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Multimodal hyperscanning reveals that synchrony of body and mind are distinct in mother-child dyads



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ABSTRACT

Hyperscanning studies have begun to unravel the brain mechanisms underlying social interaction, indicating a functional role for interpersonal neural synchronization (INS), yet the mechanisms that drive INS are poorly understood. The current study, thus, addresses whether INS is functionally-distinct from synchrony in other systems – specifically the autonomic nervous system and motor behavior. To test this, we used concurrent functional near-infrared spectroscorpy - electrocardiography recordings, while N = 34 mother-child and stranger-child dyads engaged in cooperative and competitive tasks. Only in the neural domain was a higher synchrony for mother-child compared to stranger-child dyads observed. Further, autonomic nervous system and neural synchrony were positively related during competition but not during cooperation. These results suggest that synchrony in different behavioral and biological systems may reflect distinct processes. Furthermore, they show that increased mother-child INS is unlikely to be explained solely by shared arousal and behavioral similarities, supporting recent theories that postulate that INS is higher in close relationships.

1. Introduction

Historically, the mind and the body are considered distinct in Western philosophy. This dualism however does not hold true in modern sciences (Damasio, 1994). The brain is an interoperable system which is embedded in the human body and influenced by other biological systems. In accordance with this, neuroimaging studies in individual subjects have shown that fluctuations in the autonomic nervous system (ANS) are coupled with changes in brain activity (Barber et al., 2020; Breeden et al., 2017; Critchley et al., 2005; Lane et al., 2009).

While these studies have examined single subjects, humans are social species, who continuously affect each other. During social interaction people synchronize on many different levels, including their behavior, ANS and neural signals (Davis et al., 2018; Hari et al., 2015; Hasson et al., 2012). While interpersonal neural synchrony (INS) has been robustly demonstrated in a variety of interactive tasks, the manifold factors which may lead to or affect INS are still poorly understood. Although several studies have provided first important insights (e.g., Baker et al., 2016, Cheng et al., 2015, Leong et al., 2017, Nguyen et al., 2020, Pan et al., 2017, Piazza et al., 2020), these studies often do not consider that synchrony may be established in different behavioral and biological systems. In particular, very little is known about the relationship between INS and synchrony in other biological systems, such as the ANS.

Although not measured concurrently, synchrony in either ANS or brain signals has been found in emotional tasks, such as cooperative and competitive games (Balconi and Vanutelli, 2017; Chanel et al., 2012; Järvelä et al., 2014; Reindl et al., 2018). Further, in a recent hyperscanning study, significant synchrony was observed in brain signal (measured by electroencephalography, EEG), cardiac and electrodermal signals within single subjects as well as between subjects of a dyad when they cooperated with each other (Sciaraffa et al., 2021). Thus, while

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neural and ANS synchrony may co-occur, it is not yet clear whether and under which conditions they are related to each.

To investigate this, the present study uses a well-established hyperscanning paradigm (Baker et al., 2016; Cheng et al., 2015; Cui et al., 2012; Kruppa et al., 2021; Pan et al., 2017; Reindl et al., 2018), in which adult and child either had to cooperate (to synchronize their reaction times to respond as simultaneously as possible to a signal) or to compete (to try to respond faster than their partner to a signal). Participants were 10-18-year-old children and adolescents (all female) who completed the tasks both with their biological mothers (mother-child dyads) and with a previously unacquainted female adult (stranger-child dyads). Previous research shows significant synchrony across the dorsolateral prefrontal and frontopolar cortex when 5-9-year-old children cooperated with their mother, but not in other conditions (mother-child competition, stranger-child cooperation and competition) (Reindl et al., 2018). Consistent results have been observed with adults (Cui et al., 2012; Pan et al., 2017). With older children and adolescents (8-18-yearolds), however, research using the same paradigm additionally identified significant INS during parent-child competition (Kruppa et al., 2021). One untested possibility is that developmental changes in adolescence may be associated with more emotional arousal and associated ANS synchrony during competition with the parents, potentially leading to increased INS.

To measure neural synchrony, we used functional near-infrared spectroscopy (fNIRS) which captures the brain's local hemodynamic response with a high temporal resolution and provides spatial information to locate brain regions which drive INS (e.g., Dai et al., 2018). To obtain comparable information about the temporal relationships between two person's ANS signals, metrics with a high temporal resolution are necessary. The interbeat interval (IBI), that is the time between consecutive heart beats, provides an overall index of arousal, reflective of both sympathetic and parasympathetic activity, which can be assessed reliably within short time windows (Davis et al., 2018; Helm et al., 2018). Thus, in the current study, we extend fNIRS hyperscanning by using concurrent fNIRS - electrocardiography (ECG) recordings, to measure synchrony in the dyad's brain signals and IBIs simultaneously. We examined INS in the frequency range of 0.08–0.5 Hz, which is outside the range of 1–3 Hz that may be contaminated by ECG artifacts.

To analyze INS and its relationship to synchrony in other modalities, we developed a new analytical approach based on bipartite graph analyses (described in more detail in Gerloff et al., 2021). Since complex human behavior and cognition is not localized to a single circumscribed brain region but is organized in functional brain networks, INS may be more accurately modelled as the bidirectional links between the brain networks of interacting subjects (see also Ciaramidaro et al., 2018, Santamaria et al., 2020). These functional networks can be expressed as graphs. While global graph metrics provide a scalar value, which can be easily compared to synchrony measures in other modalities, nodal metrics provide increased topological detail. Because ANS synchrony might impact INS in very specific brain regions while other nodes might be less affected (see also Chen et al., 2020), to fully understand whether INS is functionally-distinct from synchrony in other systems, an analysis on both levels may be necessary.

Here, we explored whether INS, measured at both global and nodal levels, goes beyond synchrony in the ANS, as measured by the dyad's IBIs. We also examined the relationship of INS to behavioral synchrony (indexed as the mean of the absolute differences in response times), and we measured how trial-by-trial adaptations in response times, contingent on feedback during the task, related to INS. To this end, we first compared the different biobehavioral synchrony markers (INS, ANS and behavior), and tested whether synchrony differed on each measure: (i) between mother-child and stranger-child dyads, and (ii) between cooperation and / or competition compared to a non-interactive baseline condition, in which the mother-child / stranger-child dyad watched a relaxing video together (Research Question 1). Based on our previous studies (Kruppa et al., 2021; Reindl et al., 2018), we expected an increased INS during cooperation and possibly during competition compared to baseline, as well as higher INS for mother-child dyads than for stranger-child dyads. For behavioral synchrony, we expected either no partner differences (Reindl et al., 2018) or higher synchrony for stranger-child dyads (Kruppa et al., 2021), while participants should react more synchronously during competition than during cooperation (Kruppa et al., 2021; Reindl et al., 2018). For ANS synchrony, higher synchrony is expected for cooperation and competition compared to baseline (see Chanel et al., 2012; Järvelä et al., 2014), however, no predictions were possible regarding partner effects. Second, we explored whether INS was related to ANS synchrony and / or behavioral synchrony (Research Question 2). Given the sparsity of research in this field, no hypotheses were formulated with respect to the relationship of INS and ANS synchrony. For behavioral synchrony, several studies indicate that higher INS is associated with a better cooperative performance (however, mostly measured in joint wins during cooperation and in adults; Baker et al., 2016; Cheng et al., 2015; Cui et al., 2012; Pan et al., 2017).

2. Material and methods

2.1. Participants

The initial sample consisted of 41 female children, aged between 10 and 18 years, who participated in the study with their biological mothers (mother-child dyads). In addition, each child performed identical tasks with a previously unacquainted female adult (stranger-child dyads). Because INS has been shown to be influenced by the participant's gender (Baker et al., 2016; Cheng et al., 2015), the current study focused on female children and female adults only. Participants were recruited via previous studies, postings in the intranet of the University Hospital RWTH Aachen as well as flyers. None of the participants had any severe cardiac, neurological or psychiatric conditions.

From the initial sample, one child was excluded because of an attention deficit disorder, two children were excluded due to insufficient fNIRS data quality, one child because of a heart condition and three children because of missing ECG data due to technical errors or insufficient ECG data quality. Thus, the final sample consisted of 34 children (M age = 14.26 years, SD = 2.206 years, range: 10-18 years) and 34 mothers (M age = 45.32 years, SD = 4.953 years, range: 37–56 years). Moreover, a total of 29 female adults served as strangers in the study (M age = 23.07 years, SD = 2.086 years, range: 19–29 years). Of these, 26 adults participated once, one adult twice and two adults three times. Strangers were significantly younger than mothers (t(61) = -22.532, p < 0.001). For some participants, ECG / fNIRS data were missing in specific experimental conditions mainly due to insufficient data quality or technical errors. Thus, samples sizes varied between N = 31 and N = 34 for the experimental conditions and measures (for more information see Supplementary Text 1, Table S1).

Participants were reimbursed for study participation. The study was approved by the Ethics Committee of the Medical Faculty, University Hospital RWTH Aachen (EK 151/18). All adults, including children of legal age, gave written informed consent for their own study participation as well as, in the case of mothers, for the participation of their children. Children below the age of 18 gave written informed assent.

2.2. Procedures

Prior to the experiment, participants were instructed not to exercise, not to drink alcohol or energy drinks and not to smoke at least one hour before the lab visit, as these factors may influence ANS measurements. After arriving in the lab, first, ECG electrodes were attached to the participants' bodies. This was done in the beginning of the testing session to give participants enough time to habituate to the procedure. Afterwards, the cooperative and competitive tasks were explained and five practice trials were provided for each. fNIRS optodes were placed on



Fig. 1. Illustration of the experimental design. During cooperation (A), the task was to react as simultaneously as possible to a signal via button press, while during competition (B), the task was to react faster than the other partner to win. Each cooperative / competitive trial was organized in the following way: (i) wait screen showing the two dolphins for 2 s, (ii) display of 'ready' signal (black hollow circle) for a randomly sampled time interval of 0.6-1.5 s, (iii) display of 'go' signal (colorful ball), (iv) feedback screen for 1.5 s / 1 s (cooperation / competition). RT = response time of the slower participant / faster participant (cooperation / competition).

the participant's heads shortly before the start of the measurements to reduce wearing times.

Each experiment began with the baseline condition, followed by the cooperative and competitive tasks. During the experiment, participants were seated next to each other, facing a single computer screen. They were instructed to rest their heads still on a chin rest, in order to reduce movement artifacts, and to refrain from talking to each other. To reduce the participants' ability to perceive each other's movements, a towel was placed over their hands (for a video showing the set-up and fNIRS data collection, see Reindl et al., 2019).

A total of 17 children (50%) first completed the three measurements (baseline, cooperation, competition) with the mother and after a short break with the stranger. For 17 children it was the other way around. The order of the cooperative and competitive task was kept constant for both dyads each child was part of but was balanced across children. A total of seven children (20.6%) started with mother-child cooperation, ten children (29.4%) started with mother-child competition, nine children (26.5%) with stranger-child cooperation and eight children (23.5%) with stranger-child competition.

2.3. Experimental tasks

2.3.1. Baseline

For the baseline condition, a three-minute excerpt from a relaxing aquatic video (Coral Sea Dreaming, Small World Music Inc.) was presented. The aquatic video has been effectively used in previous studies with children to obtain baseline ANS measurements (Piferi et al., 2000; Prätzlich et al., 2018) and served as a low-level control condition to account for the possibility that observed synchronous hemodynamic and physiological changes were due to shared sensory input.

2.3.2. Cooperation and competition task

Adapted versions of the cooperative and competitive computer game tasks of Cui et al. (2012) were implemented, which have been found appropriate for children (Kruppa et al., 2021; Reindl et al., 2018). Each player manipulated the on-screen movement of a dolphin towards a ball by pressing a computer key with the goal to either catch the ball together (cooperation) or win the ball for themselves (competition). Each task was composed of two task blocks with 20 trials each and three 30 s rest blocks in alternating order: rest1, task1, rest2, task2, rest3. In line with previous publications (Kruppa et al., 2021; Reindl et al., 2018), only the two task blocks were considered in the analyses. The trial organization is depicted in Fig. 1.

During *cooperation*, the goal was to "catch the ball together" by reacting as simultaneously as possible. In the beginning of each trial, two dolphins appeared and remained on the screen. After 2 s, a black circle appeared above the dolphins ('ready' signal) and was replaced by a colorful ball ('go' signal) after a variable time interval (0.6–1.5 s). Dyads were asked to respond as simultaneously as possible after the 'go' signal had appeared via pressing a computer key. If the difference in response times was below a predefined threshold, both dolphins jumped to the ball (feedback screen, 1.5 s), caught the ball (result screen, 1.5 s) and earned a point. If the difference between the response times was above the threshold, only the faster dolphin jumped towards the ball (feedback screen), none of the dolphins caught the ball (result screen) and both participants lost a point. The temporal threshold was individually adjusted to the response times of the dyad (set to T = 1/8 [RT1 + RT2], where RT1 and RT2 indicate the response times of the two participants). If one of the players reacted too early, that is before the 'go' signal, the trial started again from the beginning and both players lost a point.

During *competition*, the goal was to "catch and win the ball by oneself" by pressing the response key faster than the other partner after the 'go' signal had appeared. Only the faster dolphin jumped to the ball (feedback screen, 1 s), caught the ball (result screen, 1 s) and earned a point while the slower participant lost a point. If both reacted equally fast with an error margin of 50 ms, both dolphins jumped to the ball (feedback screen), caught the ball (result screen) and gained a point (joint win). Again, if one of the players reacted too early, the respective player lost a point and the trial started from the beginning.

2.4. Multimodal data acquisition

2.4.1. ECG data acquisition

ECG data were acquired with the Vrije Universiteit Ambulatory Monitoring System (VU-AMS; Netherlands) at a sampling rate of 1000 Hz. In addition, impedance cardiography data were acquired, which is not reported here since it is beyond the scope of the paper. After cleaning the skin with disinfection solution, H98SG, ECG Micropore electrodes (Covidien, Germany) were attached to the participant's upper body: one slightly below the right collar bone, one on the right side between the lower two ribs and one approximately at the apex of the heart. Prior to the experiment, the internal clock of the VU-AMS device was synchronized to the clock of the stimulation computer, to ensure a temporal synchronization of ECG and fNIRS devices.

2.4.2. fNIRS data acquisition

fNIRS data were acquired in both subjects simultaneously using a single fNIRS device with a sampling rate of 10 Hz (ETG-4000, Hitachi Medical Corporation, Japan). A " 3×5 " probe holder grid was mounted to a modified EEG cap (Easycap GmbH, Germany) and probes were inserted into the appropriate holder sockets on the grids. In each grid,



Fig. 2. Multimodal data analysis workflow. To examine the relationship between different biobehavioral synchrony measures in a single multivariate generative model, we proposed a symmetric data fusion approach, analyzing synchrony in fNIRS and ECG signals concurrently. Top: After motion artifact correction and detrending of the fNIRS signals, the salient wavelet coherence was calculated as the connectivity estimator. Subsequently, for each dyad and condition, individual bipartite graphs were constructed by defining the salient wavelet coherence as weighted edges connecting different regions (nodes) from adult and child. To avoid spurious connections, the graphs were reduced by a block-wise permutation procedure comparing individual graphs with the graphs of shuffled adult-child pairs. The number of surviving connections between brains was calculated for the network (global density) as well as for each node / fNIRS channel (nodal density). To reduce the dimensionality of the nodal metrics while preserving interpretability, nodal density vectors were encoded via non-negative matrix factorization. Bottom: ANS synchrony was calculated by the cross-correlation of the participant's IBI time series after R-peak correction and ARIMA modeling. Subsequent analyses were performed using (multivariate) Bayesian hierarchical models.

eight emitters and seven detectors were positioned alternatingly in three rows, resulting in 22 measurement channels. The source-detector distance was fixed at 3 cm. The caps were placed symmetrically over the participants' foreheads so that the middle optode of the lowest probe row was placed on the Fpz point of the 10-20 system, and the middle probe column aligned along the sagittal reference curve. The most probable spatial locations of the channels were estimated by the virtual registration method (Singh et al., 2005; Tsuzuki et al., 2007), using the Talairach Daemon (Lancaster et al., 2000). The brain regions covered by this optode set-up include Brodmann Areas (BAs) 8, 9, 10 and 46. Due to restrictions with respect to the number of available optodes, our recordings concentrated on the prefrontal cortex, since these regions have been frequently found to show significant INS (e.g., Pan et al., 2017, Piazza et al., 2020), and to keep the set-up comparable to previous studies with the same experimental tasks (Kruppa et al., 2021; Reindl et al., 2018).

2.5. Behavioral data analysis

The response times of both participants were recorded during the cooperative and competitive task (Supplementary Text 2, Table S2). As a measure of behavioral synchrony, the mean of the dyad's absolute differences in their response times (Mean-DRT) was calculated, with smaller values indicating higher synchrony. To further quantify the participants' task behavior, the number of joint wins during cooperation as well as the number of child's wins and joint wins during competition are reported in Supplementary Text 2 and Table S3. In addition, as an index of how strongly the dyad adapted their response times, we calculated the difference between the mean-DRT of the present trial and its subsequent trial, whereby larger values indicate a stronger adaptation of the dyad. This was calculated for all trials in which participants received feedback showing who responded more quickly and was then averaged across all 'feedback' trials of each block (in case of cooperation: only in trials in which the dyad had failed to achieve a win; Table S4).

2.6. ECG data analysis

The ECG and fNIRS data analysis workflow is depicted in Fig. 2. For the ANS synchrony analyses, we adopted previously used methods (Feldman et al., 2011; Suveg et al., 2016) (for further information see Supplementary Text 3). First, R peaks were detected in the raw ECG signal using an automated algorithm. If necessary, R peaks were manually corrected and artifacts removed. Afterwards, for each condition, the IBI time series were resampled at 10 Hz, and the samples were divided into epochs of 2000 ms with fixed on- and offsets to enable an accurate temporal synchronization of the adult's and child's IBI values (Berntson et al., 1995). An epoch length of 2000 ms was chosen based on minimal amount of time needed to reliably estimate the heart rate (Helm et al., 2018). For each epoch, the mean IBI was computed, resulting in a time series of epoch means for each participant. Artefact removal in the initial IBI series resulted in missing values. Missing values were interpolated with a cubic spline interpolation. If more than 5% of the values were missing of either adult or child in one recording, the respective experimental condition of the dyad was excluded from further analysis (Supplementary Text 1, Table S1).

A second order polynomial regression was computed for each epoch means time series in order to remove linear and quadratic trends from the data (Suveg et al., 2016). To examine the autocorrelative properties of the signals, partial autocorrelation functions (PACF) of the IBI time series after detrending, i.e., on the residuals after polynomial fitting, were plotted. The partial autocorrelation measures the signal's autocorrelation at lag k after removing effects of autocorrelations due to shorter lags. PACF results, averaged across participants, showed a strong autocorrelative component at lag = 1, likewise for adult (stranger / mother) and child and for all experimental conditions. At none of the other lags, the average autocorrelation exceeded the upper or lower confidence bounds. These results indicate that an ARIMA model with a lag = 1 is appropriate to effectively reduce the signals' autocorrelations. It is important to remove this autocorrelation, because otherwise spurious correlations may be detected in two independent but autocorrelated time

series (Dean and Dunsmuir, 2016). In order to do this, and following an approach used previously (Feldman et al., 2011; Suveg et al., 2016), the residuals of the polynomial regression were subjected to Autoregressive Integrated Moving Average (ARIMA) modeling, with one autoregressive term, one moving average term, and integrated noise, and the residuals from this analysis were entered into the cross-correlation calculations. Finally, the cross-correlation at lag = 0 was calculated between the two time series of residuals after ARIMA modeling. Cross-correlations were computed for each condition, and, in case of the cooperation / competition task, for each of the two task blocks. These served as our primary outcome value for ANS synchrony.

In addition, and in order to ensure that the validity of our findings was not specific to the exact measure used to calculate synchrony, we also calculated ANS synchrony by calculating the wavelet coherence, using a method that was as far as possible identical to the method used for calculating INS (Supplementary Text 4).

Of note, both measures used to calculate ANS synchrony do not measure how far individual heart beats occur at the same time across the dyad. Rather, they measure how changes in heart rate between consecutive 2000 ms epochs, co-fluctuate across the dyad.

2.7. fNIRS data analysis

2.7.1. fNIRS data preprocessing

fNIRS signals were preprocessed by first converting the raw intensity data to optical density data. Second, motion artifacts were detected and reduced by a cubic spline interpolation (Scholkmann et al., 2010). Third, optical density was converted to HbO and HbR concentration changes. The differential pathlength factor was estimated based on the wavelength and the participant's individual age (Scholkmann and Wolf, 2013). Finally, data were detrended. Noisy channels were identified based on a semi-automated procedure using several objective criteria in combination with visual inspection and excluded from all subsequent analysis (as described in Kruppa et al., 2021). If more than 25% of the channels of a participant in a specific experimental condition was identified as noisy, the complete fNIRS recording was excluded, resulting in missing values (Supplementary Text 1, Table S1). For further information on fNIRS data preprocessing see Supplementary Text 5 and Table S5.

2.7.2. Connectivity estimator

After signal preprocessing, the statistical dependencies between the dyad's fNIRS signals were quantified via the bivariate wavelet coherence (WCO). The WCO is a widely applied non-directional functional connectivity estimator, which localizes the signals' dependencies in the time-frequency space (Daubechies, 1990) and is thereby able to distinguish neural signal components from ANS related frequencies, such as the heart rate. For each signal pair, i.e., for each dyad in each condition and channel combination, the WCO yields a two-dimensional time - 'frequency' matrix. These coefficients were then aggregated to a single value, representing the connectivity estimator. To increase the robustness of the estimator, we only considered salient WCO coefficients that are higher than a cut-off value, since these are less affected by noise (Reindl et al., 2018). Specifically, we calculated the percentage of salient values across each task block and within a task-related frequency band between 0.08 and 0.5 Hz (period length: 2.02-12.80 s). The task-related frequency band was chosen based on previous studies (Kruppa et al., 2021; Reindl et al., 2018). It includes the trial duration (~7 s for cooperation, \sim 6 s for competition) and importantly lies outside the frequency band of the heart rate (3-1 Hz, period length 0.33-1 s). For further information on the WCO and the cut-off calculations see Supplementary Text 6.

2.7.3. Bipartite graph analysis

The complete bipartite graph, $G = (V_1 \cup V_2, E)$, was constructed, whereby the fNIRS channels of participant 1, V_1 , and of participant 2,

 V_2 , represent the nodes. These two disjoint sets of nodes are connected by edges, $E \subseteq V_1 \times V_2$, whose weights W are defined by the connectivity estimator (see 2.7.2). Consequently, in hyperscanning the edges can be interpreted as the interpersonal links between the brain regions V_1 of one participant and the brain regions V_2 of another participant. Edges connecting a 'noisy' channel were excluded.

In network analysis, it is common practice to exclude edges in order to reduce spurious links and to ensure a more robust network topology. To determine these thresholds, the WCO was calculated for all possible combinations of independent mother/stranger - child dyads, termed 'shuffled pairs', assuming exchangeability of the participant ID while holding the condition and channel combination fixed. Using this blockwise permutation, a shuffled-pair distribution was derived individually for each condition and channel-combination, and the threshold was set to its 95% quantile (for more information see Supplementary Text 7). Thus, only edges were considered which were related to the 'true' interaction of the dyad rather than related to random or systemic similarities between brain signals due to the same experimental condition. Based on these reduced graphs both global and nodal graph metrics were calculated.

Global (inter-brain) density is defined as the total number of edges, i.e., the interbrain links that survived permutation, relative to the maximum number of possible edges, after noisy channels were excluded (Santamaria et al., 2020). Nodal (inter-brain) density is the number of survived edges for each node that survived permutation, again, relative to the total number of possible edges for the respective node. Thus, nodal density estimates how strongly the temporal activation patterns of a given node are coherent to the temporal activation patterns of the other partner. Thereby it allows to determine the individual contributions of brain regions to this overall connectivity.

2.7.4. NMF

When analyzing task effects for each node individually, this may result in multiple comparison issues (N = 44 nodes; Gerloff et al., 2021). To circumvent this, we reduced the dimensionality of the nodal metrics by using a non-negative matrix factorization (NMF) and then analyzed the resulting components in one multivariate model (see 2.9). The NMF allows to encode the nodal topologies in a low-dimensional vector while preserving the contributions of the individual nodes to each component, thereby yielding localized and interpretable insights into the interbrain networks (Gerloff et al., 2021). The nodal densities are represented in a matrix $V \in R_{\perp}^{m \times n}$, with m = 44 nodes, 22 of child and adult, and n observations for each dyad in each condition. This matrix was then approximately factorized into a basis matrix $W \in \mathbb{R}_+^{m \times r}$ and a coefficient matrix $H \in \mathbb{R}_{+}^{r \times n}$, whereby the rank, r, is chosen to be smaller than *n* or m (Lee and Seung, 2001). The basis matrix W provides the assignment of nodes to components, thus, can be understood as dictionary to look up the contribution of each node to each component, which is constant across dyads and conditions (Fig. 4). The coefficient matrix H encodes the nodal densities as features for each component, which are later used in the result analysis.

To obtain stable results, we performed each NMF with 10,000 iterations. For both HbO and HbR, the rank was chosen to be four, based on the reconstruction error of the original data matrix compared to a shuffled data matrix (for further information see Supplementary Text 8). The nodes which contribute to the four components in term of their weights are depicted in Fig. 4 (HbO) and Fig. S1 (HbR).

2.8. Validation by shuffled pair analysis

To account for similarities in the dyad's IBI and fNIRS signals as well as behavioral responses not related to the social interaction, we examined whether synchrony of the actual dyads was higher than synchrony of independent participants involved in the same experimental condition ('shuffled pairs'). To this end, interpersonal synchrony measures were calculated for all possible shuffled mother /stranger - child pairs. Specifically, sets of shuffled pairs were constructed for each child by varying the adult partner and for each adult by varying the child partner, while holding the condition fixed. Next, a dyad- and condition- specific mean shuffled pair synchrony value was derived by averaging across the synchrony values of the child's and adult's shuffled pair sets in each condition. Thus, for each dyad, we obtained one actual synchrony value and one mean shuffled pair synchrony value (Kruppa et al., 2021).

While shuffled pairs performed the same cooperative / competitive task, the timing of the trials and length of task blocks differed between subjects due to the variable inter-trial interval and the subjects' responses. Since ANS and neural synchrony analysis requires an equal length of the signals, the longer task block was cut at the end to have the same length as the shorter task block.

2.9. Bayesian result analysis

To derive an estimate of how the experimental conditions affect interpersonal synchrony in different systems and the factors influencing INS, Bayesian Hierarchical Models (BHMs) were used (Fig. 2). This method is gaining increasing importance for neuroscience and brain network analyses (Bzdok et al., 2020). The Bayesian framework comes with several advantages compared to the classical 'frequentist' approach. Bayesian models allows to incorporate prior knowledge about the parameters in the models and to specify different response distributions. This is particularly important since nodal density follows a non-Gaussian distribution with long tails towards high values, thereby violating assumptions of many classical frequentist tests (e.g., ANOVA) (Bullmore and Sporns, 2009). Furthermore, the BHM does not rely on p-values but derives a probability statement for each of the parameters of interest. In the result section, we report the mean of the parameter's estimated marginal posterior distribution as well as its two-sided 90% CI, which is defined as the probabilistic interval that is believed to contain a given parameter (Aczel et al., 2020). For discussion purposes, two-sided 90% CIs which do not include zero are interpreted as statistical evidence for a given effect. For directional hypotheses (higher INS for mother-child compared to stranger-child dyads, and for cooperation/competition compared to baseline), additionally, one-sided 90% CIs (= two-sided 80% CIs) are reported. Prior to the BHM analyses, missing INS and ANS synchrony values were imputed by multiple imputation. For more information on the imputations, BHM implementations and quality checks see Supplementary Text 9.

2.9.1. In which systems does interpersonal synchrony occur?

First, to examine whether actual pairs differed from shuffled pairs, we calculated individual BHMs for (i) INS, (ii) ANS and (iii) behavioral synchrony with global density, ANS cross-correlations and Mean-DRT as the response variable, respectively. The models included pair (0 = shuffled, 1 = actual), experimental condition as well as their interaction as predictors. Reported are the effects of pair for each experimental condition.

Second, to directly compare the different experimental conditions, BHMs were again calculated for (i) INS, (ii) ANS and (iii) behavioral synchrony. For models (i) and (ii), predictors included: competition (0 = baseline, 1 = competition), cooperation (0 = baseline, 1 = cooperation), partner (0 = stranger, 1 = mother), as well as the two-way interactions between competition / cooperation and partner. For model (iii), task was coded with 0 = cooperation and 1 = competition. If there was no evidence for an interactive effect, unconditional main effects were reported. For nodal density, we calculated multivariate BHMs, which included all four NMF components as response variables. It should be noted that BHMs integrate all effects into one model and thereby address multiple comparison issues (Gelman et al., 2012). Adding the child's age (in years) as an additional predictor to these models did not change any of the main findings (for correlations between study variables and child's age, see Tables S6–S11).

2.9.2. Which factors are related to interpersonal neural synchrony?

Second, we examined whether ANS and behavioral synchrony were related to INS. To this end, we calculated univariate or multivariate BHMs for global and nodal density, respectively. First, we estimated the effects of ANS synchrony, competition (0 = baseline, 1 = competition), cooperation (0 = baseline, 1 = competition), partner (0 = stranger, 1 = mother), as well as their two-way interactions with ANS synchrony. To calculate cross-level interactions, in this case with ANS synchrony (level 1) nested in task and partner (level 2), it is advisable to conduct a group-mean centering of the level 1 predictor prior to the analysis (Enders and Tofighi, 2007). Thus, ANS synchrony values were groupmean centered by subtracting the mean value in the respective experimental condition. Equivalent (multivariate) BHMs were formulated for behavioral synchrony, estimating the effects of task (0 = competition,1 = cooperation), partner (0 = stranger, 1 = mother), behavioral synchrony (group-mean centered) and their two-way interactions on global and nodal density. Again, adding the child's age to the models did not change any of the main findings.

3. Results

3.1. In which systems does interpersonal synchrony occur?

For the first research question, we examined task (baseline vs. cooperation / competition) and partner (mother vs. stranger) differences in (i) INS, (ii) ANS synchrony and (iii) behavioral synchrony. The subsections are organized as follows. First, we compared mother / stranger-child synchrony to the synchrony of shuffled adult-child pairs, who performed the same task independently of each other. Second, we directly compared the experimental conditions.

3.1.1. INS

Neural synchrony was assessed over the prefrontal cortex using global and nodal inter-brain density (short "density"). Since many fNIRS hyperscanning studies focus on oxy-hemoglobin (HbO) signals (Baker et al., 2016; Cheng et al., 2015; Cui et al., 2012; Pan et al., 2017; Reindl et al., 2018), the results for HbO are presented in the main text and then compared to the results for deoxy-hemoglobin (HbR) to validate the findings and reduce the risk of false positives (Tachtsidis and Scholkmann, 2016) (Supplementary Text 10, Table S12).

To obtain a more robust network, the graph's edges were reduced by a block-wise permutation procedure comparing individual graphs with the graphs of shuffled adult-child pairs. Since we reduced the graphs via the 95% quantile of shuffled pairs, consequently, shuffled adultchild pairs had a global density of ~ 5%. To investigate whether global density of actual pairs was actually higher, we estimated the effects of shuffled vs. actual pair per condition within a single BHM. Descriptive results are presented in Table S13. For HbO, results showed an increased density only for mother-child competition (posterior mean (μ) = 0.11, 90% credible interval (CI) = [0.02, 0.20]), while no sufficient evidence was found for increased density in the other conditions. However, actual pairs had a lower density in the stranger-child baseline condition (μ = -0.17, CI = [-0.29, -0.05]).

Next, a BHM was calculated for the effects of baseline vs. competition, baseline vs. cooperation and stranger vs. mother as well as their two-way interactions on global density (Fig. 3; Table S12). Compelling statistical evidence was found for increased density of mother-child compared to stranger-child dyads ($\mu = 0.13$, CI = [0.04, 0.23]) and of competition compared to baseline ($\mu = 0.13$, CI = [0.02, 0.25]). Furthermore, weaker evidence was found for a positive effect of cooperation compared to baseline ($\mu = 0.08$, CI = [-0.02, 0.18], posterior samples above zero: 91.38% > 0), while insufficient evidence was observed for an interaction between competition / cooperation and partner.

Since smaller and / or more localized effects may not be detected by global graph metrics, we additionally examined nodal density, having the further advantage of an increased topological detail. The dimen-



Fig. 3. Differences between interpersonal neural and autonomic nervous system (ANS) synchrony as a function of task and partner. To examine the systems in which synchrony occurs, marginal posterior distributions were derived for the effects of stranger vs. mother, baseline vs. competition and baseline vs. cooperation on HbO and HbR global density and ANS synchrony. Forest plots show the 99 and 90% two-sided credible intervals (thin and thick black lines) as well as the posterior mean (black dot). 90% credible intervals which do not cover zero were interpreted as evidence for an effect. For both HbO and HbR, evidence was found for a higher density of mother-child compared to stranger-child dyads and of competition compared to baseline. In contrast, for ANS synchrony, there was no evidence for a partner effect, while strong support was found for both task effects, with increased synchrony for competition and cooperation compared to baseline. Together, these results indicate that synchrony in neural and ANS signals was clearly differentiable.

sionality of the nodal metrics was reduced via NMF to four components, each of which contains a collection of nodes with varying contributions (Fig. 4).

Nodal density results validated the global results, showing evidence for a partner, competition and cooperation effect (Fig. 5; Table S12). Specifically, we observed higher density for mother-child compared to stranger-child dyads across tasks in component 3 ($\mu = 0.09$, CI = [0.00, 0.18]) and component 4 ($\mu = 0.15$, CI = [0.04, 0.27]) as well as some evidence for an effect in component 1 ($\mu = 0.08$, CI = [-0.02, 0.18], 91.31% > 0). Additionally, in component 2, higher density was found for mother-child compared to stranger-child dyads in the baseline condition (competition x partner interaction: $\mu = -0.27$, 90% CI = [-0.49, -0.05]; cooperation x partner interaction: $\mu = -0.21$, CI = [-0.44, 0.02], 6.32% > 0). Evidence for a competitive task effect was observed in components 1 (μ = 0.16, CI = [0.04, 0.29]) and 4 (μ = 0.13, CI = [0.02, 0.25]) and analogously, evidence for a cooperative task effect was observed in component 1 ($\mu = 0.12$, CI = [0.02, 0.22]) and to a weaker degree in component 4 (μ = 0.09, CI = [-0.02, 0.20], 90.92% > 0). Component 3 and 4 mainly comprise orbitofrontal brain regions of adult and child as well as right superior prefrontal brain regions of the adult, while component 1 and 4 mainly comprise left and right lateralized prefrontal brain regions of adult and child. The brain regions which contribute most to each of the components can be found in Fig. 4.

Our main neural results (HbO) were further validated by comparing them to the results for HbR, which showed a mostly consistent result pattern (Supplementary Text 10). Again, increased global density was found for mother-child compared to stranger-child dyads and for competition compared to baseline (Fig. 3). Furthermore, in line with the HbO findings, HbR nodal density results confirmed the global findings, indicating increased density for mother-child dyads and for competition. In addition, increased density was found for mother-child cooperation in one component.

Together, these results indicate that INS was increased for motherchild dyads, for competition and for cooperation. Yet, the effects for cooperation were smaller and driven by a subset of nodes as indicated by the NMF results.

3.1.2. ANS synchrony

ANS synchrony was analyzed by calculating the cross-correlation of the IBI time series after reducing the time series' autocorrelations. In a preliminary step, we inspected the signal's autocorrelation via its PACFs (Fig. 6). Of note is a stronger autocorrelation at lag = 3 for the cooperative / competitive task relative to baseline (child PACF, baseline vs. cooperation: $\mu = 0.12$, CI = [0.08, 0.16], baseline vs. competition: $\mu = 0.10$, CI = [0.05, 0.14]; adult PACF, baseline vs. cooperation: $\mu = 0.16$, CI = [0.11, 0.20], baseline vs. competition: $\mu = 0.14$, CI = [0.10, 0.18]). Since the structure of our task was that, in the cooperation and competition conditions, trials were presented roughly once every six seconds (i.e., every three epochs given that a 2000 ms epoch was used), this likely reflects that both adult and child heart rate became entrained to the task structure. Furthermore, for the adult's time series, we found an interaction between competition and partner at lag = 3 ($\mu = -0.08$, CI = [-0.15, -0.00]) (and some evidence for an interaction between cooperation and partner), indicating that strangers had higher PACF values at lag = 3 than mothers.

Descriptive results for the mean IBI and ANS synchrony per condition and player are presented in Table S14. When compared to shuffled pairs, increased ANS synchrony was found for mother-child cooperation ($\mu = 0.08$, CI = [0.04, 0.12]), mother-child competition ($\mu = 0.11$, CI = [0.08, 0.14]), stranger-child cooperation ($\mu = 0.08$, CI = [0.04, 0.11]) and stranger-child competition ($\mu = 0.11$, CI = [0.06, 0.15]). However, no increased ANS synchrony was found for mother-child baseline ($\mu = 0.03$, CI = [-0.02, 0.08]) or stranger-child baseline ($\mu = 0.01$, CI = [-0.04, 0.05]).

Directly comparing the conditions, we found very strong evidence for both task effects with higher ANS synchrony for competition ($\mu = 0.11$, CI = [0.07, 0.14]) and cooperation compared to baseline ($\mu = 0.07$, CI = [0.04, 0.11]), although this effect was stronger for competition than for cooperation ($\mu = 0.03$, CI = [0.00, 0.06]). In contrast, no evidence was found for a partner effect and the μ was close to zero ($\mu = 0.01$, CI = [-0.02, 0.04]) (Fig. 3; Table S12). Thus, we can conclude with a high certainty that there was increased synchrony for cooperation and competition compared to the non-interactive baseline condition, but no meaningful difference between mother-child and stranger-child dyads.

To ensure that differences between neural and ANS synchrony cannot be attributed to difference in the synchrony estimators, i.e., crosscorrelation vs. WCO, we validated our results by calculating the WCO on the IBI signals (Supplementary Text 4). In line with the results for the cross-correlation, an increased synchrony was observed for cooperation across dyads. Further, a competition x partner interaction indicated



Fig. 4. Mapping of NMF components (HbO) to brain regions. Channels and their positions, projected on a 3D glass brain, are depicted on the top left. The basis matrix is visualized as a heat map, showing the contribution of each fNIRS channel of child (C) and adult partner (P) (x-axis) to the corresponding component (y-axis). The fNIRS channels of child and adult partner which contribute most to each of the components in terms of their nodal densities, with weights above the 80% quantile (min = 0, max = 1), are depicted on the brains below the heatmap.



Fig. 5. Effects of partner and task on interpersonal neural synchrony measured by global and nodal graph metrics (HbO). In addition to the analyses of global density (Fig. 3), marginal posterior distributions were derived for the effects of stranger vs. mother, baseline vs. competition and baseline vs. cooperation on nodal densities, encoded by the coefficients of the four NMF components. Forest plots show the 99 and 90% two-sided credible intervals (thin and thick black lines) as well as the posterior mean (black dot). Evidence of a partner and competition effect was found both globally and in components 3 and 4 (partner) / components 1 and 4 (competition). Further, evidence for a partner effect was found in component 2, however only for the baseline condition. In addition, a cooperative task effect was found in the same components as the competitive task effects, although with weaker evidence. These results show that nodal graph metrics may provide further information on the brain regions which support INS.

that stranger-child dyads had a higher ANS synchrony for competition compared to baseline, while no task effect was observed for motherchild dyads. In addition, stranger-child dyads had a higher ANS synchrony than mother-child dyads in the competition condition, while no partner effect was observed for cooperation or baseline. These results further demonstrated that increased neural synchrony of mother-child compared to stranger-child dyads was unlikely to be explained by increased ANS synchrony alone.

3.1.3. Behavioral synchrony

Task performance was quantified by first calculating how mean response time differed between conditions (Supplementary Text 2 and Table S2). Behavioral synchrony was then measured by calculating the dyad's mean of the absolute differences in response times (Mean-DRT) during cooperation and competition (Table S3). In all conditions, actual pairs were more synchronous than shuffled pairs, although effects were larger for the cooperation conditions (mother-child cooperation: $\mu = -0.43$, CI = [-0.56, -0.31]; stranger-child cooperation: $\mu = -0.45$, CI = [-0.57, -0.33]; mother-child competition: $\mu = -0.25$, CI = [-0.34,

-0.16]; stranger-child competition: μ = -0.21, CI = [-0.30, -0.11]). Thus, these findings showed that reaction times of mother / stranger and child were not independent of each other, i.e., Mean-DRTs of actual pairs were smaller than of shuffled pairs.

Directly comparing the conditions, BHM results yielded strong statistical evidence for a task x partner interaction ($\mu = 0.23$, CI = [0.05, 0.41]) (Table S12). Breaking down the interaction, we found that both mother-child and stranger-child dyads were more synchronous during competition than during cooperation (mother-child: $\mu = -0.32$, CI = [-0.48, -0.17]; stranger-child: $\mu = -0.55$, CI = [-0.65, -0.45]). Yet, stranger-child dyads were more synchronous than mother-child dyads during competition ($\mu = 0.24$, CI = [0.11, 0.37]), while no partner differences were found for cooperation ($\mu = 0.02$, CI = [-0.14, 0.17]).

In a supplementary analysis, we examined whether participants adapted their response times after receiving feedback on who had responded more quickly or more slowly. As expected, dyads adapted their RTs more strongly during cooperation than during competition (task effect: $\mu = 0.36$, CI = [0.32, 0.41], Table S4). No associations between



Fig. 6. PACF averaged across the (A) child's and (B) adults resampled IBI time series after detrending in the conditions: mother-child competition (CompM), mother-child cooperation (CoopStr), stranger-child cooperation (CoopStr), mother-child baseline (BaseM) and stranger-child baseline (BaseStr). Error bars represent standard errors. Across participants, the following mean lower and upper confidence bounds were found: CompM: \pm 0.22; CoopM: \pm 0.24; CompStr: \pm 0.26; CoopStr: \pm 0.21; BaseStr: \pm 0.21.

mean-DRT, the number of adaptations and joint wins with INS or ANS were observed (Supplementary Tables S6–S11).

3.1.4. Summary

To summarize, for research question 1, we analyzed whether interpersonal synchrony was observed in multiple systems: in neural signals, ANS, and motor behavior. While results indicated that synchrony was established in all three systems, they also showed that these different synchrony markers were differentially responsive to experimental manipulation. Importantly, only at the brain level mother-child attunement was observed, while no evidence was found for specific attunement in the mother-child dyads' movements or ANS responses.

3.2. Which factors are related to interpersonal neural synchrony?

For the second research question, we examined whether task and partner effects on neural synchrony were moderated by ANS and behavioral synchrony. Non-parametric Spearman correlations between the different measures are presented in Supplementary Tables S6–S11.

To examine the relationship to ANS synchrony, BHMs were calculated with the main and interactive effects of baseline vs. competition, baseline vs. cooperation as well as stranger vs. mother with ANS synchrony as predictors and INS as response variable (Fig. 7; Table S12). Evidence was found for an interaction of ANS synchrony with baseline vs. competition on global density ($\mu = 0.91$, CI = [0.13, 1.68]), but no sufficient evidence was found for interactions with baseline vs. cooperation and stranger vs. mother. Further analyses of this interaction revealed evidence for an effect of ANS synchrony on global density only for competition ($\mu = 0.65$, CI = [0.21, 1.08]), but not for baseline or cooperation, showing that during competition, higher ANS synchrony predicted increased INS. This should however not be interpreted as a casual or directional effect but rather as an association which could possibly be bidirectional in nature. In line thereof, we also checked the reverse relationship, confirming that increased INS also predicted increased ANS synchrony during competition in our statistical model (Supplementary Text 11).

To examine whether this effect was localized, we conducted a multivariate BHM for nodal density. Evidence for an interaction between ANS synchrony and baseline vs. competition was observed in component 1 ($\mu = 0.88$, CI = [0.05, 1.70]) and component 4 ($\mu = 1.19$, CI = [0.40, 1.99]) (Fig. 7). Again, for baseline vs. cooperation, no interactions were observed with ANS synchrony in any of the components, indicating that increased INS during cooperation was not predicted by increased ANS synchrony. Furthermore, no interactions with partner were found, supporting the notion that increased INS for mother-child compared to stranger-child dyads cannot be attributed to differences in ANS synchrony.

Results for HbR were consistent with the results for HbO, speaking to the validity of the findings (Supplementary Text 10, Table S12). Strong and widespread effects of ANS synchrony on global and nodal density were observed for competition, while no effects were found for baseline or cooperation.

For *behavioral synchrony*, BHMs were calculated with task (cooperation vs. competition), partner (stranger vs. mother) and behavioral synchrony (Mean-DRT) as well as the two-way interactions between task / partner and behavioral synchrony as predictors. For HbO, the BHM showed an interaction between task and behavioral synchrony on global density (μ = 4.02, CI = [1.87, 6.20]; Table S12). Breaking down this interaction, evidence for an effect of behavioral synchrony on INS was found only for competition: less synchronous responses were asso-



Fig. 7. Influences of interpersonal synchrony in the autonomic nervous system (ANS) on neural synchrony (HbO). To investigate whether ANS synchrony predicted increased neural synchrony of mother-child dyads, of competition or cooperation, we examined the interaction effects of ANS synchrony with stranger vs. mother (Partner:ANS), baseline vs. competition (Competition:ANS) and baseline vs. cooperation (Cooperation:ANS). Marginal posterior distributions are depicted for the interaction effects on global and nodal density in the four NMF components. Forest plots show the 99 and 90% two-sided credible intervals (thin and thick black lines) as well as the posterior mean (black dot). Evidence was found for an effect of Competition:ANS on global and nodal density in components 1 and 4. Subsequent analyses showed that only during competition higher density was predicted by higher ANS synchrony. These collective results may indicate that increased interpersonal neural synchrony during competition is related to synchronized arousal, while during cooperation it may go beyond synchrony in ANS signal.

ciated with higher INS (μ = 4.52, CI = [2.45, 6.58]). However, because no evidence was found for an effect of behavioral synchrony on HbR (no interactions with task or partner; Table S12), this finding is not further interpreted. For the effects on nodal density please refer to Table S12.

To summarize, in line with the findings for research question 1, our results showed that increased INS of mother-child dyads was not related to increased behavioral or ANS synchrony. Furthermore, while no relationships between INS, behavioral and ANS synchrony emerged for cooperation, INS and ANS synchrony were positively related during competition.

4. Discussion

In this paper we investigated interpersonal synchrony as a multimodal phenomenon (Hari et al., 2015; Semin, 2007) by applying concurrent fNIRS-ECG hyperscanning recordings as well as including behavioral assessments of motor responses in mother-child and strangerchild dyads. For the first research question, our results showed an increased INS and ANS synchrony during competition and cooperation compared to baseline. However, increased mother-child compared to stranger-child synchrony was found only on the neural level while no partner effects were found for ANS and behavioral synchrony. Further, dyads adapted their response times more strongly and reacted less synchronously during cooperation than during competition. For the second research question, our results indicate that increased INS during cooperation cannot be fully explained by ANS and behavioral synchrony, while during competition a positive relationship between INS and ANS synchrony emerged. Together, these results indicate that synchrony occurs across different systems, that the different biobehavioral synchrony markers are clearly differentiable, and that their relationship may be dependent on context.

Our neural findings are generally consistent with those in a sample of 8-18-year-old male children and adolescents, although analytical methods differed (Kruppa et al., 2021). In this previous study, we found an increased, widespread INS for parent-child competition and more localized effects for parent-child cooperation, however, no increased INS for stranger-child dyads. Here, we examined whether increased INS during competition and cooperation can be attributed to behavioral synchrony, ANS synchrony or other factors. INS was examined at both the global and nodal level.

Based on our results, we are able to rule out a number of possible drivers of the INS that we observed. For example, since INS was higher than in the baseline condition, in which dyads watched a relaxing video together, we rule out the possibility that task-related increases in INS were fully explained by a shared sensory environment. Although we are not able to provide conclusive evidence for a single cause, several possibilities are discussed and evaluated in the light of the present findings.

The first possibility is that aspects of INS may reflect shared social cognitive and attentional processes, including processes of mutual prediction and adaptation (Hamilton, 2020), and the exchange of social ostensive signals (Leong et al., 2017; Wass et al., 2020). In line thereof, we found that dyads adapted their response times based on feedback more strongly during cooperation than during competition. Thus, during cooperation, both adult and child may become entrained to each other as they pay attention to the partner's behavior, continuously predicting the other's actions and adapting their own response times based on feedback provided. However, of note is that we found no associations between INS and mean-DRT, number of adaptations or joint wins during cooperation. Thus, in contrast to previous studies using a similar cooperative game mostly in adults (Baker et al., 2016; Cheng et al., 2015; Cui et al., 2012; Pan et al., 2017), INS was not correlated with cooperative task performance. While during competition, no prediction or adaptation processes are required to successfully complete the task, social comparison processes likely take place during competition, which may potentially lead to an increased INS (see also Kruppa et al., 2021).

Specifically, lateral frontopolar cortex regions have been shown to be involved in relational integration, i.e., comparing and integrating information about self and others (Raposo et al., 2011). Based on the results of the NMF, showing task-related increases mainly in lateralized nodes for HbO, it can be speculated that comparison processes between one's own responses and those of the partner facilitate INS. In addition to such cognitive explanations, in animal studies, INS has been found to emerge from two neuronal populations that separately encoded one's own and the social partner's behavior (Kingsbury et al., 2019). Thus, both top-down and bottom-up processes may potentially play a role in facilitating INS.

The second possibility is that aspects of INS may arise due to synchronized emotional responses. Here, we found evidence that adult's and child's ANS responses become entrained to the task structure (as indicated by the increased autocorrelation every 6 s). Further, consistent with previous studies (Chanel et al., 2012; Järvelä et al., 2014), a significant arousal (ANS) synchrony was observed during both tasks, indicating that cooperation and competition elicit emotional responses in adolescence. However, while no associations between ANS and INS synchrony were found for cooperation, both were positively related during competition, indicating that their association may be dependent on context. In line with our findings for cooperation, a recent hyperscanning study found no relationship between INS and ANS synchrony, measured by respiratory sinus arrhythmia, between 4-6-month-old infants and their mothers when they played with each other (Nguyen et al., 2021). Moreover, a hyperscanning study in adults found significant links between brain, cardiac and electrodermal signals during cooperation within and between the time series of individual subjects within a dyad, yet this study did not explicitly test how different types of synchrony are related to each other (Sciaraffa et al., 2021). Further, both studies did not include a competitive task as a comparison. In the current study, the observed relationship for competition may either be explained by the 'true' relationship between neural and ANS responses (e.g., Critchley et al., 2005, Lane et al., 2009) or by false positives, i.e., influences of the ANS on non-neural hemodynamic changes (e.g., Chen et al., 2020, Tachtsidis and Scholkmann 2016). Speaking against the latter are the arguments that (i) the relationship was only observed during competition and not during cooperation, (ii) it was present both in the HbR and HbO signals even though HbR has been found to be less affected by the systemic physiology and cardiac oscillations are less prominent in the HbR signal (e.g., Kirilina et al., 2013; Tachtsidis and Scholkmann, 2016), and (iii) the fNIRS connectivity estimator did not include the frequency band of the heart rate.

A third possibility is that INS arises as a result of factors unrelated either to shared cognitive and attentional processes, or to shared ANS entrainment to the task structure. The most consistent aspect of our results was the partner effect (mother-child INS > stranger-child INS). This was observed across conditions (baseline, cooperation, competition), but not found for behavioral and / or ANS synchrony. Instead, shared experiences (Gabard-Durnam et al., 2016; Lee et al., 2017), social affiliation (Zheng et al., 2020), as well as genetic influences (Glahn et al., 2010) may lead to higher similarity in brain signals with the mother compared to a stranger. However, since significant heritability is also found for cardiac activities (Muñoz et al., 2018) and motor reaction times (Kuntsi et al., 2006), the latter explanation seems less likely. Further, this finding is consistent with an increasing number of studies showing higher INS in close relationships, including parent-child dyads and romantic partners (Kinreich et al., 2017; Kruppa et al., 2021; Pan et al., 2017; Reindl et al., 2018), and studies indicating that INS may be related to affiliative bonding (Zheng et al., 2020). In line thereof, increased similarities in the resting state network connectome of parent-adolescent child dyads have been related to the dyad's day-to-day emotional synchrony (Lee et al., 2017).

Linked to the question of the source of interpersonal synchrony is the question of the function of different types of synchrony. Interpersonal synchrony is already observed within the first months of life between infant and caretaker when they coordinate their affect, gaze and vocalizations and is accompanied by a coupling in ANS and neural signals (Feldman, 2007; Feldman et al., 2011; Piazza et al., 2020). Thus, it has been hypothesized that one of the main functions of synchrony is to promote social bonds and connections (Wheatley et al., 2012). Further, it may give us access to each other's internal states and allow us to predict each other's behavior, thereby facilitating emotional sharing and corregulation, social understanding and cooperative actions (Semin, 2007; Valdesolo et al., 2010; Wheatley et al., 2012). Yet, the specific and possibly interrelated functions of different types of synchrony (neural, ANS and behavioral synchrony) remains an open question. Of note is that the

question whether INS and ANS synchrony between two interacting persons indeed facilitates certain functions, such as promoting social bonds, cannot be answered by hyperscanning studies alone, but will require study protocols including e.g., multibrain stimulation (Novembre and Iannetti, 2021).

Our study demonstrates the feasibility and utility of multimodal hyperscanning to provide a more holistic view on the neurobiological underpinnings of social interactions. However, one limitation of the study is that our fNIRS set-up focused on the prefrontal cortex, which is important both for social-cognitive and for emotional processing (Amodio and Frith, 2006; Decety et al., 2004; Goldin et al., 2008). With recent improvements in fNIRS hardware, future studies may extend the measurements to cover most of the cerebral cortex. Yet, the inability of fNIRS to measure brain activity in subcortical regions, such as limbic areas, will remain a limitation of the technique. This should also be kept in mind when talking about "global" density, which should not be misinterpreted as an effect across the whole brain. Furthermore, with more recent technological developments it is possible to deduct cardiovascular influences originating from the superficial layers of the head (e.g., the skin) from the fNIRS signal by including short distance measurement channels (Nozawa et al., 2016). Another limitation which should be noted is that mothers were significantly older than strangers, who were mostly university students. Due to the smaller age difference, the stranger may be perceived by some adolescents more as an unfamiliar peer than an unfamiliar mother. To deal with such age differences, we used a block-wise permutation procedure when thresholding the graph holding the condition (task and partner) and channel combination fixed. Future studies may further evaluate the influence of a familiar vs unfamiliar parent and familiar vs unfamiliar peer. Finally, it should be noted that the question of how far findings generalize across tasks, but also across analysis methods and INS/ANS measures, remains an open avenue for future hyperscanning research.

In conclusion, multimodal hyperscanning using concurrent dualbrain and dual-ANS recordings may allow us to validate findings and gain a better understanding of the sources of INS. While our results provide support for models which view interpersonal synchrony as a multimodal phenomenon, they also show that synchrony in different behavioral and biological systems should not be considered as a single common factor or unified construct. Instead, results suggest that synchrony in different systems does not necessarily co-occur; rather, that it may reflect distinct processes and that their functional meaning is likely dependent on context. Importantly, we found that increased INS was observed in mother-child compared with stranger-child dyads across conditions (including baseline), and appeared unrelated to increased ANS or behavioral synchrony. Thus, these findings provide support for recent theories which postulate that INS is higher with "significant" others (Gvirts and Perlmutter, 2020).

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Data and code availability statement

Full results of the Bayesian hierarchical models, including \hat{R} and effective sample size (ESS) indices, are presented in Supplementary Table S12. The raw fNIRS and ECG data of our study may be made available upon request after a confirmation from the ethical committee of our institution. To analyze the data, we used several publicly available Mat-

lab, Python and R functions and toolboxes, which are reported in the Supplementary Information.

Declaration of Competing Interest

The authors declare no competing financial interests.

Credit authorship contribution statement

Vanessa Reindl: Conceptualization, Funding acquisition, Methodology, Investigation, Formal analysis, Writing – original draft. Sam Wass: Methodology, Writing – original draft. Victoria Leong: Methodology, Writing – review & editing. Wolfgang Scharke: Methodology, Writing – review & editing. Sandra Wistuba: Investigation, Methodology, Writing – review & editing. Christina Lisa Wirth: Investigation, Writing – review & editing. Kerstin Konrad: Conceptualization, Funding acquisition, Writing – review & editing. Christian Gerloff: Conceptualization, Funding acquisition, Methodology, Formal analysis, Writing – original draft.

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Supplementary materials

Supplementary material associated with this article can be found, in the online version, at doi:10.1016/j.neuroimage.2022.118982.

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