Advantages and Challenges of Small Animal Magnetic Resonance Imaging as a Translational Tool

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Abstract
The utilization of magnetic resonance imaging (MRI) methods in rodent models of psychiatric disorders provides considerable benefits for the identification of disease-associated brain circuits and metabolic changes. In this review, we discuss advantages and challenges of animal MRI and provide an overview of the major structural (voxel-based morphometry and diffusion tensor imaging) and functional approaches [resting-state functional MRI (rs-fMRI), MR spectroscopy (MRS), regional cerebral blood volume measurement and arterial spin labelling] that are applied in animal MRI research. The review mainly focuses on rs-fMRI and MRS. Finally, we take a look at some recent developments and refinements in the field.

Animal Magnetic Resonance Imaging in Psychiatric Research

Introduction
Preclinical animal models make valuable contributions to improving our understanding of human diseases, ranging from insights into the molecular and cellular underpinnings of a disorder to the development of novel pharmacotherapeutic treatment approaches. One of the main advantages of animal studies is group homogeneity, which cannot be easily achieved in clinical (human) studies. Yet, establishing valid animal models of psychiatric disorders has proven to be an undertaking of substantial proportions [1, 2]. The pronounced heterogeneity of mental disorders and a strong reliance on phenomenology have impeded any precise evaluation of the validity of respective animal models [3]. One way to confront this problem has been to reverse-translationally adapt the concept of endophenotypes [4, 5] to preclinical research.

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thereby emphasizing the dissection of functional circuits related to animal behaviour [2]. In this regard, neuroimaging is a highly valuable method in animal research; conceptually, it is informative with respect to the investigation of experimental manipulations shaping functional circuits and behaviour. Methodologically, the application of analogous imaging technology to animals and humans positively affects the translational potential of animal research (fig. 1).

Moreover, the last decade has seen the critical appraisal of a combined impact of genetic and environmental factors upon disease susceptibility [6]. Due to its complexity, the investigation of gene-by-environment-by-treatment interactions calls for an intermediate level of analysis. Since risk genes as well as environmental and pharmacological manipulations specifically influence brain functioning, high-resolution animal imaging provides a highly suitable technique to investigate these aspects in isolation or in their interaction. Finally, functional magnetic resonance imaging (fMRI) drives the progress of in silico technologies as are used in the Blue Brain Project (http://www.humanbrainproject.eu/), the Human Brain Project (http://www.humanbrainproject.eu/) or the Bernstein Centres for Computational Neuroscience.

Advantages of Non-invasive Imaging Research: Repeated Measurements and Multimodality

Neuroimaging provides an excellent way to investigate structural and functional aspects of disease-associated pathology in animal models and clinical populations alike. Apart from this methodological advantage, non-invasive imaging as a research tool implements Russell and Burch’s [7] classic ‘3R’ principles of ‘reduction’, ‘refinement’ and ‘replacement’ for the humane treatment of research animals. Reduction can be achieved through smaller group sizes (reduced variance due to intra- and interindividual measurements). In addition, due to the possibility of performing repeated intraindividual measurements, the number of experimental groups can be reduced. High-resolution morphological and functional analyses as well as the use of methods causing as little distress as possible contribute to refinement.

An important feature of non-invasive procedures concerns the possibility of performing multimodal investigations on one and the same animal, allowing for a simultaneous or step-wise analysis of structure and function. In theory, a specific brain region may be investigated by voxel-based morphometry (VBM), resting-state fMRI (rs-fMRI), arterial spin labelling (ASL), diffusion tensor imaging (DTI) and MR spectroscopy (MRS). Impressive studies combining spectroscopy and fMRI have recently shown the great potential of a multimodal investigational approach to imaging research [8].

Challenges and Limitations

One of the biggest challenges of animal MRI concerns its resolution: while the human brain has an average volume of about 1,450 ml, that of a mouse is only 0.5–0.6 ml.
Accordingly, while resolutions of about 1 mm suffice for most human brain imaging needs, the investigation of a rodent brain requires voxel sizes to be 0.1% of those required for human imaging. For MRS, in humans, the typical hippocampus voxel would contain approximately 3,000–4,000 μl, compared with 3–4 μl in mice and 12–15 μl in rats. Since the signal-to-noise ratio (SNR) is directly proportional to the voxel size, this difference should be balanced in order to produce high-quality images.

Limitations

The SNR of a typical imaging application (e.g. MRS of the hippocampus) in a 3-Tesla human scanner exceeds that of a comparable animal measurement in a 9.4-Tesla scanner by a factor of 5, which can only be balanced by repeated signal acquisition, thus necessitating longer durations of measurements; doubling the SNR, for instance, requires quadrupling the time of measurement.

Physiological noise is a much greater problem in animal imaging as well. Basically, the amount of movement a small animal induces through breathing (even under anaesthesia) is higher in relation to the voxel size in humans. Also, at a higher signal strength, physiological noise (breathing and cardiac pulse) fraction increases [9]. Due to the higher field power and resolution of animal scanners, physiological parameters may cause relevant disturbances. At best, they contribute noise to the acquired data; at worst, they cause strong artefacts, thereby seriously impairing image quality. Very sensitive DTI sequences in fact may be triggered by breathing in order to avoid or minimize artefacts.

For fMRI, a continuous recording of physiological activity is required at a sufficient temporal resolution of about 100 Hz sampling frequency. Since fMRI, especially in animals, always undersamples physiological noise, the frequencies of respiratory and cardiac noise may appear anywhere in the frequency band of interest due to aliasing effects. Using physiological curves recorded with high temporal resolution allows retrospective methods such as RETROICOR [10] to assess the influence of these distortions on the fMRI time series and to remove them by using regression methods.

For most applications of animal MRI, anaesthesia is a necessity. Even though it places restrictions on fMRI, experiments with awake animals also have their limitations [11]. For example, even after habituation to the apparatus, animals are still stressed by the fixation, which may subsequently have an impact on the investigated brain functions. So far, there has been a relative paucity of studies looking at the effects of different types and depths of anaesthesia on cortical and subcortical functioning. Commonly, anaesthesia is achieved through the inhalative application of isoflurane, which is advantageous in different respects: it is usually well tolerated, and animals breathe spontaneously and do not require endotracheal intubation (fig. 2). Moreover, anaesthesia can be easily controlled by adjusting the isoflurane concentration. Rats usually receive 1.5–2% isoflurane, 30–50% oxygen and 50–70% air or N2O; mice require slightly lower doses of isoflurane. The concentration may initially be higher and drop during the course of a longer experiment. However, isoflurane has major disadvantages for rs-fMRI studies, which are discussed in detail in the section titled 'Resting-State fMRI'.

Field Strength

Animal imaging systems often possess higher field powers and stronger gradient systems than human scanners. Higher-intensity fields are more favourable regarding the SNR (roughly linear) and allow for a higher spectral resolution in MRS. A 9.4-Tesla system increases the SNR by about a factor of 3, which is actually countered in part by a disadvantageous change in relaxation parameters for imaging. The advantages of the higher field strength become really important for MRS, where not only the gain in SNR but also the higher spectral resolution improve the data. This means that overlapping resonance lines move further apart on the frequency axis and are easier to quantify. For imaging applications, this higher chemical shift is rather a disadvantage, since it could produce artefacts in the data (e.g. lipid artefacts in fast imaging sequences).
Apart from the changes in relaxation parameters, the most important disadvantages of a higher field strength are a higher magnetic field inhomogeneity and an increased specific energy absorption rate. High-field-strength scanners are more prone to the influence of disruptive artefact-generating factors. Due to physical limitations, homogeneous static fields (B0) or radio frequency fields of the high-frequency coils (B1) are increasingly difficult to provide with field intensities of 7 T and above. However, a homogeneous B0 field is required for high-quality spectra and artefact-free images, and B1 inhomogeneities impede the quantification of MRS and imaging data. In order to balance B0 inhomogeneity, shim coils are applied. Field distortions depend on the measured object and require repeated field adjustments, which often do not yield the desired effect. MRS and fast imaging sequences such as echo planar imaging (EPI) used in fMRI and DTI are particularly sensitive in this regard. The increased specific energy absorption rate means that for the same application more energy will be absorbed by the tissue, which is a problem in human rather than in animal imaging.

**MR Coils**

The maximum increase in SNR in animal imaging is achieved by dedicated transmit-and-receive coils. Small radio frequency coils placed in close proximity to the animal can highly improve the quality of the acquired data. Recently, manufacturers introduced cryogenic coils cooled to low temperatures around 20–30 K. This greatly reduces the noise generated by the coil itself and the pre-amplifier electronics and can thus improve the SNR by another factor of about 3 for protons and by much higher factors for other nuclei. In the case of larger animals and humans, the benefit from a cooled coil would be much smaller, since most of the noise would come from the much larger object of interest.
The emergence of high-resolution scanning technology and sophisticated image processing has significantly advanced structural analysis of the rodent brain. Endophenotype mapping as well as the assessment of the structural effects of ageing and environmental or pharmacological manipulations would considerably benefit from improved structural data.

Morphometric analysis can generally be carried out in two different ways. Regions of interest (ROIs) can be selected, which requires an a priori hypothesis or knowledge about structures potentially affected by an experimental factor. Methods like VBM or DTI, on the other hand, are based on semi-automatic procedures of tissue separation.

Voxel-Based Morphometry

In VBM, different anatomical attributes can be analysed and compared based on tissue contrast mapping due to MRI intensity differences. VBM is an unbiased method allowing a fine-grained structural analysis — which, however, requires extensive processing of data. High-resolution $T_1$- or $T_2$-weighted 3D datasets are partitioned into tissue classes (grey matter, white matter and cerebrospinal fluid) by segmentation algorithms. In this process, images are matched onto a common template with non-linear spatial normalization procedures, and the tissue probability maps are modulated with the local non-linear volume change. The segmented and normalized images containing a semi-quantitative measurement of the local tissue concentrations are then smoothed so that each voxel has the average of grey matter from itself and its surroundings. Smoothing is done to reduce the impact of residual misregistration between subjects/images and to lower the possibility of non-gaussian data.

Voxel-wise statistical testing is performed using generalized linear models, whereby local structural differences (in volume or concentration) can be identified. Figure 3 depicts the steps required for the creation of a group template for segmentation and subsequent individual segmentation and statistics.

The output of VBM is statistical map displaying regions with significant differences in volume or concentration of the investigated tissue (fig. 4). VBM is mainly a qualitative assessment of local differences in volume between groups. A semi-quantitative absolute volume measurement of a tissue class in a region can be conducted by summation of the signal intensities in the normalized segmented images. Each voxel has a probability of containing one of the three tissue classes multiplied by the Jacobian determinant of the vector field used to non-linearly normalize each individual dataset to the group template. This is a factor greater or smaller than 1 if the local volume is compressed or inflated during the normalization, respectively. The sum of all voxels over a region of the anatomical atlas in a normalized grey matter image thus approximates the total amount of grey matter in this region. However, depending on the normalization parameters, there could still be residual misregistrations in the final images; hence it has to be taken into account that there is a possible inaccuracy of quantitative volume assessment by VBM.

Diffusion Tensor Imaging

DTI is a technique for visualizing the (micro-)structural brain organization. Movement of water molecules is more rapid in the direction of fibre tracts and slower in the perpendicular plane; the measurement of this difference (diffusion anisotropy) can be used to make inferences about the presence, position and integrity of white matter tracts.
The diffusion tensor provides information about restricted and directed movements of water molecules with high spatial resolution. It is calculated from a comparison of MR signals acquired with diffusion gradients along at least 6 non-collinear, non-coplanar directions with MR signals without diffusion gradients. In practice, not only 6 but 32 or more diffusion measurements are taken in order to calculate a tensor of sufficient quality. This makes the technique time-consuming and prone to movement artefacts.

Essentially, two different processing methods are used for DTI. The simpler one is a calculation of the fractional anisotropy (FA) coefficient in every voxel, which states how directed the movement of water is in the current position compared with freely moving water. Since movement is restricted to one direction along white matter fibres, the resulting FA maps are particularly useful to measure changes in white matter. Therefore, DTI is often referred to as a white matter imaging method.

In the second method, the information from the calculated diffusion tensor is used to reconstruct white matter pathways in vivo. Tractography algorithms use this information to track the neural pathways based on the assumption that the dominant direction of water motion aligns with the fibre orientation in an imaged voxel. While this method produces striking visual presentations on fibre tracts, the results are not necessarily always accurate, since most algorithms are based on probability functions and are easily thrown off course by the presence of noise or artefacts in a single voxel along the tract. Also in voxels with fibre crossing the calculation of the diffusion tensor would represent a mean of the movement along all fibres present in the given voxel, thus impeding an estimation of the dominant direction in the given voxel by tractography algorithm.

Since the measurement of 32 or more diffusion directions is time-consuming, fast imaging sequences such as EPI are used for DTI. However, they are prone to geometrical artefacts and are highly sensitive to movement. Hence, most mouse brain DTI reports use data from ex vivo analysis of the brain, with only few attempts to perform in vivo DTI [13]. The movement problem can often be successfully solved by triggering the sequence to the animal’s breathing cycle, which prolongs the time of measurement of one DTI dataset to about 1.5 h. Figure 5 displays an example of data from a murine DTI experiment.

**Translational Functional Methods**

**Arterial Spin Labelling**

The translational application of functional imaging methods requires acknowledgement of the fact that most classic sensory stimulation paradigms cannot be used because of the need for animals to be anaesthetized. The blood oxygen level-dependent (BOLD) technique has long been the closest approximation to a functional analysis. Though well established and possessing a high degree of sensitivity, the BOLD effect is complex, and despite its dependence on cerebral blood flow, it does not give a direct measure of it. The direct and absolute quantification of cerebral blood flow requires a diffusible tracer. The most effective approach, called ASL, uses arterial blood water that is magnetically (spin) labelled by radio frequency irradiation. The change in tissue magnetization due to the labelled arterial blood is proportional to cerebral blood flow. In general, ASL has a rather poor SNR but does not require the application of contrast agents and is directly translational. It allows the assessment of task-independent measures of brain activation via the measurement of resting-state cerebral blood flow. Several translational ASL techniques exist which rapidly measure blood flow with improved SNR and high retest reliability [14], one of them being continuous ASL [15–17].
Regional Cerebral Blood Volume

Regional cerebral blood volume (rCBV) measurements make use of the fact that the acute passage of a bolus of non-diffusible contrast agent through the brain causes a change in signal intensity that correlates with the regional blood volume. The mechanisms of contrast agents rely on their changes in local longitudinal relaxation time ($T_1$) and/or transverse relaxation time ($T_2$).

Several kinds of contrast agent are generally available for rCBV measurement in the rodent brain. On the one hand, there are gadolinium-based contrast agents with particle sizes smaller than 1 nm, and on the other hand, several kinds of superparamagnetic iron oxide (SPIO) nanoparticles, which are based on magnetite ($\text{Fe}_3\text{O}_4$) or maghemite ($\gamma\text{-Fe}_2\text{O}_3$) and have particle sizes between 10 and 180 nm. The large particle size originates from the coating of the paramagnetic molecules with polysaccharide or synthetic polymers or monomers. The smaller SPIO nanoparticles with a particle size below 50 nm – also called ultrasmall SPIO or USPIO particles – have a longer half-life in the blood after intravenous injection and are thus better suited for rCBV measurements than are larger particles. All contrast agents usually prolong the longitudinal $T_1$ relaxation constant and shorten the transverse $T_2/T_2^*$ relaxation times. USPIO particles have a stronger difference in these changes with an emphasis on shorter transverse relaxation times, and they have an excellent relaxivity as well as stronger effects than gadolinium-based contrast agents on a per-atom basis. Quantification of the different effects produced by a contrast agent on the signal is complex and depends on vasculature size, field strength and pulse sequence; the $T_2$ contrast from spin echo sequences, for instance, is sensitive to small vessels, while changes in $T_2^*$ which can be detected by gradient echo-based sequences are attributed more to larger vessels. When a contrast agent reaches a uniform distribution, relative CBV maps can be measured from steady-state $T_2$-weighted images with the dependence $\text{CBV} \propto \Delta R_2 = \ln(S_{\text{pre}}/S_{\text{post}})/TE$, with the relaxivity $R_2 = 1/T_2$, and $S_{\text{pre}}$ and $S_{\text{post}}$ as the signal intensities before and after contrast agent application.

While USPIO particles have a comparably strong effect on relaxation time and a long circulating half-life, they are more costly than gadolinium-based contrast agents and can lead to organ toxicity, which makes them not ideally suited for longitudinal studies. Gadolinium-based agents, on the other hand, can repeatedly be applied intraperitoneally and have minimal toxicity [18]. They typically reach a steady state about 30–60 min after intraperitoneal injection, which normally lasts for approximately 1 h. The kinetics depends on the nature of the contrast medium and its way of application. Regional blood volume measurements are well suited for pre-post comparisons in the context of pharmacological interventions [19]. Voxel-wise calculation of differences in rCBV intensity between the presence and absence of a pharmacological challenge allows a direct and non-invasive calculation of rCBV maps.

Different types of sequences can be used for quantifying changes in signal intensity after the application of a contrast agent. Figure 6 displays EPI, fast low-angle shot and rapid acquisition with relaxation enhancement sequences, for instance, is sensitive to small vessels, while changes in $T_2^*$ which can be detected by gradient echo-based sequences are attributed more to larger vessels. When a contrast agent reaches a uniform distribution, relative CBV maps can be measured from steady-state $T_2$-weighted images with the dependence $\text{CBV} \propto \Delta R_2 = \ln(S_{\text{pre}}/S_{\text{post}})/TE$, with the relaxivity $R_2 = 1/T_2$, and $S_{\text{pre}}$ and $S_{\text{post}}$ as the signal intensities before and after contrast agent application.

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Initial studies have been able to impressively demonstrate the potential of the rCBV method to boost high-quality translational research: Small’s group [20], for example, showed that baseline abnormalities in rCBV in the CA1 region of the hippocampus of prodromal patients were predictive of the development of psychosis. In the same study, rCBV mapping was performed on mice receiving antipsychotics, and the antipsychotic medication could be excluded as a potential confounder of the study results [20].

Resting-State fMRI

Although the first report on human brain functional connectivity in the absence of any stimulus or task was already published as far back as 1995 [21], only recently have resting states also been described for the rodent brain. One of the main reasons why it took longer to establish them in animals is the requirement for anaesthesia and physical restraint in the scanner. However, low-frequency (<0.1 Hz), temporally synchronized BOLD oscillations reflecting functionally connected brain networks analogous to humans were demonstrated in anaesthetized rodents [22–28]. The observed networks relate to sensorimotor, hippocampal, prefrontal, retrosplenial, visual, auditory, caudate-putamen and thalamic networks. It has been suggested that anaesthetized rat brains also possess a default mode network that is comparable to that of humans and includes hippocampal, prefrontal, cingulate, posterior parietal and retrosplenial cortices [29]. Therefore, these coherent BOLD fluctuations may represent a principal intrinsic property of a functioning brain and can be used as a valuable preclinical tool in animal models of disease to investigate diagnostic markers in the form of changes in functional connectivity. In addition, the effect of pharmacological manipulations of functional connectivity can be studied preclinically (fig. 7) [30].

Resting-state networks have also been reported in non-anaesthetized rats [31–35]. Before conducting experiments, the animals are habituated to the restrained position; typically, they are lightly anaesthetized and placed into a mock scanner with scanner noises simulating the future experimental set-up. The number of acclimation sessions has varied in different studies (from 2–3 days [31, 34] to 8 days [32, 35]). However, these studies are still few in number due to their main limitation, i.e. the stress animals have to undergo in the restrained position in the scanner and its consequential effect on brain functional connectivity patterns, and also due to their need for several habituation sessions. Interestingly, a recent study has shown that although an anaesthetized rat brain (under isoflurane) displays a weaker connectivity, it still preserves global network properties of the awake brain (in terms of graph theory: global clustering coefficient, mean shortest path length, small-worldness and modularity), and long-distance connections are not preferentially reduced [33]. A recent study on anaesthetized human brains (under propofol) exploring functional connectivity in terms of graph theory has also shown that long-range and short connections are not differentially affected by anaesthesia, and the small-world architecture of the brain networks persists (is in fact increased); clustering and modularity are elevated, possibly indicating an increase in localized processing and lower integration [36].
The usual choice of anaesthetic has been isoflurane, because narcosis can be easily adjusted and kept stable for hours. However, the α₂-adrenoreceptor agonist medetomidine was demonstrated to provide functional connectivity maps more comparable to what is seen in humans [37]: under medetomidine, more localized correlations were observed, in contrast to the widely spread connectivity across the cortex seen under isoflurane – which is possibly due to the latter’s interference with neural activity, which medetomidine does not affect [38]. Medetomidine allows longitudinal experiments and does not require invasive procedures, in contrast to α-chloralose, which necessitates intubation and mechanical ventilation of animals. Functional connectivity maps from rats anaesthetized with α-chloralose are more similar to the ones observed under medetomidine than those seen under isoflurane; however, they exhibit localization to a lesser extent [37]. Therefore, medetomidine is currently the preferred choice [23, 26, 28, 39–41].

Once an animal is anaesthetized and placed into the scanner, it undergoes a typical rs-fMRI experiment similar to the one on humans. Preferably, fast EPI sequences acquire data during a period of approximately 10 min with a repetition time of 1–3 s. Since geometrical artefacts and the influence of physiological noise are bigger at a higher field strength, the B0 field maps and physiological signals (respiration and cardiac pulse) are sampled during the experiment. The cardiorespiratory cycle and variations in respiratory volume/rate may inflate ROI-ROI correlations and bury the real ones. Cardiac noise mostly affects brain regions bordering blood vessels, whereas respiratory motion can be detected in areas near the edges of the brain or near the ventricular system due to shifts in B0 from abdominal movement. Since both cardiac and respiratory motion usually occur at frequencies higher than the fMRI sampling frequency, their effects on the signal are aliased to lower frequencies possibly within the range of the resting-state fluctuations and cannot be simply filtered out by band-pass filtering. Thus, during data preprocessing, retrospective methods estimate the effect of the noise on the fMRI signal and regress it out of the data [10, 42]. A resting-state network analysis usually concentrates on the spontaneous BOLD fluctuations in a frequency range between 0.01 and 0.1 Hz [21, 43]. However, recent studies have demonstrated that meaningful information is also contained at higher frequencies (at 0.17 and 0.25 Hz and higher) [44–46]. In the emerging field of rs-fMRI, numerous methods for detecting and representing these fluctuations have become available. One of the common methods is independent component analysis, which looks for independent spatio-temporal modes to detect areas belonging to a synchronized network (fig. 8). In seed-based analysis, temporal correlations between distinct brain regions are calculated voxel- and/or ROI-wise, offering a way of functional connectivity quantification. This analysis may be expanded by graph-theoretical methods to investigate network models over the selected brain areas, allowing a characterization of network-intrinsic properties. Further analytical approaches include clustering, pattern classification and several ‘local’ methods; their pros and cons are thoroughly reviewed e.g. by Margulies et al. [47].

**Translational MRS**

MRS is a unique in vivo method for investigating biochemical processes by detecting and quantifying in situ concentrations of certain chemical substances in the brain. MRS involves the detection of signals of compounds other than water by making use of the fact that certain atomic nuclei, if placed in a strong, static magnetic field, can interact with an externally applied, pulsed magnetic field, resulting in a frequency-specific resonance. Nuclei of different molecules possess distinct resonance frequencies and can be identified and distinguished on this basis.

**Single-Voxel MRS**

For single-voxel MRS, a cuboid-shaped target ROI is activated, and a single spectrum representing the biochemical composition of the ROI is measured. Spatial localization is achieved by sequential activation of three orthogonal layers, the intersection volume of which equals the ROI. An ideal ROI contains only tissue of interest and is of small size. The spatial resolution, however, is limited because MRS signals are generally quite weak and may easily disappear in the background noise if in vivo concentrations are very low. It is generally possible to compensate for signal loss by repeated signal acquisition; however, a 50% reduction in ROI size requires an extension of the duration of a measurement by a factor of 4 in order to achieve a similar SNR. Established sequences for single-voxel MRS analyses are the point-resolved spectroscopy sequence, consisting of one 90-degree and two 180-degree high-frequency pulses, and the stimulated echo acquisition mode sequence, consisting of three 90-degree high-frequency pulses.

Figure 9 displays an exemplary 1H spectrum from a murine hippocampus indicating characteristic peaks of different metabolites. The area under a metabolite reso-
Fig. 8. Hippocampal-prefrontal resting-state functional connectivity network in mice. This network was identified by independent component analysis. PFC = Prefrontal cortex; pHC = posterior hippocampus.
nance in the spectrum is proportional to the concentration of the metabolite within the measured volume. The integration of peak areas is usually sufficient in high-resolution MRS, since the peaks are clearly separated and the baseline does not fluctuate. In vivo MR spectra, however, often do not meet these prerequisites, and both the dominant water resonance in $^1$H-labelled spectroscopy and signals of slowly moving molecules may lead to an overestimation of the peak areas. In addition, the spectral resolution, i.e. the ability to distinguish adjacent peaks, is limited. Moreover, scalar coupling of different nuclei, indicative of molecular connectivity, frequently occurs and causes a splitting of the nuclear MR signal into multiplets with different intensity ratios.

As a consequence, there is a multitude of methods for in vivo MRS whereby the time or frequency signal is sought to be manipulated by mathematical operations [48, 49]. For example, more recent procedures utilize a priori information about the composition of the spectrum to be measured which is derived from simulations or measurements of each substance in model solutions. The signal intensity of the resonance lines can then be calculated from parameters of the adapted model functions. The resulting intensity values, however, are not true representations of metabolite concentrations, due to the influence of hardware- and sequence-specific parameters and the metabolites’ relaxation times. Moreover, the volumes feeding into the spectra usually do not encompass just one tissue, eventually leading to the measurement of metabolite concentrations of different types of tissue. Segmenting the MRS volume elements into grey matter, white matter and cerebrospinal fluid and entering their values into the analysis as covariates can be helpful in this regard [50]. Most problems can be avoided by reporting intensity ratios rather than absolute values. For example, the MR signal of creatine is often used as a reference in $^1$H-labelled spectroscopy. However, this technique is only valid and useful if the concentration of one metabolite is assumed to be constant [51, 52].

Animal MRS benefits in two ways from the high field intensities used in modern animal scanners: the SNR and spectral resolution are higher, which means that overlapping resonance lines can be separated more easily. Particularly the distinction and identification of glutamate, glutamine and γ-aminobutyric acid (GABA) in animal scanners is easier and of superior quality compared with the performance of human scanners, which usually have lower field intensities of 1.5–3 Tesla.

**In vivo $^1$H-MRS**

Besides tritium, a hydrogen isotope only rarely found in nature, protons are the most sensitive nuclei in MRI.
They are easy to measure due to their high prevalence and high gyromagnetic ratio. With a concentration of up to 40 mol/l, water is the most frequently occurring proton-containing molecule in live tissue. Even though its MR signal is the basis for MRI, it is a noise generator in MRS due to its much (up to 1,000-fold) higher intensity in comparison with other MR-detectable metabolites, requiring either suppression of the water signal during data acquisition or removal from the spectra in postprocessing [53].

Protons are part of most organic molecules, and the range of chemical shift is rather small. Thus, one would assume that in vivo $^1$H spectra consist of a large number of overlapping lines. However, the resonances of only a few metabolites are directly detectable in $^1$H spectra, for two reasons: (1) only small and mobile molecules generate a signal of sufficient intensity for detection and (2) a metabolite’s concentration must be high enough (1–2 mmol/l) to be separable from the background noise.

Figure 9 displays the most important metabolites detectable by $^1$H-MRS. N-acetylaspartate (NAA) is a neuronal marker, and the intensity of the NAA signal is a highly sensitive marker for morphologically intact and functioning neurons. The in vivo resonance of NAA consists of overlapping signals of NAA and, to a smaller extent, N-acetylaspartylglutamate. Both signals can be separated only in a highly homogeneous field.

Even though occurring in smaller concentrations than NAA, choline-containing molecules have a high resonance at 3.22 ppm due to their 3 methyl groups containing 9 magnetically equivalent protons. Choline emits further signals with a multiplet structure at 3.54 and 4.05 ppm – which, however, cannot be separated from the signals of other metabolites with lower intensity in vivo. The choline signal represents the choline pool. Acetylcholine, glycerophosphocholine and phosphatidylcholine – the latter two being integral cell membrane components [54] – are the functionally most important molecules of this pool. Increases in the choline signal indicate increases in membrane turnover, neuronal plasticity or reactive glia due to neuronal repair or degeneration.

Creatine and phosphocreatine form another group of important resonance signals. Creatine is partly ingested through food and partly synthesized in the liver, kidneys and pancreas from arginine, glycine and 5-adenosylmethionine. Phosphocreatine is the storage form of chemical energy in muscle, brain and nerve tissue [52]. Resonances of creatine and phosphocreatine are indistinguishable in in vivo $^1$H spectra. Protons of the CH$_3$ group are represented by a resonance signal at 3.02 ppm.

Myo-inositol (ml) is most easily detectable through its resonance at 3.54 ppm. The substance participates in intracellular signalling via inositol phosphate; however, the pathophysiological relevance of increased cerebral ml is largely unknown. In newborns, ml concentrations are significantly higher and decrease over several weeks. Further, ml is a sensitive marker for Alzheimer’s disease [55].
Glutamate is the most important excitatory amino acid. After synaptic release, it is metabolized to glutamine in the glia. Due to their complex multiplet structure, the resonances of glutamate and glutamine can only be separated in high field intensities (from 7 Tesla onwards). GABA can also be quantified at 7 Tesla. A change in the glutamate/GABA ratio may indicate a change from neurotoxicity to neuroprotection [54].

Non-Translational Methods

Optogenetic fMRI

An essential focus of psychiatric research lies on the investigation of functional networks within the brain. For example, prefrontal/hippocampal network connections are relevant in a variety of disorders such as depression or schizophrenia [56–58]. It is hitherto unknown how interregional interaction or functional connectivity alters under conditions of activation of specific groups of neurons. The combination of high-field MRI with optogenetic methods makes a pioneering move into investigating this question.

Optogenetics is a type of neuromodulation technology based on the specific expression of light-sensitive transmembrane conductance regulators such as channelrhodopsin-2, a monovalent Na conduction ion channel [59]. The activity of neurons expressing channelrhodopsin can be regulated directly in the range of milliseconds by application of blue light of 472 nm wavelength, which can be directed to a specific region in vivo by laser beam-transducing glass fibres. Both interneurons and excitatory neurons can be activated in a highly selective manner. Lee et al. [60] were the first to successfully describe the combination of optogenetics and functional imaging (opto-fMRI): rhodopsin was specifically activated in CaMKIIa-expressing excitatory neocortical and thalamic neurons and elicited BOLD signals not only locally but also in downstream targets of the stimulation sites. An example from our own group demonstrating the effect of CaMKIIa-expressing excitatory hippocampal neurons is displayed in figure 10.

Opto-fMRI offers a huge potential regarding the causal identification and analysis of functional networks and endophenotyping of dysfunctional disease-associated circuits. It could further facilitate understanding of the therapeutic effects of non-specific deep brain stimulation.

Further Refinements: Cryogenic Coils, X Nuclei, Manganese-Enhanced MRI

A promising avenue regarding the improvement of SNR and thus image quality is the use of cryogenic coils in animal scanners. Both the coil and the preamplifier of the signal receiver are helium cooled at approximately 20 K, which eliminates a large part of the thermal noise of the coil and electronic equipment. Cryogenic coils are mainly used in mouse scanners as bigger animals are a significant source of noise themselves. In mice, with $^1$H-labelled imaging techniques, an improvement in SNR by a factor 2.5 or higher is possible (fig. 11). In nuclei with lower frequencies (e.g. $^{13}$C), the improvement can be even more substantial.
MRS allows the measurement of a growing number of different isotopes (X nuclei) besides protons, such as phosphor (31P), carbon (13C), lithium (7Li), fluoride (19F) or sodium (23Na), each possessing specific technological requirements (coil, high-frequency generator and sequences) and preferred areas of use. For example, lithium, with its scarce natural occurrence, is particularly suitable for the analysis of brain metabolic or pharmacokinetic features. Through quantifying ADP and ATP, phosphorus spectroscopy can be employed in the investigation of energy metabolism. Carbon spectroscopy could play a relevant role in the future of tracer technology as 13C is a stable, non-radioactive isotope occurring in nature in minimal quantities (1.1%). Compared with micro-PET, theoretically there is a much larger repertoire of tracers, since no radiochemical synthesis is needed.

Manganese-enhanced MRI (MEMRI) has been used for nearly two decades in the investigation of biological systems with the aims of contrast enhancement, assessment of neuronal activity and tract tracing. Manganese is a paramagnetic calcium analogue; depending on the place of injection (intracerebral, intraventricular or intraperitoneal), the anatomical disposition and excitability as well as the neural networks and their functional connectivity can be displayed in a unique and distinct manner. MEMRI is advantageous for animals as they can freely move 24–48 h after injection; neuronal activity recorded by MEMRI, provided by accumulation of manganese, is anaesthesia free. On the other hand, manganese is relatively neurotoxic, which is why often only individual injection attempts are made, making the use of automated analysis difficult.

Conclusion

To conclude, MRI methods have proven their high reliability and sensitivity in translational research and build an important link between basic and clinical research. A number of discussed issues and pitfalls are to be considered and improved, which can then further amplify the potential of rodent translational imaging studies.

References

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