Summary

Recent research using multimodal neuroimaging has revealed that focal-onset epilepsies are accompanied by pathological changes in brain structure and function that extend far beyond the site of primary injury. These findings lend support to the notion that epilepsies are in fact network disorders, and that their clinical presentation, time course and treatment responsiveness might critically depend on network-wide effects. Although the network hypothesis is not new, current advances in neuroimaging technology have made it directly quantifiable - in vivo, non-invasively, with high spatiotemporal resolution and minimal inconvenience for the patient. In this article, we provide an overview of recent studies using advanced morphometric and functional magnetic resonance imaging (fMRI), show how these modalities can be combined with one another and/or electroencephalographic (EEG) data and present clinical examples of how these techniques might be applied in practice.

Key words: Temporal lobe epilepsy, lesion localization, functional connectivity, network effects

Bildgebung bei Temporallappenepilepsie: von der Läsion zum Netzwerk

Neuroimaging of Temporal Lobe Epilepsy

Introduction

It has been known since the work of Ramón y Cajal (1852-1934) that the brain is basically a vast, dynamic, densely connected biological network of breathtaking complexity [1]. Much of clinical neurology to date, including epileptology, has mainly focused on identifying critical nodes within network structures in order to identify lesions that e.g. cause hemiparesis or generate seizures. However, it is well known that a lesion can exert distributed effects across large-scale brain network, leading to metabolic dysfunction or structural alterations of remote brain areas connected to the primary lesion site [2, 3]. An example of these effects is crossed cerebellar diaschisis, classically described as a reduction in regional cerebellar blood flow contralateral to a supratentorial ischemic lesion [4-6], but also apparent during focal epileptic seizures and status epilepticus [7, 8]. Indeed, using advances multimodal neuroimaging techniques, network-wide effects have been increasingly recognized in focal epilepsies [3, 9]. It has therefore been proposed to view epilepsies as a “network disorder” and to integrate local and remote effects into a systems-level model of disease [3, 9 - 11] that would expand the well-known zonal concept commonly used in epilepsy [12]. In this article, we provide a brief overview of structural and functional imaging findings that support the network hypothesis in epilepsy. We focus on temporal lobe epilepsies (TLE), because they represent the most prevalent form of focal epilepsies and one of the most frequently investigated with neuroimaging [13, 14]. Also, they can be used as a paradigm for network-wide effects of focal lesions, given their often (at least visually) restricted pathology to structures of the temporal lobe.

Lesion Analysis in TLE

The hallmark of mesial TLE (MTLE) is hippocampal or mesial temporal sclerosis (MTS), which is detected by MRI in 75-85% of the patients [15]. Current clinical MRI protocols for temporal lobe abnormalities incorporate high-resolution T1w, T2w and FLAIR imaging, with at least one set of T2w or FLAIR coronal slices perpendicular to the long axis of the hippocampus [16]. Interpretation should focus on hippocampal volume, increased signal intensity on T2-weighted imaging, and disturbed internal architecture. The presence of MTS on preoperative MRI is predictive for good outcome after epilepsy surgery [17], whereas preoperative temporal lobe volume differences provided equivocal results [18, 19]. In non-lesional MTLE 18F-fluorodeoxyglucose-positron emission tomography (FDG-PET) is predictive for good surgical outcome; whereas an added value of 11C-Flumazenil-PET (FMZ-PET) and proton magnetic resonance spectroscopy (MRS) has not been established [20]. Subtle focal cortical dysplasia may coincide in the temporal lobe or in extratemporal areas ipsilateral to the MTS [21]. While clinical applications of automated voxel-based analyses and cortical thickness analysis for individual patients with MTLE are still experimental, improvements within a range of 10-30% in the detection rate of focal cortical dysplasias have been reported [22 - 24]. In addition to conventional MRI, advanced imaging techniques as functional MRI and tractography lateralize eloquent brain areas (language and motor function), predict effects of temporal lobe resection on memory and visualize the topography and composition of cerebral white matter tracts [25].

Network Analysis in TLE

While lesion analysis and interpretation is the essential task for the neuro radiologist in clinical practice, a substantial body of epilepsy research has shown that focal lesions influence brain areas beyond the epileptogenic lesion, across ensembles of functionally and anatomically brain areas [26 - 30] characterized by coherent physiological activity, such as high-frequency oscillations on EEG or low-frequency (< 0.1Hz) blood oxygen level dependent (BOLD) signal fluctuations. The BOLD signal is the most commonly used contrast mechanism in functional MRI and is thought to reflect the hemodynamic response of the brain to exogenous stimuli (such as an image or a tone) or endogenous neural activity (such as interictal epileptiform discharges). Covariance analysis of these fluctuations determines areas of the brain that form functionally connected networks. More than twenty independent networks have been identified that correspond to intrinsic and extrinsic systems, and are associated with internal- and external-oriented processing, respectively [31]. Many studies that investigated alterations of gray matter volume and concentrations using MRI-based automated analysis techniques as voxel-based morphometry and cortical thickness analysis on a whole brain level have shown that structural changes in mTLE patients are not restricted to the hippocampus or the elements of the mesial temporal lobe [32], but show widespread abnormalities that extend into the temporal pole, temporolimbic and frontocentral regions [33, 34], the cerebellum and the thalamus [35] and longitudinal increases in volume loss [36]. Neuronal network damage in TLE with hippocampal atrophy and in MRI-negative TLE has been reported to be more widespread in patients with a left-sided seizure focus [35]. This may be explained by more extensively connected temporo-frontal networks in the dominant hemisphere, due to their involvement in language function with more intense seizure propagation in the left hemisphere re-
sulting in a prominent neuronal loss in LTLE [37]. Furthermore, an association between residual seizures and atrophy in temporopolar and insular cortices in TLE with hippocampal atrophy and in the posterior quadrant in MRI-negative TLE has been reported [38].

Our group has recently examined the spatial relationship between widespread cortical atrophy and functionally connected networks linked to interictal epileptiform discharges, using simultaneous EEG-fMRI in a cohort of patients suffering from mTLE [39]. In EEG-fMRI, both modalities are recorded at the same time using MRI-compatible electrodes. After postprocessing, the time course of interictal epileptiform discharges found on the EEG trace can be used to indentify spatiotemporal correlates of the BOLD signal, i.e. regions of the brain whose hemodynamic response is closely coupled to the EEG-activity. It is thought that these areas represent the irritative zone and its associated network. In the aforementioned study, we detected hemodynamic correlates to interictal epileptiform discharges beyond the seizure onset zone in the ipsilateral insula, the temporal pole and temporo-lateral neocortex, in the cerebellum, along the central sulcus and bilaterally in the cingulate gyrus. Equally widespread reductions in grey matter volume were detected in the middle and inferior temporal gyrus, the uncus to the hippocampus, the insula, the posterior cingulate and the anterior lobe of the cerebellum. These findings were in line with the cortical thickness changes described above. Previous ictal connectivity studies in mTLE exhibited patterns of bilateral increases of cerebral blood flow in the temporal lobes (predominantly the middle and superior temporal gyrus including the temporal pole, the posterior temporal lobe and the cerebellum) and decrease cerebral blood flow in the inferior temporal gyrus, the inferior parietal lobe and posterior cingulated [40].

Clues from morphometric analyses and combined EEG/fMRI recordings implicate that the widespread structural damage linked to abnormal hemodynamic responses is suggestive of TLE being a system rather than a focal disorder leading to a disruption of structural networks. Both functional network analysis (derived from a temporal correlation of neurophysiologic signals in different brain regions) to illustrate brain dynamics on a system level and structural network analysis (using advanced neuroimaging techniques as diffusion tensor or diffusion spectrum imaging) provided further advances in knowledge about interregional network disruptions in mTLE [41]. While graph-theoretical analyses revealed a small-world organization of the cerebral cortex in healthy individuals [42], global network organization of patients revealed increased path length in TLE [41]. Several authors demonstrated altered connectivity in patients with mTLE. As a key finding, they reported increased connections within the mesial temporal lobe and decreased connectivity along extratemporal areas, including contralateral temporal regions [43 - 45]. Others suggested altered bitemporal connectivity patterns in patients with mTLE [46]. These findings suggest a deleterious impact of the epileptic lesion and the epileptogenic zone on the whole brain, potentially impacting multiple cerebral networks [47].

Network Relationships across Modalities

What is the relationship between the functional and structural changes reviewed above? A recent study by Voets et al. examined the association between resting-state functional networks, brain atrophy and changes in white matter microstructure in 35 patients with TLE [48]. Interestingly, Voets et al. reported reduced functional interactions between the hippocampus and anterior temporal and sensorimotor cortices that covaried with the extent of grey matter atrophy, i.e. reduced integration of the hippocampus with the rest of the network could be explained by the degree of grey matter volume reduction in the same areas. Of note, these areas parallel those that we found to correlate with IEDs in simultaneous EEG-fMRI. Additionally, Voets et al. discovered that functional integration outside this network, e.g. between frontal and temporal areas, was dependent on white matter microstructure rather than grey matter atrophy.

Network Relationships across Modalities: Clinical Examples

We present two cases where we used visual and automated methods to identify the epileptogenic lesion, EEG-fMRI to map the interictal network, and functional connectivity to define the lesion-dependent network.

Case 1

We used surface based morphometry (SBM) with FreeSurfer [49, 50] as a tool for comparison of cortical properties like thickness, curvature and sulcal depth between subject groups. As mentioned above, in [51] it was shown that SBM may also be useful for quantitative assessment of cortical malformations in individual epilepsy patients during pre-surgical evaluation. Here we compared cortical thickness of our patient (female, 34 years) to a gender matched control group of 16 females (age 30 to 40 years, mean 33.4, standard error of the mean 0.9) who received T1 weighted imaging with the same MR sequence (MPRAGE, TE=2.2ms, TR=1950ms, TI=900ms). While on the left hemisphere no significant deviations from the control group were found, several regions survived false discovery rate (FDR) correction on level 0.05 in the right hemisphere.
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Figure 1. Patient example: lateral TLE. A: Lesion in the right temporo-parietal cortex leads to extended deactivations of the default-mode network during interictal epileptiform discharges (IED), and shows pathological resting-state connectivity to the same areas. A: surface-based cortical thickness analysis (increase in hot colors, decrease in cool colors). Color bar represents z-scores compared to healthy controls. Insert shows a magnification of the largest cluster of cortical thickness increase in the right temporo-parietal junction, corresponding to an extended cortical heterotopia. B: Upper row, (rectified) time course of IED, derived from a multivariate analysis of the EEG inside the scanner during simultaneous EEG-fMRI. Lower row, time course of BOLD-signal fluctuations from the same examination, extracted from the cortical thickness cluster seen in Panel A. Time courses are normalized to unit standard deviations of each time series (y-axis). For clarity, only the first 200 of 460 time points (scans) have been plotted (x-axis). C: BOLD-correlation maps superimposed on 3D renderings of the patient’s brain (peak threshold p < .001, uncorrected, extent threshold p < .05, FDR-corrected). Upper row: positive (IED+) correlation between BOLD-signal and IED time course in the lateral and inferior temporal gyrus on the right. Middle row: negative (IED-) correlations between BOLD-signal and IED time course in an extended network including (i) on the right the lateral and inferior temporal gyrus, the parietal operculum, the inferior frontal cortex (ii) on the left the anterior cerebellum, and (iii) bilaterally the precuneus and the inferior parietal cortex. Lower row: positive resting-state functional connectivity (CON+) between the cortical heterotopia and a network overlapping with IED- above, including (i) right inferior temporal cortex and (ii) bilateral inferior parietal cortex and precuneus. (*) indicates approximate center of mass from Panel A.

(Figure 1, Panel A). These included notably two large clusters of increased cortical thickness in the temporo-parietal junction and the middle temporal gyrus, corresponding to an extended cortical heterotopia, but also significant atrophy in the fusiform gyrus (not shown). It has recently been shown that these malformations might still be functionally connected to healthy cortex [52], which might have epileptogenic effects. We therefore calculated the functional connectivity using cross-correlations between the BOLD-signal from the heterotopic cortex (Figure 1, Panel B, lower row) and the rest of the brain. We found a bilateral functional network (Figure 1, Panel C, lower row) that included the inferior parietal cortex and the precuneus bilaterally, areas known to participate in the DMN. Interestingly, a broadly similar network was found to be anticorrelated with, or deflected during, IED (Panel C, middle row), which extended into right frontal areas and the contralateral anterior cerebellum. Note that this pattern of deactivation has already been shown in a group of TLE patients using cerebral blood flow measurements with SPECT [40], underscoring the validity of our single-subject results. Note also that the region that correlated positively with the IED time course (Panel C, upper row) was restricted to a localized region close to the heterotopia, in strong contrast to the network-wide, possibly pathological functional connectivity effects (Panel C, lower row). Due to the volume of heterotopic cortex, no surgery could be performed.

Case 2

This 21 years old female suffered from longstanding pharmacoresistant epilepsy (from 5 years of age) due to left MTS (Figure 2, Panel A). She underwent presurgical EEG-fMRI to identify BOLD-correlates linked to IED (Figure 2, Panel B, upper row) that could be found...
in the depth of the left posterior temporal lobe (semi-transparent rendering in Figure 2, Panel C, upper row). Similarly to the 1st case, there was a widespread deactivation including the bilateral posterior parietal cortex, cuneus and precuneus and the left sensorimotor cortex. We used the BOLD-time series from the epileptogenic lesion to generate brain-wide maps of functional connectivity (Figure 2, Panel C) and detected a bilateral mesio-temporal connectivity network including bilaterally the hippocampus, insula, anterior cingulate and parietal operculum, and the lateral temporal cortex on the left (Figure 2, Panel C). In contrast to Case 1, there was no apparent overlap between IED- and functional connectivity networks. Of note, the left polar and lateral temporal regions visible on the functional connectivity map were included in the surgical resection. The patient has remained seizure-free (current follow-up 2.5 years).

Conclusions

An increasing body of literature — further illustrated by two clinical cases — indicates that epileptogenic lesions have widespread effects on the network structure of the brain. Here, we provided two examples of a network perspective on epilepsy that could further implicate clinical research and practice in future. One practical implication of the network concept is that it may radically alter our current classification of epilepsies [53]. Further, a concept of network organization along strategic nodes and connections may influence surgical planning. Several outcome studies indicated that seizure freedom can be attained in 60-70% of TLE patients with different resective approaches (anterior, medial or lateral temporal lobectomy and combinations thereof) [17, 54].

Despite the promises of a network-based perspective in epileptology, translation to the bedside remains challenging. From a methodological point of view, integrating multimodal data sets into a coherent analysis framework is a non-trivial problem, and will certainly necessitate advanced statistical modeling techniques. Much research is needed in this area. On the other
hand, there are practical questions for the daily clinical routine that need to be answered, before these methods can be brought to fruition. For instance, what is the sensitivity and specificity of neuroimaging techniques, alone or in combination [55]? How can results be reported in a rapid, easily understandable manner? What is the added value of multimodality neuroimaging and network analysis for patients and doctors; do they really inform clinical decision-making and improve treatment outcomes, or just add another layer of complexity to an already challenging situation? As we have shown with practical examples, methods are already in place to examine these questions and might help to advance epilepsy into a truly network-based medical discipline [53, 56, 57].

References

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