Review Article

A Review of Theoretical Perspectives in Cognitive Science on the Presence of $1/f$ Scaling in Coordinated Physiological and Cognitive Processes

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Time series of human performances present fluctuations around a mean value. These fluctuations are typically considered as insignificant, and attributable to random noise. Over recent decades, it became clear that temporal fluctuations possess interesting properties, however, one of which is the property of fractal $1/f$ scaling. $1/f$ scaling indicates that a measured process extends over a wide range of timescales, suggesting an assembly over multiple scales simultaneously. This paper reviews neurological, physiological, and cognitive studies that corroborate the claim that $1/f$ scaling is most clearly present in healthy, well-coordinated activities. Prominent hypotheses about the origins of $1/f$ scaling are confronted with these reviewed studies. It is concluded that $1/f$ scaling in living systems appears to reflect their genuine complex nature, rather than constituting a coincidental side-effect. The consequences of fractal dynamics extending from the small spatial and temporal scales (e.g., neurons) to the larger scales of human behavior and cognition, are vast, and impact the way in which relevant research questions may be approached. Rather than focusing on specialized isolable subsystems, using additive linear methodologies, nonlinear dynamics, more elegantly so, imply a complex systems methodology, thereby exploiting, rather than rejecting, mathematical concepts that enable describing large sets of natural phenomena.

1. Introduction

The presence of $1/f$ scaling in human performances is arguably one of the most puzzling, yet lawful phenomena in cognitive science (see [1–7]). $1/f$ scaling represents fractal, self-similar processes nested across multiple scales of measurement. Its occurrence implies that the rescaling of a time series leaves the distributional properties of the time series unaffected. Intriguingly from both a statistical and theoretical point of view, fractal scaling is widespread across the central nervous system, motor behavior, cognitive performances, and well beyond. As will be discussed, many physical and physiological signals exhibit scale-invariant features. Because it is now well known that the presence of $1/f$ scaling has a profound impact on cognitive and physiological activities, the phenomenon warrants serious attention.

Despite the fact that $1/f$ scaling is a mathematical concept that allows for descriptions of large sets of natural phenomena, the study of this relatively simple lawful description is relatively new. $1/f$ scaling, nonetheless, has the potential to reveal unexpected congruities among neurological, physiological, and cognitive activities. A general back draw is that the observation of $1/f$ scaling in itself runs against standard statistical intuitions. That is, successive observations of a repeated behaviors are typically assumed to represent measurement values that are independently drawn from a Gaussian distribution and thus to fluctuate randomly from trial-to-trial. Over recent decades, however, it has become clear that movement variability rarely equates with random, Gaussian noise and that temporal variability is usually structured and reveals specific details of the system dynamics [2, 5, 6, 8, 9].
In fact, structured variability appears to be the rule rather than the exception and is often as revealing as aggregate information in terms of unpacking the nature of the system organization [3, 10, 11].

$1/f$ scaling implies long-range dependence in the signal, often called long-range correlations or long memory. The associated serial correlations from trial to trial decay very slowly as the number of intervening trials increases, indicating persistent serial correlations, in contrast with the traditional view that they are transient (see e.g., [2]). This sets $1/f$ scaling apart from random noise, which lacks this serial dependence.

One way of revealing $1/f$ scaling is by translating dependencies in the time domain (i.e., a pattern of change in response time over trials) as simple features in the frequency domain using an operation called a Fourier transform, which decomposes the data series containing changes in response over trials into its constituent frequencies. Next, the power
(the square of the amplitude) of each contributing wave at that frequency in the decomposed signal is plotted in a log-log power spectrum. Also called power spectral density function, such a log-log power spectrum for a random data series (white noise, see Figure 1(a)), is shown in Figure 1(b).

White noise is not long-range correlated, as represented by a flat slope in a log-log power spectrum. $1/f$ scaling, in contrast, is expressed as an inversely proportional relation between log power and log frequency (see Figure 1(d)). This implies a nested sequence effect spanning over the entire time course of a measurement and even beyond, encompassing undulating "waves" of relatively longer and then shorter response times travelling across the series. In particular, faster (high-frequent) changes in response time are typically small and embedded in overarching, slower (lower-frequent) changes of higher amplitude (see Figure 1(c)). A third class of variability is Brownian noise (see Figure 1(e)), which can be generated by adding successive observations generated by a white noise process. Brownian noise is nonstationary, which means that variance increases over time. The log-log power spectrum of Brownian noise has a slope of $-2$ (see Figure 1(f)).

Typically, repeated human behaviors show a scaling exponent $\alpha$ in the range of 0 and 1, in between random noise and $1/f$ scaling. Examples of cognitive tasks include mental rotation, lexical decision, and visual search [2], simple noise and reaction time and word-naming [3], forearm oscillation [12], bidaily reports of self-esteem [15], and movement times (see Figure 1(f)).

The aim of this paper is to review the fascinating linkage between $1/f$ scaling and healthy, well-coordinated activities. Next, the most pertinent hypotheses about the origins of $1/f$ scaling in human activities are discussed, followed by a critical assessment of their relative strengths and weaknesses. Any of these hypotheses are able to successfully explain the presence of $1/f$ scaling but postulate opposing underlying mechanisms. The proposal put forwards is that questions concerning potential mechanisms and models that produce $1/f$ scaling should be rephrased. What matters is which explanation accounts best for the general linkage between observed fractal dynamics and coordinated physiological and cognitive processes. After all, each distinct account will have to deal with this broad linkage.

2. Fractal Dynamics and System Coordination

Recent years have witnessed increasing empirical support for widespread $1/f$ scaling in time series of physiological and behavioral processes. A main theme of this review focuses specifically on the association of $1/f$ scaling with healthy and well-coordinated behaviors. That is, deviations from $1/f$ scaling are often related to pathologic disorders, aging, external perturbations, high workload, or other situations where the system was not fully functional or coordinated. Rather than an incidental finding, this relation appears to be global and extends from small spatial and temporal scales (e.g., neurons) to the larger scale of human behavior itself.

That said, the growing body of research has further complicated this picture, showing that changes in the scaling relation occur simultaneously, and sometimes independently, on shorter (i.e., sleep-wake cycles, rest versus exercise, or circadian phases) and longer time scales, such as learning or aging (e.g., [21–26]). Observing deviations from $1/f$ scaling in shorter scale processes (e.g., transitions from one physiological state to another) under young and healthy conditions invite for a critical review of the relation between system performance and the presence of $1/f$ scaling on the longer behavioral scales, to (a) evaluate whether the current state of knowledge allows arriving at a coherent theoretical framework, and (b) pinpoint the questions that remain to be answered.

One biological example that hints at the generality of the claimed relation between coordinated system behavior and $1/f$ scaling is the vibratory motions of the membrane of red blood cells. The human body continuously produces new red blood cells (approx 2.4 million a second, cf. [27]), which compose a third of the cells in the human body [28]. Red blood cells are renewed after approximately 120 days, however. Interestingly, the membrane of red blood cells spontaneously vibrates, or "flickers," revealing $1/f$ scaling. Costa et al. [29] revealed, in addition, that the dynamical properties of this flickering behavior change with in vivo aging. Older cells emit less clear $1/f$ scaling, compared with newer cells that carry more oxygen.

This example is meaningful, since red blood cells constitute an important, basic system component, which is strongly related to an adequate functioning of one's physiology (and, hence, by extension human behavior). By reviewing functional benefits of fractal dynamics for living systems, an attempt is made to offer a vehicle for theoretical progress on the topic of $1/f$ scaling in cognitive performances. The steadily growing body of reliable positive evidence contradicts and grossly surprises conventional thinking about the architecture of cognitive systems. For one, a common scale-free “language” among perceptual-motor or cognitive tasks and (neuro-)physiology are not expected from classical componential models, which typically posit domain-specific control structures.

2.1. Pervasive Fractal Scaling in the Central Nervous System.

Brain activity is being investigated across a multitude of embedded scales of analysis. The finest level is the molecular scale which makes up cells, neurotransmitters, and so forth. Courser levels of brain activity include cell membranes with their synapses, microcircuits of dendritic trees, whole neurons, local cortical circuits consisting out of nearby neurons, entire cortical regions, and interactions among cortical regions, and pathways connecting them. At the coarsest scale one finds the central nervous system as a whole, embedded in a body, history, and environment. Evidence of fractal scaling at all of these levels would suggest common dynamical constraints across these embedded scales of organization,
begging the question whether the presence of fractal scaling provides support to this multiscale organization.

Anatomically out first, a mammalian brain reveals fractal properties in branching dendrite patterns, thereby maximizing functionality for a fixed dendrite cost [30–32], suggesting that fractal scaling in morphological specializations at the cellular level has a functional role (see also [33–35]). As a different example, fractal characteristics of microglia differentiate between healthy and pathological brains [36, 37]. And, also at more global organizational levels of the brain, fractal morphology yields functional advantages [38, 39].

Apart from the spatial fractal structure of brain anatomy, temporal brain dynamics are also reminiscent of complex behavior across many scales: $1/f$ power-law scaling in temporal activation patterns has been observed at all levels of neural organization, from ion channels opening and closing times to cortical networks [40]. The question of primary interest is how the dynamics of ion channels [41–45] interact with, rather than concatenate to fractal spike intervals [46–49] and the functional characteristics of larger scale neural ensembles [50–56].

Important clues may come from pathological brains. For instance, deviations from $1/f$ scaling, as typically observed in healthy controls, have been found with major-depressive disorder [57], mania [58], autism [59], epilepsy [60], and Alzheimer's disease [61]. These results suggest that $1/f$ scaling is prevalent in healthy, coordinative behavior of a healthy brain, and less so in the global activity of the brain with disorders and disease. Following on this suggestion, it was shown that the presence of $1/f$ scaling in brain activation correlates with the severity of depression symptoms [57] and the success rate of recovery from traumatic brain injury [62].

The scale complexity inherent to the study of brain activity makes it an enormous challenge for neuroscience to arrive at a universal theory of brain function. One first complication comes from the requirement of different methods at each level or scale of neural investigation. Each of the methods available yields a compromise between spatial and temporal resolution and, consequently, yields a priori choices in the organizational level of interest. No current method in isolation is likely capable of revealing the full-blown complexity of the brain, from the decimeter to the micrometer scale and from milliseconds to minutes up to developmental timescales. The discussed findings, however, should be taken as a strong suggestion that the inherent scale complexity of the human brain is a (if not the) key feature in itself.

2.2. Pervasive Fractal Scaling in the Body. A substantial amount of fractal applications in cognitive research have been motivated by initial successes in physiology, which had related $1/f$ scaling to the coordination, adaptivity, and flexible stability of the involved regulatory processes. Deviations from $1/f$ scaling in physiology have been found to relate to the degradation and decoupling of integrated systems, pathological conditions, severe disease, and increased mortality risk.

For instance, heartbeat interval series show $1/f$ scaling, with the clearest examples in healthy subjects. Deviations from $1/f$ scaling, either associated with excessive order (pathologic periodicity) or uncorrelated white noise (lack of temporal organization), indicate pathological conditions like congestive heart failure and ventricular arrhythmia [63, 64], and thus predict mortality [65]. Smaller deviations from $1/f$ scaling have been observed in heartbeat intervals in obese children [66] and adults with Down syndrome [67].

However, although alterations in the fractal properties of physiologic signals have been found to be reliable markers of changes in physiologic control, with healthy aging scale-invariant and nonlinear properties of heartbeat dynamics remain unchanged, with heart rate variability is significantly reduced healthy elderly, nonetheless [68, 69]. These findings suggest that healthy aging may not result in a continuous gradual change in scaling properties of heartbeat intervals, and implies that alterations in the cardiac control mechanism with advanced age differ conceptually from the mechanistic changes in the autonomic regulation associated with pathologic conditions.

Also breathing rhythm reveals $1/f$ scaling behavior but shows deviations thereof with aging [70, 71]. The contrary goes for development. Fetal breathing movements show more pronounced $1/f$ scaling with gestational age [72]. In contrast, deviations from $1/f$ scaling are observed in asthma patients, out of which those with more pronounced $1/f$ signatures in breathing rhythm showed better recovery after treatment [73]. These findings indicate together that fractal dynamics increase the overall efficiency of the respiratory system (cf. [74]).

Also fluctuations in blood pressure are found to scale as $1/f$ [75, 76]. Diabetic patients, however, show reduced $1/f$ noise specifically in glucose fluctuations compared with healthy controls [77, 78]. Another example are temporal fluctuations in colonic pressure. Yan et al. [79] observed $1/f$ fluctuations in the colonic activity of the healthy subjects, while patients hospitalized for slow transit constipation showed colonic pressure fluctuations deviating from $1/f$ noise towards Brownian noise; a condition yielding hardly bearable levels of pain sensation.

In physiology and medicine it is being increasingly acknowledged that a disease not only changes an average measure, such as heart rate or breathing rate, but is manifest in departures from fractal variability. Deviations from $1/f$ scaling are taken to imply a loss of physiologic control, which is often visible at very early stages of pathological development. The change in fractal dynamics with disease and, in some cases, aging suggested the new definition of disease as a loss of complexity, rather than the loss of regularity (e.g., [80]).

2.3. Pervasive Fractal Scaling in Motor Control. As in neuroscience, medicine, and physiology, there has been an increasing interest in fractal dynamics in human movement science. For instance, time series of postural sway differ reliably from random noise, revealing fractal properties (e.g., [18, 81, 82]).
Moreover, the clearest examples of $1/f$ scaling are found in young participants, while elderly participants show less clear fractal scaling in their postural dynamics [83], indicating degraded balance control [84–88]. This interpretation was further supported by Manabe et al. [89], who demonstrated less clear $1/f$ scaling in postural sway dynamics for patients suffering from Parkinson's disease and spinocerebellar ataxia, when compared to a control group. Interestingly, fractal measures of postural sway were even found more reliable than more traditional measures [90].

Also in gait intervals ([91, 92] are reviews)—the time intervals between successive steps in locomotion—the clearest examples of $1/f$ scaling are observed in young and healthy participants, whereas with aging, gait interval series become more random. It must be noted, however, that, while cardiac and respiratory dynamics show a more pronounced $1/f$ scaling and also higher degree of nonlinearity with gestation age and maturation, in locomotion deviations from $1/f$ scaling towards more random scaling are observed throughout maturation from childhood to adulthood (cf. [93, 94]).

In an elderly population, however, the presence of $1/f$ scaling successfully differentiates between fallers from non-fallers, and between Huntington's patients and control participants [91]. Moreover, the fractal dynamics of stride intervals produced by Huntington's patients correlates strongly with the severity of the illness ($r = -0.78$, cf. [91]), suggesting that deviations from $1/f$ scaling suggest impaired control of locomotion. Hausdorff [92] describes similar findings in Parkinson's disease patients and observed that among patients Parkinson's disease, the $1/f$ scaling relation in gait intervals "breaks down" and the stride-to-stride fluctuation in gait becomes very similar to white noise; each stride starts a new process, unlinked and unrelated to the previous stride. After successful treatment with proper medication, however, $1/f$ scaling becomes more prominent again in Parkinson patients' gait dynamics [95].

Using a different motor task, Wijnants et al. [17] presented college students with a very challenging precision-aiming (i.e., Fitts) task. The instruction was to move as fast and as accurately as possible between two circular targets with an inkless stylus. Participants were presented with five training blocks of 1100 back-and-forth movements. Very small targets were used, positioned wide apart, while participants were instructed to use their nondominant, least-practiced hand. A learning effect was observed after the extensive number of practice trials. Participants reached the narrow targets much faster after practice, while accuracy was maintained. A reliable effect of practice was found in the presence of $1/f$ scaling in movement time fluctuations; the presence of $1/f$ scaling became more pronounced with motor learning.

In a later study, Wijnants et al. [20] examined $1/f$ scaling in both spatial (movement amplitude) and temporal (movement time) time series in the challenging precision-aiming task, so notorious for revealing speed-accuracy trade-off (cf. [96]). Because of the difficulty of the task, participants were required to either emphasize the speed or the accuracy side of the trade-off (while equally being instructed to move as fast and as accurately as possible between targets) simply because the dual task constraints were so incompatible. The emphasized task requirement (temporal or spatial) directly affected the presence of $1/f$ scaling. Faster participants produced clearer $1/f$ scaling in movement times, but more random dynamics in movement amplitudes, as they performed the task less accurately. Conversely, more accurate participants produced more random dynamics in movement time sequences, as they performed the task more slowly, and clearer $1/f$ scaling in movement amplitude series. This effectively led to a trade-off between spatial and temporal $1/f$ scaling ($\rho = -0.64$), contingent on the well-established speed-accuracy trade-off.

In the same study, Wijnants et al. [20] established a strong relation between the fractal dynamics of movement time versus movement amplitude and the biomechanical constraint of minimizing the dissipation of mechanical energy. Faster participants, who produced clearer $1/f$ scaling in their movement time series also better capitalized on the elastic properties of the muscular system to recyle the kinetic energy of the approaching hand, arm, and shoulder in potential form, which is energy-related to the benefit of the next movement. Together, this amounted to a strong coupling among measurement scales in human performance: the biomechanical details emerged within the timeframe of a single movement, while speed and accuracy are determined by entire movements. The third timescale included the fractal dynamics that extend up to minutes of performance. The strong coupling between these multiple levels of motor performance provided further exquisite support for the nested multiscale properties in the organization of the cognitive system.

In contrast, externally driven performances have been found to minimize intrinsic fractal fluctuations. For instance, Chen et al. [13] observed no $1/f$ scaling when tapping to a metronome ($1/f$ scaling was observed in the asynchronies to the metronome, nonetheless), while $1/f$ scaling is clearly present in self-paced intertap intervals (cf. [97]). Similarly, in consecutive gait-cycles the external drive of a metronome breaks down the long-time correlations of the natural pace and generates random variability [91].

In a bimanual force production task, Wing et al. [98] observed random or close to random variability when feedback was presented to the participants (another form of external drive), while much clearer $1/f$ scaling was observed when no feedback was provided. Kello et al. [11] and van Orden et al. [3] interpreted these findings, altogether, as a confirmation that feedback provided constitutes a type of external perturbation that decorrelates the intrinsic fluctuations of $1/f$ scaling.

Perhaps the epitome of exercising external perturbation to motor control is to add a cognitive task on top of an initially simple motor task. Dual-tasking paradigm has been found to induce reduced $1/f$ scaling in the primary motor task [99]. This finding was replicated in a different task by Hausdorff [92] who invited Parkinson's patients to walk while performing a challenging secondary task. The gait dynamics revealed a reduced presence of $1/f$ scaling, compared with a no dual-tasking condition.
2.4. Pervasive Fractal Scaling in Human Cognition. The presented results from neuroscience, physiology, and movement science naturally extend to cognitive performances. Consider Figure 2, which depicts the average $1/f$ scaling exponents observed in four different experiments: word-naming [3], choice reaction [11], simple reaction [3], and precision aiming [17]. It is clear that the presence of $1/f$ scaling gradually increases over these tasks. Note that in precision aiming and simple reaction tasks, each experimental trial is identical. Each trial yields the same stimuli and the same type of response. This means that external sources of variation are minimized in these task performances. Consequently, the observed variation must largely reflect internal sources, which can be seen as a clearer presence of $1/f$ scaling in those tasks. One can readily notice the difference between precision aiming and simple reaction times. Precision aiming is a cyclic task, while simple reaction times become perturbed by a discrete signal to respond at each trial, and requires a discrete response at each trial. It can be seen that the extent of this perturbation effectively leads to a reduced presence of $1/f$ scaling.

The other end of the scale shows the $1/f$ scaling exponents from a word-naming task and a choice reaction task. In these tasks, each experimental trial differs, but to a different extent. In the choice reaction task, four different signals to respond were presented, with each signal requiring a different response. This procedure introduces more external sources of variation, which is revealed by a more whitened $1/f$ signal. Word-naming reveals the most reduced example, a task in which every experimental trial introduces a unique signal to respond. Hence, external sources of variation are maximized in this procedure, and the measured values likely reflect the intrinsic sources of variation to a lesser extent, requiring a more flexible and adaptive, perhaps less organized, control strategy. Similarly, the presence of $1/f$ scaling is typically reduced under high workload conditions [14, 100, 101] and when a sequence of presented stimuli becomes less predictable [11].

A closer look at one of the tasks, choice reaction, reveals a similarly interesting finding. Ward [102] discusses an experiment that was set up to manipulate the number of stimulus and response alternatives. In that study, Ward and Richard [103] presented participants with a choice reaction task that consisted out of either one (simple reaction), two, or four stimulus-response alternatives. The authors effectively showed that the presence of $1/f$ scaling is reduced as the number of stimulus-response alternatives increased. The scaling exponents were .60 for the simple reaction task, .37 for the two-choice reaction task, and .24 for the four-choice reaction task. (Note that the scaling exponents in this four-response choice reaction task are somewhat reduced, compared with the findings of Kello et al., 2007 [11]. Note however, that these differences might be due to the fact that Ward and Richard fitted the spectral slope over all estimated frequencies, while Kello et al. [11] only incorporated the lowest 25% of frequencies in their linear fit, as suggested by Eke et al. [104] and Holden [105]. All results in Figure 2 were obtained by fitting the spectral slope over the lowest 25% of frequencies only).

Another example of systematic perturbations attenuating the presence of fractal dynamics comes from [106], who presented participants with a temporal estimation task, in either of five conditions of accuracy feedback. In one condition, feedback was provided at every estimate, in another condition no feedback was given at all. In the remaining conditions, feedback was displayed only if the temporal estimate deviated from the target interval by more than 50, 100, or 200 ms, respectively. The results revealed an increase of $1/f$ scaling in conditions where less feedback (i.e., intermittent sources of perturbation) was provided, and hence, where the intrinsic cognitive fluctuations were revealed most clearly.

Gilden and Hancock [107] compared the performance of adults who reported Attention Deficit Hyperactivity Disorder (ADHD) symptoms with a control group of adults who did not, in a mental rotation task. The instruction was to press a key if the stimulus (a letter rotated by 0, 60, 120, 180, 240, or 300 degrees) was mirror-inverted and another key if the stimulus was not. Distinctive dynamical patterns of response in the reaction times were observed in both groups. More consistent participants revealed fractal scaling exponents close to $1/f$ scaling, while the more variably performers revealed random walk dynamics with a spectral slope close to $-2$. Hence, the more efficient system responses yielded a relatively clear $1/f$ signal, while the less coordinated, attention-deficit responses deviated from that pattern.

Wijnants et al. [108] inquired fractal scaling properties in the context of another learning disability. Response latencies of young children diagnosed with developmental dyslexia were compared with response latencies of nondyslexic children. The authors revealed that $1/f$ scaling measures reliably differentiated between both groups of readers when reading aloud single words sequentially. The dyslexic readers revealed more random response times, while average readers revealed clearer $1/f$ scaling. Furthermore, in the dyslexic group the $1/f$ scaling exponents of the response time series strongly
correlated with the severity of the reading impairment, as determined by average response time ($r = -0.56$), the negative correlation indicates that slower responses were associated with less clear $1/f$ scaling and standardized reading scores ($r = 0.77$), the positive correlation indicates that lower reading scores are associated with less clear $1/f$ scaling).

The summarized findings from research incorporating behavioral responses to repeated stimuli are very similar to the findings in neuroscience, physiology, and movement science. These findings raise the suggestion that many measurements related to how humans respond to repeated stimuli reveal signatures of complexity akin to the scaling behavior observed in simultaneously occurring processes (e.g., in the brain and the body). Importantly, these “cognitive” processes are traditionally treated as independent and self-supporting “directive systems” (i.e., the brain modules “in charge”) or “executive systems” (i.e., the autonomous nervous system “doing what it is directed to do”). The very observation that they may not be independent and self-supporting, but rather massively interactive, and complex in a true complex systems sense, could however cast doubt on the validity of the common sense distinction between the directive and executive front seemingly involved in the coordination of human action [55, 109].

3. Theoretical Perspectives on $1/f$ Scaling in Human Performance

$1/f$ scaling relations are systematically manipulable and indicate adaptive and flexible performances where no single timescale dominates coordination. Cognitive systems thus appear to ongoingly maintain a balance between competitive and cooperative processes in a flexible coupling across brain, body, and cognition. By confronting the most prominent, yet competing, hypotheses with the empirical observations presented above, an attempt is made towards a single conclusive theoretical account.

3.1. Multiscaled Randomness. One account for widespread $1/f$ scaling in human cognition conceives the phenomenon as reflecting many independent processes, each acting individually on their own timescale (e.g., [98, 102, 110–113]). For example, consider a time series where each measured response $X$ constitutes the sum of three different processes $Y_1$, $Y_2$, and $Y_3$, each evolving at their own independent timescale, in the additive form; $X(t) = Y_1(t) + Y_2(t) + Y_3(t)$. If one then assumes that $Y_1$ is a quickly random changing process, $Y_2$ is a random intermediate process, and $Y_3$ is a slowly randomly changing process, the additive series of all $Y$’s may yield $1/f$ scaling under specific circumstances (see for instance, [III, I13]).

Figure 3 shows schematically a power spectrum from repeated responses that combine the activity of a multitude of independent processes, each at their own timescale. In this schematic representation, indeed, the schematic distributions of time scales lead to $1/f$ behavior (see, e.g., [I14], for a more detailed description). This suggests that the complex fluctuations and $1/f$ scaling observed in many biological systems do not convey anything “special” about the mechanism generating these dynamics. From this perspective, $1/f$ scaling is a rather coincidental event that reflects nothing more than an artifact of information processing.

The main benefit of explaining $1/f$ scaling by aggregation of component processes is that such an explanation is conceptually simple and seemingly transparent and may therefore “demystify” the widespread occurrence of $1/f$ scaling (cf. [113]). That is, a componential model, consisting of many independent components, is effectively able to produce transient short-range correlations that, together, by chance may closely mimic the $1/f$ scaling behavior that is so ubiquitously observed across neurology, physiology, motor behavior, and cognition (e.g., [7, III–I13]).

Consider for instance the often cited dice-throwing algorithm in Gardner [II5]. Given three dice, when the first is thrown only rarely, the second intermittently, and the third at every observation. The sum of the values, when taken over a range of observations, will fluctuate in a $1/f$-like fashion, under specifically constrained conditions.

Nonetheless, there are a number of drawbacks. For one, independent multiscaled processes must exactly match the power and frequency associated with each involved component in order to coincide with a $1/f$ scaling pattern. However, if all parameters of the model are free, it is very unlikely that a system would, by chance, choose the “proper” parameters necessary to consistently generate $1/f$-like noise, let alone accumulate consistent relative changes across so many manipulations in neuroscience, physiology, and cognitive science. That is, “simulations demonstrate that if model parameters are unconstrained, the likelihood of generating $1/f$ noise is quite small. Thus, while the model can be used to generate $1/f$ noise with various scaling exponents, it is unlikely that the $1/f$ behavior observed in many biological systems is due only to the fact that these systems are regulated by many different inputs acting on different time scales.” [III, page 2154].

A second important drawback from multiscaled randomness is the lack of parsimony in the model. For every independent $1/f$ signal observed new components need to be
asserted [3, 4, 11, 116]. If not, that would mean the components are shared between processes, meaning each component is not strictly interdependent. Components of perception and action would hinge on synergies of physiological and cognitive component activities.

If one, in addition, considers a nongeneric 1/f process measured over a time range of ten minutes, or twenty minutes, regardless of its nature, multiscaled randomness would imply scaling relations to bend off at the lowest frequencies in the log-log power spectrum when sampling extends over longer periods of time (e.g., [117]). In the counter case, infinitely more short-range processes are to be invoked to keep the spectrum from flattening at the low frequencies (e.g., [118], see [4]), which would not be very parsimonious. Surely, the scaling relation may break down at extreme sides of the spectrum, but the criterion held in physics, which is observing 1/f scaling over at least two decades of frequencies, is generally well met in cognitive performances.

Thirdly, notwithstanding that the idea of identifying processes at different time scales is interesting and seems viable, the theoretical interpretation of the different time scales and the identification of their source still remain quite speculative. That is, the theoretical underpinnings for how and why exactly the hypothesized processes fluctuate in that specific manner are often underspecified in multiscaled randomness approaches. The critical question is not whether multiple independent processes could cause 1/f scaling in a measured signal (cf. [7, 113]), but rather the specific manner in which those processes must fluctuate. In particular, for any observed 1/f time series, which are many, one needs to examine the number of reasonable time scales involved in the specific process, the approximate values for those time scales, and the relative magnitudes of each influence. The involved processes that evolve at distinct time scales may themselves be as general as corresponding to conscious, preconscious processes, and unconscious processes [102], neural, behavioral and cognitive events [112], planning and control [16], or automatic, conscious, and sustained attention [113].

Despite the simplicity of the model itself, the examination of the data is limited to post hoc accounts for how neural, behavioral and cognitive events, attentional processes, or consciousness components are supposed to overlap (see [4, 118]).

When confronted with the experimental results reviewed above, revealing consistent changes in scaling exponent in different performances, from the fastest neural scales down to the slower behavioral processes, the “demystifying” multiscaled randomness approach of 1/f in human cognition, in fact, becomes an extraordinary hypothesis in itself. That is, the assumption that the observed behavior is jointly determined by many independent groups of neurons, each with their own different relaxation rate (determined by an autoregressive decay parameter), carries a massive theoretical load. The challenge would be to seek for deficient system components in depression symptoms, retiring red blood cells, as well as in severe constipation, asthma, dyslexia, or heart failure, among the many other examples, all happening to line up as a reduced 1/f signature.

3.2. Regime Switching. A second perspective on 1/f scaling in human cognition assumes that response processes show discrete transitions from one mode of operation (i.e., a specific mean or variance) to the next. These so-called regime-switching models propose that shifts in the mean or the variance of the observed process express shifts in strategy, fatigue or attention. First, it is assumed that, over the course of an experiment, participants repeatedly change strategies. These shifts are employed for only a limited period of time each. During these periods, particular threshold levels determine the criterion amount of accumulation of information required for a response. These changing threshold criteria present themselves as different plateau-like variations (see Figure 4(a)). Secondly, the speed with which the accumulation process approaches the current threshold is assumed to vary between successive measurements as a first- (or sometimes higher) order autoregressive process (see Figure 4(b)).

Under well-specified parameterizations regime-switching models are able to effectively generate 1/f-like fluctuations (e.g., [7, 113]), as shown in Figure 4(c). Regime-switching models may also account for nonstationarity (i.e., large criterium switches) and sudden switches in performance mode. Cognitive experiments designed to measure the degree of 1/f scaling are generally lengthy, as they usually require 2^9, 2^10, or more trials. In those experimental set-ups measured values are sometimes indeed susceptible to
undesired effects of learning, fatigue, shifts in strategy and attention, and the like. This means that it is well possible that many short-range dependencies happen to line up as 1/f scaling, showing “that 1/f noise is by no means ubiquitous in psychology” [119, page 740]. In particular, “something that looks like” 1/f scaling may simply cause a coincidental by-product of cognition through fluctuations in fatigue, attention, or strategy.

It is admittedly mysterious then, that 1/f scaling is so genuinely observed across the broadness of time scales of neurology, physiology, motor behavior, and cognition, simultaneously: Adding the coincidences (as it applies to independent events in probability theory) associated with the discussed studies underscores that the critical question is not whether strategy, attention, or fatigue fluctuations cause 1/f scaling, but rather the specific manner in which the component activities must fluctuate to cause exactly 1/f scaling, and its variations in the face of experimental manipulation. The associated coincidence renders regime-switching models as poor theoretically viable candidates to account for 1/f scaling in human cognition, lacking any physical motivation.

In addition, one could consider the fact that the broad family of autoregressive moving-average models can reproduce exactly any spectral function after the fact, as long as the spectrum has an asymptotic white region at low frequencies, because the model family has so many free parameters. A first inconvenience, as mentioned before, is that in psychological data the low frequency region does not bend off, suggesting genuine long-range dependence.

Secondly, the reduced 1/f scaling in repeated cognitive performances revealed in some of Wagenmakers et al’s experiments, which would indeed be expected given the postulated multifarious origins, could be explained by the random intertrial intervals used in these experiments, which was later shown to disrupt scaling (see [120]).

Third, the apparent success of the model is determined by goodness-of-fit per se, and again, realistic data can only be described by a rather narrow set of possible parameter variation (see [121]). The question is thus whether a model is a true representation of psychological processes or whether the model itself is so flexible that it is simply able to bend with the measurement error in producing good scores on goodness-of-fit. As Gilden [121] notes “Regardless of how small the minimum chi-square is for a particular set of parameter values, one will eventually have to reckon with the fact that the model did not predict that specific outcome; it predicted a range of outcomes, one of which may have happened to look like the data.”

Wagenmakers and colleagues claim that their componential models are “specified in enough detail to allow a wide range of data to be successfully described and, more important for scientific rigor, predicted” [7, page 114]. The first part of the sentence (i.e., the descriptive part) is obviously true, but disappointedly has little to do with the second part (i.e., prediction). As aptly summarized by Gilden, Wagenmakers et al’s approach describes so much, it in turn explains very little, and only offers a vehicle for mathematical formalism aimed at post hoc data fitting, lacking supporting theoretical literature. Therefore, the theoretical predictions mentioned in the second part of the sentence remain unclear. How do the ARMA model parameters corroborate and predict such consistent changes in the presence of 1/f scaling across such a variety of experimental manipulations? What is the theoretical ground for exactly the fitted parameters? Why is 1/f scaling so allied with coordination in complex systems, being only an artifact? The requisite list of post hoc explanations required to dismiss 1/f scaling as being functional, across all scales of observation, from the neuron up on to behavior itself, is simply too long, rendering the regime-switching model a rather speculative hypothesis.

### 3.3. A Blend of Short and Long-Range Correlations

A second prominent theory about 1/f scaling in human performance is the two-source model [2, 97, 121–123]. This model is motivated by the fact that a log-log power spectrum often does not exactly follow a straight line and may reveal a flattened (hence, whitened) slope at the highest frequencies.

The high- and low-frequency range can thus be modeled in terms of a constrained mixture of two distinct families of variability: white noise and 1/f scaling. These blended sources of variability are referred to as fBmW, where fBm stands for fractional Brownian motion and W for whitened, to emphasize the hybrid structure of whitened 1/f noises.

A fBmW response series has following form: 

$$R_T = (1/f^a)_n + \beta N(0,1)$$

where $R_T$ is the $n$th data point in a 1/f signal with zero mean and unit variance, and $N(0,1)$ denotes a random sample from the normal distribution with zero mean and unit variance. $\beta$ denotes a free parameter that determines the relative contributions of the white noise component. Accordingly, the power spectrum is built from the correlated and uncorrelated parts in a fBmW signal: 

$$S(f) = N(a)f^a + \beta^2$$

where $\beta^2$ is the fraction of variance in the random process attributable solely to white noise (see Figure 5).

A difference between Gilden et al’s account and the two models previously described is that 1/f scaling is conceived of as residing from a functional part of the cognitive system, rather than as a statistical artifact. In the approach of Gilden and colleagues, the presence of 1/f scaling represents a genuine fractal process, indicating that cognitive processes are complex across multiple temporal scales. A natural prediction is that 1/f scaling is generic to the behavior of cognitive systems and an intrinsic property of the system.

At the other hand one could note that 1/f scaling is not exactly seen as a dynamical signature of an entirely integrated system either. 1/f scaling is rather hypothesized to reflect the self-organization of a component within the system, a component that is associated with elementary aspects of cognition, including the goals, intentions, and representations of a participant. In particular, Gilden [2] postulated that intrinsic fluctuations of a memory module, which serves the purpose of continuity of “mental set,” are potentially causal in the formation of 1/f scaling. Other processes, for example, motor processes, constitute a source of contaminating white noise, and human performances constitute an additive blend of a complex 1/f process and
Figure 5: (a) Shows a white noise signal, which was added to the $1/f$ scaling in (b) to construct the power spectrum in (c), which reveals $fBmW$ noise, as indicated by the flattened slope at the highest frequencies, shown as a solid line. The dashed line represents the ideal $1/f$ scaling slope.

The pure white noise signal, in varying degrees in different performances.

The two-source approach fits in with more traditional models, where cognitivist dual system models are common. The mental set hypothesis may explain why the presence of $1/f$ scaling is reduced in cognitive performances when the task is not constant across trials. For instance, when task parameters change unpredictably, mental set is interrupted, which will induce trial independence in responses, resulting in scaling properties closer to white noise. An example given by Gilden [2] is task switching. This interpretation appears viable if one considers the varying degree of $1/f$ scaling in the four tasks presented in Figure 2. In particular, Thornton and Gilden [123] presented a detailed account for Figure 2 based on their model, by showing that temporal estimation tasks (comparable with the precision aiming task in Figure 2 for the present purpose) not only reveals clear $1/f$ scaling, but also a contribution of the white noise component, while a choice reaction task shows less clear $1/f$ scaling, with a much stronger white noise contribution, and word-naming data reveal the least clear presence of $1/f$ scaling and the strongest contribution of white noise. It thus appears that the model adequately describes a subset of the discussed data.

Nonetheless, the model does not specify specifically which of the dual components in the model would change specifically, and specifically why or how the error term gives complementary information. The presented results also suggest that an independent system component attributed to mental set [2] or vigilance [107] might be too limited in scope to account for all shared instances between $1/f$ scaling and coordination in brain, body, and behavior, across the measurement scales involved.

A related approach to $1/f$ scaling originates from Delignières and colleagues and has been applied primarily to rhythmical movements. These authors have presented a series of concrete models, each specific to a well-defined domain of motor performance, which effectively mimic empirically observed scaling properties. Like Gilden, their approach is to insert a local source of $1/f$ scaling in statistically well-defined, local parts of traditional cognitive models, to account for the relative presence of $1/f$ scaling and specific spectral deviations thereof. As an example, Delignières et al. [12] adapted the Wing-Kristofferson model [124] for finger tapping, by inserting fractal properties, into the cognitive timekeeper module assumed by the traditional model, using the regime-switching model previously discussed. Their model accounted for the scaling properties of self-paced tapping. Other examples include synchronized tapping [125] and forearm oscillations [12].

Delignières et al. point out that their purpose is to account for the workings of a particular, domain-specific component encapsulated in the system and use $1/f$ noise as a constraint for modeling. Sources of $1/f$ scaling are taken to represent complex timekeeping processes that can be statistically localized in components within the system. Thus from this perspective, $1/f$ scaling represents a functional aspect of human performance related to cognitive timekeeping, but the different source of $1/f$ scaling in each domain-specific application has the form of a fractal generator that together with nonfractal components, makes up the dynamics of repeated responses.

Mechanistic (i.e., two-source) modeling has the advantage to be experimentally testable and thus falsifiable and allows establishing links to current models of sensorimotor behavior. There is no a priori reason why long-range and short-range dependence should be mutually exclusive (cf. [4, 7]), and thus the observed serial correlations are possibly the result of both, and mechanistic models provide a route
to separate the long-range from the short-range components. Separation of variance components led to good fits to empirical data in specific tasks, under the assumption that different system components are responsible for the differences in performance (e.g., [126]).

Interestingly, the timekeeping hypothesis proposed by Delignières and colleagues appears to be consistent with the lack in $1/f$ scaling when participants synchronize to a metronome (i.e., an external timekeeper) evidenced in tapping and walking experiments as discussed previously.

The concern, again, is the approach of post hoc data fitting. While good data fits may be compelling, agreement between model and data is not proof that the model is correct (cf. [121]). The pitfall of mechanistic modeling is that one may mistakenly believe that a good quantitative model fit equals qualitative or theoretical insight. A good fit to the data is a necessary, but not a sufficient criterion for a model's usefulness [127]. A consideration of a model's usefulness involves, for instance, also a consideration of the theoretical foundations, the extent to which the model points to new research directions, and the generalizability of the model.

Advanced data fitting may lead to an indefinite number of fractal generators throughout brain, body, and cognition, possibly leading to multiple competing models capable of equally compelling fits to the data [121], which would question in itself the tenability of domain-specific theoretical explanations of $1/f$ scaling [11]. In particular, a full account for the reviewed studies would require domain-specific models for each of the observations across scientific domains and scales of observation. Although this could arguably lead to a well-fitting model for each separate phenomenon, that approach would lack the coherence of a solid theoretical framework over brain, body, and cognition necessitated.

3.4. Interaction-Dominant Dynamics. $1/f$ scaling is a typical behavior of self-organizing systems, which reflects a fundamental aspect of all physiological and cognitive functions: their emergence in the balance of independent versus interdependent component activities. Interaction-dominance postulates that the cognitive system actually uses this “flexible stability” to its own advantage by exploiting the interdependencies among ongoing processes over a multiplicity of interdependent scales, allowing for a coordinative basis of cognitive function (e.g., [11, 17, 128, 129]), and suggesting a common control mechanism among brain, body, and cognition, across the full range of scales involved.

This perspective conceives $1/f$ scaling as the natural outcome of complex, living systems. In the true physical sense, complex systems consist of a set of interrelated and interdependent parts with an almost infinite amount of degrees-of-freedom that cohere into a coordinated functional system. The system components dynamically interact in nonlinear ways at multiple embedding time scales, so that the intrinsic dynamics of the components matter less than the mutual interactions among components. This position is a departure from the more traditional view on human cognition that conceives human performance as caused by a number of discrete components (i.e., independent regions of the brain, internal clocks, or other information-processing devices), whose internal dynamics, when integrated, account for the observed performance. This convention can be referred to as component-dominant dynamics because the intrinsic activities of the components are held to be much more influential, much more dominant in determining the observed performance, than the interactions among the components.

The starting point of the interaction-dominance approach is that the conventional way of thinking about cognitive processes underestimates the number of temporal scales on which cognitive activity is actually assembled. The interaction-dominant perspective claims that cognition is more than a collection of independent processes operating in a modular cognitive system and entails that the same processes govern cognitive performance in very short (i.e., neural) and very long time frames (e.g., behavior). Any measured behavior nests processes at faster time scales, and in turn, is nested within processes at even slower time scales, and the behavior of any one process at any one time scale is susceptible to, and reflective of, the behaviors of all processes evolving over shorter scales. This means that the many processes involved interact so completely, up to the periphery of the nervous system and that one can no longer parse the individual activity of any component apart, which reveals itself through widespread scaling relations, as the essential outcome the interdependence among components across multiple scales.

In interaction-dominant systems control is distributed rather than localized as in specialized devices that form efficient causes for behavior. Coordinated system performance is therefore taken to emerge through the interdependence and cooperation of processes that operate at many timescales simultaneously. The tight coupling of mutually constraining timescales of performance, as in $1/f$ scaling, through self-organization (the spontaneous organization that coordinates system behavior in the absence of a central controller) and emergence (the appearance of features that are not implicit in the parts of the system), in an a priori prediction.

Interaction-dominance is a viable explanation for the presence, and the relative changes in the presence, of $1/f$ scaling in the activity of brain, body, motor system, and cognition. Self-organization parsimoniously explains that $1/f$ scaling is so generic to the behavior of such a wide variety of embodied processes. To give rise to $1/f$ scaling, a system does not depend on the behavior of a specific subsystem, but rather coordinates its behavior system-wide. As a consequence, one cannot dissect functionally entangled phenomena into component processes. Coordinated system consists of entangled processes evolving over a multitude of temporal scales, observed as a clear fractal signal. A further natural prediction from interaction-dominance, confirmed empirically, is that external perturbations to the system reduce or decorrelate the $1/f$ signal.

Nonetheless, some authors remain reticent and cautious to embrace an interaction-dominant position to the study of cognitive phenomena. For one, Wagenmakers et al. [7] have argued that $1/f$ noise is not a unique property of interaction-dominant systems, nor do all interaction-dominant systems
display $1/f$ scaling under all circumstances. Wagenmakers
et al. [130] have taken a pile of salt as an example. It
remains unclear, however, whether this abstract example
truly invalidates the broad linkages between $1/f$ scaling and
human coordination presented here. In contrast, what this
argument underscores is that all model systems, like rice
piles, need to be tuned to elicit the emergent behavior. This
illustrates just how hard the scaling by coincidence hypothesis
is to maintain for complex living systems.

Another issue is whether concepts and mathematical
tools from nonlinear dynamical science to cognitive phenomenen
ca lead to a notable advance in the understanding of
those phenomena, if such applications are not accompanied
by concrete mechanistic models. That is, some scientists have argued that, in order for interaction-dominance to be testable and falsifiable, the equations that govern the system should be
written out [1, 7]. As it stands, however, reproducing $1/f$ by
a model is not sufficient, since many mechanisms can lead to
the same scaling behavior, but with different nonlinear and
multifractal properties that do not match the behavior of the
studied system.

van Orden et al. [4] have spelled out some of the difficulties that arise at this point. For one, “there is presently no workable entry level below the level of the emergent phenomenen themselves” (page 121). While it seems doable to postulate adequate models that describe separately the functioning of, for instance, red blood cells, motor behavior, or brain activity, it remains unclear as to at which of these levels one should start in a reduction of an interaction-dominant system in the true physical sense. In other words, “emergent macrolevel behavior is antithetical to the conventional reductive pursuit
of cognitive mechanisms.” [4, page 121]. Therefore, support for interaction-dominance must come from either empirical results (i.e., observing consistent changes in the presence of $1/f$ scaling following careful experimental manipulations) or other phenomena that reveal analogous self-organizing properties (i.e., convection rolls or ecosystems, [3, 131, 132]), and neural network models [133, 134].

In the context of models that generate $1/f$ scaling, there have been important developments after the proposition of the multiscaled randomness and the two-source model mechanism that complement earlier analogies with simplified physical models (i.e., an Ising model, see [135], or a sandpile model, see [136]). Some of these efforts have been successfully applied in cognitive science (e.g., [137, 138]) to generate scaling that holds for a range of model parameters. However, no single model family currently seems able to coherently
capture the widespread scaling relations empirically observed
across the fast neural scales down to the slower scale of
human behavior and cognition.

4. Conclusion

In this paper a discussion is provided about the linkage between the presence of $1/f$ scaling and coordinated brain,
motor, and cognitive activities. In particular, it has been argued that $1/f$ fluctuations govern healthy, flexibly-stable
systems, regardless of the scale of observation, from the level
of the cell up to the level of brain, body, and cognition. $1/f$ scaling has triggered a lot of controversy over recent years as an empirical phenomenon. Therefore, the goal of
the present paper was to advance this debate, by focusing on
a broad range of empirical observations that transcend the boundaries between scientific domains. The discussion focused on different perspectives that seek theoretically
motivated accounts for these findings.

The question posed was how to understand the close
linkage between $1/f$ scaling and system coordination. The
goal was to promote answering the harder type of questions based on this linkage. What do the systematic fluctuations observed in the performance of so many different studied systems signify? Do these fluctuations constitute errors or
do they play a functional role? How and why is long-term
stability associated with short-term flexibility when system behavior is governed by $1/f$ scaling? While all of
the discussed approaches may potentially account for the
presence of $1/f$ scaling per se, the specific suggestion here
was to rephrase the question as to which of the approaches
accounts best for the so general linkage between observed fractal dynamics and coordination in the analyzed system.

Wagenmakers et al. have laid out an argument showing that
the presence of $1/f$ scaling (per se) should not be taken as evidence for self-organization and complexity. The argument refutes the enterprise of observing (some extent of)
$1/f$ scaling in some measure in some task and concluding,
based upon that observation, that the system under scrutiny is complex. The goal of the present paper was actually to step beyond a discussion about the presence versus nonpresence of genuine $1/f$ scaling, however, and to question instead
how and why it is that $1/f$ scaling changes so consistently
across different manipulations of task, condition, and level of
analysis. For example, why would cognition be short-range?
Why this exception in the context of fractal scaling at all other scales of the system?

In their respective approaches, Gilden and colleagues as
well as Delignières and colleagues, have focused specifically
on observed changes in the extent of $1/f$ scaling, including
changes that may emerge from nonlinearity in the power
spectra, but if cognitive components are independent and
domain-specific, why is fractal scaling evident throughout
the entire system?

Furthermore it remains unclear whether two-source models can include other complexity measures as well (e.g.,
[21, 23, 139, 140]).

While the perspectives presented by Wagenmakers and
colleagues and Delignières and colleagues are yet to account for the generality of the association between $1/f$ scaling and coordinated human activities, Gilden’s model is in fact much
more widely applicable and generalizable (see [121]). It is
unclear, however, whether the distinction between cognitive
$1/f$ noise and random motor noise remains tenable given the
thin line separating motor behaviors from cognitive behav-
iors (i.e., the radical embodiment thesis). Also the accom-
panying phenomenological account (hence, breakdown of
mental set or vigilance) might be to narrow to account for
the totality of presented changes in the extent of $1/f$ scaling.
The perspective of interaction-dominance, however, does
provide one general explanation for the reviewed studies, using principles of self-organized control.

Embracing interaction-dominance requires stepping beyond the traditional modular view of human cognition. If one considers the unexpectedness of the observations presented, however, based on a modular approach it is clear that the post hoc explanations required are massive. In addition, the modular defense has not gone any much further than simply expressing the opinion that studies that include 1/f scaling are too “limited, superficial, and overly general” [130, page 4] to be of any use, which simply ignores the empirical outcomes and the broadness of scales to which they apply. Under the current state of affairs, therefore, it seems unwarranted to categorize studies addressing the functional role of 1/f scaling in behavior as “mostly speculation, wrapped in jargon, inside wishful thinking” [130, page 5]. Quite to the contrary, although many questions remain to be answered and important challenges lie ahead, studying 1/f scaling and allied concepts of interdependence in human performances arguably has the potential to answer the tougher type of question about the scales and modes involved in coordinated human activities.

Conflict of Interests

The author declares that there is no conflict of interests regarding the publication of this paper.

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