

# Navigating Pubertal Goldilocks: The Optimal Pace for Hierarchical Brain Organization

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Adolescence is a timed process with an onset, tempo, and duration. Nevertheless, the temporal dimension, especially the pace of maturation, remains an insufficiently studied aspect of developmental progression. The primary objective is to estimate the precise influence of pubertal maturational tempo on the configuration of associative brain regions. To this end, the connection between maturational stages and the level of hierarchical organization of large-scale brain networks in 12-13-year-old females is analyzed. Skeletal maturity is used as a proxy for pubertal progress. The degree of maturity is defined by the difference between bone age and chronological age. To assess the level of hierarchical organization in the brain, the temporal dynamic of closed eye resting state high-density electroencephalography (EEG) in the alpha frequency range is analyzed. Different levels of hierarchical order are captured by the measured asymmetry in the directionality of information flow between different regions. The calculated EEG-based entropy production of participant groups is then compared with accelerated, average, and decelerated maturity. Results indicate that an average maturational trajectory optimally aligns with cerebral hierarchical order, and both accelerated and decelerated timelines result in diminished cortical organization. This suggests that a “Goldilocks rule” of brain development is favoring a particular maturational tempo.

## 1. Introduction

Similar to early childhood development, adolescence is a time-bound event with an onset, tempo, and duration. While the interaction between genetic and environmental factors is established as a fundamental contributor to diversities in adult behavioral consequences, the temporal aspect, specifically, the speed of maturation, remains an insufficiently explored feature of developmental progression.<sup>[1,2]</sup> In the current study, we focus on the alterations of hierarchical brain organization, related to different timings of pubertal maturation.

Adolescent cerebral maturation involves intricate sequences of neurodevelopment,<sup>[3,4]</sup> interwoven with alterations by age and stress.<sup>[5-7]</sup> Normative trajectories delineating volumetric<sup>[8]</sup> and synaptic density<sup>[9]</sup> shifts underscore the ongoing developmental dynamics of the adolescent brain. It is important to acknowledge, however, that the comprehensive volumetric transformations and mean synaptic density reflect the cumulative outcome

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of anatomical reconfiguration within the human brain, where the different components undergo heterochronous maturation.<sup>[10,11]</sup> Increasingly earlier onset of puberty referred to as worldwide secular trends,<sup>[12–14]</sup> or delayed onset due to, e.g., malnutrition or eating disorders<sup>[15,16]</sup> might have significant consequences on developmental trajectories, impacting a wide range of physiological and psychological processes. For instance, puberty-associated reductions in cortical thickness<sup>[17]</sup> or synaptic density<sup>[18]</sup> may manifest prematurely or belatedly, deterring the adaptability of regions maturing at later stages to the atypical developmental milieu. Premature or postponed puberty can also generate challenges in behavioral adjustment, potentially leading to social isolation and heightened anxiety, particularly in the absence of adequate coping mechanisms,<sup>[19]</sup> thereby emphasizing the critical role of adolescence in mental health.<sup>[8,20]</sup> Consequently, it is essential to investigate the impact of pubertal timing on the developing brain.

Uncertainty around the impact of atypical pubertal timelines arises from ineffective maturity measures and difficulty dissociating chronological age from pubertal maturity.<sup>[21,17,22–24]</sup> While puberty onset and its potential impact on developmental trajectories have long been a topic of interest, with significant individual variability suspected,<sup>[23,25,26]</sup> a precise assessment of the relationship between maturity and brain development remains noticeably absent from the literature, which is due to the lack of a reliable method to assess maturity levels. The Tanner Scale,<sup>[27–29]</sup> considered as the benchmark, relies on subjective evaluations of physical attributes such as breast and testicle size, rendering it susceptible to evaluator subjectivity. This scale originates from a post-war longitudinal study carried out within an orphanage from 1949 to 1971, thus failing to account for contemporary nutritional conditions and secular trends in growth.<sup>[13,28,30–32]</sup> Self- and parent-report versions of this method have also been deemed unreliable.<sup>[28,29,33]</sup>

Recently, we have introduced ultrasonic bone age as a promising alternative for maturity assessments in human developmental research,<sup>[34]</sup> with evidence for selective maturity-dependent effects in cognitive,<sup>[34,35]</sup> motor,<sup>[36]</sup> and emotional development.<sup>[37]</sup> Operating without harmful radiation, this technology gauges acoustic conductivity at hand and wrist growth zones to estimate bone age, demonstrating robust correlation with pubertal hormone levels,<sup>[28,33,38,39]</sup> X-ray estimations,<sup>[40]</sup> and displaying high

reliability.<sup>[40]</sup> This method overcomes traditional pubertal staging accuracy issues by providing an objective, continuous measure with unprecedented accuracy. In the current study, skeletal maturity serves as a proxy for individual pubertal maturity status. Addressing the impact of pubertal timing on brain development, the degree of maturity of each participant is calculated based on the difference between their bone age (BA) and chronological age (CA). It is widely acknowledged that the timing of puberty can differ significantly among individuals and follows a normal distribution,<sup>[41–44]</sup> therefore, to allow for statistical comparison across groups with different maturity levels, in our cross-sectional study, we selected an equal number of decelerated (BA < CA), average (BA = CA) and accelerated (BA > CA) maturity participants after screening them for bone age (**Figure 1a**, and please also consult Section Participants in Experimental Section for details). This approach not only disentangles chronological age from pubertal maturity (**Figure 1b**), but also provides insights into how adolescents with varying maturity levels navigate pubertal goldilocks.

Our primary objective is to investigate the precise influence of pubertal maturation on the configuration of associative brain regions, which manifest the most substantial developmental shifts during adolescence.<sup>[10,45,46]</sup> For this purpose, we assess resting-state brain activity and extract the degree of hierarchical arrangement among participant groups exhibiting diverse levels of physical maturation.

Recent progress in neuroimaging has enabled the creation of normative brain-growth charts,<sup>[8,47]</sup> similar to those applied for anthropometric attributes like height and weight. Despite their reliance solely on chronological age and their predominant cross-sectional nature, these investigations reveal the spatiotemporal dynamics of developmental plasticity and have even identified a sensorimotor-to-associative sequence of refinement.<sup>[10,11]</sup> The latter findings align with the progress of phylogenetic brain development from less variable unimodal areas toward heteromodal association cortices.<sup>[48–50]</sup> It is important to recognize that the most recently evolved associative networks within the brain might be particularly vulnerable to diverse pubertal timelines, given their ontogenetic maturation coincides more closely with the adolescent phase. Irregular timelines could also relate more to alterations in the finely tuned intrinsic activity of these networks, as opposed to extensive anatomical reconfigurations. While functional Magnetic Resonance Imaging (fMRI) has proven useful in assessing intrinsic resting-state activity, it is sensitive to neural, vascular, and respiratory influences. Therefore, direct measures of neural activity might offer better options for investigating this intrinsic behavior. Resting-state EEG provides direct measures of intrinsic cortical activity, allowing for the detection of changes in oscillatory activity and functional connectivity that are not captured by fMRI. In this study, we rely on resting-state EEG in the alpha frequency range (see Section Resting State Data Recording and Section Data Analysis in Experimental Section) to compare the progression of hierarchical brain organization across groups with early, on-time, and late maturing groups as defined by the difference between their bone age and chronological age.

To reveal any relationships between pubertal pace and the advancement of brain organization, we analyze how skeletal age-based maturity levels link to high-level features of resting-state

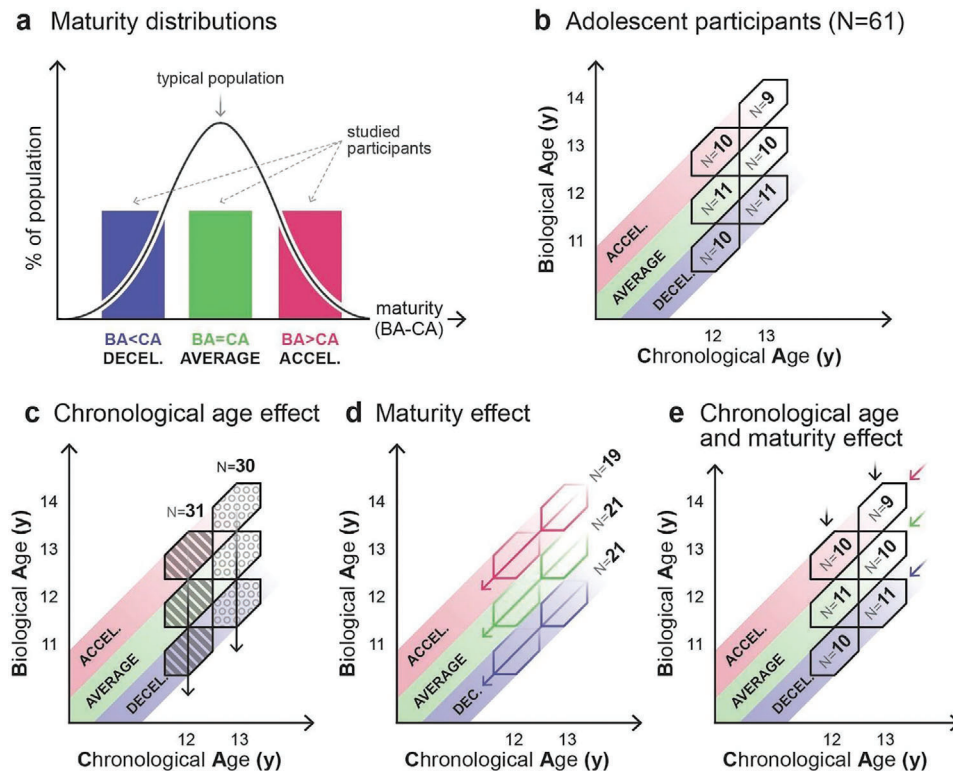
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**Figure 1.** Participant grouping by adolescent age and maturity. a) The black curve illustrates the normal distribution of diverse maturity levels in a standard population, where maturity is represented along the horizontal axis as the discrepancy between biological age (BA) and chronological age (CA). Positive differences indicate accelerated, while negative differences indicate decelerated maturity. In the present study, to explore the influence of varying maturity levels, we initially assessed bone age using an ultrasonic device to estimate the biological age of participants. We then selected an approximately equal number of participants with decelerated, average, and accelerated maturity levels. b) Dissociation of biological and chronological age is achieved through the assignment of participants into one-year-wide hexagonal bins which ensures the absence of overlap between maturity groups. The data were then analyzed through various grouping approaches: c) The effect of chronological age is analyzed by averaging across different maturity groups within specific age groups; d) The effect of maturity is analyzed by averaging across age groups within distinct maturity groups; e) By comparing data at the bin-level, the interplay between age and maturity effects is revealed.

EEG-dynamics. Specifically, we introduce “entropy production” to capture the degree of functional hierarchical organization of the brain. In thermodynamics and systems biology, the asymmetry and directionality of flow in the state space of the components of a living system is known as the “breaking of the detailed balance” and gives rise to non-equilibrium states which can be captured by the level of non-reversibility in time.<sup>[51,52]</sup> In other words, different levels of hierarchical organization allow the orchestration of the whole-brain dynamics accordingly in different ways. Hierarchy is built up by the underlying asymmetry in the directionality of information flow between different regions. Thermodynamics establishes a direct link between hierarchy, non-equilibrium, and time irreversibility. This comes from the fundamental idea of the second law of thermodynamics, explicitly stating that a system will go from order to disorder over time. We capture these different levels of organization and non-equilibrium through the level of irreversibility, i.e., through the “arrow of time.”<sup>[51,53]</sup>

Asymmetric flow can be observed in state spaces at various scales, from the molecular<sup>[54]</sup> to the macroscopic brain dynamics.<sup>[52]</sup> In general, asymmetric flow reflects a degree of non-reversibility in the dynamics of the system. In the case of

the brain, such non-reversibility is thought to reflect the hierarchical organization of brain activity.<sup>[51,55]</sup> Entropy production is a direct measure of non-reversibility and indirectly captures the functional hierarchy expressed by brain activity<sup>[51,53,56]</sup> (see Section Data analysis in Experimental Section). The validity of this approach using combined resting-state EEG – fMRI recordings has been recently confirmed on adult human patients with a neurodegenerative disorder.<sup>[57]</sup> The study demonstrated that compromised brain networks show a breakdown of temporal irreversibility which is accompanied by cognitive decline. An out-of-sample validation procedure additionally confirms these results.<sup>[57]</sup> The fact that broken detailed balance can characterize large-scale brain dynamics as it is related to function has been shown in a study discovering entropy production signatures of different levels of conscious awareness.<sup>[53]</sup> In terms of anatomical considerations, recent studies converge on finding increased entropy production in the default mode network (DMN) of the brain being active during resting states.<sup>[53,55,57]</sup>

The purpose of our study is to investigate the relationship between entropy production, as a proxy of hierarchical brain activity, and the level of physical maturation. We hypothesize that during

the development and maturation of the brain, there is a surge in hierarchical organization which will be reflected by the measured level of entropy production within chronological and maturational age groups.

## 2. Results

As explained in Figure 1b, we dissociated biological and chronological age through the assignment of participants into one-year-wide hexagonal bins to ensure the absence of overlap between maturity groups with approximately equal numbers of participants. We then analyzed the data through various grouping approaches illustrated in Figure 1c,d, allowing for the analysis of chronological age and maturity levels independently. Results are discussed according to those groupings. We have obtained and analyzed closed-eye resting state high-density EEG recordings in the alpha frequency range (8–12 Hz) in 61 adolescent, and 26 emerging adult female participants (see Figure 1 and Experimental Section/Participants). To measure the level of hierarchical brain activity in resting-state EEG recordings, we parsed the recorded neural data into 12 topographically distinct, phase-based activity patterns, referred to as “brain states” (see Figure 6), utilizing a modified *k*-means clustering algorithm.<sup>[58]</sup> Each brain state captures patterns where two major electrode sets record congruent signals, but these sets are in opposition to one another. After extracting the brain state patterns, we studied the sequence in which these brain states occurred, specifically focusing on the asymmetrical nature of their transitions. A sequence of consecutive brain states is illustrated in Figure 7a as a representative example in Section Data analysis in Experimental Section, while Figure 7b presents a table that counts the observed occurrences of each transition within this example, highlighting their asymmetric quality.

### 2.1. Chronological Age-Related Entropy Production

To evaluate the effect of chronological age on entropy production, we averaged across maturity groups within a 12-year-old ( $N = 31$ , mean age = 12.48 y,  $SD = 0.27$  y) and a 13-year-old ( $N = 30$ , mean age = 13.58 y,  $SD = 0.29$  y) group. In order to have normative data from young adults, we also included 22-year-old participants ( $N = 26$ , mean age = 21.69 y,  $SD = 0.64$  y). For further details see Figure 1c, Figure 2c, and Section Participants in Experimental Section. Importantly, adult data should only be compared to age-averaged results of adolescents as the adult cohort has not been selected with respect to maturity. Given that bone formation is completed by adulthood, bone age assessment is irrelevant in adults. Therefore, adult data were included in the statistical analysis only where the basis of the analysis is chronological age, irrespective of maturity.

Figures 2a and 3b provide a detailed look at brain state dynamics within each age-group. Logarithmic irreversibility transition matrices in Figure 2a illustrate a contribution to entropy production of brain state transitions in age groups, derived from the log ratio of the forward and reverse transition probabilities, estimated from transition counts. Matrices were generated by averaging individual matrices within each age group, followed by

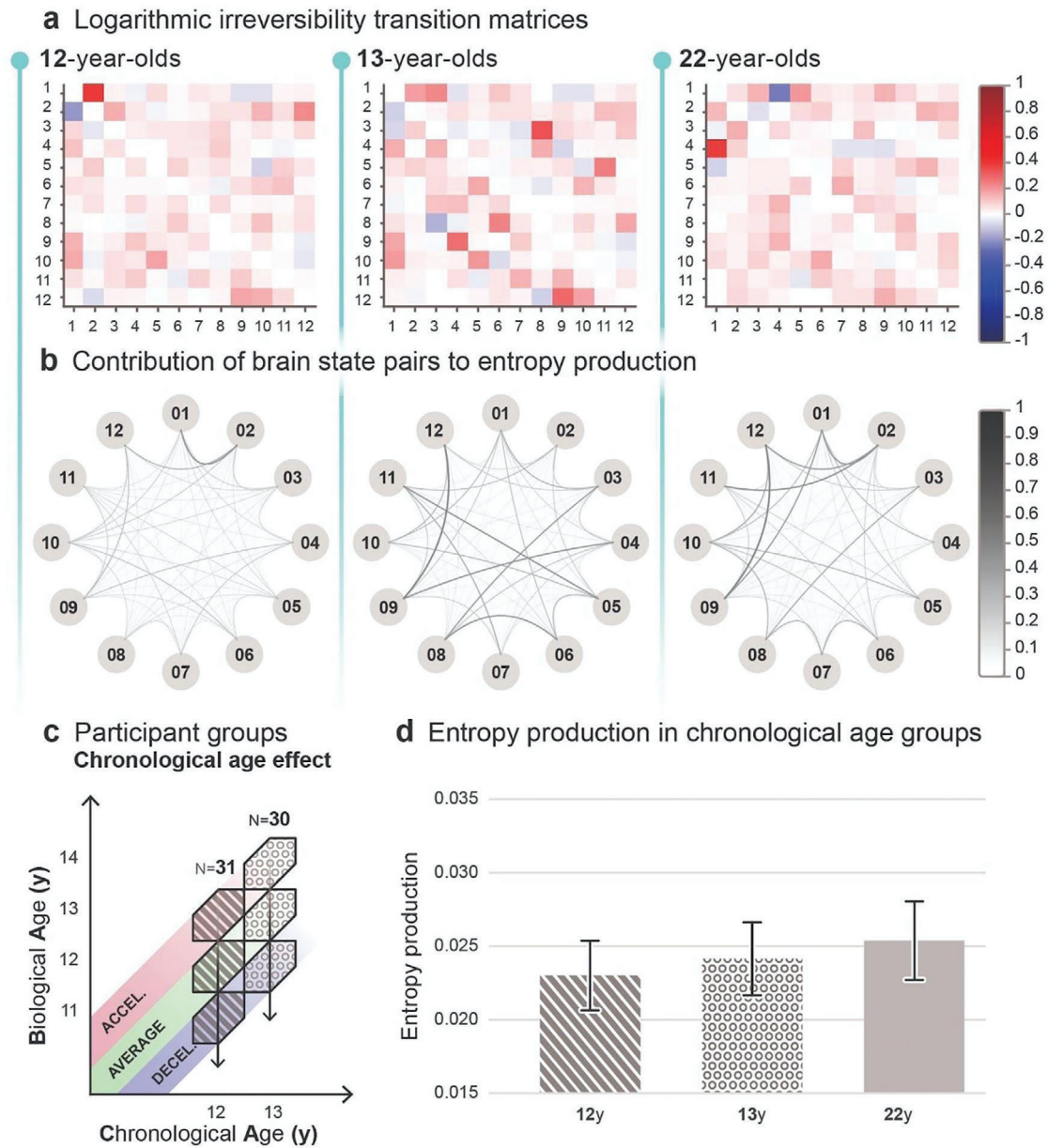
normalization of values to a  $-1$  to  $+1$  range (using the largest absolute value of the global extremum), indicated by the color scale. Each matrix holds  $12 \times 12$  values, with the starting brain states on the horizontal and the subsequent states in the brain state switching on the vertical axis. Log ratios of the forward and reverse transition probabilities are non-symmetric. Positive and negative values indicate a preference for forward and reverse transitions, respectively. The relative similarity in the variability in hue and intensity indicates comparable entropy production between age groups, with a slight indication of age-related improvement. Graphs in Figure 2b show the total entropy production contributions of brain state pairs, computed by summing the log ratio values of forward and reverse transition probabilities within each pair, creating symmetric matrices. Graph values are generated by averaging the individual symmetric matrices within age groups, and then normalizing them between 0 and 1. The nodes represent the 12 brain states, with the most dominant state (highest percentage of total activity) positioned at the 12 o'clock position, and subsequent states are arranged clockwise in order of descending dominance. The edges indicate pairwise contributions to entropy production, where increasing thickness and darkness represent greater contribution. The slight increase in line thickness and darkness across age-groups may indicate a subtle rise in contributions to entropy production with age.

Group averages and variances of entropy production were computed via inverse-variance weighting to minimize the biasing effect on the weighted arithmetic mean introduced by variance. Ninety-five percent confidence intervals of group-level entropy production were also calculated using the weighted group average. Weighted group averages revealed a tendency to increase entropy production with age, as it is demonstrated in Figure 2d. Specifically, the 12-year-old group exhibited an average mean entropy production of 0.02300 bit (95% CI: 0.02064–0.02537), while the average mean entropy production of the 13-year-old group was 0.02415 bit (95% CI: 0.02166–0.02663), and the average mean entropy production of the 22-year-old group was 0.02538 bit (95% CI: 0.02271–0.02805). To determine whether the group differences are statistically significant, we utilized two-sample *z*-tests to compare the group averages (see Section Statistics in Experimental Section for further details). While a slight increase in groupwise average entropy production with chronological age is observed, pairwise comparison of group averages was not statistically significant ( $p > 0.05$  in each comparison). Hedges' *g* effect size estimates revealed small effect sizes in all cases: between the 12-year-old and 13-year-old ( $g = 0.17$ ), and between 13-year-old and 22-year-old ( $g = 0.18$ ) groups, and in the comparison of the 12-year-old and 22-year-old group ( $g = 0.35$ ).

Our statistical results indicate that, while there is a slight increase in entropy production with increasing chronological age, this variable independently does not have a substantial impact on entropy production, further warranting the inspection of maturity in adolescence.

### 2.2. Maturity-Related Entropy Production

To analyze the relationship between maturity and group-level average entropy production, we sorted adolescent participants according to their level of maturity, irrespective of

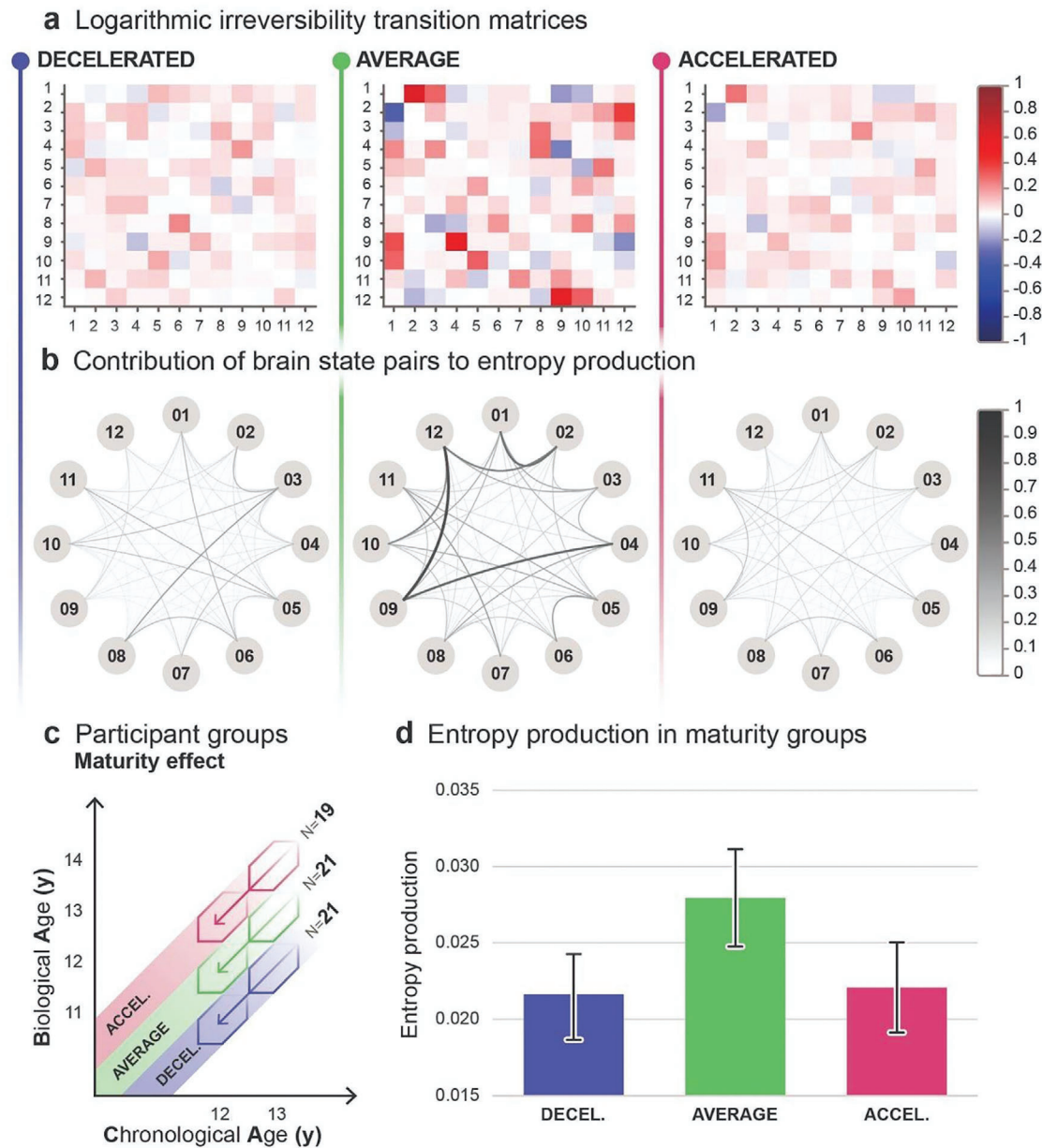


**Figure 2.** Chronological age-related entropy production. a) Logarithmic irreversibility transition matrices illustrate two distinct, non-symmetric values in each brain state pair, derived from the log ratio of the forward and reverse transition probabilities. The matrices inform about the preference for directionality in brain state switching. The extent of deviation from 0 reflects the strength of directional preference (forward transitions in red, backward transitions in blue), indicating the level of irreversibility and hierarchical organization in the brain. The relative similarity in the variability in hue and intensity indicates comparable entropy production between age groups. b) Graphs represent the cumulative contribution to entropy production of both forward and reverse transitions and reveal the level of irreversibility in the dynamics of each brain state pair within age groups. Nodes represent the 12 brain states, and edges indicate pairwise contributions to entropy production, where increasing thickness and darkness represent greater contribution and, therefore larger irreversibility. There seems to be a subtle rise in contributions to entropy production with age. c) The effect of chronological age is analyzed by averaging across different maturity groups within specific age groups. d) Bar plots illustrate the average entropy production and respective 95% confidence interval for each age group calculated via inverse-variance weighting. Differences among groups were statistically assessed using two-sample z-tests, accompanied by Hedges' *g* effect size calculations, yielding no statistically significant difference among the age groups in pairwise comparisons ( $p > 0.05$ ) and small effect sizes in all comparisons.

chronological age, forming three distinct groups: decelerated maturity ( $CA-1.5 \leq BA < CA-0.5$ ;  $N = 21$ , mean age = 12.98y,  $SD = 0.60y$ , mean  $BA-CA = -0.92$ ,  $SD = 0.24$ ), average maturity ( $CA-0.5 \leq BA \leq CA+0.5$ ;  $N = 21$ , mean age = 12.97y,  $SD = 0.60y$ , mean  $BA-CA = 0.03$ ,  $SD = 0.29$ ), and accelerated matu-

riety ( $CA+1.5 \geq BA > CA+0.5$ ;  $N = 19$ , mean age = 13.12y,  $SD = 0.68y$ , mean  $BA-CA = 0.88$ ,  $SD = 0.31$ ). Participant groups are illustrated both in Figures 1d and 3c.

Figure 3a,b offers detailed insight into brain state dynamics within each maturity group. As it is described above, with



**Figure 3.** Maturity-related entropy production. a) Logarithmic irreversibility transition matrices illustrate two distinct, non-symmetric values in each brain state pair, derived from the log ratio of the forward and reverse transition probabilities. The matrices inform about the preference for directionality in brain state switching. The extent of deviation from 0 reflects the strength of directional preference (forward transitions in red, backward transitions in blue), indicating the level of irreversibility and hierarchical organization in the brain. The larger hue and intensity variation in the matrix of the average maturity group indicates higher entropy production relative to the decelerated and accelerated groups. b) Graphs represent the cumulative contribution to entropy production of both forward and reverse transitions and reveal the level of irreversibility in the dynamics of each brain state pair within maturity groups. Nodes represent the 12 brain states, edges indicate pairwise contributions to entropy production, where increasing thickness and darkness represent greater contribution and, therefore larger irreversibility. Graphs indicate higher entropy production in the average maturity group, apparent in the thicker and darker lines. c) The effect of maturity is analyzed by averaging across age groups within distinct maturity groups. d) Bar plots illustrate the average entropy production and respective 95% confidence interval for each maturity group, calculated via inverse-variance weighting. Using two-sample z-tests, we compared the average group to both decelerated ( $p = 0.0028$ ) and accelerated ( $p = 0.0082$ ) groups, observing significant differences in each comparison with large effect sizes (Hedges'  $g$ ).

respect to Figure 2a, logarithmic irreversibility transition matrices in Figure 3a illustrate a contribution to entropy production of brain state transitions, in this case within decelerated, average, and accelerated maturity groups among adolescent participants. The larger hue and intensity variation in the matrix

of the average maturity group indicates higher entropy production relative to the decelerated and accelerated groups. Graphs in Figure 3b represent the total contribution of both forward and reverse transitions to entropy production (see the description related to Figure 2b above), indicating higher entropy production

in the average maturity group, apparent in the thicker and darker edges of the graph compared to that of the decelerated and accelerated groups.

Using inverse-variance weighting, we calculated the group-level average and variance of entropy production to minimize the distorting effect of variance and determined the 95% confidence intervals based on the weighted group average. The inverse-variance weighted group average and confidence interval of the decelerated maturity (mean = 0.02146 bit, 95% CI: 0.01866–0.02426), average maturity (mean = 0.02794 bit, 95% CI: 0.02475–0.03113), and accelerated maturity (mean = 0.02208 bit, 95% CI: 0.01913–0.02503) groups are illustrated in Figure 3d. It is clear that the average maturity group expresses higher average entropy production than both accelerated and decelerated maturity groups. We utilized two-sample z-tests to determine the statistical significance of the apparent differences, reported with Hedges' *g* effect size tests. Results showed that the average maturity group expresses significantly higher Groupwise average entropy production than both accelerated ( $p = 0.0082$ ,  $g = 0.83$ ) and decelerated ( $p = 0.0028$ ,  $g = 0.92$ ) maturity groups. Effect sizes were large in both pairwise tests.

The average maturity group consistently expressed significantly higher ( $p < 0.05$ ) average entropy production than both decelerated and accelerated groups with additional clustering approaches, specifically, working with 6, 8, and 10 brain states and 12 brain states that are specific to each unique group; see Section Extended Analyses under Supporting Information for further details.

Our findings show that maturity significantly influences entropy production, irrespective of chronological age, with the group-level entropy production being significantly higher in the average maturity group.

### 2.3. Chronological Age and Maturity-Related Entropy Production

To study the interplaying effect of chronological age and maturity, we divided adolescents into two age brackets, namely 12-year-olds ( $N = 31$ , mean age = 12.48 y,  $SD = 0.27$  y) and 13-year-olds ( $N = 30$ , mean age = 13.58 y,  $SD = 0.29$  y), and within these brackets, we further assorted participants into three distinct maturity groups, forming six groups, referred to as bins, in total.

In the 12-year-old group, we identified three maturity subgroups: decelerated maturity ( $CA-1.5 \leq BA < CA-0.5$ ;  $N = 10$ , mean age = 12.41 y,  $SD = 0.26$  y, mean  $BA-CA = -0.86$ ,  $SD = 0.21$ ); average maturity ( $CA-0.5 \leq BA \leq CA+0.5$ ;  $N = 11$ , mean age = 12.48 y,  $SD = 0.28$  y, mean  $BA-CA = 0.04$ ,  $SD = 0.31$ ); and accelerated maturity ( $CA+1.5 \geq BA > CA+0.5$ ;  $N = 10$ , mean age = 12.54 y,  $SD = 0.27$  y, mean  $BA-CA = 0.91$ ,  $SD = 0.34$ ). Similarly, in the 13-year-old group, we established three maturity subgroups: decelerated maturity ( $CA-1.5 \leq BA < CA-0.5$ ;  $N = 11$ , mean age = 13.49 y,  $SD = 0.24$  y, mean  $BA-CA = -0.97$ ,  $SD = 0.27$ ); average maturity ( $CA-0.5 \leq BA \leq CA+0.5$ ;  $N = 10$ , mean age = 13.52 y,  $SD = 0.31$  y, mean  $BA-CA = 0.03$ ,  $SD = 0.28$ ); and accelerated maturity ( $CA+1.5 \geq BA > CA+0.5$ ;  $N = 9$ , mean age = 13.76 y,  $SD = 0.26$  y, mean  $BA-CA = 0.85$ ,  $SD = 0.28$ ). Participant bins are illustrated both in Figures 1e and 4c

Figure 4a,b offers detailed insight into brain state dynamics within each bin. As described above, with respect to Figure 2a,

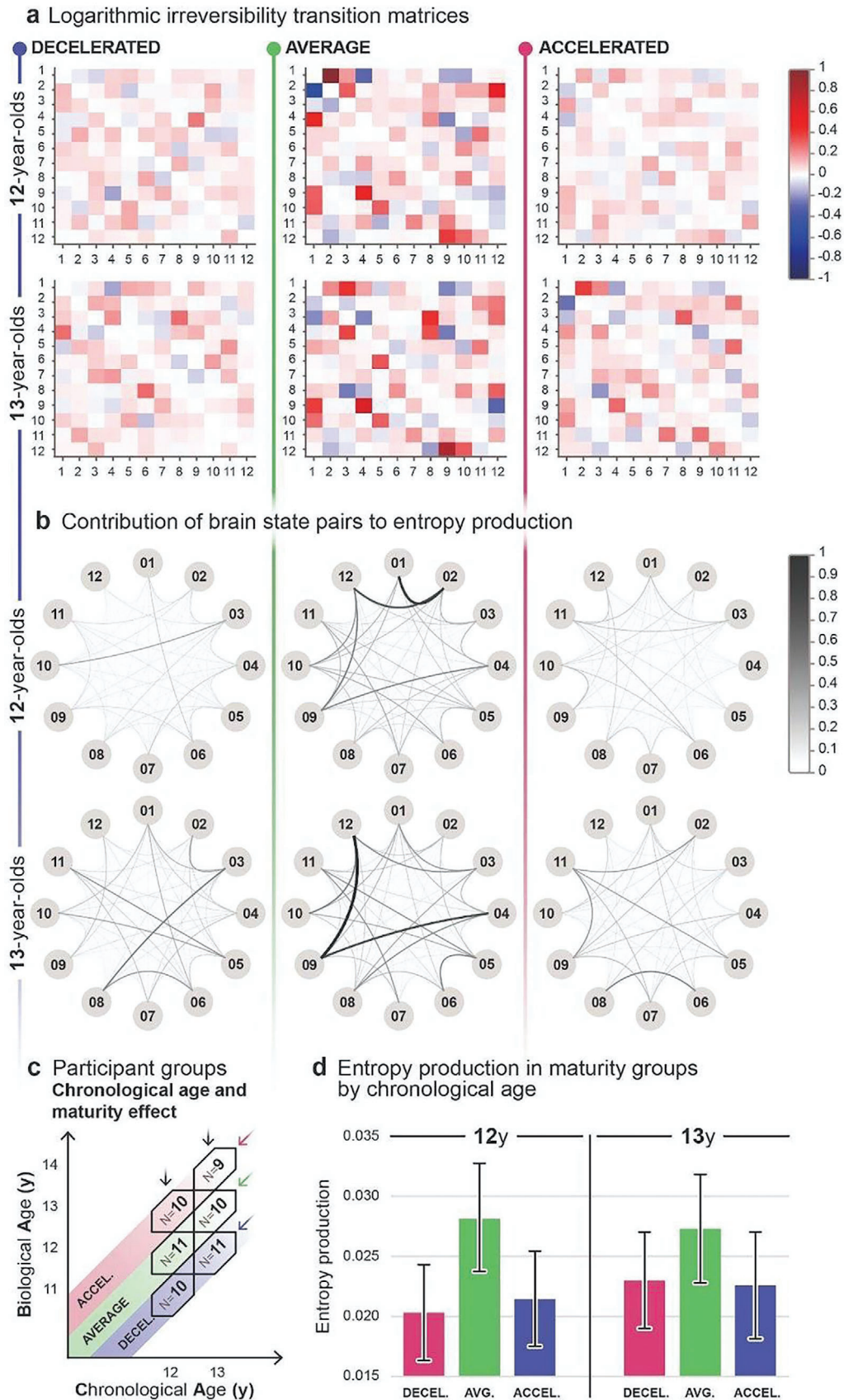
logarithmic irreversibility transition matrices in Figure 4a illustrate a contribution to entropy production of brain state transitions, in this case within decelerated, average, and accelerated maturity groups within age groups. In both age brackets, the larger hue and intensity variation in the matrix of the average maturity group indicates higher entropy production relative to the decelerated and accelerated groups. Graphs in Figure 4b represent the total contribution of both forward and reverse transitions to entropy production (see the description related to Figure 2b above), indicating higher entropy production in the average maturity group in both age groups, apparent in the thicker and darker edges of the graph compared to that of the decelerated and accelerated groups.

We utilized inverse-variance weighting to calculate group averages and variances of entropy production, minimizing the biasing effect that variance introduces to the weighted arithmetic mean. Additionally, we determined the 95% confidence intervals for group-level entropy production based on the weighted group average. As it is suggested in Figure 4d, the average maturity group expresses higher average entropy production at the group-level than decelerated and accelerated groups in the same chronological age category. We performed two-sample z-tests to determine the statistical significance of differences. Two-sample z-tests, accompanied by Hedges' *g* effect size estimations, confirmed statistically significant differences among the 12-year-old maturity bins (see Figure 4d); we compared the average entropy production of the average maturity (mean = 0.02823 bit, 95% CI: 0.02374–0.03271) bin to both decelerated (mean = 0.02032 bit, 95% CI: 0.01633–0.02431;  $p = 0.0098$ ,  $g = 1.120$ ) and accelerated (mean = 0.02145 bit, 95% CI: 0.01750–0.02540;  $p = 0.0261$ ,  $g = 0.963$ ) bins, revealing significant differences and large effect sizes in these comparisons. Performing two-sample z-tests on the maturity bins of the 13-year-olds did not yield a significant result ( $p > 0.05$  in all comparisons). Effect size measurements revealed a medium effect size ( $g = 0.61$ ) between the average maturity (mean entropy production = 0.02728 bit, 95% CI: 0.02277–0.03179) and decelerated (mean entropy production = 0.02299 bit, 95% CI: 0.01899–0.02700) bins, and large effect size in the comparison of average versus accelerated (mean entropy production = 0.02257 bit, 95% CI: 0.01813–0.02701;  $g = 0.67$ ) bins.

Our results show that maturity strongly influences entropy production among 12-year-old participants. While we did not find significant differences between the maturity groups in the 13-year-old age bracket, the higher group-level entropy production in the average maturity group is still present in the logarithmic irreversibility transition matrices (Figure 4a), in the graphs of Figure 4b, and in the effect size calculations, indicating a persistent pattern of increased entropy production associated with average maturity.

## 3. Discussion

A novel question was addressed regarding the role of adolescent maturation in brain development. While individual differences in maturity among teenagers are well-established, it has been unclear whether there is a difference in brain development between groups of adolescents with diverse maturity levels. The introduction of skeletal age into research now allows for conclusive answers to this question in a field obscured by





methodology issues around maturity assessments. Our results suggest that an average maturational trajectory might be optimal in terms of hierarchical brain organization. The significance of this study stems from the possibility that developmental plasticity may not fully compensate for accelerated or decelerated timelines, and a heightened risk of behavioral problems and psychiatric disorders may arise from such alterations.

To address how well adolescents with different maturational paces can navigate pubertal goldilocks, we relied on skeletal age as a proxy for pubertal progress and resting-state EEG-derived entropy production as a gauge of hierarchical brain organization. As illustrated in Figure 1, we employed the strategy of selecting an equal number of participants with accelerated, average, and decelerated maturity levels. We have also completed our sample with a group of emerging adult participants who were not selected by maturity levels since bone age assessments are not informative at this age. The dissociation between biological age and chronological age illustrated in Figure 1b allowed us to independently analyze the impact of age and maturity on the extent of resting-state EEG-derived entropy production.

First, we analyzed the relationship between chronological age and entropy production by averaging across the three maturity groups within the 12 and 13-year-old participant groups (Figures 1c and 2c) and comparing them to the emerging adult group. Despite a noticeable increasing trend in entropy production with age (Figure 2d), chronological age in itself did not have a statistically significant effect on entropy production. The similar logarithmic irreversibility transition matrices of the age-groups (Figure 2a), and the relatively similar dispersion of entropy production among brain state pairs (Figure 2b) support the statistical results. These findings suggest that the level of hierarchy in the adolescent brain may not be meaningfully influenced by chronological age alone. It is important to note that we lack information about the adolescent maturational trajectories of emerging adult participants, while adolescents have been screened and selected for the maturity groups in approximately equal numbers. Therefore, the distributions of individuals with different maturity levels might be different between emerging adults and adolescents (Figure 1a), and their direct comparison should be approached with caution.

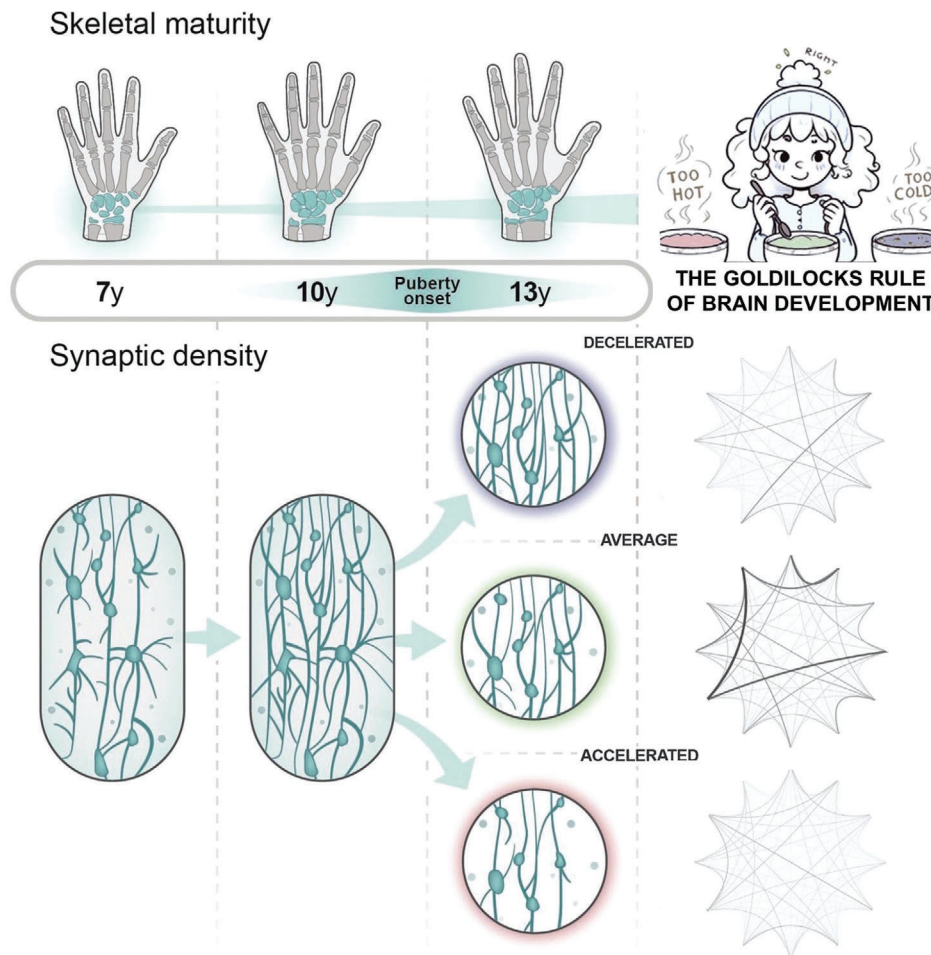
To investigate the influence of maturity on entropy production, we categorized adolescent participants into three distinct groups based on their level of maturity (decelerated, average, and accelerated maturity groups in Figures 1d and 3c.). Statistical analysis of group averages (Figure 3d) revealed significantly higher entropy production in the average maturity group with large effect sizes,

pointing to a considerably higher level of hierarchical organization when chronological age and bone age align. A closer look at the logarithmic irreversibility transition matrices (Figure 3a) illustrating the strength of directional preference in switching among the 12 states provides further insight into the role of maturity in hierarchical organization. The average maturity group has a relatively higher number of brain state pairs among which a strong directionality is established. The contribution of brain state pairs also reflects an elevated level of hierarchy in the average maturity group (Figure 3b). The dynamics between brain states underlying the entropy production levels in each group indicate well-established causal interactions, and hierarchical brain organization in the average maturity group, while these relationships seem relatively less dense in the decelerated and accelerated groups. These findings indicate that physical maturity is inherently linked to the shaping of irreversible, causal interactions within the cortex<sup>[51]</sup> during adolescence. Furthermore, it suggests that the optimal maturational pace for fostering these relationships is one in which chronological and biological age are closely matched.

To see whether the maturity-related effect is stable across different chronological ages, we compared the three maturity groups within the 12- and 13-year-old age groups (Figures 1d and 4c). The analysis revealed that the average maturity group displays significantly higher entropy production compared to both the decelerated and accelerated maturity groups among the 12-year-olds (Figure 4d). While this tendency is still apparent in the 13-year-old group, statistically significant differences between maturity groups were not detected. However, the logarithmic irreversibility transition matrices of each bin (Figure 4a) make a case for the optimal pace of maturation even in the absence of statistical significance, as the average maturity groups in both age brackets convey a larger number of brain state pairs where a strong preference of switching direction is established. The contribution of each brain state pair to entropy production (Figure 4b) also highlights a larger number of well-established irreversible interactions in the average maturity groups. Taken together, the statistical results (Figure 4d) may suggest that the clear advantage of an on-time maturational pace is limited to a specific chronological age window with a declining tendency in older age groups, however, this possibility may need further investigation as the graphical analyses presented in Figure 4a,b suggests a persistent pattern of increased entropy production associated with average maturity.

The difference in entropy production between the three different maturity groups, and the increased entropy in the average

**Figure 4.** Chronological age and maturity-related entropy production. a) Logarithmic irreversibility transition matrices illustrate two distinct, non-symmetric values in each brain state pair, derived from the log ratio of the forward and reverse transition probabilities. The matrices inform about the preference for directionality in brain state switching. The extent of deviation from 0 reflects the strength of directional preference (forward transitions in red, backward transitions in blue), indicating a higher level of irreversibility and hierarchical organization in the brain. The hue and intensity diversity in the matrices in the average maturity groups across both age groups indicates higher entropy production values compared to decelerated and accelerated maturity groups. b) Graphs represent the cumulative contribution to entropy production of both forward and reverse transitions and reveal the level of irreversibility in the dynamics of each brain state pair within age groups. Nodes represent the 12 brain states, edges indicate pairwise contributions to entropy production, where increasing thickness and darkness represent greater contribution and, therefore larger irreversibility. Graphs indicate higher entropy production in both average maturity groups, apparent in the thicker and darker lines. c) The effect of maturity is analyzed by averaging across age groups within distinct maturity groups. d) Bar plots illustrate the average entropy production and respective 95% confidence interval for each maturity group, calculated via inverse-variance weighting. Using two-sample z-tests, we compared the average group to both decelerated ( $p = 0.0098$ ) and accelerated ( $p = 0.0261$ ) groups, observing significant differences in each comparison with large effect sizes (Hedges'  $g$ ) in 12-year-olds, however, the analysis of 13-year-olds did not yield significant results.



**Figure 5.** Summary of findings. To reveal the relationship between pubertal pace defined by skeletal maturity and the advancement of brain organization, the connection between skeletal age-based maturation stages and hierarchical organization in the temporal dynamics of resting-state EEG recordings was analyzed. Not only skeletal growth but brain development is also profoundly influenced by the dynamic shifts in pubertal hormone levels. Puberty onset initiates synaptic pruning (reduction in dendritic spine density) and pruning then continues until adulthood throughout the nervous system. However, as the conceptual figure suggests, synaptic pruning may not adhere to its expected course, but it may undergo dysregulation in cases of accelerated or decelerated pubertal maturation. Untimely maturation with dysregulated synaptic pruning (under- or over-pruning) may lead to the emergence of alterations in hierarchical brain organization (illustrated by the graphs in the right column – please see the caption for Figure 3b). These findings indicate that an average maturational trajectory optimally aligns with cerebral hierarchical order and both accelerated and decelerated timelines result in a reduction of cortical organization. This suggests that a Goldilocks rule of brain development might exist, favoring a particular maturational tempo.

maturity group in particular raises the issue of the functional relevance of these results. As mentioned earlier, increased entropy may reflect heightened hierarchical organization during intrinsic brain activity within the resting state network. In terms of behavioral consequences, modern theories of consciousness might assist interpretation. The Temporo-spatial Theory of Consciousness (TTC) of Northoff,<sup>[59–61]</sup> for example, brings the intrinsic or spontaneous activity of the brain into focus and assumes that the temporal dynamic and spatial topography of neural activity constitute the “common currency” of neural and mental states. In this context, intrinsic activity prepares the organism to interact with external stimuli, which means that a higher level of organization during intrinsic activity will lead to a more adaptive interaction. One may also speculate that an atypical shift in the spatial-temporal pattern of intrinsic activity – such as in the case of accelerated or decelerated development – may affect conscious states

such as metacognitive functions.<sup>[62]</sup> Currently, it is not known whether adaptive developmental plasticity fully compensates for accelerated or decelerated timelines, or increases the risk of behavioral problems and psychiatric disorders, therefore it is important to further investigate these issues.

The refinement of brain structure and bone development are both part of the anatomical changes associated with puberty and adolescence as illustrated in Figure 5. One of the significant maturational events during this phase is the targeted elimination of synapses, known as synaptic pruning<sup>[45,63–65]</sup> (Figure 5), which occurs in various species<sup>[66,67]</sup> and across the nervous system<sup>[45]</sup> during development to adapt to environmental conditions.<sup>[65,67]</sup> This process is vital for optimal brain function and shaping a healthy adult brain.<sup>[64,68]</sup> The timing of synaptic pruning has been linked to puberty onset in rodents<sup>[69,70]</sup> and humans,<sup>[18]</sup> suggesting that early or late puberty may lead to a correspondingly early

or delayed initiation of synaptic pruning. However, it is not clear whether synaptic pruning follows a normal course or becomes dysregulated when it begins earlier or later (Figure 5). As insufficient synaptic pruning or “underpruning” has been tied to neurodevelopmental disorders<sup>[4,22]</sup> and excessive pruning or “overpruning” to psychiatric conditions such as schizophrenia,<sup>[71,72]</sup> it is important to ask whether untimely maturation may lead to the emergence of alterations in brain organization, thereby contributing to the current psychological challenges faced by adolescents.<sup>[73–77]</sup> The temporal aspect of synaptic pruning holds significance within hierarchical neurodevelopment as well. Typically, transmodal association regions undergo peak synaptic density later than sensory zones, experiencing pruning throughout adolescence.<sup>[11,45,46]</sup> This design permits a broad experience-dependent refinement window but also introduces the prospect of prolonged vulnerability.<sup>[11,78]</sup> Untimely maturation with dysregulated synaptic pruning (under- or over-pruning) may lead to the emergence of alterations in hierarchical brain organization and may provide the anatomical basis of our findings.

In terms of limitations, the current project is confined to the examination of adolescent females, a limitation that might be apparent. However, a selection was essential when aiming to distinctly disentangle the impacts of maturation and chronological age on brain development. Involving both sexes would have made it difficult to carry out the detailed analysis within the narrow temporal windows of biological and chronological age depicted in Figure 1b. As male pubertal onset occurs  $\approx 1\text{--}1.5$  years later than that of females,<sup>[32,79]</sup> this analysis would necessitate a considerably larger sample size. We opted for the inclusion of females since menarche age provides valuable supplementary data affirming the accuracy of bone age assessments (see Section Procedure in Experimental Section). Given the conclusive findings with females, underscoring a substantial role of biological age in brain development, it is plausible to assume a comparable effect might manifest in males. Indeed, we would expect that maturity is a relevant factor in the general brain development of males as well; however, owing to the distinct gonadal hormone levels governing female and male developmental trajectories,<sup>[21,80]</sup> disparities are also anticipated, including the timing of the impact of maturity levels.

An additional limitation of our study pertains to its cross-sectional design within a relatively narrow age spectrum, which prevents exploration of potential long-term consequences due to variations in pubertal pace. The evolutionarily expanded heteromodal association cortices, mentioned in the introduction, hosting a range of higher-order functions<sup>[11,48–50,81]</sup> may have the potential to compensate when confronted with divergent maturational paces. Nevertheless, the consequences of accelerated or delayed development on relevant and precisely timed adolescent brain alterations such as cortical volume reduction,<sup>[82,83]</sup> gray matter thinning,<sup>[17,84]</sup> and synaptic pruning<sup>[18,45]</sup> remain unknown. The human brain could conceivably adapt to distinct maturational tempos, and with only transient disparities, developmental trajectories might still converge within the normative range. While the distinctions in hierarchical brain organization uncovered in our study may be transient, even these potentially short-term developmental discrepancies are essential to discuss since they could temporarily place the child outside the typical range and cause heightened stress levels. Subsequent investiga-

tions must undoubtedly address the longitudinal trajectories of brain development within cohorts characterized by diverse rates of maturation. It needs to be considered, however, that any such study faces a serious trade-off: observing long-term alterations due to accelerated puberty onset is more likely with age, but direct assessment of pubertal timing becomes less reliable with age. Although a long-term follow-up between puberty onset and adulthood seems necessary to complement our work, in the current study we have chosen to work with a cross-sectional sample to get a first glimpse of the impact of different maturational tempos on brain development.

## 4. Conclusion

To conclude, we would like to re-emphasize that maturation plays a pivotal role in shaping both brain structure and behavior. This process ensures that the brain efficiently coordinates distributed computations across its entirety, allowing not only survival but also optimal functioning. This efficiency hinges on a hierarchical orchestration of state-dependent, self-organized brain activity over time. This orchestration facilitates near-optimal information processing and transfer while minimizing energy consumption. Therefore, a paramount concern of a comprehensive theory of brain function is the exploration of hierarchy. This entails unraveling the dynamic and evolving self-organization of brain states, spanning from stability to transitions, as they navigate through probabilistic and sometimes chaotic shifts. Here, we assumed that the inherent hierarchy of spontaneous brain activity might be reflective of the level of maturation. Harnessing the principles of thermodynamics, we quantified hierarchy by computing the level of irreversibility in brain signals during resting state conditions. Specifically, we captured the level of irreversibility through entropy production<sup>[51,53,56,57,85]</sup> as computed through a sophisticated clustering strategy. Note that resting states cover a rich range of the dynamical repertoire of the brain and thus those are ideal to characterize the most general and wide variety of interaction across the whole brain, i.e., the underlying hierarchical orchestration.

With respect to the broader relevance of our findings, although the participants enrolled in the present study were deliberately chosen to fall within a normative range of maturational speed and were further screened for favorable socioeconomic circumstances, as well as the absence of developmental or neurological disorders, the findings bear more general implications, especially for clinical domains. We found an advantage of on-time maturation with respect to hierarchical brain organization within our non-clinical cohort, indicating that significant deviations toward either accelerated or delayed maturational speeds might amplify disparities, leading developmental trajectories into the clinical spectrum. The acceleration of pubertal development is a growing concern today, particularly given the surge in medical referrals for young girls experiencing precocious puberty globally, with some instances manifesting as early as 6 or 7 years old.<sup>[86–97]</sup> Premature puberty carries the risk of compromised adult stature, psychosocial challenges, and potential health complications later in life.<sup>[44,98,99]</sup> The sudden peak in the lower left segment of the puberty onset age distribution curve (akin to that depicted in Figure 1a), corresponding to instances of early puberty onset, suggests that there may be significant implications for the

broader population involving a rapid secular trend. The possibility that developmental plasticity may not fully compensate for these changes is highly relevant. Secular trends in pubertal onset age persist across both developing and modern societies. While advancements in living conditions primarily account for trends in developing countries today,<sup>[13–15,100,101]</sup> negative anthropogenic influences, such as endocrine-disrupting chemicals, have shaped the trend in developed nations in the last decades.<sup>[100]</sup> The recent pandemic has emerged as a novel potential factor influencing the age of pubertal onset, and it is essential to further investigate its impact on the general population.

In present times, within the most affluent nations, important developmental events, such as birth and puberty can be artificially timed, inducing intraspecies heterochrony which denotes variations in the timing of developmental events. Genetic-induced heterochrony introduces natural variability throughout evolution.<sup>[102–104]</sup> It remains uncertain whether anthropogenic-induced heterochrony (e.g., accelerated trends in pubertal timing, or artificially delayed puberty) will yield compromised mental functioning and health, or foster adaptive diversity in successive generations. Therefore, our study seems to be very timely, and can be considered as a first step toward investigating the potential outcomes of “anthropogenic heterochrony”. Mental health problems have surged following successive lockdowns, demonstrating the psychological toll on adolescents worldwide.<sup>[74–77]</sup> The findings of our study will aid in devising interventions more attuned to developmental contexts, thus lowering the risk of developing disabling psychopathologies in future generations.

## 5. Experimental Section

**Participants:** The study is limited to investigating adolescent and emerging adult females, a deliberate choice made to disentangle the impacts of maturation and chronological age on brain development. The inclusion of both genders would have hindered in-depth analysis within the narrow temporal windows of biological and chronological age depicted in Figure 1b. Given the delayed male pubertal onset,<sup>[32,79]</sup> a significantly larger sample would be required. The inclusion of females since menarche age that provided valuable supplementary data affirming the accuracy of bone age assessments in adolescents was opted for.

Participants were recruited via school contacts and online advertisements from Budapest, Hungary. Parents and/or participants provided demographic, educational, and medical history details. One participant with attention deficit disorder was excluded.

In order to determine the optimal sample size, power calculations for a two-sample z-test for means were performed. A sample size of  $N \geq 25$  was found to be required for a power of at least 0.8 to detect a 0.8 effect at  $p < 0.05$ . A sample size of  $N = 30$  in the adolescent and emerging adult chronological age groups was aimed for. This yielded 20 subjects in the adolescent maturity groups with a power of at least 0.7. Ultrasonic bone age screening was conducted, and adolescent participants were categorized into BA/CA-defined bins, ensuring non-overlapping maturity groups: average ( $CA - 0.5 \leq BA \leq CA + 0.5$ ), accelerated ( $CA + 1.5 \geq BA > CA + 0.5$ ), and decelerated ( $CA - 1.5 \leq BA < CA - 0.5$ ). A pilot study revealed that  $\approx 20\%$  of adolescents exhibited accelerated or decelerated maturation. Thus, 200 participants were invited (100 per CA group) for BA screening, aiming for  $\geq 10$  participants per bin. Extreme cases ( $BA > CA + 1.5$ ,  $BA < CA - 1.5$ ) were excluded to avoid endocrinological complexities. This initial process resulted in a total of 65 participants between 12 and 14 years of age. However, during Section EEG Preprocessing (Experimental Section), 9 recordings were excluded from the study due to excessive noise. To compensate for this, targeted bone age screenings were conducted further and EEG data

of participants who fit into the aforementioned bin criteria were recorded, aiming to maintain a nearly equal distribution of participants across bins. From this additional recruitment, five new EEG recordings were successfully added. Thus, 61 girls aged 12–14 were included in the final cohort as depicted in Figure 1b. The cohort of emerging adult participants consisted of 30 subjects. EEG data of 26 participants was included after the preprocessing procedure, as 4 recordings were removed due to noise.

The percentage of postmenarche participants in both age groups was the lowest in the decelerated groups (see the Open Science Framework link, Supporting Information), suggesting that bone age is indeed a reliable gauge of maturity in adolescence. Mean menarche age of adolescent participants was 12.06y (SD = 0.72). Twenty seven of sixty one participants did not provide data on menarche as they have not reached that stage yet. Mothers (88.52%) and fathers (83.61%) of adolescent participants had a university degree demonstrating high socioeconomic status among the participating families. The EEG data collection was part of a large-scale study, within which each adolescent participant was also administered the Wechsler Intelligence Scale for Children, 4th edition (WISC-IV),<sup>[34]</sup> carried out a fine motor task,<sup>[36]</sup> and a Stroop test.<sup>[35]</sup>

The Hungarian United Ethical Review Committee for Research in Psychology (EPKEB) approved the study (reference number 2017/84). Written informed consent was obtained from all subjects and their parents. Participants were given gift cards ( $\approx$ EUR 15 value each) for their attendance. All research described in this paper was performed in accordance with the Declaration of Helsinki.

**Procedure—Bone Age Assessment:** Body measurements were taken according to the protocol laid down in the International Biological Programme<sup>[105]</sup> using standard instruments. Skeletal maturity assessment began with an anthropometric procedure that includes the measurement of full body and sitting height (GPM Anthropometer, DKSH, Switzerland Ltd, Zurich, Switzerland), as well as body weight (Seca digital scale). Ultrasonic bone age estimation was conducted using the Sunlight BoneAge System (Sunlight Medical Ltd, Tel Aviv, Israel), scanning the speed of sound (SOS) between the distal radius and ulna epiphysis.<sup>[106]</sup> This bone structure reflects the level of maturation as well as it changes significantly during stages of physical growth. The BoneAge device calculates bone age in years and months using software that accounts for ethnicity and gender. Using ultrasound for skeletal maturity assessment is safe for the subjects as opposed to ionizing radiation-based techniques, while results obtained from these two types of procedures are highly correlated.<sup>[40]</sup>

Bone age assessment was performed on the left hand and wrist of participants in each case. Participants were instructed to position their hands between the transducers on the unit's armrest. The device is then attached to the wrist, applying a pressure of  $\approx 500$  g. Transducers transmitted 750 kHz frequency ultrasound through the wrist to measure SOS. The transducer on the ulnar side emits ultrasound while the other transducer acts as a receiver. Measurement protocol was repeated five times for accuracy, each measurement lasting  $\approx 20$  s. Measurements were completed either at the high schools of participants or at the Research Centre for Sport Physiology at the University of Physical Education, Budapest. The entire procedure took 5–10 min per person. Trained assistants conducted the measurements, and a biological anthropologist oversaw the measurement procedure and data analysis.

Bone age assessment results are considered valid for 3 months. Subjects were invited for the resting state electroencephalography (EEG) recording session within this timeframe. If a subject participated in laboratory testing 3 months after the bone age assessment, the procedure was repeated.

**Procedure—Resting State EEG Data Recording:** Continuous EEG data were recorded using the HydroCel GSN 130, a 128-channel high-density EEG (HD-EEG) system (Electrical Geodesics, Inc., Canada), with the Net Station Acquisition software (version 5.4) in a windowless, air-conditioned room insulated from external noise and electromagnetic interference. The physical setting of each EEG recording session was uniform. All participants underwent the same procedure during recording, minimizing the variability caused by external stimuli. Before recording, participants were asked to remove all electronic devices, e.g., phones and smartwatches from their pockets and bodies. Following a brief guided relaxation, two

5-min (300 s) long resting state recordings were collected at a 1000 Hz frequency sampling rate. Impedances were aimed to be kept below 5 K $\Omega$  on all channels. EEG data recording took place at the Laboratory for Psychological Research, Pázmány Péter Catholic University, Budapest.

Two 5-min-long resting state conditions were recorded according to standard protocol.<sup>[107–110]</sup> In the first 5-min-long recording segment, participants were instructed to keep their eyes open, blink minimally, and maintain posture to minimize artifacts generated by muscle movement.<sup>[111]</sup> During the second 5 min of recording, participants followed the same protocol, but were instructed to keep their eyes closed. The two conditions are separated since they exhibit different characteristics, namely a mismatch in topography and activity levels.<sup>[112]</sup> EEG spectral analysis of closed eyes HD-EEG resting state recordings can be found in Supporting Information, under the title of “Age-related changes in the power spectrum” and confirms that the dataset shows the known and expected<sup>[113–115]</sup> age-related changes, thereby supporting the reproducibility of the measurements. In the present study, all statistical analyses were performed on closed-eye recordings. The selection of the closed-eye condition was motivated by the known reduction of alpha oscillations in the presence of visual stimuli and visual attention. Interested readers can find the power spectra of both open eyes and closed eyes conditions in Supporting Information (“Alpha power reduction in “open eyes” condition”) demonstrating the obvious reduction in alpha power when the eyes are open. In spite of this reduction, all the following computations were also carried out for the latter condition and no significant maturity-related variations (Supporting Information “Entropy production under “open eyes” condition”) were found.

**Procedure—EEG Data Preprocessing:** EEG preprocessing was performed in MATLAB (The MathWorks, Inc.; version: R2021a Update 4), using the EEGLAB toolbox<sup>[116]</sup> (version 2021.1). First, a high-pass filter was applied at 1 Hz (transition bandwidth 1 Hz; passband edge 1 Hz; cutoff frequency –6 dB 0.5 Hz; zero-phase non-causal filter) to eliminate low-frequency noise and direct-current offset, and a 40 Hz low-pass filter (transition bandwidth 10 Hz; passband edge 40 Hz; cutoff frequency –6 dB 45 Hz; zero-phase non-causal filter) to remove 50 Hz line noise and high frequency components, using EEGLAB’s basic finite impulse response (FIR; version 2.4) filter. Channel locations were then imported into EEGLAB, the channel layout map was supplied by Electrical Geodesics, Inc. Following filtering steps, noisy channels, determined by two independent raters via visual inspection, were excluded manually. As the last steps, noisy epochs were removed from the recordings using the Clean Rawdata (version 2.4) plugin, then data were re-referenced to average reference. The same preprocessing procedure was implemented for all recordings.

**Data Analysis—Leading Eigenvector Dynamics Analysis:** After preprocessing, Leading Eigenvector Dynamics Analysis (LEiDA),<sup>[58]</sup> adapted to EEG data by the group, was employed to analyze dynamic functional connectivity<sup>[51,55]</sup> in the developing brain. The steps described below were performed on individual EEG recordings and were uniform in all cases.

Although EEG recordings offer an opportunity to analyze data across a wide range of oscillation frequencies, further analyses within the alpha (8–12 Hz) frequency band was chosen to be carried out. This choice is largely based on the fact that both adolescent<sup>[113,114]</sup> and adult<sup>[117,118]</sup> resting-state brain activity is dominated by the alpha rhythm. The relevance of the alpha band in resting state studies is further emphasized by more recent findings demonstrating the overlap between fMRI-based large-scale networks and resting state connectivity in the alpha band.<sup>[119,120]</sup> It might also be relevant to mention that oscillations below the alpha range tend to engage the entire cortex and mostly contribute to long-range temporal integration,<sup>[121]</sup> while oscillations beyond the alpha range tend to be more local, preventing a meaningful analysis of global hierarchical brain organization. However, for interested readers, the entropy production results of delta, theta, beta, and sigma bands on the OSF website (under Supporting Information) have been included. These analyses did not result in any significant maturation-related differences in entropy production.

First, preprocessed EEG recordings were filtered with a 6th-order Butterworth bandpass filter for the alpha band (8–12 Hz) without introducing a phase shift, then downsampled to 100 Hz (intervals of 10 ms). Subsequently, signals were detrended and demeaned, followed by a Hilbert

transformation to obtain the phase of all channels at all timepoints  $t$ . The nominal recording duration of 300 s yielded a temporal sequence of up to 30 000 phase vectors per subject. However, the EEGLAB toolbox typically excised small parts of the sequence, so that the average sequence length was 28 500 vectors (SD = 1600). During preprocessing, a minority of electrodes with interrupted EEG signals during some or all of the recording (“bad channels”, range: 0–12; see Section EEG Data Preprocessing in Experimental Section) was observed. To obtain full-size (128  $\times$  128) coherence matrices at all timepoints and for every subject, a two-step approach termed “patching” was developed. In the first step, group-level lists of “bad channels” by combining noisy channels from all subjects in a group, specifically, the 6 bins of adolescent participants (depicted in Figure 1e) and the group of adult subjects were formed. For  $n$  remaining “good channels” shared by all subjects of the group, instantaneous phase coherence  $PC_{xy,t}$  was computed at every time point for each electrode pair as

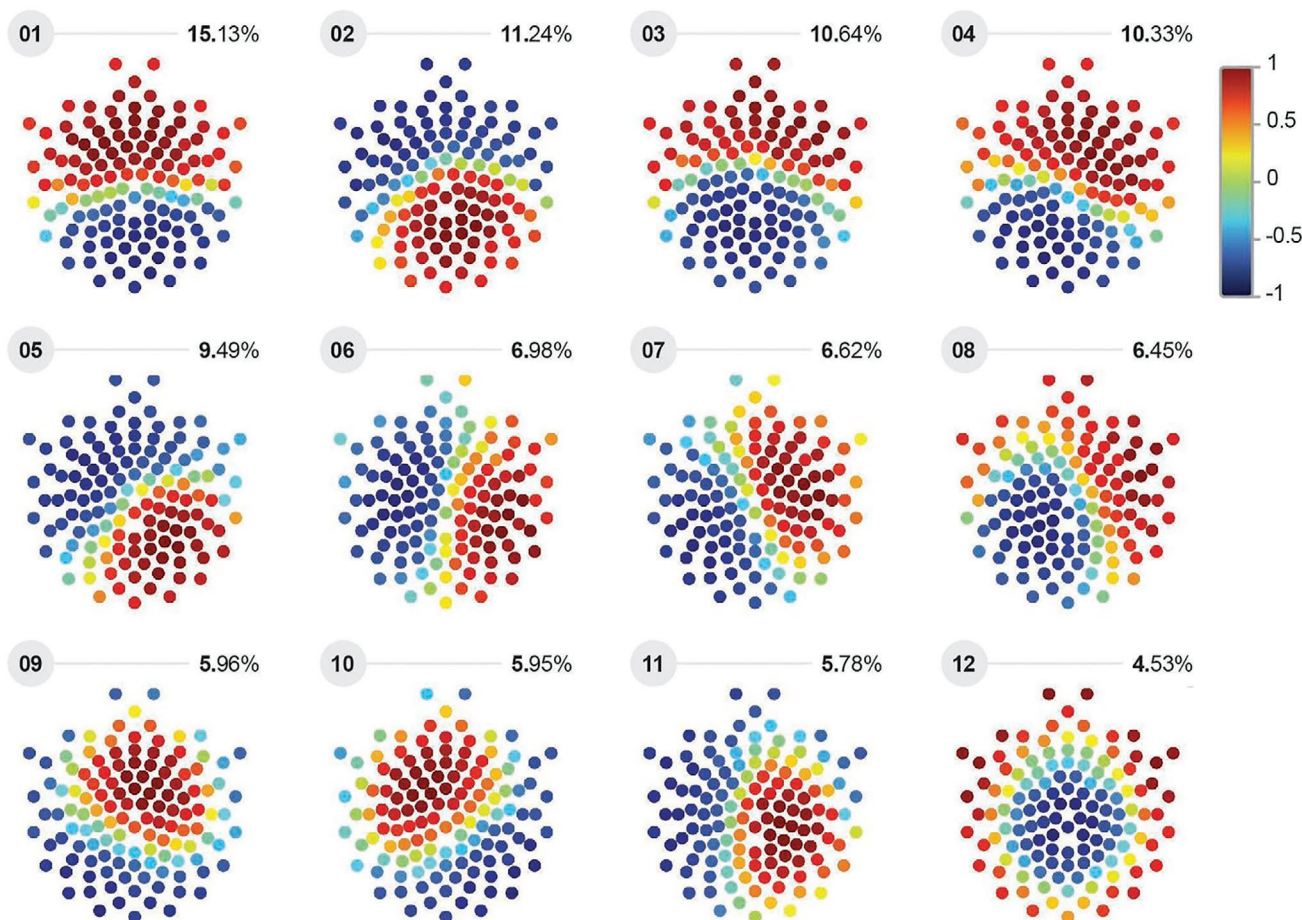
$$PC_{xy,t} = \cos \left( \phi \left( E_{x,t} \right) - \phi \left( E_{y,t} \right) \right) \quad (1)$$

where  $\Phi(E_{x,t})$  and  $\Phi(E_{y,t})$  denote the instantaneous phase of electrode  $x$  and  $y$ , respectively, at each time point  $t$ . This resulted in a temporal sequence of  $n \times n$  phase coherence matrices, sampled at 10 millisecond intervals, for each subject. The leading eigenvector ( $1 \times n$ ) of each matrix was used to summarize the instantaneous pattern of phase differences between electrodes at every time point. Then, these vectors were clustered with Matlab’s built-in  $k$ -means clustering algorithm, assigning a cluster label to all timepoints in all subjects. In the second step, the missing phase values of “bad channels” in all subjects were patched. To this end, the missing phase values of a “bad channel”  $x$  at a time point  $t$  assigned to a particular cluster was replaced by randomly sampling a phase value from the ensemble of values recorded in other subjects of the same bin at the same electrode  $x$  and at time points assigned to the same cluster. Performing this procedure for all groups, subjects, and “bad channels” permitted us to reconstruct full-size phase vectors ( $1 \times 128$ ) and full-size phase difference matrices ( $128 \times 128$ ). Full-size leading eigenvectors ( $1 \times 128$ ) were obtained from full-size phase difference matrices. It should be noted that eigenvectors had the same dimension as the original phase vectors, but represented relative rather than absolute phase.

To identify patterns of phase differences that were common to all subjects – also referred to here as “brain states” and depicted in Figure 6 –, the leading eigenvectors at all timepoints and from all subjects were pooled and clustered via a modified  $k$ -means algorithm<sup>[122]</sup> such as to obtain polarity invariant topographical maps. Through the modified  $k$ -means clustering algorithm, each vector is assigned to one of  $k$  clusters (or brain states) and is labeled accordingly (see Figure 6), allowing us to represent the continuous EEG activity as a trajectory of, and transition between, brain states. LEiDA was carried out in Matlab (version: 2021a). While the results presented here are based on 12 clusters, analyses were also conducted using 6, 8, and 10 clusters and 12 clusters specific to each unique group (see the Open Science Framework link under Supporting Information). The results of these additional analyses were consistent with those presented in Section 2. Results regarding maturity-related entropy production. The results based on 12 brain states since it provides the most detailed description of the data were chosen to be presented.

A representative example of a sequence of successive patterns of phase differences (brain states) is shown in Figure 7a. The average duration of brain states was 125 ms, with a standard deviation of  $\pm 0.4$  ms over groups (range: 12–13 ms) and  $\pm 1.5$  ms over subjects (range: 100–170 ms). However, the durations of individual brain states were quite variable (range: 50–500 ms), with the average standard deviation over subjects being  $\pm 9$  ms (range: 7–14 ms).

**Data Analysis—Entropy Production Calculation:** To quantify asymmetry in the temporal dynamic of state space trajectories, the temporal sequence of brain states (Figure 7a) of each subject was transformed into a table of transition counts  $N_{ij}$  by removing diagonal entries (remaining in the same brain state) and by counting the observed number of forward and reverse transitions between pairs of different brain states ( $i, j$ ). A representative example table is shown in Figure 7b. To avoid singularities in the subsequent



**Figure 6.** Brain state topographies. After initial preprocessing, EEG signals were analyzed via Leading Eigenvector Dynamics Analysis (LEiDA). First, Hilbert transform was applied to each recording to acquire the phase of each electrode. Then, the instantaneous phase coherence between all electrode pairs was computed, yielding a  $128 \times 128$  phase distance matrix for each time point  $t$ . Leading eigenvectors were extracted from the matrices via eigenvector decomposition. Following LEiDA, leading eigenvectors of all subjects were pooled, then clustered via modified  $k$ -means clustering to obtain topographically distinct connectivity patterns, which is referred to here as “brain states”. As leading eigenvectors from all subjects are combined, the twelve activity patterns are rendered uniform in each group. Brain states are ranked according to their dominance duration, expressed in percentages accompanying rank labels, with brain state number 01 accounting for the largest percentage of total activity, and brain state number 12 the smallest. The color scale denotes cosine similarity, with similarly colored dots indicating electrodes with congruent signal phases. Accordingly, values of 1 (dark red) and  $-1$  (deep blue) represent similar and dissimilar (opposite) phases, respectively. Note that electrodes with similar and dissimilar phases form two spatially contiguous subsets.

computation of entropy production, all counts were incremented by one,  $N''_{ij} = N_{ij} + 1$ . Any effect of this change was nullified by the noise floor correction (see below). The corresponding transition probabilities were then obtained as  $P_{ij} = N''_{ij} / \sum_{ij} N''_{ij}$ . To quantify the asymmetry in the pattern of state transitions, deemed indicative of the level of hierarchical organization, entropy production (also termed relative entropy)<sup>[52]</sup> was computed for each subject as the Kullback–Leibler (KL) divergence:

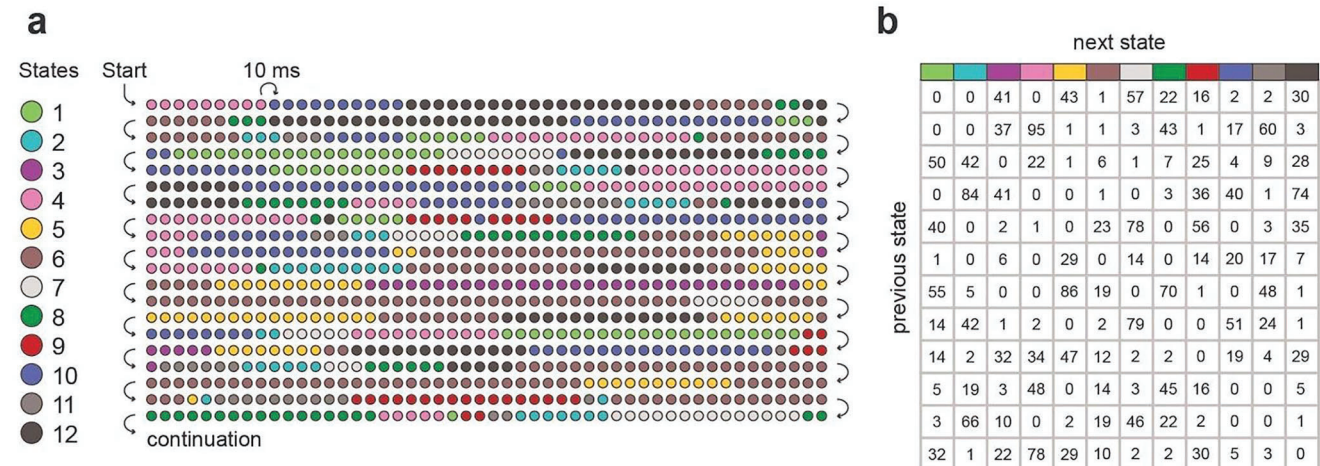
$$H = D_{KL} = \sum_{i,j} P_{ij} \log \frac{P_{ij}}{P_{ji}} \quad (2)$$

where  $D_{KL}$  and  $H$  denote the total entropy production (i.e., the asymmetry between forward and reverse transition probabilities in brain state switching),  $P_{ij}$  denotes the transition probability from brain state  $i$  to brain state  $j$ , while  $P_{ji}$  expresses the probability of switching from brain state  $j$  to brain state  $i$ .

To estimate the variance of the observed entropy values, 1000 synthetic transition trajectories as Markov chains were generated from the transi-

tion probability table observed for each subject via MATLAB’s two built-in functions, one of which creates discrete-time Markov chains and the other simulates Markov chain state walks. Synthetic trajectories had the same length as observed trajectories. Converting synthetic chains into entropy production values with Equation (2) produced a distribution of 1000 entropy production values for each subject.

As the finite length of trajectories inflates entropy production values, a noise floor correction was additionally performed.<sup>[52]</sup> To this end, 1000 synthetic trajectories of the same length from a symmetric transition table (obtained by averaging the observed table with its transpose) were further synthesized. Although perfectly symmetric transition probabilities should not produce any entropy, the resulting value distribution was positive, revealing inflation due to finite trajectory length. To obtain corrected entropy production values, the mean symmetric-table entropy was subtracted from the all asymmetric-table entropy values. From the distribution of corrected values, a mean and estimated variance of entropy production for every subject were obtained. It should be noted that the procedure kept intact the temporal order of trajectories, taking subject-specific brain dynamics into account, and improving on the simpler approach of Lynn



**Figure 7.** Transitions between brain states. a) Brain states numbered 1–12 are represented here by colored disks. Only the initial 10 s of a recording lasting 284 s are shown, specifically, the initial 1000 leading eigenvectors (observed at 10 ms intervals) in a sequence of 28 400 eigenvectors. In this particular example, the average duration of brain states was 98 ms (i.e., 9.8 consecutive eigenvectors were assigned to the same state, on average). b) Number of observed transitions between different states in the entire sequence of 28400 states. Previous states are represented by rows, and the next states by columns. Table entries indicate how often each particular transition was observed. Entropy production quantifies the degree of asymmetry in this matrix.

and colleagues.<sup>[52]</sup> All calculations were performed in Matlab (version: 2021a).

**Data Analysis—Calculating Group Averages and Confidence Intervals:** Group averages and variances of entropy production were computed by inverse-variance weighting. This was done to address the subject heterogeneity and to attenuate the influence of subjects with high variance on the weighted arithmetic mean. The following formulas were used:

$$\bar{H} = \frac{\sum_i H_i / \sigma_i^2}{\sum_i 1 / \sigma_i^2} \quad (3)$$

$$\text{var}(\bar{H}) = \frac{1}{\sum_i 1 / \sigma_i^2} \quad (4)$$

where  $\bar{H}$  is the inverse variance weighted group average entropy production,  $H_i$  denotes the mean observed entropy production for subject  $i$ ,  $\sigma_i^2$  denotes the observed variance for subject  $i$ , and  $\text{var}(\bar{H})$  denotes the inverse variance weighted group variance. 95% confidence intervals for group-level mean entropy production were also computed based on  $\pm 1.96 \cdot \sqrt{\text{var}(\bar{H})}$ . Group averages and confidence intervals were calculated in Matlab (version: 2021a).

**Data Analysis—Statistics:** Statistical assessment of group differences was carried out using two-sample z-tests,<sup>[123]</sup> using the following formula:

$$Z = \frac{\bar{H}_1 + \bar{H}_2}{\sqrt{\text{var}(\bar{H}_1) + \text{var}(\bar{H}_2)}} \quad (5)$$

The significance level ( $p$ ) in the two-sample z-test is determined as in the following:

$$p = 2\Phi(-|Z|) \quad (6)$$

where  $\Phi$  is the standard normal cumulative distribution function.

Results obtained from two-sample z-tests were reported with Hedges'  $g$  effect size estimations. All statistical tests were carried out in Microsoft Excel Professional (version 2021).

## Supporting Information

Demographic and anthropological datasets, EEG analyses, and further analyses are available at the Open Science Framework (OSF) platform at this address: [https://osf.io/eqmwd/?view\\_only=0c48de5f396d4cc681aa894ad093b0fc](https://osf.io/eqmwd/?view_only=0c48de5f396d4cc681aa894ad093b0fc).

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## Conflict of Interest

The authors declare no conflict of interest.

## Data Availability Statement

The data that support the findings of this study are available from the corresponding author upon reasonable request.

## Keywords

bone age, brain development, electrophysiology, entropy production, thermodynamics

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