Introduction: The cerebral cortex consists of many structurally distinct areas. All cortical areas are interconnected through fibre tracts of the white matter, i.e., each cortical area receives projections from and sends projections to other cortical areas [1] [2] [3]. Such structural organisation can be modelled [4] [5] in a way that BAs are represented by a series of vertices, their connections by lines, and the lines are weighted by the strengths of the interareal connection. Vertices and lines then constitute the human anatomical cortico-cortical connectivity (CCC) network. The cortical network offers a quantitative description of cortical connectivity, which is still largely missing in the human brain [6]. It may also represent a link to studies employing tracers and electrophysiological methods of nonhuman primates as well as mathematical models of primate brain connectivity [1] [2], and, therefore, overcome the often problematic extrapolations from primate to human brain connectivity. The understanding of networks may contribute to our knowledge about how information is processed, disseminated and integrated among multiple specialised cortical areas. Here we attempt to propose a method for building the human anatomical CCC-matrix and investigate the topological properties of the cortical networks using graphical theory [7].

Methods: This study was carried out in 14 right-handed healthy males (mean age 31 ± 7 years). Written, informed consent was obtained from all subjects. MR data were acquired using a Siemens Tim-Trio 3T scanner with a 12-channel head coil. DWI data were acquired using a twice-refocused, spin-echo diffusion-weighted EPI sequence (60 volumes with b = 800s/mm² and 7 volumes with b = 0, voxel-size = (1.8mm)³, TE = 88ms, 75 transverse slices). DWI scan was repeated four times. 3D anatomical images were obtained using a T1-weighted MP-RAGE sequence. A brain mask, extracted from the averaged b = 0 volume, was taken as diffusion space and the 3D anatomical images as structural space.

Data analysis was carried out using FSL. Transformation matrices were calculated between diffusion, structural and MINI-152 standard space using FLIRT. BAs were defined using the template of a flattened Brodmann map which contains 41 discrete volume masks (= regions-of-interest, ROI), most of them representing one, but some also representing several BAs. An example of the latter was IPI – the insular and perinsular cortex [8]. Each BA was registered into every individual structural and diffusion space.

Using FDT [9], we performed fibre tracking for all possible pairs of BAs for each subject in diffusion space. Fig. 1 illustrates the steps for constructing the cortical networks: (a) calculating the fibre projection dαβ’s from α- to β- BAs, (b) computing the target-based fibre density from α- to β- BA as Cαβ = dαβ / Vα, (c) constructing the atlas-based ICC connectivity matrix, C, (d) generating the symmetric matrix, S, and antisymmetric matrix, A, for each subject and grading matrix elements. Each element of the symmetric matrix was scaled from 5 to 0 indicating the decrease of the strength of ICC connection and each element of the antisymmetric matrix was scaled from 3 to 0 indicating the difference of fibre densities. The group network was constructed by averaging the raw cortical networks from all individuals.

Results: (I) Connectivity-based cortical subdivisions: We applied an unsupervised learning network approach, a modified k-nearest neighbour algorithm [10], onto each individual anatomical symmetric CCC-matrix. The re-ordered correlation matrix thus obtained shows clustering structures. After mapping the identified clusters back onto the cerebral cortex, seven major connectivity-based cortical subdivisions were identified. The comparison of the individual matrices showed that the borders, the locations and the number of cortical subdivisions were nearly identical for all subjects, except for BA7 and IPI. As examples, the symmetric CCC-matrix and the re-ordered correlation matrix for an individual brain are shown in Figs.2a-b, and the group connectivity-based cortical subdivisions derived from all subjects in Figs.2c-d.

(2) Cortical networks derived from the subject group: Topography of the graded group-mean symmetric cortical network is shown in Fig.3, which indicates that most BAs connect to four to seven different BAs; BA6 and IPI connect to some more BAs, both of them occupy large parts of the cortical surface. The graded group mean anatomical antisymmetric CCC-matrix is shown in Fig.4. It can be used to represent the difference of fibre densities between every pair of BAs. Actually, the distribution of the fibre densities on the cortex is not uniform. Given two linked BAs (α and β), Aαβ > 0 is represented by a directed line β-to-α, i.e. the directed line leaves an area with lower fibre density and reaches at an area with higher fibre density. Fig.4 shows that the fibre density at the ‘IPI’ area is extremely low, BA10, in contrast, is not linked to other areas in this antisymmetric network, i.e. the density of incoming and outgoing fibres is similar.

Discussion: The abstract representation of the cortical network captures several essential characteristics of the cortical network although it has to be noted that it represents a rather simple static picture. The data showed the dense connections between the neighbouring and next-to-neighbouring cortical areas which is consistent with the results of nonhuman primate studies [1] [11]. Furthermore, the architecture of the cortical network was characterized by a small number of strong connections, a large number of weak connections, and by antisymmetry. The antisymmetric connectivity matrix may further contribute to a better understanding of the dynamics of brain function in more detail [12], such as information flow. It is worth emphasising that the constructed human brain ICC connection matrix and cortical networks are influenced by several methodological aspects, including the template of cortex, the tractographic method, and the grading schemes. An alternative strategy is to define the ROIs according to the cytoarchitectonic probabilistic map [13]. A challenge for further research is to explain how the cortical connectivity pattern mediates the behaviour: can the cortical areas be classified as ‘relay areas’, ‘giver areas’ or ‘taker areas’?