Aging of population becomes an emerging health problem nowadays. One of symptoms of societal aging is an observed significant increase in the rate of dementia, which gradually excludes large group of patients from every aspect of life. Magnetic resonance imaging (MRI) is one of the most rapidly developing methods of neuroimaging. Modern MR techniques not only present structure and function of the brain, but also allow to assess metabolism, microcirculation and integrity if nervous tissue at the cellular level. The paper presents opportunities and perspectives of MRI in the diagnostics and differentiation of dementia.

Key words: magnetic resonance imaging, dementia, Alzheimer’s disease

Summary

Aging of population becomes an emerging health problem nowadays. One of symptoms of societal aging is an observed significant increase in the rate of dementia, which gradually excludes large group of patients from every aspect of life. Magnetic resonance imaging (MRI) is one of the most rapidly developing methods of neuroimaging. Modern MR techniques not only present structure and function of the brain, but also allow to assess metabolism, microcirculation and integrity if nervous tissue at the cellular level. The paper presents opportunities and perspectives of MRI in the diagnostics and differentiation of dementia.

Introduction

General improvement of living conditions and better health care systems resulted in a longer mean life span. Aging of the population is a significant social and medical problem of our times. It is estimated that in 1900, 1% of world population accounted for individuals over 65 years of age. In 1992 that percentage rate increased to 6.2% and according to prognoses, will reach as much as 20% in 2050 [1].

One of the symptoms of aging population is the witnessed significant increase in the incidence of dementia syndromes resulting in progressive exclusion of a large group of patients from every aspect of life. That is why the problem of advanced-age diseases is connected with increasing interest of researchers and also politicians (e.g. the document of the European Council from 2008, entitled Council Conclusions on public health strategies to combat neurodegenerative diseases associated with aging and in particular Alzheimer’s disease).
The main objectives of the research on degenerative syndromes of the elderly include searching for biomarkers, i.e. indicators or symptoms allowing early and reliable diagnosis of dementia. These may include aberrations in psychological tests, laboratory indicators or specific features of the brain that can be detected in imaging studies.

Searching for biomarkers of dementia focuses on such currently used imaging techniques as PET (positron emission tomography) and MRI (magnetic resonance imaging). The PET examinations, due to their limited accessibility, high price and exposition to ionizing radiation, seem to have less chance for a broad clinical use. Therefore, most examinations focus on the diagnostics of dementia with the use of MRI being a method that allows evaluation of brain morphology, chemical composition, ongoing metabolic processes and function. The aim of this work was to present possibilities and perspectives of the use of MRI in early diagnostics and differentiation of dementia syndromes.

DEMENTIA SYNDROMES

According to the definition of WHO, dementia syndrome is a set of symptoms caused by brain disease, usually a chronic or progressive one, clinically characterized by numerous disorders of higher cortical functions such as memory, thinking, orientation, reasoning, counting, ability to learn, language, and evaluation [1]. Moreover, disturbances of cognitive functions are frequently accompanied or even preceded by emotional, behavioral and motivational disorders. A more precise definition is given by DSM-IV (Diagnostic and Statistical Manual-American Psychiatric Association), which defines dementia as a set of symptoms of disturbed cognitive processes including, apart from disturbed memory, deficits of at least two functions from the following: speech (aphasia), voluntary complex movement (apraxia), ability to discriminate and identify objects (agnosis), as well as inability to plan, initiate, control and correct complex behaviors (disturbed executive functions) [2]. Moreover, cognitive deficits should be serious enough to disturb professional activity, social functioning and execution of everyday activities.

It seems that dementia is not a normal process of body aging. It can result from over 50 pathologies which cause primary or secondary disorders of the central nervous system (CNS). To systematize the background of the problem, we may distinguish 6 main etiological groups in the literature: vascular, degenerative, metabolic, infectious, toxic, and CNS injuries. The statistics and demographic data clearly show that the incidence rate of dementia is determined by age, i.e. the increasing rate of elderly individuals in the population is directly connected with increased incidence of dementia. Studies have shown that among individuals between 65 and 85 years of age the incidence is the highest and the main neurodegenerative disease is the Alzheimer’s disease, and among individuals over 85 y.o.a., the vascular dementias are the most prevalent [3, 4].

The most numerous group includes vascular and degenerative dementias. The remaining conditions, although sporadic, require insightful differential diagnostics, as they are potentially reversible. In Europe, the dominant form of dementia is the Alzheimer’s disease (AD), accounting for 50-60% of all cases. It is followed by DLB dementia (dementia with Lewy bodies) and VaD (vascular dementias) [3, 4].

The Alzheimer’s disease is a degenerative disease of the CNS, with progressive deficits of cognitive function and cortical disturbances – apraxia, aphasia, memory and behavior disorders, and psychotic symptoms. Early phase of AD reveals non-specific symptoms with dominant disturbances of memory. The disease is strictly age-determined. In individuals aged between 65 and 85 years, the incidence doubles every 5 years [4]. People with advanced AD are not able to function independently in their everyday life.

Dementia with Lewy bodies is characterized by cognitive disturbances with features of Parkinson’s disease of changing type with delusions and hallucinations. Typical for this disease is hypersensitivity reaction and poor response to neuroleptics. The accompanying symptoms include syncope, repeated falls and transient disturbances of consciousness [5, 6].

Vascular dementia reveals a picture typical for subcortical disturbances: memory disturbances, changes of emotions and personality, executive function disturbances and slower information processing. Usually, these symptoms result from numerous minor strokes. Risk factors of VaD include mainly age, diabetes, atherosclerotic lesions of internal carotid, vertebral and cerebral arteries, hypercholesterolemia and stroke in past [7].
Diagnostics of dementia underscores 4 basic clinical features of the disease: the profile of cognitive disturbances, the onset and time sequences of appearing clinical symptoms, psychopathological disorders (psychotic syndromes and disturbed behaviour) and neurological symptoms [8]. These components are included in detailed criteria differentiating VaD from degenerative diseases of the brain. A thorough analysis of criteria of dementia supported by advanced MR imaging facilitates the proper diagnosis.

DEMENTIA SYNDROMES IN IMAGING STUDIES – CURRENT IMAGING PRACTICES

The main symptom of brain degenerative diseases includes disturbed cognitive functions. It is estimated that in half of the patients suffering from dementia, the disease was not diagnosed and according to NDS (National Dementia Strategy), dementia is diagnosed too late and too rare. Development of neuroimaging techniques facilitates clinical diagnosis of dementia, mainly by excluding the organic causes of brain disorders. Imaging methods used in the diagnostics of dementia may be divided into structural (CT, morphological MRI with volumetry, Doppler ultrasonography) and functional: evaluation of diffusion, diffusion tensor in MRI, evaluation of perfusion with MRI and CT, functional magnetic resonance (fMRI), as well as SPECT, i.e. single-photon emission computed tomography, and PET.

Owing to the availability of CT, this is frequently the first examination performed in a patient with dementia or suspected dementia. However, it should be remembered that its diagnostic capabilities are quite limited and the examination detects only some pathological lesions typical for dementia. In CT it is easy to evaluate brain atrophy (cortical, subcortical or mixed). Their location may suggest the type of dementia, e.g. AD-related atrophy of hippocampal area mostly, FTD-related atrophy of frontal and temporal lobes, subcortical atrophy in Huntington’s disease and atrophy of frontal lobes and cerebellum in alcoholic dementia [9]. The evaluation of the degree of atrophy should be evaluated regarding the age of the patient. Moreover, an important advantage of atrophy evaluations is the possibility of a precise control of dementia changes in time [9].

Over the past years, morphological imaging of dementia with MRI has gained on importance. This tendency results from a higher contrast resolution of MRI as compared to CT, which enables precise evaluation of particular structures of the brain, e.g. hippocampus, typically undergoing atrophy in AD, or the gray matter of the hemispheres selectively. Increased interest in dementia imaging with this method was a result of studies carried out by Award et al. [9, 10]. They showed that in MRI images of the studied patients, the incidence of lesions in the white matter reaches 22% in a group below 40 years of age, 51% among individuals aged 41-60, and 92% among individuals over 60 y.o.a. To compare with, CT detects focal lesions in 7% of individuals aged 41-60, and in less than 1% of individuals under the age of 45 [10]. The advantage of MRI over CT is also visible in the evaluation of ischemic lesions. This follows from a significantly higher sensitivity of T2-weighted images in detecting small ischemic lesions within basal ganglia, white matter, brain stem, and cerebellum. Specific MRI image allows also for diagnosing rare neurodegenerative diseases, such as Hallevorden-Spatz syndrome, Wilson or Creutzfeldt-Jakob disease [9].

ADVANCED MRI TECHNIQUES IN DEMENTIA IMAGING

Morphometry

VBM morphometry is a measurement of volume of the grey matter, comparing signal intensity within particular pixels of an image of a selected anatomical structure of the brain. The measurement of the volume of tissue structures or fluid spaces is carried out at sites strategic for a given type of dementia. On the basis of information thus received it is possible to evaluate the advancement of the disease as well as clinical prognosis of disease progression [9, 11].

However, it should be remembered that due to inter-individual variability of the size of the skull as well as insufficient specificity, the value of single linear measurements in the diagnostics of CT imaging may raise doubt. MRI imaging is characterized by high accuracy in volume evaluation – up to 0.1 cm³. Such a
precision in measurements guarantees high
repeatability and sensitivity of the method [10]. In case
of high-resolution MRI imaging, software packages
required for data processing generate planimetric and
volumetric measurements. Planimetric measurements
are obtained with area segmentation technique. This
method uses the difference in signal intensity between
the brain cortex, subcortical structures and the white
matter [11]. Values obtained in such a way are
normalized with regard to so called total intracranial
volume [11]. The evaluation is made with the use of
semi-automatic technique.

In order to obtain images, the IR (inversion
recovery) sequences are used or typical T2-weighted
images, in FSE (fast spin echo) sequence. Examinations with the use of VBM showed that in AD
there is not only a typical atrophy of structures of the
superior-medial part of the temporal lobes, but also
atrophy within the insular cortex and caudate nuclei
[11, 12].

Jack et al. [13] in their longitudinal studies showed
that in the group of study patients with diagnosed AD
and healthy, elderly individuals, a higher index of
hippocampal atrophy was found in patients with AD
(4-6% a year). A similar interrelation was also shown
for the entorhinal cortex [13]. This finding
acknowledge volumetric measurements of the
entorhinal cortex as dedicated indicator of clinical
deterioration in patients with AD [14]. For evaluation
of progression of brain atrophy, it is recommended to
use a method which allows for “superimposition” of
subsequent images and subtraction of the baseline
image. As a result, the acquired image will include
marked in color (usually red) areas of brain atrophy

Magnetic Resonance Spectroscopy (MRS) in the
literature called „window for tissue metabolism”. In
MRS it is possible to non-invasively assess the amount
of selected chemical compounds in tissues and
indirectly draw conclusions on metabolism or cellular
integrity of the examined area [15]. The spectroscopy
uses signals of chemical compounds including nuclides
not only 1H but also 13C, 19F, 31P, 15N and 23Na. The
result of the test is the resonance spectrum of the tested
sample, shown in coordinates. The spectrum results
from converting signals of free precession FID
registered within a selected area.

The most frequently used method of localization is
the SVS (single voxel spectroscopy). It allows to
ascribe a different resonance frequency to every
examined area. Undoubtedly, the advantage of this
method is high homogeneity of the study area, short
examination time (4-8 min), high signal-to-noise ratio
and easiness in selective attenuation of water signal.
Other localization methods include ISIS (Image-
selected in vivo spectroscopy), PRESS (point-resolved
localized spectroscopy) and STEAM (Stimulated echo
acquisition mode). Differences between them involve
changing RF impulse and coordinates of time
sequences [10]. We should also mention the
MultiVoxel technique. It allows for chemical shift
imaging (CSI) and allows for simultaneous registration
of signal from a few adjacent voxels. The acquired
image is a map of selected metabolites with their
spatial position in the examined layer of the organ
[10, 16].

MRS has long been applied in studies on dementias
[9]. Proton spectroscopy is the most frequently used
one (hydrogen, 1HMRS). More rarely used, mainly
due to equipment requirements, are phosphorus
spectroscopy (31P-MRS), fluorine spectroscopy
(19F-RS) and carbon spectroscopy (13C-MRS). The
1HMRS allows for testing the following brain
metabolites: NAA (N-acetylaspartate), choline and its
metabolites, myoinositol, creatinine and creatinine
phosphorane, as well as the glutamate-glutamine
complex. NAA in immature oligodendrocytes is found
mostly intracellularly. Thus, it is held for a marker of
neuronal density and as an indicator of metabolic
activity of nervous cells [15, 17] and NAA decrease is
connected with neuronal atrophy [18]. In a normal
spectroscopic image of the brain, it shows the highest
peak.

MRS creates new possibilities of diagnostics of AD
in living individuals. A typical picture of AD is as
follows: lower NAA concentration, higher myoinositol
concentration [9, 10]. Intensification of lesions is
especially significant in associative centers. In a test
differentiating healthy middle-aged individuals and
patients with AD, MRS showed sensitivity of 83% and
specificity of 98%. In a test differentiating AD and
other forms of dementia, MRS showed specificity of
64% and sensitivity of 82% [10].

In case of dementia with Lewy bodies, diagnostics
with spectroscopy has been incidental so far. The
reason was technical problems which can appear
during the examination (parkinsonism, no cooperation
of the patient and fluctuations of cognitive functions).
So far, the first and the largest project aimed at
spectroscopic examination in DLB was the study by
Molina et al. [18]. It included 12 patients meeting the criteria for DLB. All included individuals had fluctuations of cognitive functions, visual hallucinations, and parkinsonism. The authors tried to evaluate brain metabolites in the temporal lobes and basal ganglia but the examination of the mentioned areas ended in failure due to no homogeneity of the brain in those areas. Significant differences in the concentration of metabolites between DLB and the control group were found only in the white matter.

**Diffusion tensor imaging (DTI)** is a technique of MRI used to visualize the course and to evaluate the integrity of the pathways of the white matter. DTI is an advanced diffusion technique which, apart from quantitative evaluation of free diffusion of water in the extracellular space of living tissues, expressed as MD (mean diffusivity) allows also for qualitative and quantitative evaluation of the direction of diffusion [17].

A natural barrier for free diffusion in the tissues of the cerebral and spinal white matter is the cell membrane and myelin sheath of axons. This is where the movement of water particles is limited and anisotropic – it is expressed by limited diffusion in perpendicular directions with regard to the white matter fibers. The degree of diffusion anisotropy is shown by FA (fractional anisotropy) parameter. FA is also called an index of white matter integrity [11]. A decrease in its value signals damage and degradation of white matter pathways or widening of the extracellular space in the course of vasogenic edema [19]. An additional option of DTI is tractography of diffusion tensor. This method visualizes in 3D the anisotropy of water diffusion in the white matter, which corresponds to the course of white matter fibers. Tractography uses an algorithm of drawing the course of continuity of fibers from a chosen point. It should be remembered that the registered course of fibers is averaged for every voxel [11].

DTI is widely used in the evaluation of dementia disorders [9]. In dementia there is a secondary degeneration of the fibers of the white matter, probably due to Waller degeneration, which concerns the associative pathways (longitudinal upper and lower bundles, fronto-occipital bundles, and posterior part of the cingulum), callosal pathways (genu and splenium corporis callosi) [9]. Sasiadek et al. showed the highest lowering of FA within the posterior cingulum, with a strong difference between AD and mild cognitive impairment (MCI), and between MCI and healthy population. DTI performed in the area of the posterior cingulum was the best in differentiating those groups of patients, showing superiority over MRS and PWI (perfusion weighted imaging), or even PET [9].

**Functional MRI** is a method presenting activation of the brain cortex in response to specific stimuli or at rest. The intensity of MRI signal is related to the degree of hemoglobin oxidation. Activation of neurons in response to a stimulus results in a decrease in oxyhemoglobin level in local microcirculation and increase of CO2 and deoxyhemoglobin level. As a result of autoregulation, the local blood flow rises after 2-6 sec, which causes an increase in the amount of oxyhemoglobin and washout of the deoxygenated form. Both forms of hemoglobin present different magnetic properties, which causes a change in MR signal intensity of 1-5% in the BOLD technique [20].

fMRI has a high linear and temporal resolution. It is carried out in two phases. The first one is the morphological (structural) examination. Next, the areas of the brain cortex are subjected to a series of active or passive (stimulus removal) stimulation separated by periods of rest [11]. After comparing images of activation obtained in control situation of rest and task, a map of brain activity is obtained, correlated with the influence of a selected component [21].

The fMRI examination in case of cognitive disorders gains on its importance. An essential role is played by the possibility of comprehensive evaluation of structural changes and brain function [11].

Scientific research with the use of fMRI are usually conducted on good-cooperating volunteers. The results of neuropsychological or movement tests are much better in those individuals than in patients. Patients with dementia have their cognitive functions tested. Therefore, when preparing the paradigms – tasks – it is necessary not only to include neuropsychologist’s help but also to simplify stimuli adequately [11]. Cognitive testing of dementia is based on evaluation of working memory (remembering pictures, associating face with name), arithmetic functions (counting forward and backward, addition and subtraction), visual-spatial functions (comparing objects seen from different sides, identifying the angle between the arms of the clock) [10].

In AD there is a decreased activity of the cortex during performance of cognitive tasks, especially in parietal lobes and hippocampus [9]. It was also shown that specific changes in fMRI appear even before the occurrence of clinical features typical for AD:
increased activity of the praecuneus and decreased activity within the middle part of temporal lobes and prefrontal cortex [10, 11]. One should remember that the area of increased or decreased activity would point to disturbances of interaction between certain centers and not to their damage [11].

This method allows for an assessment of the influence of procognitive medications on memory functions during long-term therapy. Increased activity was found in the areas of the cortex stimulated during memorization [21]. fMRI may also be helpful in differentiating AD from other pathologies associated with dementia, e.g. the Parkinson’s disease, vascular dementia or fronto-temporal dementia [22]. In case of AD, it is extremely important to select patients at an early stage of the disease. AD is frequently preceded by the MCI. Distinguishing MCI patients with increased risk of development the AD is clinically challenging and fMRI seems to facilitate this.

**Perfusion-weighted imaging (PWI)** permits evaluation of brain microcirculation on the basis of changes in MR signal intensity. Depending on the technique applied, PWI allows for quantitative and qualitative evaluation (color maps, regional blood flow (CBF), volume of blood in the vascular bed (CBV), and mean time of blood flow through microcirculation (MTT)). Examination of perfusion is used in detecting and differentiating the type of dementia and in monitoring of its progression.

PWI can be performed with or without administration of contrast media. Perfusion examination without the use of gadolinium preparations is carried out with the ASL (Arterial Spin Labelling) method, i.e. marking of blood spins. This technique includes saturation or inversion of arterial blood spins at the level of the neck, and then their detection in the measured volume. The marked spins, moving to the vascular bed of the brain, change magnetization of brain tissues, which is registered as dynamic change in signal intensity [9, 10, 17]. The second method requires i.v. injection of paramagnetic contrast agents and is based on the evaluation of a simple change in relaxation time T1 or uses the effect of magnetic susceptibility of the bolus, and it influences the relaxation time T2 (DSC). DSC is a more precise method but requires repeated acquisition of data during the first pass of the bolus, and thus poses high requirements for the equipment [10].

MR perfusion is valuable method in differentiating VaD and AD [11]. It can be also used in monitoring the progression of dementia. Walecki et al. showed that there is a correlation between CBF and CBV proportional to the MMSE score [14]. Decreased perfusion in cortex indicate dementia [9]. Location of abnormalities seen in PWI in different forms of dementia correlates well with SPECT results [9]. According to Sąsiadek et al., the most intensified perfusion abnormalities in dementia can be found within the cingulate and this may be the cue for differentiating AD patient from MCI patient, and MCI patient from a healthy person [9].

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Diagnostic opportunities of new techniques of magnetic resonance imaging in detecting and differentiating dementia syndromes

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