Traveling cortical netwaves compose a mindstream
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ABSTRACT
The brain creates a physical response out of signals in a cascade of streaming transformations. These transformations occur over networks, which have been described in anatomical, cyto-, myeloarchitectonic and functional research. The totality of these networks has been modelled and synthesized in phases across a continuous time-space-function axis, through ascending and descending hierarchical levels of association via changing coalitions of traveling netwaves, where localized disorders might spread locally throughout the neighboring tissues.
This study quantified the model empirically with time-resolving functional magnetic resonance imaging of an imperative, visually-triggered, self-delayed, therefor double-event related response task. The resulting time series unfold in the range of slow cortical potentials the spatio-temporal integrity of a cortical pathway from the source of perception to the mouth of reaction in and out of known functional, anatomical and cytoarchitectonic networks. These pathways are consolidated in phase images described by a small vector matrix, which leads to massive simplification of cortical field theory and even to simple technical applications.

INTRODUCTION
On a first sight it seems to be self-evident that a local perturbation within the cortical sheet should spread locally. The underpinning networks (see methods table II), dictating pathways, have been developed using different technologies and have varying numbers of nodes: anatomical50-38 cards are underpinned by fasciculi and smaller subcortical connectomes48-52 and structured by cytoarchitectonic39-47,94-96,98-100,104,109,111,121-124. These biological networks carry functions which, originally descriptively43,55 defined in psychological terms, lead, with integrated systems perspective56,57, to the low-dimensional multifocal cards from dynamic functional magnetic resonance imaging (dfMRI) seen in resting states58,61,62,64-89.
The networks are subject to a vectored dynamic system: in its multiple primary areas influxes of information flow inside and intermingle before running towards the only outlet in the primary motor area. This produces well defined temporal, spatial and functional boundary conditions. The system, even taking into account systemic feedforward-feedback-loops, yields an orientated direction and thus, using Hebbian1 principles as a basis, Mesulam2,3,4 develops a model (figure I) where a linear thought stream is represented via four levels of association. A hierarchical vector is created across the four levels, integrating the above-mentioned boundary conditions. This four levelled phase space illustrates not only effective but also causal and structural connectivity by using orthogonal, inter- and intra-associative projections:
A cycling phase might be started with the perception of a stimulus in a primary, idiotypical, unimodal association area, as e.g. within the visual cortex (ai); discrimination occurs further upstream in a secondary, still modality-specific association area (aii); heteromodal and often conscious integration follows in a tertiary, high-order association area (aiii); and as highest association area, a limbic, emotional response will upspring (l). Further ahead in time, the chain of ai-aii-aiii in reverse, falls down in descending order via a tertiary multimodal level of association, as in prefrontal thought processes (aiii); the anticipatory planning of movements, as in the vast secondary unimodal, supplementary-motor area (aiii); to the execution of the movement in the primary level of association (ai), the motor cortex – a process of execution which ultimately triggers the ai-aii-aiii chain once again: this time through self-perception which takes place in the primary somatosensory area (ai) over unimodal discrimination in secondary parietal regions (aii) to multimodal parietal regions (aiii), resulting in temporal models as shown in figure I and more detailed in the methods figures IV – VII and movie I.
Independent of complex, dispersed processes, the order of the sequence ai-aii-aiii-l-aiii-ai should be retained like a palindrome as a quasi-periodic pattern and be continually repeated in parallel and with varying phase lengths, just as the perception-reaction sequence involved in systematic thought processes also remains chronological and recurs rapidly throughout the day and gradually over the course of a lifetime in an ever-repeating self-similar loop.
Figure I. Hierarchies, longlasting waves and sensorimotor phases.

Above - The phasic hierarchy model of a visuo-motor task, adapted from Mesulam2, figure 2, stretched over time. The nodes named (within circles) following Mesulam (on the left), Jubrain45-47,63 respectively (on the right), 2π from visuo-perception to auto-perception. AIT - anterior inferotemporal cortex; f - area specialized for face encoding; L - hippocampal-entorhinal or amygdaloid components of the limbic system; LIP - lateral intraparietal cortex; MST - medial superior temporal cortex; P - heteromodal posterior parietal cortex; Pf - lateral prefrontal cortex; s - area specialized for encoding spatial location; PIT - posterior inferotemporal cortex; T - heteromodal lateral temporal cortex; TF - part of medial inferotemporal cortex; V1 - primary visual cortex; V2, V3, V4, V5 - additional visual areas; W - Wernicke’s area; wr - area specialized for encoding word-forms; 7a - part of dorsal parieto-occipital cortex.

Red lines – ventral streams; green lines - dorsal streams.
Details see methods, figures IV-VII and movie I.

Bottom - Event-related potentials ordered over time demonstrate like the rapid monosynaptic transmission-times in the range of 10 ms (from V1 to V5) to 50 ms (from V1 to AIT), as between Mesulam’s hierarchical levels, add up to longlasting waves in the order from 2 Hertz down to 0.25 Hertz. Note that Hebbian monosynaptic feedforward-backward loops occur in parallel to the slow frequency oscillations in the range of some 10 ms.

SPN - Stimulus-preceding negativity, awaiting an informative stimulus; N100 – preattentive perceptual processing; N170 - faces; P200 – preattentive perceptual processing; N200 – stimulus detection; N250 – face identity; P300 – stimulus categorization and memory updating; N400 – semantic and conceptual processing; P600 – syntactic processing; SEP – Sensory evoked potential; ERN NE - Error negativity, false acts; Pe - Error positivity, false acts.
The model raises a series of prognoses regarding the where, what and when of a transformation whose content is ultimately arbitrary and of the likelihood of such a linear, successive combination occurring. Regulated by feedforward-feedback-loops, interferences appear over longer time and space periods as static or moving waves, described by Crick & Koch as

... a forward-traveling, ... propagating net-wave of neuronal activity, (but is not the same as a wave in a continuous medium) ... These net-waves represent at each point in time 'coalitions' ... There are coalitions, which achieve winning status and hence produce [a] conscious experience ... In the conscious mode, it seems likely that the flow is in both directions so that it resembles more of a standing net-wave.

Deco & al. ground this dynamic into a neural field model where the above described neural masses rise out of spiking neurons, concluding in summary, for any model of neuronal dynamics, specified as a stochastic differential equation, there is a deterministic linear equation that can be integrated to generate ensemble dynamics.

The corresponding empirical spatio-temporal matrix \( M \) and its physical node parameters (such as location, volume, point in time, phase, frequency, amplitude) and edge parameters (such as distance, direction, duration, hierarchy, functions) are determined, using time-resolved functional magnetic resonance imaging (fMRI), both separately and as arrays in tasking, i.e. double-event-related brains, where a visual instruction was followed by a considered motor-reaction (see methods).

RESULTS

The appearance of the resulting phase image (figure II, details supplementary figures VIII & IX, movie II) unfolds before the view a wave train, which reflects the flow of the visual stream, the dorsal and the ventral one, fulminant, and reproduces at first glance in great clarity the sensorimotor stream, ventral and dorsal stream well separated too.

A right-left hemispheric symmetry of the activation pattern allows to limit the detailed analysis to one hemisphere, which will be the right one in the following detailed descriptions.

There is a medio-dorsal cluster in the visual phase image, the medial supplementary motor area SMA (figure II, pink oval), the latest cluster in time to have been activated. The same cluster is the first to be activated in the motor pictures through the movement. The two clusters in the two different phase images overlap in the right hemisphere over 356 mm\(^3\) (table I) and are constructed out of the same microanatomical, functional and temporal elements. Activation begins in the two conditions in pre-SMA (visual 3.1 s post, motor 2.1 s pre) and flows over the SMA proper to the medial upper border of the cortical mantle (visual 3.9 s post, motor 1.6 s pre). A joint time period is therefore consolidated in which a time arrow of 5.2 s represents the time from one perception over cognition to action and back to autoperception, the phase 2\(\pi\) in Mesulam’s model out of figure I, with wave fronts covering expansive, anatomical, cyto- and myeloarchitectonical, functional and hierarchical areas, parallel and perpendicular, as shown in table I:

i) Anatomical network architecture is quantified (table I, column 3; both hemispheres supplementary table II) with Automated anatomical labeling (AAL)\(^{32}\) over 15 out of the 44 AAL-nodes (calcarine, temporal, fusiform, lingual, occipital inferior, parietal inferior, cuneus, parietal superior, supplementary motor area, middle cingulum, precentral, postcentral, Rolandic operculum and once more parietal inferior in each hemisphere). The AAL-algorithm gives activated volumes accumulating in total to almost 10000 mm\(^3\) in each hemisphere; with an outer layer thickness\(^7\) of 2.3 - 2.8 mm a neural coverage of about 250 cm\(^2\) for a surface\(^8\) of 1200 cm\(^2\). Identifying first, extreme and last point in time of correlation allows modelling of the regional standing netwaves, the frequencies moving with 0.3 – 0.5 Hz in the lower delta band. The entire set of standing netwaves results in a stream of continuously phase-shifted waves, where the individual regions develop in parallel as winning coalitions and form a forward traveling netwave across time and space.

ii) The cytoarchitectonic network (table I, columns 4 - 7) resolution augments the number of nodes up to 26 out of the 64 nodes shown in figure V. The ventral visual stream is here composed of 17\(^{39}\), FG1\(^{91}\), ventral 18\(^{93}\), hOc4\(^{95}\), hOc3\(^{v}\), FG2\(^{91,98}\), BA 36, with a midtemporal branch hOc5\(^{96}\), BA 21\(^{97,100}\). The dorsal visual stream
Figure II. Phase images of a self-triggered response task.

Phase images from time series (supplementary figures VIII and IX, movie II) show the dynamic of an optically triggered, imperative, self-delayed response task.

Maximal-intensity-projections of the point-in-time of extreme activities as indicated by the colour bars. Left image event-related to the signal, and the two right pictures event-related to the respective reactions; stimulus-related from 1 second pre-stimulus (red) to 4 seconds post-stimulus (blue); reaction-related from 4 seconds prior to the reaction (blue) to 2 seconds after the reaction (red). The colour bar codes hierarchies\textsuperscript{3,4} too, separating unimodal (red/brown/green) from heteromodal (green/blue) areas.

Note the spatio-temporal gradient spanning from the calcarine sulcus over the medial, lateral and inferior extrastriate cortex to the precuneus and superior parietal cortex, and from the deep mid-frontal pre-SMA to the upper SMA proper (purple circle), where a spatio-temporal gradient starts in the motor images, followed within the lateral convexity by gradients from the central sulcus to anterior and to posterior areas, 7PC/mip short cutting to the latter visual superior parietal side (green circle).
is composed of 17, dorsal 18, hOc3d\textsuperscript{98}, ip\textsubscript{1}\textsuperscript{109}, mip\textsuperscript{104}, 7PC, 7A, and 7P\textsuperscript{105-108,124} and finally, a new observation, BA 6, pre-SMA\textsuperscript{63,110-116}.

The corresponding volitional motor response is made of a medial part from paleocortical aCMA\textsuperscript{113} over pre-SMA, SMA proper to 6aβ; and a lateral part from 4p\textsuperscript{121}, a dorsal stream to prefrontal 6aα and to postcentral 3, to 1\textsuperscript{122,123} and further on to 7P\textsuperscript{124}, as well as a ventral stream to 6aα inferior\textsuperscript{63} and to caudal PFt, Pfop\textsuperscript{100}.

iii) These macro- and microanatomical architectures port dMRI-correlations (table I, column 2). Their temporal architecture is obvious in a sequence over seven networks with sixteen nodes, representing their hypothesized functional succession: At first, consecutive visual networks\textsuperscript{62,64,66,69,71,78,80} turn up - the occipital, lateral and medial visual network as per Smith & al.\textsuperscript{64} with the latter two forming the high-level visual network, as per Zhang & al.\textsuperscript{78} including the six hierarchical nodes of Yeo’s & al.\textsuperscript{62} visual hierarchies. Within the higher association areas it is the dorsal attention network\textsuperscript{62,70,72,75,78-81}, which follows parietal and ultimately the medial-precentral part of the sensory motor network\textsuperscript{64,65,78,81} originating with the latter. This and the anterior cingulate cortex, a part of the ventral attention/salience network\textsuperscript{62,69,72,78,79}, is also the origin of the motor branch which subsequently employs the ventral motor network\textsuperscript{78}, and flows posteriorly and anteriorly back into the dorsal attention network, and inferiorly towards the Roland’s motor operculum in the direction of the insular lobe, spreading in an almost elliptical fashion from the sulcus centralis.

### Table I

Hierarchical, functional, anatomical and histological visuo-motor nodes of the right hemisphere forming a mindstream.

| M | dMRI | AAL | BA | Subregion | mm\textsuperscript{2} | s |
|---|------|-----|----|-----------|----------------|---|
| al, V1 | caudatus | 17 | 20 | 0.0 |
| all, X T | temporal std. | 21 | 199 | 0.1 |
| all, V4 | frontalis | 31 | 716 | 0.1 |
| all, V2 | lingualis | 14 | 816 | 0.3 |
| all, vm | occipitalis inf. | 10 | 149 | 1.1 |
| all, V2 | temporal std. | 21 | 122 | 1.2 |
| all, V3 | lingualis | 19 | 448 | 1.5 |
| all, vm | frontalis | 31 | 272 | 1.8 |

| M | dMRI | AAL | BA | Subregion | mm\textsuperscript{2} | s |
|---|------|-----|----|-----------|----------------|---|
| all, Ld | dat par sup. | 7 | ip | 111 | 0.5 |
| all, V2 | cuneus | 10 | 200 | 1.5 |
| all, V2 | cuneus | 19 | 170 | 2.4 |
| all, Ld | dat par sup. | 7 | 7PC | 52 | 2.6 |
| all, Td | dat par sup. | 7 | 1A | 200 | 2.8 |
| all, P | dat precuneus | 7 | TP | 666 | 2.8 |

| M | dMRI | AAL | BA | Subregion | mm\textsuperscript{2} | s |
|---|------|-----|----|-----------|----------------|---|
| all | cingulum medi. | 32,24 | aCMA | 220 | 2.2 |
| all | pre-SMA | 314 | 2.1 |
| all | SMA | 6 | pre-SMA | 302 | 2.1 |
| all | SMA | 6 | SMA proper | 42 | 3.9 |
| al | precentral | 4 | ip | 122 | 2.6 |
| al | postcentral | 3 | l, b | 110 | 0.2 |
| al | postcentral | 5 | 5 | 110 | 0.2 |
| all | SMA | 6 | 45 | 1.3 |
| all | precentral | 6a sup | 366 | 1.6 |
| all | postcentral | 7 | 7PC | 207 | 1.7 |
| all | Ld | dat postcentral | 7 | FP | 253 | 1.8 |
| all | par inf. | 40 | FP | 202 | 1.9 |

Columns: M - Mesulam\textsuperscript{2,3} (abbreviations as in figure I); dMRI - dynamic functional resonance imaging (networks: vp – visual primary; vl – visual lateral; vm – visual medial; dat – dorsal attention; vat – ventral attention; sm – sensorimotor; vm – ventral motor); AAL, automated anatomical labeling\textsuperscript{6}; BA – Brodmann\textsuperscript{40-41} areas; JuBrain\textsuperscript{42-47}, but the differentiation within the large surface region 18 between 18v (18 ventral, gyrus lingualis) and 18d (18 dorsal, cuneus and gyrus occipitais); also the as yet in JuBrain uninvestigated Brodmann areas 36 and 21; also aCMA, a small portion of BA 32 and the differentiation of the large surface volume of SMA according to myelographic classification\textsuperscript{63}; mm\textsuperscript{2} – activated volume; s - the average points in time of maximal correlations for a given node in seconds related to the event.
iv) The hierarchical architecture (table I, column 1), including dynamic ideas, is of particular importance and shows the phenomenon of a linear traveling netwave over the visual perception, discrimination and higher association areas as delineated by Mesulam\textsuperscript{2,3}. This demonstrates the starting point in early idiotypical, primary areas and its evolution over time via secondary unimodal to multimodal high association areas, located midparietal. These midparietal high-association areas are omitted in a speeded response task, demonstrating their high hierarchical position (supplementary figure X).

The reversed hierarchy is present in the movement-linked development from limbic regions over secondary to primary unimodal motor, to primary, secondary unimodal somatosensory and tertiary heteromodal sensory areas. Over the course of time the process recruits all the four Mesulam hierarchies over nine nodes.

The complex concordance from i) - iv) is grouped into large clusters, which radically simplifies the description to the determination of local and temporal extreme values (figure III). Vectors (black in figure III) are constructed between these extremes over the extension of those clusters, which in their orientation cover the above described, ordered sets of anatomies, cytoarchitectures, functions and hierarchies. Putting it together leads to a vector matrix bridging the combined visual occipito-temporo-parietal-midprefrontal, visual occipito-lingual-fusiform and sensorimotor prefrontal-precentral-postcentral-parietal networks. The matrix spans a field from the source of the visual perception through the conscious reflection to the action output into the newly emerging autoperception. By way of the constructed vectors, a wave develops like Hayne’s stream of consciousness\textsuperscript{9}.

Thus, the wanted matrix $M$ is well approximated.

DISCUSSION

Waves are built from their amplitude and their duration in space and time, oscillate in different frequencies, appear time- and space-locked to events, can spread locally and are believed to be fundamental mechanisms of neuronal computation and interregional communication\textsuperscript{7}.

Neurons are the elemental building blocks of such cortical waves, spiking predominantly in the form of propagating, traveling waves, while at the meso- and macroscopic levels, mainly stationary waves have been described so far. Such stationary signals have been explored regarding the significance of their physical properties, as especially frequency. Slow cortical potentials, $< 4$ Hz, so in the time range of the here observed signals, have been seen by He & Raichle\textsuperscript{10} as optimally positioned for carrying out large-scale information integration in the brain, have even been linked by them to conscious processing, and have been inverted by neurofeedback\textsuperscript{11}.

In fMRI the temporal properties of oscillations could be used to describe the standing, anticorrelated bumps on unimodal networks superficially and on multimodal networks very profoundly. Time-locked event-related standing oscillations are fundamental in electrophysiology (see examples of event-related potentials in figure I) and functional MRI, and even more information seems to be linked to phase-coded signals. Montemurro & al.\textsuperscript{12} calculate in macaques that at low field potentials frequencies, the phase of firing conveyed 54% additional Shannon information beyond that conveyed by spike counts. Glierean & al.\textsuperscript{13} demonstrate by using instantaneous phase synchronization as a measure of dynamic, time-varying functional connectivity in the visual cortex of man the same regions to be activated as in the present study but the mid-precentral SMA.

Lakatos & al.\textsuperscript{13} show even – again in the same wave-length as in the present study - that stimulus-locked phase-coding in the delta frequency range are present when attended stimuli are in a rhythmic stream. These oscillations ... entrain to the rhythm of the stream, resulting in increased response gain for task-relevant events and decreased reaction times.

However, traveling netwaves have only since recently, with the dawn of methods as multielectrode array, electrocorticography, voltage-sensitive dye, low field potential and intracellular recordings, been in the focus of interest. Traveling netwaves over some hundred’s milliseconds, observed in the cat’s visual cortex, voltage-sensitive dye registration in V1, V2 and V4 of the awake monkey, traveling beta oscillations during movement planning in
Figure III. Dynamic topology of a visuo-motor vector field.

The right hemispheres from figure II combined to an entire stream, surface-rendered and the histological architecture as per the definitions of JuBrain\textsuperscript{45-47}, with the exception of BA 21 and 36, which are not yet published. The subclassification of SMA/BA 6 as per Zilles & al.\textsuperscript{63}. 18m – medial (cuneus), 18l - lateral part (gyrus occipitalis) of dorsal JuBrain-area 18.

The colour bars provide the time scale in reference to visual perception and movement in seconds (supplementary movie III). Temporal extreme values were determined within the regions and vectors extrapolated (black arrows). The table to the left provides the temporal extreme values (t in seconds, the time-arrow of the reaction transposed by -5.2 seconds to visual perception) and their point in space (x, y, z – Talairach\textsuperscript{28}-coordinates in mm). The table to the right provides the vectors with their Euclidian distances in time (dt, in seconds) and space (ds, in mm).
macaques, and especially the electrocorticographic slow oscillations. 0.1 - 1.0 Hz, orchestrating large-scale neural spiking in human slow-wave sleep, traveling anterior-to-posterior with 1.2 - 7.0 m/s, lead Muller & al.\textsuperscript{14} to the conclusion that stimulus-evoked travelling waves are a general and important feature of dynamics in the visual system ... so ... that dynamic wave fields introduce a general conceptual framework.

All these individual biophysical properties - on standing waves, on frequencies, on rhythmic oscillations, on timelocking, on phase synchronization, on wandering waves – as observed in those studies are continuously and in overwhelming totality and simplicity present in the vector-matrix constructed in fig. III. The propagation speeds are from 0.1 to 0.25 m/s, the frequencies of 0.5 Hz, traveling across several centimeters, and adding up to a complete stream of about 50 cm length and 10 s duration. These waves develop parallel and dense in temporospatial continua, so that a current grows out of them. The phases of the current waves are temporally synchronized in a continuum along spatio-hierarchical-functional axes.

Methods to exploit the information encoded in time-varying connectivity have been developed in recent years in order to reach beyond the static measures of resting fMRI\textsuperscript{15,16}. Task positive paradigms are, in their nature, even more proximately linked to time-varying phenomena. While resting state might resemble testing an orchestra preparing for the concert, a task positive brain would be like observing a playing orchestra, where time synchronization will occur as the result of supervised Hebbian learning. Finally, it is the task-positive space where we look for orbits and attractors\textsuperscript{17}. However, the solution space shown here can now be used as seed-point series for dfMRI.

In the course of these considerations, the phases cluster were understood as Hebbian layers, which adhere to each other spatially. The synchronized stream that was observed here would, however, not represent the Hebb's error minimization but it's result. Thus high synchronization, sharing information between entities of a system, might be, far away from concrete, local functions as the perception of colour, intention or emotion, the essential Hebbian supervisor.

**CONCLUSION**

Slow cortical potentials are ordered to a spatio-temporal matrix $M$ out of a series of vectors, forming together a stream of consciousness\textsuperscript{9}.

Well evaluated neuro-biological networks are reflected in $M$ in a linear formation: The temporal order is reproduced i) between the dynamic functional networks visual, dorsal and ventral attention, and sensorimotor; ii) between the anatomical regions contained within these networks; iii) between the Brodmann and Zilles areas also found in these regions; iv) inside these areas and v) via functions and hierarchies.

Geometric waves are observed ‘hovering’ in the event of high over time synchronization\textsuperscript{18,19} between input and output, in a similar fashion to a taut bow string. Crick & Koch’s\textsuperscript{5} concept of consciousness as a forward-traveling, propagating net-wave of neuronal activity holds true and its quantification as phase-shifted synchronisation\textsuperscript{20} includes the simplification of biophysical dynamic causal models\textsuperscript{21,22}, as well as cortical field theory\textsuperscript{6}.

The Mesulam and more deeply stacked hierarchical models can be linearly parametrised in task-positive networks accordingly.

The discussed models of Hebb\textsuperscript{1}, Libet\textsuperscript{14}, Kornhuber & Deecke\textsuperscript{115}, Birbaumer\textsuperscript{108}, Crick & Koch\textsuperscript{5}, Mesulam\textsuperscript{2,3}, He & Raichle\textsuperscript{10}, Kjaer & al.\textsuperscript{106} and Haynes\textsuperscript{9} are essentially about the formalization of consciousness. It seems on the first sight evident that a complex system like consciousness should be composed out of even few and simple elements like other natural phenomena are constructed from simple elements. These elements, vectors, compose $M$, which can now be inverted\textsuperscript{23} in a computational form leading, like Birbaumer & al.\textsuperscript{108} did in neurofeedback with standing slow cortical potentials, to dynamic applications as multi-channel transcranial stimulation, inducing a virtual reality – spectra of cortical cycling phase continua forming a mindstream.
METHODS

The paradigm required a visual signal which prompted a test subject to push a button using either their left or right hand following a period they felt was “up to around 15 seconds”, with this process monitored using an fMRI scan:

Ten subjects (age range from 21 to 59 years, average age 30 years, 4 female, 6 male, all right-handers, as determined by the Edinburgh Inventory of Handedness) were studied. Informed written consents and approvals by the local ethic commission were obtained. The subjects held a button in each hand and watched visual stimuli presented from a video beamer onto a screen. The stimuli consisted of a short white flash (150 ms) on a black screen, followed by the appearance of a white arrow, which pointed in a pseudo-randomized sequence either to the right, or to the left. The arrow remained visible over the whole trial. The subjects were instructed to press the button with the corresponding thumb (left or right). They were furthermore instructed to choose, for each button press, varying temporal delays from zero up to approximately 15 seconds following the instruction flash. The whole experiment required 20 minutes, during which 87 trials of this delayed response task were presented with pseudo-randomized interstimulus-intervals of 14 +/- 2.5 seconds. The delayed task was contrasted with a speeded response task.

Event-related functional Magnetic Resonance Imaging was performed with a 1.5 Tesla MRI scanner (Siemens Vision, Erlangen, Germany). Twenty-four axial slices with 5 mm thickness and 3x3 mm² in-plane resolution, 64³ matrix, 8/8 192 mm² FOV covering the whole brain were acquired using a gradient recalled single shot echo-planar-imaging sequence. To shorten examination time TE was set to 40 ms. The measurement time for one whole brain acquisition was 2.32 seconds and volume repetition time was 2.52 seconds. Thus, for each experiment, 480 whole brain measurements were performed during a time interval of 20 minutes resulting in a mean of 5.5 volume measurements for each trial.

The image evaluation was performed with the Statistical Parametric Mapping programme package (SPM99, Wellcome Department of Cognitive Neurology, London). Preprocessing of the single subject time series images consisted of i) slice-timing, referring to the first slice (top), ii) realignment with sinc interpolation and adjusting for sampling errors, iii) non-linear normalisation to the canonical echo planar target image in Montreal Neurological Imaging space, interpolated with 2 x 2 x 2 mm³ voxel size, in order to achieve maximal spatial signal-resolution in the group statistics, and iv) spatial smoothing with a Gaussian filter with full-width half-maximum of 6 mm.

To generate time-resolved statistical maps, a series of models was calculated with time-shifted event delays. For each delay, the BOLD response was modelled using a generic hrf (global scaling, specified high-frequency-filter, using the hrf as low-pass-filter, which corresponds to a Gaussian filter with a cut-off level of 4 seconds full-width half-maximum, and without correcting for further internal correlation), resulting in separate t-maps for time points prior to the signal and the self-paced movements (-2 s for the signal, -3.5 s for the movements) and following (6 s for the signal, 2.5 s for movements). T-values were scaled for each volume proportionately from threshold (p < 0.001) to individual volumes extreme t-value at point in time t. The resulting values were assigned to a baseline corrected activity I. Spatial segmentation used the algorithm of Tzourio-Mazoyer, leaving only regions which excised 10% volume activity, except supplementary, figure X, insert i.

Table II shows cortical networks that served for comparison with the classification of the results. Figure IV shows the Jubrain-architecture, aligned with the Mesulam model of hierarchies, simplified in figure V, used to sculpture in figures VI and VII models of cortical streams that served for comparison with the classification of the results. Movie I shows the Mesulam model unfolded over time.
Table II. Nodes and networks in parcellations of the human brain.

The space, the anatomy, the cytoarchitecture, the myeloarchitecture, and the functions supported by the aforementioned structures, for the whole cortex as well as for the visual and motoric subsystems. Given are the nodes (per hemisphere) and, if investigated, their organization in networks and hierarchies.

i) Space – constituent Talairach [28] and MNF [29] spaces.

ii) Macroanatomy - Different anatomical subdivisions, as per old observations such as Ecker [30], Smith [31] and newer ones as Tzourio-Mazoyer [32], Smith [33], Desikan [34], Harvard–Oxford Atlas Makris [35], Desikan & al. [36], Klein & al. [37], Au 

iii) Cytoarchitecture – Campbell [39], Brodmann [40], and its transcription by the Yale Bioimage Suite Package Lacardie & al. [41], von Economo & Koskinas [42], Sarkisov & Filimonoff [43], Bailey & von Bonin [44], Zilles & al. [45], [46], [47].

iv) Myeloarchitecture - as per Vogt & Vogt [48], Flechsig [49], Geschwind [50], Catani & Ffytch [51] and the ongoing efforts of the Human-Connectome-Project [52] on cortical parcelling.

v) Function descriptive – Gall [53], Zilles & Rehkaemper [54], Van Essen & al. [55], the two latter ones consolidating Brodmann’s atlas.

vi) Function visual – Van Essen & al. [56] (12 hierarchies), Ungerleider [57] (6 hierarchies), Gleran & al. [58] (dynamic regional phase synchrony), Naser & al. [59], Grandinetti & al. [60], Pitrimos & al. [61], Yeo & al. [62].

vii) Function motoric – Zilles & al. [63] (integrating architectonic, transmitter receptor, MRI and PET data).

viii) Function dMRI – resting state functional magnetic resonance imaging observing networks (default mode network (consisting of a core, a dorsal medial and a medial temporal subsystem), executive control network, frontoparietal network, dorsal and ventral attention network, salience network, several visual and sensorimotor networks): Smith & al. [64] (ICA, 7000 subjects), Andrews-Hanna & al. [65,66] (hierarchical relations within the default network), Bellec & al. [67], Power & al. [68], Yeo & al. [62] (clustering ICA, 1000 subjects, separation of the early sensory and late motor cortices), Buckner & al. [69,70] (seed-based ICA), Choi & al. [71,72] (caudal-rostral parietal and rostro-caudal frontal hierarchies), Craddock & al. [73] (spectral clustering), Shen & al. [74], Spreng & al. [75] (hierarchical relations between three networks), Wang & al. [76] (interindividual), Joliot & al. [77], Zhang & al. [78] (directionality, subdividing two visual networks), Fan & al. [79] (eigenentropy), Gordon & al. [80] (individual), Braga & al. [81] (within individual), Schaefer & al. [82] (clustering and local boundary detection), Kong & al. [83] (individual, character-predictive).

ix) Other – Huth & al. [84] (multimodal, language), Fan & al. [85] (multimodal, structure, connectivity, function), Glasser & al. [87] (multimodal), Varfok & al. [88].

See Eickhoff & al. [89] for a detailed discussion.
Figure IV. Temporal and spatial phasic hierarchies in Mesulam’s\textsuperscript{2} anatomical/histological/functional model (centre, adapted from Mesulam\textsuperscript{1}) translate to the cytoarchitectures published in JuBrain\textsuperscript{45-47} (see methods movie I). The right hemisphere, unfolded.

Waves might find their way in any direction, as, e.g., the top-down effect of the Bereitschaftspotential\textsuperscript{115-116} or the top-down effect of attention in early sensory cortex, largely invisible to spike recordings, but readily seen in the fMRI signal of monkeys\textsuperscript{13,90}. 

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Figure IV: Temporal and spatial phasic hierarchies in Mesulam's anatomical/histological/functional model (centre, adapted from Mesulam's) translate to the cytoarchitectures published in JuBrain (see methods movie I). The right hemisphere, unfolded.

Waves might find their way in any direction, as, e.g., the top-down effect of the Bereitschaftspotential or the top-down effect of attention in early sensory cortex, largely invisible to spike recordings, but readily seen in the fMRI signal of monkeys.
Figure V. Mesulam\textsuperscript{2,3} hierarchies applied to the Brodmann\textsuperscript{40,41} /Zilles\textsuperscript{45-47}-cytoarchitecture.

The right hemisphere of a normalized\textsuperscript{28,29} brain, unfolded. Monofocal histological regions named following Brodmann & Zilles: FG3, FG4, 2, hip3, Te2.1, Te2.2, TPI, TI, HATA, amygdala, ID2-3, OP5-7, p24a, p24b, pd24sv, pd24cv, pd24cd, pv24c, s24b, s24a, 25p, 25a, p32, CH1-4 announced but not published.

8, 9, 38, 25, 34, 29, 26, 20, 35, 30, 23, 31, 24, 46, 47, 52 Brodmann-areas only.
The points give the centers of gravity and the lines give the long axes of the areas volumes out of figure IV.
The colors represent hierarchical hebbian layers, mono- or bisynaptic apart from each other.
Figure VI. The architecture of the ventral and dorsal visual stream.

Sculptured out of figure V.

Paragraphs: top row - concordance between Brodmann40,41/Zilles45-47/Mesulam2/dfMRI62,64-66,70,72,75,78-81/other; following rows - presumed functions2,5,6,57,59,60,62,64,58,91,97,104,108,105, all using different number and localizations of nodes, as well as different numbers of hierarchical levels; bottom row - deficit-syndroms.

**Red - The ventral stream ... shows a considerable functional heterogeneity from early visual processing (posterior) to higher, domain-specific processing (anterior)**91, an ascending hierarchy with the sequence V1 – V2 (V3) – V4 – inferior temporal cortex (IT), starting with simpler features posteriorly (TEO/PIT) that increase in complexity as processing moves anteriorly (TE/AIT) to perform object recognition92,93; asymmetrical, tending to the right hemisphere93; 1794 - V1, primary visual, retinotopic two-dimensional picturing of the external world; 1894 - gyrus lingualis, occipitalis inferior; hOc3v95 - lingual cortex; FG1/FG292,96 - fusiform cortex, orientation selective cells concerned with form, and direction selective cells concerned with motion. The fusiform gyrus hosts several "high-level" functional areas. It is involved in a core network subserving different cognitive tasks, that is, object recognition, visual language perception, or visual attention. It processes high-level visual information associated with faces, bodies, words, and places. FG2 is even a higher-order one96.

**Green - The dorsal stream comprises the sequence V1 – V2 (V3) – middle temporal cortex, MT, V5, always retinotopic – medial superior temporal cortex, MST – and finally intermediate object representations used by the decision-making circuitry further down the dorsal stream in lateral intraparietal sulcus, LIPS97; hOc3d98, cuneus, V3, V3A. There is a medio-temporal component consisting of hOc599,100 - temporo-occipito-parietal TOP-junction, related to the deficit symptom of neglect101; and BA 21 - medial temporal gyrus, it has been connected with processes as different as contemplation, recognition of known faces, and accessing word meaning while reading102;**

There is evidence to suggest that there is integration of both dorsal and ventral stream information into motion computation processes, giving rise to intermediate object representations, which facilitate object selection and decision-making mechanisms103. The areas within the intraparietal sulcus, IPS, in particular, serve as interfaces between the perceptive and motor systems for controlling arm and eye movements in space104. This region is functionally and histologically complex in so far as several association areas, default mode, executive control and dorsal attention105,106 overlap in here as well as the cytoarchitectonic areas 7A, 7PC, mIP, Ip1, Ip2, PF and PFM, sulcus intraparietalis, concerned with the integration of multimodal...
information for constructing a spatial representation of the external world in relation to the body or parts thereof, and planning and executing object-centered movements\textsuperscript{108}; 7P - precuneus, integration to a complex space out of proprioceptive and visual sources, information on objects in depth which are translating in space, because of the self-motion, object distance in a dynamic condition (as that created by self-motion) to orchestrate the eye and arm movements necessary to reach or avoid static and moving objects in the environment\textsuperscript{107}; self-perception\textsuperscript{108}. Premotor area engaged in the mental operation, the precuneus aids monitoring the operation in terms of internally represented visual images. Interfaces between the perceptive and motor systems for controlling arm and eye movements in space. Visuomotor tasks comprising target selections for arm and eye movements, object manipulation and visuospatial attention, deficit syndrome of apraxia, neglect, Bálint’s syndrome\textsuperscript{109}.

![Diagram](image)

Figure VII. The sensorimotor architecture.

sculpted out of figure V.
Paragraphs: top row - concordance between Brodmann\textsuperscript{40,41}/Zilles\textsuperscript{43-47}/Mesulam\textsuperscript{2}/dfMRI\textsuperscript{62,64-66,70,72,75,78-81}/other; following rows - presumed functions\textsuperscript{12,60,104,110,118}.

aCMA - cingulum anterior, its proposed role in cognition and its relationship with the arousal/drive state of the organism placing it in a unique position to translate intentions to actions\textsuperscript{110}, and being dedicated to the limbic portion of the cortex; 6\textsuperscript{111} - pre-SMA and SMA proper\textsuperscript{112,113}, intention to act, generation of readiness potential\textsuperscript{114,115,116,117}, action planning and gesture recognition, the operation of the motor monitoring, namely detecting the mismatch between the actual motor output and intended action goal\textsuperscript{117}, SMA proper and pre-SMA are active prior to volitional movement or action, as well as the cingulate motor area (CMA). This is now known as ‘anterior mid-cingulate cortex (aMCC)’. It has been shown by integrating simultaneously acquired EEG and fMRI that SMA and aMCC have strong reciprocal connections that act to sustain each other’s activity, and that this interaction is mediated during movement preparation according to the Bereitschaftspotential amplitude\textsuperscript{112}; 6\textbeta - the supplementary eye field (SEF), which has a special role in high level aspects of saccade control, such as complex spatial transformations, learned transformations, and executive cognitive functions\textsuperscript{118,119}; 6\textbeta superior - visuosomatomotor area hosting the frontal eye field (FEF)\textsuperscript{120}, which constitutes together with the intraparietal sulcus (IPS) ... an important role in the control of visual attention and eye movements\textsuperscript{104}, the coding of object positions\textsuperscript{59}; 4p\textsuperscript{121} - central sulcus, primary area, involved in kinematic and dynamic parameters of voluntary movements\textsuperscript{122}; 3a/3b,1\textsuperscript{122} - primary area; 2\textsuperscript{123} - the projection emphasising size and shape, secondary unimodal; 5\textsuperscript{124} - secondary, unimodal.
Movie I. Traveling cortical Netwave, hypothesis.\cite{hypthesis}

Jubrain\textsuperscript{45-47} regions aligned over time and neighbourhood from the visual perception to the voluntary motor reaction, and back over the autopereception and heteromodal association, assigned to the cortical hierarchies (coloured box bottom, right) of Mesulam\textsuperscript{2}, unfolded right cortical hemisphere.
Movie II. Traveling cortical Netwave, results\(^{126}\), [https://youtu.be/efEaO79gBkM](https://youtu.be/efEaO79gBkM)

Part i - The time series for the visual perception, 2 seconds ante until 5 seconds post perception, perception at \( t = 0 \) s; last picture the resulting phase image.

Part ii - The right and left hand motor reaction, 3.5 seconds ante until 2 seconds post reaction, movement at \( t = 0 \) s; last picture the resulting phase image.

The red-white boxes at the bottom represent the time axis in seconds.

Activity strength (scaled \( t \)-values of hrf-correlation) following the colour bar at the bottom, but in frame 12 and 27, the resulting phase images, where colour codes time.

Figure VIII. Perception-related time series.

Time steps in second distances (2 seconds ante until 5 seconds post perception, white numbers). Perception at time step three. Activity strength following the colour bar (see supplementary movie II, part i).
Figure IX. Reaction-related time series.
Right hand - above; left hand - below. Time steps in half second distances (3 seconds ante until 2.5 seconds post reaction, white numbers). Reaction at time step eight. Activity strength following the colour bar (see supplementary movie II, part ii).
Table III. Right and left hemisphere in the AAL\textsuperscript{32}-network.

|                         | mm$^3$ | $t_{\text{mean}}$ | $t_{\text{max}}$ |
|-------------------------|--------|-------------------|------------------|
| Parietal.Inf.L          | 416    | 5.0               | 0.3              |
| Rolandic.Oper.L         | 317    | 5.0               | 0.4              |
| Postcentral.L           | 2608   | 5.0               | 0.3              |
| Precentral.L            | 1657   | 4.6               | 0.3              |
| Temporal.Sup.L          | 156    | 4.6               | 1.0              |
| Insula.L                | 158    | 4.5               | 0.3              |
| HeschL                  | 15     | 4.0               | 1.0              |
| Cingulum Mid.L          | 233    | 3.8               | 0.3              |
| Supp_Motor.Area.L       | 408    | 3.0               | 0.3 right hand   |
| Precuneus.L             | 353    | 2.4               | 0.3              |
| Cuneus.L                | 488    | 1.8               | 0.3              |
| Parietal.Sup.L          | 268    | 1.2               | 0.5              |
| Lingual.L               | 356    | 0.4               | 0.5              |
| Calcarine.L             | 745    | 0.3               | 0.5              |
| Fusiform.L              | 439    | -0.2              | 0.5 visual signal|
| Calcarine.R             | 819    | 0.1               | 0.2              |
| Temporal.Mid.R          | 324    | 0.5               | 0.3              |
| Fusiform.R              | 1108   | 0.6               | 0.3              |
| Lingual.R               | 1334   | 0.7               | 0.3              |
| Occipital.Inf.R         | 140    | 1.1               | 0.5              |
| Parietal.Inf.R          | 309    | 1.5               | 0.5              |
| Cuneus.R                | 370    | 2.0               | 0.3              |
| Parietal.Sup.R          | 511    | 2.0               | 0.2              |
| Precuneus.R             | 686    | 2.8               | 0.3              |
| Supp_Motor.Area.R       | 356    | 3.2               | 0.3 left hand    |
| Cingulum_Mid.R          | 220    | 3.0               | 0.5              |
| Precuneus.R             | 1589   | 4.3               | 0.5              |
| Postcentral.R           | 2141   | 5.2               | 0.3              |
| Rolandic.Oper.R         | 253    | 5.3               | 0.5              |
| Parietal.Inf.R          | 202    | 5.9               | 0.3              |
|                       | 10160  |                   |                  |
| Supp_Motor.Area.L lha   | 709    | 4.0               | 0.3              |
| Supp_Motor.Area.R lha   | 220    | 4.0               | 0.3              |
| Supp_Motor.Area.L rha   | 1074   | 4.6               | 0.2              |
| Supp_Motor.Area_R lha   | 877    | 4.6               | 0.2              |

Visual signal, moving left and right hand (blue digits); AAL - Anatomical regions following Tzourio-Mazoyer\textsuperscript{32}; mm$^3$ – activated volume; $t_{\text{mean}}$ – mean-time of maximal activation over all voxels within the region in seconds, relating to the signal = 0 s (standard deviation 0.5 to 1 s); blue numbers for the movement, red for the signal. Black numbers indicate that this region activated under all conditions (the results for the SMA of the visual branch in the table, the results for the other conditions below); $f$ – frequency in Hertz; lha – left hand; rha – right hand; nw – netwave. The colouration of the waves corresponds to Mesulam\textsuperscript{2}.

The individual standing netwaves are approximated by

$$I_r = e^{-(t-t_{\text{max}})f}$$

where $r$ is the coordinate, its base-normalised regional nodes activity $I_r$, the time $t$, the mean time point $t_{\text{max}}$ of the maximum regional activities, and frequency $f$.

Phase-shifted waves develop in parallel as winning coalitions and form a forward traveling netwave across time and space.
The unique hierarchical position of the complex made of the parietal Brodmann area 7, a core area of the dorsal attention network\(^6\), becomes clear in contrast imaging of speeded and delayed-response tasks. The speeded responses (right) activate all of the same regions as the sum of the delayed responses (the visual phase image on the left, next to it both motor phase images) – with the exception of the high-association superior parietal regions (precuneus and parietalis superior, BA 7, pink rings). The high-association areas are apparently used only when they are involved in long-lasting feedforward-feedback-loops or, in short, in the case of accelerated responses, there is no time left for daydreaming.

Movie III. Traveling cortical Netwave, interpretation\(^{127}\). [https://youtu.be/fvLWPrw2Ro0](https://youtu.be/fvLWPrw2Ro0)

The fMRI time series for the visual perception and the following the left hand motor act, fusioned out of sup. movie II, using the medial supplementary motor area (resulting time axis 2 seconds ante until 7.5 seconds post perception, 7 seconds ante until 2.5 seconds post reaction),

The red-white boxes at the bottom represent the time axis, scaled in seconds above with respect to the visual signal and below with respect to the motor reaction.

Activity strength (scaled t-values of hrf-correlation) following the colour bar at the bottom.

The anatomical regions labelled following JuBrain\(^{45,47,91,99,100,109}\).

The evolution of different standing netwaves, the resulting travelling netwave and winning coalitions\(^5\) following AAL\(^{32}\) assigned to the hierarchies of Mesulam\(^2,3\) in the insert above left (see sup. tab. III).

Frame 20 gives the surface rendered phase image, right hemisphere, as well as the extreme values in space and time (left table) and the resulting vector-matrix (right table).

Colour codes time, with respect to the signal (upper timescale) and to the reaction (lower timescale). Labelling JuBrain. Temporo-spatial cortical vectors red. Their position in space \((x, y, z,\) Talairach-space (blue boxes) in mm) and time \((t\) in s) as shown in the left table. Their length in time \((dt\) in s) and space \((ds\) in mm) in the right table.
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