parameter map is used first to detect interesting or suspicious regions, whereas TICs in selected regions are analyzed later.

Summary. Basic visualization techniques for exploring perfusion data were described by Behrens et al. [10] and König et al. [11]. Meanwhile, the techniques described above have been integrated into commercial software packages (recall Sec. 2). The diagnostic quality achieved by using these techniques often depends on appropriate data processing (recall Sec. 3).

5 ADVANCED VISUALIZATION AND ANALYSIS TECHNIQUES

Advanced visualization techniques for exploring perfusion data are motivated by three essential drawbacks of basic visualization techniques. They do not support the integration of:

• several parameter maps in one image,
• information derived from perfusion data with morphologic information from another dataset,
• extracted features.

The following sections 5.1 to 5.3 show how these problems may be tackled. Data analysis techniques which support the visual exploration by means of classifying tissue according to perfusion characteristics are described in section 5.4. The combination of data analysis and advanced visualization techniques is presented in section 5.5.

5.1 Multiparameter Visualization

The integrated analysis of several perfusion parameters in a suspicious region is essential for various diagnostic tasks [12]. To support the analysis, we discuss the appropriateness of integrated multiparameter visualizations. These visualizations are based on pre-computed parameter volumes, where the corresponding perfusion parameters are represented for each voxel of the original dataset.

In principle, color may be employed for two or three parameters as well. Among the wide-spread color spaces, the HSV space (describing a color by its Hue, Saturation, and Value component) is the best choice since it is perceptually roughly linearized [13]. To be compatible with expectations of users, the most suspicious parameter combinations may be mapped to a red color (Hue) with high saturation and high intensity (Value), whereas normal parameter values are mapped to lower saturation and intensity values and a bluish hue component. With this approach, the viewer’s attention is directed to suspicious regions. However, the simultaneous visualization of three quantitative values relating to data with high spatial frequency is in general very hard to interpret. The correct interpretation of two or even three perfusion parameters by means of one color cannot be achieved by preattentive vision. Therefore, Oeltze et al. [14] investigated methods where color (for one perfusion parameter) is combined with another visualization technique for displaying a second parameter. Isolines, height fields, or orientations of textures might be employed to combine several parameters within a single image [15]. In particular, the combination of isolines and colors is effective and can be easily interpreted. Oeltze et al. [14] also discussed the use of color icons [16].

Combining isolines and color-coding. Isolines connect regions where the investigated perfusion parameter has a certain value. Isolines are easily computed by the Marching squares algorithm [17]. Noise removal is important in order to prevent that many irrelevant small and distracting isolines or relevant but jaggy lines result (see Fig. 6). In contrast to color-coding isolines are not interpreted at a glance but allow a more quantitative interpretation.

Colored height fields. Colored height fields enable the integrated visualization of two parameter maps. A 3D elevation profile is generated based on the pixel-values of the first parameter map. In a next step, the resulting profile is colored according to the pixel-values of the second parameter map and an arbitrary color look-up-table. The profile may be freely rotated such that initially occluded parts become visible. The mapping to height is scalable. It is initially adapted to the domain of the first parameter (see Fig. 7). As a natural mapping, PE should be mapped to the height parameter.

Fig. 5. Parameter maps TTP, MTT, and the relative cerebral blood volume (roughly corresponding to the general perfusion parameter Integral) of a cerebral MRI perfusion dataset are depicted. The delayed blood flow in the right hemisphere (left part of the images!) becomes obvious. (Images are courtesy of Jonathan Wiener, Boca Raton Community Hospital)
Exploration of multiple parameter images with lenses. Lenses are used to explore conventional images. Digital lenses—working as pixel magnifiers—are also required in digital image exploration to analyze small scale phenomena within enlarged visualizations. The interaction with movable viewing lenses (Magic Lenses) is useful for exploring multidimensional data [18], where lenses do not magnify information but show different information in the lens-region. For parameter maps, lenses may show information relating to one parameter either in the context of the original perfusion data (see Fig. 8) or in the context of a map of another parameter (see Fig. 12).

With this interaction style, the user starts by selecting a foreground and a background parameter (for example TTP and MTT) and then moves a lens (a rectangle or an ellipse) to select either of the parameter set. Inside the lens region, displays of the foreground and background parameter are combined by means of alpha-blending (Fig. 12) thus combining an opaque background and translucent foreground to imitate transparency. With an alpha value equal to 1, only the foreground parameter is represented (Fig. 8).

Glyph-based visualization of multiple parameters. Glyphs represent a standard technique in the visualization of multi-field data. A glyph is a simple geometric primitive which is positioned with respect to the original data points in space and whose attributes, e.g., color, extension, size, and orientation, are modified according to some represented values. The integrated glyph-based visualization of multiple perfusion parameters is presented by Oeltze et al. [19] with a focus on an intuitive mapping of perfusion parameter values to glyph shape. Intuitive mapping here refers to the generation of an easy to learn glyph coding of TIC shape, e.g.,
by mapping parameter $Integral$ (area below the curve) to size, and $Slope$ (steepness of the ascending curve) to orientation. Several 2D glyph shapes, e.g., circular discs, ellipses, rectangles or toroids, with different visual attributes besides color have been implemented in slice-based visualizations. Inspired by the work of Doleisch et al. [20], a feature definition component has been integrated to speed up the evaluation of the complex multi-field parameter data. It is based on the tight integration of the glyph display with multiple statistical representations, connected by (smooth) brushing facilities applied to scatterplot representations. Initial tests showed that the placement of one glyph per data point results in too small glyphs which is due to the limited screen space (see Fig. 9 (left)). Hence, a multi-resolution glyph display has been proposed to improve the readability of the glyph attributes (see Fig. 9 (right)). The display incorporates different resolution layers which can be interactively explored by the user.

**Summary.** Multiparameter visualizations enable the simultaneous visualization of two or more perfusion parameters. Height fields are visually attractive and might be used to present diagnostic results, e.g., in educational settings. However, the use of height fields poses interaction problems due to occlusion and is probably not an optimal choice for efficient routine diagnosis. The combination of color and isolines is more promising to support the integrated visualization of two parameters. The use of lenses is based on ideas of clinical users and was considered useful in informal discussions with a neuroradiologist and a cardiologist from the medical faculty of the University of Magdeburg. Both appreciated that they can explore two parameter maps simultaneously instead of having to mentally integrate the information from spatially separated visualizations.

### 5.2 Integrating Dynamic Information and Morphology

Relevant perfusion parameters are often only extracted and visualized for a restricted region in the entire dataset, e.g., a tumor or the myocardium. However, other constituents such as the bony structures might provide substantial information for displaying the diagnostically relevant regions in their anatomic environment. Also other surrounding tissues that are not enhancing, can be of indispensable diagnostic value. Therefore, it is useful to add spatial reference information in the regions not containing relevant dynamic information. As a simple example, the display of an original slice together with the height field in Fig. 7 supports the spatial orientation which is hampered by the circular shape of the myocardium. A reasonable strategy to add spatial reference information is to color-code dynamic information and to display the reference data in the background using a gray scale. Depending on the resolution of the image data, the integration of dynamic and morphologic information should be carried out in 2D slice visualizations or 3D renderings. For DCE-MRI mammography data with more than 50 slices, 3D renderings are appropriate, whereas cerebral and myocardial perfusion data provide a too small number of slices.

The assessment of perfusion data might benefit from segmentation information, for example, concerning suspicious breast lesions. In this case, the visualization of dynamic information might be restricted to the segmented region.

### 5.3 Probing and Annotating of Perfusion Data

Another way to depict temporal curves specified at every position of the data is to render them outside of their spatial location. The most common way is to show a set of time-dependent graphs for a set of pre-selected spatial locations, respectively. With this approach one might loose the correspondence between the spatial position of the measured data and the curve data itself.

Mlejnek et al. [21], [22] proposed the Profile Flag, an intuitive tool for browsing and annotating of temporal data. It enables the visualization of spatial or temporal curves closely connected to the rendering of the anatomic structure of the data without removing any parts thereof. The Profile Flag looks like a board-pin-like glyph which consists of a banner, a marker, a range selector and a set of needles (see Fig. 10 (left)). The Profile Flag can be positioned on and dragged along the surface or inside of the inspected anatomical object. For probing of the underlying data, the set of needles is positioned beneath the surface of the probed structure at locations of interest. Each needle defines the position of one probed location or curve. The flag pole is a cylinder

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**Fig. 9.** Glyph-based visual exploration of cerebral perfusion parameters. The glyph display in all images has been restricted to suspicious regions by means of smooth brushing. *Left:* One circular disc is placed per data point. Changes in glyph size are hard to interpret. A magnification (inlet) improves the readability but involves a loss of context information and spatial orientation. *Right:* The application of a lower resolution layer solves the problem. (Data is courtesy of Jonathan Wiener, Boca Raton Community Hospital)
which connects the banner with the range selector. The size of the range selector can be changed by moving the marker along the flag pole. The range selector defines the size of the ROI, i.e., it encloses the set of needles. For 3D dynamic perfusion data, the probed information is taken along the time axis (TICs). The temporal development is probed at a specific 3D point location. In the simplest case, this probing position and interesting nearby locations are specified by a spherical range selector. In the more general case, the range selector can be a general polytope including all emphasized TICs. Another way of interaction with the range selectors deals with the selection of TICs based on its properties. One can define more general criteria (e.g., maximal deviation from a pre-defined reference curve) for the selection of a set of profiles. In this case, the shape of the range selector determines the size and shape of the region which includes the selected curves (see [22] for details).

The probed curve data is visualized on the banner. Several types of banners can be defined, e.g., a single-profile banner only shows the probed values at the position of a single needle. During the investigation, one or more Profile Flags can be stuck into the inspected object. They can be moved along the object’s surface, while showing the underlying probed TICs. Multiple Profile Flags can be placed to emphasize differences between areas from different spatial locations, e.g., healthy vs. suspicious regions. For the visualization of dynamic data, the horizontal axis usually corresponds to the time-axis. Therefore, for time-varying data, the banner visualizes the time steps along the horizontal axis, while the vertical axis shows the values for each measured time step. Additionally, for sparse temporal data (i.e., just a few time steps), vertical lines are included in order to facilitate reading off the values at particular time steps. Figure 10 (right) shows an annotation of a DCE-MRI visualization with four time steps by two Profile Flags. The left Profile Flag is located at a healthy region (low increase in the first post-contrast step), while the right Profile Flag illustrates the size and the shape of a suspicious region. The range selector encloses all spatial locations of profiles with similar properties. In this case the selected set of profiles defines the shape of the range selector. In order to avoid visual clutter, an averaged banner is applied, which shows one curve computed by averaging all probed curves.

The basic concept of Profile Flags has been qualitatively evaluated by the radiologists from Catharina Hospital in Eindhoven [22]. An informative survey indicated that the tool is a valuable add-on to the currently used examination procedures. Furthermore, the complete diagnosis including the anatomy as well as the annotated regions can be printed on one sheet of paper. Thus, one of the essential aspects of the Profile Flags is the efficiency of diagnosis communication between different hospital divisions.

5.4 Analysis of Perfusion Data

Another venue of analyzing perfusion data relates to a statistical analysis as well as to mining and knowledge discovery techniques. In particular, the classification of DCE-MRI mammography data by means of artificial neural networks and clustering techniques is an active research area [23], [24], [25]. As an example, Twellmann et al. [25] applied an artificial neural network architecture which combines unsupervised and supervised techniques for the voxel-by-voxel classification of temporal kinetic signals derived from DCE-MRI mammography data. Chen et al. [23] investigated and developed a fuzzy c-means clustering-based technique to automatically identify characteristic kinetic curves from segmented breast lesions in DCE-MRI mammography data. Nattkemper and Wissmueller [26] described the application of self-organized maps to time curve features of DCE-MRI mammography data and discussed how the results may be visually represented as color-coded cross-sections. Automatic classification may be useful in a screening setting in order to replace the opinion of a second radiologist or to direct a radiologist to suspicious regions.

5.5 Combining Analysis and Visual Exploration

Recently, data analysis techniques and advanced information visualization techniques have been combined in order to efficiently explore the space of perfusion parameters [27]. In particular, a correlation analysis is carried out so as to investigate which perfusion parameters strongly correlate. The remaining parameters are processed by a Principal Component Analysis in order to detect major trends. Inspired by the work of Doleisch et al. [20], the trends as well as the original perfusion parameters are displayed in 2D-histograms and scatterplots and are used for (smooth) brushing of
relevant subsets of the data (see Fig. 11). This overall strategy turned out to be useful to discriminate different tissues in cerebral perfusion data, tumor perfusion and myocardial perfusion data.

The overall analysis pipeline is probably too complex for routine diagnosis but may be essential for researchers investigating the effects of CAs, magnetic field strength and other imaging parameters on the diagnostic value of certain perfusion parameter combinations. The pipeline has been discussed with two experienced radiologists from the medical faculty of the University of Magdeburg, both familiar with perfusion imaging in the clinical routine, though not in a research context. Both argued that the data analysis is only applicable in the clinical routine–in particular in emergency cases–if carried out in the background leading to an initial suggestion for suspicious regions. Both assessed brushing as valuable for exploring a non-standardized parameter domain. They appreciated the visualization of perfusion data in 3D since it provides a good overview.

The combination of analysis techniques, linking and brushing for efficiently locating features in perfusion data has been extended with a dense visualization of TICs for all voxels of a perfusion data set. Special techniques are used to reduce clutter in the visualization of a multitude of TICs and dedicated brushes are employed to define TIC target shapes, e.g. a sudden increase and a later decrease of the signal intensity [28].

More specifically, the exploration of higher dimensional histograms for discriminating tissue in ischemic stroke diagnosis was described by Grzesik et al. [29]. They incorporate MR perfusion data and diffusion-weighted MR data in order to integrate the information from both imaging data.

6 Case Study: Cerebral Perfusion

In contrast to highly permeable vessels in malignant tumors, microvessels in normal brain tissue do not leak as a result of the blood brain barrier. Consequently, there is no enhancement in the extracellular volume. Instead, we observe the first-pass of the CA through the vessel components. About 10 seconds after the first-pass of blood circulation, a broadened second-pass can be seen. The volume of blood in each voxel is diagnostically relevant. It is measured by the Integral parameter of the TIC (see Fig. 2).

CT and MRI are primarily used to assess cerebral perfusion in clinical routine. MRI studies suffer from a lower spatial resolution compared to CT, but allow scanning of the entire brain, and are thus better suited to detect an infarction, if its location is not a priori known. CT and MRI have both been proven to be useful in diagnosing the acute ischemic stroke and in decision making for therapeutical interventions [30]. However, the feasibility of MRI studies is usually restricted due to the low availability of emergency settings in most clinical institutions, superior costs and patient-specific difficulties with obtaining MRI, e.g., claustrophobia [30].

Typical parameters for cerebral MRI perfusion datasets are listed in Table 1. In single-slice perfusion CT, one slice covering a brain area of typically 1 cm is acquired. With multi-slice CT, an area of 2 cm may be covered by two or four slices [31]. To reduce image noise, a large slice thickness (one slice with 10 mm thickness or more recently, two slices with 5 mm thickness) is used. The brain coverage may be extended to 4-5 cm by using two successive bolus administrations [32]. Multi-slice CT with greater arrays of elements, e.g., 64/256-slice CT, will pave the way for whole brain imaging in clinical routine.

Cerebral perfusion images are used for ischemic stroke diagnosis, in particular to discriminate cerebral hemorrhage and ischemic stroke. In case of an ischemic stroke, the existence and the extent of “tissue at risk” surrounding the core of the stroke has to be evaluated. While the core exhibits no significant perfusion (red curve in Fig. 1 (left)), “Tissue at risk” is characterized by a reduced and delayed perfusion (green curve in Fig. 1 (left)). Surgical and chemical interventions may salvage at least parts of the “tissue at risk” [33].

The value of combining cerebral perfusion and diffusion data for predicting stroke evolution is discussed in...
7 Case Study: Tumor Perfusion

The process of CA enhancement in a tumor can be described by the diffusion of tracer particles from the inside of blood vessels into the extravascular space and vice versa before it becomes excreted in the kidneys [37]. The permeability of the vessel walls and the extracellular volume fraction determine the amplitude and the shape of the TIC. TICs—which show a high early enhancement followed by a rapid wash-out—are especially suspicious (see red curve in Fig. 1 (right)), because they indicate strong perfusion and high permeability of vessels. Strong perfusion often results from tumor-induced vessel growth (neoangiogenesis). These newly formed vessels are highly permeable, leading to a rapid wash-out [38]. Less suspicious are curves showing a plateau later on (green curve), or regions that continue to enhance (blue curve). This is typically observed in benign tumors.

Dynamic Contrast-Enhanced (DCE)-MRI mammography has been introduced by Kaiser and Zeitler [39] in 1989. However, only recently the imaging modality gained wide-spread acceptance, which is partially due to the effective computer support [40]. The major diagnostic task is to confirm or reject the hypothesis of a tumor being malignant. Data processing, in particular motion correction is challenging (recall Sec. 3).

In DCE-MRI mammography, T1-weighted images are employed. Contrast enhancement lasts considerably longer than in cerebral blood vessels. Therefore, longer acquisition times are employed. DCE-MRI mammography data is characterized by a high spatial resolution and a low temporal resolution. Typical parameters for DCE-MRI mammography data are listed in Table 1. For more details on tumor perfusion, see [41], [42] and a recent multicenter study [43].

Substantial research to support the diagnosis of DCE-MRI mammography data has been carried out at MeVis Research, Bremen. In the work by Behrens et al. [10], the display of parameter maps, the selection of ROIs, the calculation of TICs, and the quantitative analysis of these curves was presented. Meyer et al. [44] presented a software assistant adapted to the needs of the clinical routine, in particular with respect to the support of breast cancer diagnosis. This assistant has been successfully applied in a clinical trial evaluating the DCE-MRI datasets of 46 patients suspected of having breast cancer [45]. With more advanced visualization options, a fast motion correction, and the incorporation of parameters from a pharmacokinetic model, a new research prototype, DynaVision, was presented by Alfke et al. [46]. The prototype was tested in examining the perfusion of pancreatic carcinoma xenografts in mice with severe combined immunodeficiency disease. The findings from perfusion analysis were well correlated with those from histopathological analysis and could be achieved within a reasonably fast time (15 min).

A long-term effort on visualizing DCE-MRI mammo-
graphy data has also been accomplished by Hellwig et al. [47]. They investigated virtual reality input and output technology for exploring stereoscopic 3D visualizations of the original data and derived parameter volumes. They conclude that the stereoscopic display facilitates an improved localization and differentiation of lesions in space. This is further supported by virtual input devices to adjust, e.g., orientation and transparency of the display. Recently, Coto et al. [48] explored advanced volume rendering techniques for an efficient diagnosis. They have shown that a novel workflow to analyze DCE-MRI mammography data can reduce time needed for diagnosis by utilizing the combination of interactive examination, segmentation and advanced visualization techniques. Moreover, their software MAMMOEXPLORER allows to indicate different types of lesions based on both the shape of the lesion and the temporal development of the TIC.

In the following, selected visualization techniques which have been used for computer support for breast tumor analysis are described.

**Color coding.** Two parameters describing the diagnostically significant shape and amplitude of each pixel’s TIC may be mapped to color [49]: (1) the slope of the early CA enhancement to brightness and (2) the slope of the late wash-out to the color value, encoding suspicious wash-out in red. Using continuous color values creates a smooth transition between slowly enhancing and depleting regions.

**Projection methods.** For the integration of morphologic information and perfusion parameters, 3D visualization techniques are useful. To avoid visual clutter, the visualization of perfusion parameters should be restricted to those voxels exhibiting a high dynamic range (significant signal intensity changes). Projection techniques, such as MIP and Closest Vessel Projection [50], provide a direct link between pixels and corresponding voxels with the related TIC. Thus, with a colorized projection image, morphological information can be visualized together with physiological parameters [49] (see Fig. 13).

Kohle et al. [49] suggest to apply a colorized temporal MIP (maximum value along the temporal scale). With this approach, voxels characterized by a strong dynamics (either in the wash-in or wash-out phase) are represented by a color which incorporates the wash-in as well as the wash-out behavior by mapping these values to Hue, Value and Saturation. As an improvement to MIP, closest vessel projection (CVP) also known as local MIP, was developed to add depth information to MIP images [50]. The most intense voxel along the projection ray is no longer selected; rather, the voxel which represents the first local maximum above a certain threshold is taken. The threshold has to be adjusted to display only the interesting structures. A threshold of 20% relative enhancement is appropriate for restricting the visualization to the interesting structures of DCE-MRI mammography data. Figure 13 shows a colored CVP of DCE-MRI mammography data. Both MIP and CVP are offered as whole-volume visualization techniques and as slab rendering—restricted to a portion of the data characterized by two parallel clipping planes.

**Volume Rendering and Information Visualization Techniques.** The analysis of perfusion data with a large number of slices, such as DCE-MRI mammography data, is a tedious and time-consuming task. It involves browsing through all slices and searching for suspicious areas. Afterwards, the procedure is repeated for the identified areas. An effective examination requires the extraction and visualization of diagnostically essential data.

Coto et al. [48] presented several investigation tools (e.g., scatterplots and volume rendering) for the classification and visualization of DCE-MRI mammography data. The approach combines brushing and linking interaction with effective visualization of the selected suspicious areas. For the computation of the TICs, the pre-contrast scan ($t_i$) is subtracted from all post-contrast scans ($t_{i+}$). This step emphasizes the gradient in the temporal dimension of the analyzed curve and highlights suspicious areas. An enhancement scatterplot (see Fig. 14 (left)) is calculated for each post-contrast step. It shows the relative enhancement of the pre-contrast step with respect to the post-contrast step $t_i$. In the interaction step, brushing is performed on one of the scatterplots, while the selected set is emphasized on all remaining scatterplots. If the brushing is performed on multiple scatterplots, the result of the selection is calculated by a
logical “and” operation. The selection set and the way of interaction with the scatterplots depends on the specific application. The result of the interaction can be displayed in a 2D view by highlighting the selected areas, or in a 3D view with importance-driven volume rendering (see Fig. 14 (right)).

8 CASE STUDY: MYOCARDIAL PERFUSION

Perfusion data is also crucial in the diagnosis of coronary heart disease (CHD). At an early stage, the CHD is characterized by a perfusion defect caused by a stenosis (an abnormal vessel narrowing). The localization of the perfusion defect with respect to the myocardium combined with anatomical knowledge about the supplying coronary arteries is essential in detecting stenosis as well as in early CHD diagnosis [51]. In particular, for assessing stenosed coronary arteries, imaging modalities from nuclear medicine, such as dynamic PET and SPECT are widespread. With these modalities, the regional distribution of a radioactive isotope, such as Rb-82 chloride, nitrogen 13 (N-13) ammonia, or oxygen 15 (O-15) water is represented [52], [53], [54], [55].

PET is more specific than SPECT in discriminating vital and irreversibly damaged tissue after an infarction [56]. However, both imaging modalities exhibit a lower spatial resolution and more artifacts, compared to MR and CT perfusion. Also the amount of radiation for the patient is significantly higher for PET and SPECT data acquisition. Myocardial MR perfusion offers an especially attractive alternative since measures of myocardial perfusion, viability and ventricular function can be integrated in a single scanning protocol. It has shown to have at least a similar sensitivity and specificity in comparison to PET and SPECT without exposing the patient to radiation [57], [58]. However, wide-spread use is still hampered by a variety of artifacts and the strong experience necessary to perform the examination and interpret the results. Recent technical advances in hardware, CAs and imaging sequences are described in [51].

The data acquisition is typically accomplished according to the standards of the American Heart Association (AHA) [59] (Fig. 15 (left)). In MR perfusion imaging, the data acquisition is accomplished during breathhold and it is electrocardiogram (ECG)-triggered over a period of at least 40 consecutive heart beats. The acquisition is often carried out at rest and under drug-induced stress. The stress test may even reveal marginal stenosis and is usually performed prior to the test at rest using identical imaging parameters. Typical parameters for myocardial perfusion data are listed in Table 1.

According to the AHA standard, the myocardium is divided into 17 segments based on a correspondence between those regions and the supplying coronary branch: ramus circumflex (RCX), left anterior descending (LAD) and right coronary artery (RCA). The perfusion parameters characterizing the CA distribution for each segment are derived from the corresponding averaged TIC. The latter is computed based on the TICs of all voxels in the respective segment. The perfusion parameters are visualized separately by polar coordinates in a color-coded Bull’s-Eye Plot (BEP) (Fig. 15 (right)).

For myocardial perfusion diagnosis, the parameters $PE$, $TTP$, $Up$-slope and $MTT$ have been evaluated as especially meaningful [60]. Based on the ratio of the $Up$-slope at rest and under stress, an additional perfusion parameter, the so-called myocardial perfusion reserve index (MPRI), is computed. The coronary perfusion reserve is defined as the ability of the coronary arteries to increase blood flow under stress by vessel dilation. The MPRI facilitates a more reliable detection of ischemic areas.

Fig. 15. Left: AHA conform acquisition of myocardial perfusion data in short-axis views. Schematic representation of the left ventricle which is imaged with 3 to 4 slices dissecting the left ventricle basally, centrically, apically and at the apex. Right: Bull’s-Eye Plot and associated AHA conform nomenclature. The plot is generated by projecting the myocardial segments onto a plane. The segments are colored according to the respective supplying coronary branch.
The detection and localization of a perfusion deficit as well as the assessment of the severity are directly relevant for treatment decisions. Major diagnostic tasks to be performed are: to evaluate whether the patient suffers from CHD, to evaluate the severity of the disease, and to assess the vascular supply of less perfused tissue. For an overview on MR-based diagnosis of the CHD, see Edelman [61].

The major preprocessing step for myocardial perfusion analysis is motion correction. A combination of rigid and elastic registration employing mutual information as the similarity measure works reasonably well for most datasets [62]. For parameter computation, the user examines the motion-corrected dataset and selects a ROI in healthy myocardial tissue. Utilizing these data, the parameters are calculated for each voxel of the dataset. Since the diagnostic questions mostly relate to the myocardium of the left ventricle, the visualization of perfusion parameters is restricted to this structure. The following description is based on the work by Oeltze et al. [14].

**Multiparameter visualizations for myocardial perfusion analysis.** Multiparameter visualization techniques (Section 5.1) are useful for evaluating myocardial perfusion data (see Fig. 16). The visualization of perfusion parameters by means of colored height fields should be restricted to the segmented myocardium (see Fig. 7).

**Refined Bull’s-Eye plot for rest/stress comparison.** In a rest/stress comparison, BEPs (or parameter maps) may be displayed side by side to identify areas where perfusion defects first appear or become worse with stress. In order to simplify a mental integration of rest and stress perfusion in one area, a refined BEP was introduced in [14]. Each segment ring is bisected, thus, duplicating the number of segments. The resulting outer and inner rings represent the stress and the rest state, respectively (Fig. 17). This circular bisection ensures that neighboring segments in the plot are adjacent in the myocardium as well as that they show the same state. Compared to the original variant [14], the segments are visually separated by a gap (grey ring) for better recognizability. The refined plot may also be used for comparing two different perfusion parameters of one state.

The diagnosis of CHD benefits from a link to morphologic image data, in particular MR/CT angiography data. Oeltze et al. [14] provided a link between the BEP and the 3D view of the coronaries and picking facilities for both, the plot and the 3D view. A segment of the plot exhibiting a suspicious parameter value may be selected by pointing—resulting in an animated emphasis of the corresponding vessel branch in the 3D view (Fig. 18) and vice versa emphasizing the supplied segments.

The refined BEP and the linking to morphologic data have been considered useful in informal discussions with a cardiologist from the medical faculty of the University of Magdeburg. He explained that supporting the simultaneous examination of rest and stress perfusion has been widely neglected so far. He suggested that the 3D view would benefit from mapping the perfusion analysis results to the ventricular surface hereby facilitating an exact assignment of the perfused territories to the supplying coronary branches. However, this requires the registration of the morphologic and the perfusion data.
9 GUIDELINES FOR VISUAL EXPLORATION

Over the years, we collected feedback from our clinical collaborators concerning the usefulness of the visual exploration techniques presented in sections 4-5. Based on this feedback, we propose guidelines for the use of these techniques. It turns out that some general guidelines are valid independent of the application area. Before we define the guidelines, we arrange the exploration techniques in three different classes whose separating criterion is the type of required input data:

C1 Input data: the original signal intensities (4D), techniques: cine-movies, subtraction images, and profile flags

C2 Input data: the perfusion parameters (3D), techniques: parameter maps, multiparameter visualizations and Bull’s-Eye plot

C3 Input data: the original signal intensities and perfusion parameters, technique: MIP + CVP

The second class C2 may be further subdivided into two subclasses: C2a (single parameter techniques: parameter maps) and C2b (multiparameter techniques: multiparameter visualizations). The original Bull’s-Eye Plot fits into C2a, whereas the refined version belongs to C2b. The MAMMOEXPLORER (recall Sec. 7) and the exemplary implementation of the analysis pipeline presented by Oeltze et al. [27] (recall Sec. 5.5) do not fit in either of these classes. Both rather represent a complex system comprising several techniques and integrate a sophisticated solution to link them.

Techniques of C1 are useful for an initial inspection of the original data. Tissue characterized by a very strong or by (almost) no enhancement can be roughly identified. Cine-movies require a mental tracking of signal intensity change over time. Their application in breast tumor diagnosis is limited by the low temporal resolution of the data. Profile flags show the signal intensity development plotted in a graph and embedded in a volume rendering, though only for a restricted region. Their application in cerebral and especially in myocardial perfusion is hampered due to the low slice number preventing a volume rendering. Subtraction images emphasize the highest differences in signal intensity between two time points. However, care must be taken to choose the “right” points. All of the before mentioned techniques require no time-consuming computation or complex user interaction.

After the initial inspection, techniques of C2 may be used for a more thorough analysis of the dynamic behavior in the suspicious tissue. Depending on the number of parameters that are interesting for the clinician, either techniques of class C2a or C2b should be used. The choice of a technique from C2b may be influenced by the number of parameters that can be simultaneously displayed, differences in the required computational effort and the complexity of the required interaction.

The class C3 contains techniques which exploit the original signal intensities as well as the perfusion parameters. The combined analysis of both has already been proven to be useful in a clinical trial for assessing breast tumor perfusion by Wiener et al. [45]. They could show that an analysis based on MIPs of subtraction volumes and parameter maps detected many occult cancers and thereby altered and allowed more confident treatment planning. Kohle et al. [49] proposed an integration of MIP and CVP for the combined analysis. However, this method is not applicable in cerebral and myocardial perfusion due to the low number of slices.

10 CONCLUDING REMARKS

This paper discussed general techniques for exploring perfusion data. However, many aspects of the exploration are relevant for other dynamic medical volume data. As an example, the analysis of dSPECT (dynamic Single Photon Emission Computed Tomography) data is also based on the selection of regions and the investigation of curves depicting changes over time in these regions [63]. The analysis of functional MRI data also
involves the analysis of time series [64]. The concept of application profile flags which enable an integrated view of the TICs and the underlying dynamic image data is also a general approach for exploring a variety of dynamic medical image data [22].

The basic principles of deriving, filtering, and analyzing TICs were developed for the analysis of scintigrams in the 70s and refined for the analysis of X-ray image sequences [65].

We described a variety of advanced visual analysis techniques which are motivated by discussions with medical doctors and observations of their work which revealed problems with simple visualization techniques. As an obvious drawback, we could not cite any substantial user study related to the superiority of the advanced visualization techniques with respect to specific diagnostic questions. To date, no such user studies exist. Therefore, we conclude that a systematic evaluation of techniques and their parameters as well as combinations is the most important task left open for future work. We also reviewed recent work on data analysis techniques, such as cluster analysis, feature detection and correlation analysis. Significantly more research is necessary to investigate whether these analysis techniques actually lead to a faster and more accurate interpretation of perfusion data.

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REFERENCES


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