Structural and Functional Trajectories of Middle Temporal Gyrus Sub-Regions During Life Span: A Potential Biomarker of Brain Development and Aging

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Although previous studies identified a similar topography pattern of structural and functional delineations in human middle temporal gyrus (MTG) using healthy adults, trajectories of MTG sub-regions across lifespan remain largely unknown. Herein, we examined gray matter volume (GMV) and resting-state functional connectivity (RSFC) using datasets from the Nathan Kline Institute (NKI), and aimed to (1) investigate structural and functional trajectories of MTG sub-regions across the lifespan; and (2) assess whether these features can be used as biomarkers to predict individual’s chronological age. As a result, GMV of all MTG sub-regions followed U-shaped trajectories with extreme age around the sixth decade. The RSFC between MTG sub-regions and many cortical brain regions showed inversed U-shaped trajectories, whereas RSFC between MTG sub-regions and sub-cortical regions/cerebellum showed U-shaped way, with extreme age about 20 years earlier than those of GMV. Moreover, GMV and RSFC of MTG sub-regions could be served as useful features to predict individual age with high estimation accuracy. Together, these results not only provided novel insights into the dynamic process of structural and functional roles of MTG sub-regions across the lifespan, but also served as useful biomarkers to age prediction.

Keywords: middle temporal gyrus, gray matter volume, resting-state functional connectivity, regression model, deep learning network

INTRODUCTION

The middle temporal gyrus (MTG) is suggested to be involved in various functions (Giraud et al., 2004; Sato et al., 2012) with several distinct sub-regions (Sewards, 2011). Recently, a consistent topography pattern of structural and functional delineations in MTG was identified in our previous studies (Xu et al., 2015, 2019a), resulting in four distinct clusters, namely, the anterior part of MTG (aMTG), the middle part of MTG (mMTG), the posterior part of MTG (pMTG), and the sulcus part of MTG (sMTG). Although structural and functional roles of MTG were decoded
in the healthy adults (21–25 years), accumulating evidence of significant age-related effects on structural and functional patterns of MTG was identified. Previous studies based on functional MRI (Blumenfeld et al., 2006; Chou et al., 2006) showed age-related brain activation in left MTG, whereas one similar study showed no age effect (Moore-Parks et al., 2010). Using other functional neuroimaging technologies, such as magnetoencephalography, old adults exhibited both declined cortical activities and delayed activation in MTG as compared to younger adults (Lin et al., 2018). Moreover, two graph theory studies (Cao et al., 2014; Zhao et al., 2015) also showed inverted U-shaped trajectories of nodal degree and functional connectivity strength in the MTG across the lifespan (7–85 years). Structurally, the thickness of left MTG showed a negative relationship with age (10–60 years) (Raznahan et al., 2010), and the gray matter volume (GMV) of MTG showed a dramatic decline from 6 to 26 years (Mu et al., 2017). In addition to healthy individuals, patients with reading impairment presented a delay in the development of left MTG in the children group (Luniewska et al., 2019). Moreover, our recent study revealed specific structural and functional patterns of MTG sub-regions in children and adults with autism spectral disorders (Xu et al., 2019b). Together, these studies provided enough evidence to indicate that the structural and functional roles of MTG are dynamic processes with remarkable changes across the lifespan (Fair et al., 2008; Zhao et al., 2015; Li et al., 2019). Thus, investigating age-related trajectories of MTG sub-regions across the lifespan is critical and helpful to guide the interpretation of alterations associated with these regions in many brain disorders.

To date, many studies have been performed to investigate lifespan trajectory about gray matter (Gennatas et al., 2017; Mu et al., 2017), white matter (Ziegler et al., 2012; Zhao et al., 2015), cortical morphology (Sowell et al., 2003; Khundrakpam et al., 2015), and topography of structural and functional brain networks (Fair et al., 2008; Betzel et al., 2014; Li et al., 2019), revealing that the structure and function of the human brain undergo complex changes across the lifespan. Moreover, several studies have been performed to characterize age-related effects on functional connectome (Wang et al., 2012; Cao et al., 2014; Yang et al., 2014; Jiang et al., 2019) using the same dataset (NKI sample). For example, Cao et al. (2014) observed that the human brain functional connectome not only exhibited highly preserved nonrandom modular organization, but also showed linearly decreases in modularity, inverted-U-shaped trajectories of local efficiency, and inverted-U-shaped trajectories of rich club architecture over the entire age range. Wang et al. (2012) showed a linear increase in the emotion system and a decrease in the sensorimotor system during brain development from childhood to senescence. Yang et al. (2014) observed stronger age dependence in the spatial pattern of a precuneus dorsal–posterior cingulate cortex network as compared to the default network. However, little attention was paid to the lifespan changes of particular regions at the sub-regional level. Therefore, our study aimed to investigate the structural and functional lifespan trajectory of MTG at the sub-regional level in healthy individuals.

Moreover, age prediction is considered to be an important way to understand brain developmental and degenerative processes in healthy individuals (Zhao et al., 2019). Many brain structural and functional features are sensitive to maturational processes throughout the lifespan, such as cortical thickness (Khundrakpam et al., 2015; Lewis et al., 2018), GMV (Franke et al., 2010; Mohajer et al., 2020), resting-state functional connectivity (RSFC) (Ciesielski et al., 2019; Li et al., 2019), brain network topography (Cao et al., 2014; Zhao et al., 2015), as well as multi-modal MRI features (Betzel et al., 2014; Zhao et al., 2019). However, these studies mainly focused on extracting different features to increase the accuracy of age prediction using traditional machine learning techniques (i.e., non-deep learning algorithms) rather than improving prediction models. Recently, the Long Short-Term Memory (LSTM) deep learning model (Hochreiter and Schmidhuber, 1997), an evolution over the Recurrent Neural Networks (RNN), has been widely adopted in clinical predictions, such as epileptic seizures (Tsouris et al., 2018; Wei et al., 2019), heart failure (Maragatham and Devi, 2019), cancer survival outcomes (Koo et al., 2020), and even mortality of patients (Xia et al., 2019). This method succeeded better performance than traditional ways (Tsouris et al., 2018; Wei et al., 2019; Xia et al., 2019), suggesting potential applications to increase the accuracy of age prediction.

In the current study, we aimed to (1) investigate structural and functional trajectories of MTG sub-regions across the lifespan; and (2) assess whether these features can be used as biomarkers to accurately predict an individual’s chronological age. To uncover these two questions, we used T1 structural and resting-state fMRI data from the publicly available Nathan Kline Institute-Rockland Sample (NKI-RS) dataset (7–85 years). The GMV and RSFC patterns of each MTG sub-regions were calculated and subsequently used to explore their relationship with age using linear and quadratic regression models. Finally, the structural and functional features of MTG were used as inputs for LSTM to predict an individual’s chronological age.

**MATERIALS AND METHODS**

**Participants**

Data used in this study are publicly available at the International Neuroimaging Data-sharing Initiative (INDI) (see text footnote 1) from the Nathan Kline Institute (NKI, NY, United States). The initial dataset included 207 participants. The inclusion criteria were (1) aged 7–85 years; (2) without any lesions in brain scans; (3) no problems in semi-structured diagnostic psychiatric interviews; and (4) normal in psychiatric assessments. We also excluded participants who were (1) left-handed; (2) missing demographic data; (3) with bad image integrity and quality on screening; and (4) only T1 or fMRI images. Finally, 160 subjects were included in the structural analyses and functional analyses. These subjects include 25 children and adolescents (10 female / 15 male, 22 right / 3 left-handedness, age range of 7–18 years), 48 young adults (22 female / 26 male, 43 right / 5 left-handedness, age range of 19–30 years), 37 middle-aged adults (11 female / 26 male, 31 right / 6 left-handedness, age range of

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1http://icon_1000.projects.nitrc.org/indi/pro/nki.html#LastRelease
31–45 years), 22 older adults (7 female / 15 male, 20 right / 2 left-handedness, age range of 46–60 years), and 28 elders (12 female / 16 male, 24 right / 4 left-handedness, age range of 61–85 years). This information is given in Supplementary Table 1. Informed consent was obtained from all subjects, and the study was approved by the NKI institutional review board (Nooner et al., 2012). Moreover, this study was performed in accordance with the Declaration of Helsinki.

MRI Acquisition

Magnetic resonance imaging scanning was performed using a Siemens 3.0 T Trio Tim MRI scanner. During scanning, subjects were told to keep their eyes closed, relax their minds, and to not move. T1 images were obtained using the magnetization-prepared rapid gradient echo (MPRAGE) sequence: time repetition (TR) / time echo (TE) = 2,500 / 3.5 ms, inversion time = 1,200 ms, flip angle (FA) = 8°, field of view (FOV) = 256 mm × 256 mm, voxel size = 1.0 mm × 1.0 mm × 1.0 mm, and number of slices = 192. Resting-state fMRI scans were collected using an echo-planar imaging (EPI) sequence: TR / TE = 2,500 / 30 ms, FA = 80°, FOV = 216 mm × 216 mm, voxel size = 3.0 mm × 3.0 mm × 3.0 mm, number of slices = 38, and 260 volumes.

Image Preprocessing

The T1 images were preprocessed using a toolbox for Data Processing & Analysis for Brain Imaging (DPABI)⁴. First, the quality of each image was visually checked and 30 subjects were excluded for really bad quality. Then, the remaining T1 images were segmented into gray matter, white matter, and cerebrospinal fluid, and transformed to a standard Montreal Neurological Institute space. Next, these images were modulated to preserve regional volume information. Finally, the modulated gray matter images were smoothed with a Gaussian kernel of 6 mm full-width at half maximum and used in the following analyses.

The resting-state fMRI data were also preprocessed using DPABI. The main steps included the following: (1) removing the first 10 volumes; (2) slice timing; (3) realigning (subjects with head motion exceeding 3 mm in any dimension or 3° of angular motion were removed, and 20 subjects were excluded); (4) spatially normalizing; (5) resampling to a voxel size of 3 mm × 3 mm × 3 mm; (6) smoothing using a Gaussian kernel of 6 mm full-width at half maximum; (7) removing linear and quadratic trends; (8) regressing out head motion effects using the Friston 24-parameter model (Satterthwaite et al., 2013), as well as the white matter, cerebrospinal fluid, and global signals; (9) temporal band-pass filtering (0.01–0.1 Hz); and (10) “scrubbed” two-time points before and one time point after bad images, whose frame displacement (FD) > 0.5 (Power et al., 2012).

Definition of Middle Temporal Gyrus Sub-Regions

The MTG sub-regions were obtained from our previous results (Xu et al., 2019a), which were defined as the overlap between functional and anatomical parcellation results of MTG (Supplementary Figure 1). All the MTG sub-regions were then resampled to 1.5 mm × 1.5 mm × 1.5 mm for structural analyses and 3 mm × 3 mm × 3 mm for RSFC analyses.

Calculations of Structural and Functional Indexes

Structurally, the GMV was defined as the mean value of each MTG sub-region in the modulated and smoothed gray matter map. The analysis of covariance (ANCOVA) was performed to identify between-group differences. The significant levels were set at P < 0.05 with Bonferroni correction.

The RSFC was defined as the Pearson correlation coefficients between the mean time series of each MTG sub-region and that of each voxel in the rest of the brain, respectively. Then, all correlation coefficients were converted to z values using Fisher’s z transformation to improve normality.

Age Effects on Regional Gray Matter Volume

To determine how age affects GMV of MTG sub-regions, both linear and quadric regression models were used controlling for covariates of gender and handedness. The significant level was set at p < 0.05/8 = 0.00625. The models for GMV can be expressed with the following equations:

Y = β0 + β1 × age + β2 × gender + β3 × handedness (linear);

Y = β0 + β1 × age + β2 × age² + β3 × gender + β4 × handedness (quadric).

Then, Akaike’s information criterion (AIC) was used to determine the best-fitting model. The extreme age along the quadric trajectory is important to the investigation of timing differences in maturation or degeneration across the lifespan, and could be calculated using age = -β1/2β2. The fitted GMV was defined as the residual of GMV regressed out gender, handedness, and age.

Age Effects on Resting-State Functional Connectivity

To explore the corresponding quadric relationship between age and RSFC of MTG sub-regions, we performed correlation analyses in the DPABI using age² as an independent variable, and gender, handedness, age, and FD as covariates. The significance level was determined using the Gaussian random field corrections with a voxel-level threshold of p < 0.001 and a cluster-level threshold of p < 0.05.

For the regions which showed a significant quadric relationship with age, we calculated their mean RSFC and performed quadric regression models controlling for covariates of gender, handedness, and FD. The model can be expressed by:

Y = β0 + β1 × age + β2 × age² + β3 × gender + β4 × handedness

http://rfmri.org/dpabi

Supplementary Table 1

| Age Group | Gender Distribution | Number of Subjects |
|-----------|---------------------|--------------------|
| 31–45 years | 7 female, 15 male, 20 right / 2 left-handed | 22 |
| 46–60 years | 12 female, 16 male, 24 right / 4 left-handed | 28 |
| 61–85 years | 12 female, 16 male, 24 right / 4 left-handed | 28 |

Supplementary Figure 1

The resting-state fMRI data were also preprocessed using DPABI. The main steps included the following: (1) removing the first 10 volumes; (2) slice timing; (3) realigning (subjects with head motion exceeding 3 mm in any dimension or 3° of angular motion were removed, and 20 subjects were excluded); (4) spatially normalizing; (5) resampling to a voxel size of 3 mm × 3 mm × 3 mm; (6) smoothing using a Gaussian kernel of 6 mm full-width at half maximum; (7) removing linear and quadratic trends; (8) regressing out head motion effects using the Friston 24-parameter model (Satterthwaite et al., 2013), as well as the white matter, cerebrospinal fluid, and global signals; (9) temporal band-pass filtering (0.01–0.1 Hz); and (10) “scrubbed” two-time points before and one time point after bad images, whose frame displacement (FD) > 0.5 (Power et al., 2012).

Definition of Middle Temporal Gyrus Sub-Regions

The MTG sub-regions were obtained from our previous results (Xu et al., 2019a), which were defined as the overlap between functional and anatomical parcellation results of MTG (Supplementary Figure 1). All the MTG sub-regions were then resampled to 1.5 mm × 1.5 mm × 1.5 mm for structural analyses and 3 mm × 3 mm × 3 mm for RSFC analyses.
FIGURE 1 | Quadratic relationships between gray matter volume (GMV) of middle temporal gyrus (MTG) sub-regions and age across lifespan. The GMV was fitted by controlling for gender and handedness. The analysis of covariance (ANCOVA) was performed to explore group differences (* represents significant difference).

\[ \beta_5 \times \text{FD} \] The significant level was set at \( p < 0.05/15 = 0.003 \). The extreme age was also calculated using \( \text{Age} = -\beta_1/2\beta_2 \). The fitted RSFC was defined as the residual of RSFC regressed out the gender, handedness, age, and FD.

Long Short-Term Memory Network to Predict Individual Age

We adopted a commonly used LSTM network (Tsiouris et al., 2018; Liu and Gong, 2019; Supplementary Figure 2) to predict individual age using 23 features, including GMV of all MTG sub-regions and all RSFC listed in Table 1. All features are standardized to 0–1 and served as inputs for a simple 4-layer LSTM network with 40 hidden units. Each hidden unit has self-memory and was fully connected. Especially, it is composed of an input gate \( i \), an output gate \( h \), and a forgetting gate \( f \), which were defined by the equations in the right bottom of Figure 2, respectively. The output layer was composed of a sigmoid function. In the initial model, the batch size was set to 10, and the training rate was set to 0.006 for the Adaptive Moment Estimation (Adam) optimizer, which used the Root-Mean-Square Error (RMSE = \( \sqrt{\frac{1}{m} \sum_{i=1}^{m} (y_i - \hat{y}_i)^2} \), \( m = 1, 2n \)) as loss function. Moreover, we used leave-one-out cross-validations to evaluate the model by 1,000 times. To assess prediction accuracy, the Pearson correlation coefficient and the Mean Absolute Error (MAE) between the actual and predicted ages were calculated. The LSTM networks were built using Keras 2.2.4 upon Tensorflow 1.12.0 backend in Python 3.6.

RESULTS

Age Effects on Regional Gray Matter Volume

Controlling for effects of gender and handedness, the fitted GMV of all MTG sub-regions exhibited U-shaped trajectories with troughs nearly 60 years (Figure 1).

Age Effects on Resting-State Functional Connectivity

Resting-state functional connectivity between MTG sub-regions and most cortical brain regions showed inversed quadric relationships with age, but RSFC between MTG sub-regions and
sub-cortical brain regions/cerebellum showed the opposite way (Figure 2A and Table 1).

Further quadratic regression models showed significantly U-shaped or inverse U-shaped trajectories between fitted RSFC of MTG sub-regions and age with extreme age nearly 40 years (Figure 2B). Especially, RSFC between aMTG.L and left medial frontal gyrus (MFG.L)/ right vermis (VER.R) exhibited U-shaped trajectories with age. RSFC between mMTG.L and right operculum part of inferior frontal gyrus (opercFG.R) showed inverted U-shaped trajectory with age. RSFC between sMTG.L and left anterior cingulate cortex (ACC.L)/ left putamen (PUT.L)/ right amygdala (AMG.R) showed U-shaped trajectories, whereas RSFC between sMTG.L and left post-central gyrus (PostCG.L)/ right post-central gyrus (PostCG.R) showed inverse way.

Resting-state functional connectivity between aMTG.R and right pole part of temporal gyrus (pSTG.R) exhibited U-shaped trajectory with age. RSFC between pMTG.R and left caudate (CAU.L)/ right superior temporal gyrus (STG.R) showed U-shaped trajectories with age. RSFC between sMTG.R and ACC.L/right pallidum (PAL.R) showed U-shaped trajectories,
TABLE 1 | Brain regions that showed a quadratic relationship between RSFC of MTG sub-regions and age.

| Seed regions | Brain regions                                      | Abbreviations | Cluster size | Peak coordinates  | Peak intensity |
|--------------|--------------------------------------------------|---------------|--------------|------------------|---------------|
| aMTG.L       | Left medial frontal gyrus                        | MFG.L         | 112          | (−18, 21, −6)    | 0.36222       |
| Right vermis |                                                   | VER.R         | 111          | (6, −45, −3)     | 0.3867        |
| mMTG.L       | Right operculum part of inferior frontal gyrus   | operIFG.R     | 89           | (36, 12, 27)     | −0.3259       |
| sMTG.L       | Left anterior cingulate cortex                   | ACC.L         | 82           | (−12, 30, −9)    | 0.3710        |
| Right amygdala| AMG.R                                            |               |              |                  |               |
| Left putamen  | PUT.L                                            |               |              |                  |               |
| Right postcentral gyrus | PostCG.R | 249 | (36, −15, 36) | −0.3849       |
| Left postcentral gyrus | PostCG.L | 296 | (−39, −12, 33) | −0.3575       |
| Right pole part of superior temporal gyrus  | pSTG.R                                             | 54            | (42, 18, −24) | 0.3443        |
| Right superior temporal gyrus | STG.R | 46 | (51, −21, 0) | 0.3521        |
| Left caudate  | CAU.L                                            |               |              |                  |               |
| Right pallidum| PAL.R                                             | 211           | (21, −6, −6) | 0.3789        |
| Left anterior cingulate cortex | ACC.L | 150 | (−9, 33, 0) | 0.3408        |
| Left precentral gyrus | PreCG.L | 1038 | (−36, −18, 51) | −0.4014       |
| Right precentral gyrus | PreCG.R | 516 | (60, −9, 45) | −0.3964       |

whereas RSFC between sMTG.R and left precentral gyrus (PreCG.L) and right precentral gyrus (PreCG.R) showed inverse way.

Maturation Age

We also calculated the extreme age and identified that the maturation ages of RSFC in all MTG sub-regions were nearly 20 years earlier than those of GMV (Figure 3).

Age Predictions

When all 23 features (Figure 4A) were combined as input for the LSTM network to predict individual age, a high estimation accuracy was obtained (MAE = 3.89, r = 0.95, p = 9.13 × 10^{−64}) between predicted versus actual ages (Figure 4B).

DISCUSSION

In the current study, we (1) examined GMV and RSFC of each MTG sub-region; (2) performed both linear and quadratic regression models to investigate structural and functional trajectories of MTG sub-regions across the lifespan, as well as ANCOVA to examine between-group differences; and (3) adopted LSTM to assess whether these features can be used to accurately predict an individual’s chronological age. As a result, we identified that GMV of all MTG sub-regions showed U-shaped trajectories with extreme age around the sixth decade, whereas RSFC between MTG sub-regions and most cortical brain regions showed inverted U-shaped trajectories, but RSFC between MTG sub-regions and sub-cortical regions/cerebellum showed U-shaped way. Further prediction models indicated that GMV and RSFC of MTG sub-regions could be served as biomarkers to predict individual age with extremely high estimation accuracy.

Structurally, we identified that GMV of all MTG sub-regions showed U-shaped trajectories with troughs around the sixth decade, suggesting a nonlinear decline between 7 and 60 years and a slight increase after that. These trajectories were similar to one previous study, which reported significant U-shaped trajectories across 7–85 years in whole-brain total GMV (Sowell et al., 2003), as well as gray matter density in dorsal frontal and parietal association cortices. Moreover, some developmental neuroimaging studies also supported our results by showing modest decreases in GMV in almost all cortical regions during adolescence (Sowell et al., 2004; Brain Development Cooperative Group, 2012; Gennatas et al., 2017), which is considered to be generally attributed to a combination of synaptic pruning of exuberant connections and increasing myelination, two essential aspects of cortical development (Stiles and Jernigan, 2010). However, different brain structures showed unique trajectories across the lifespan. For example, the basal ganglia generally follow linear decreases of volume over the age of 4–18 years (Brain Development Cooperative Group, 2012), 8–30 years (Ostby et al., 2009), 18–94 years (Fjell et al., 2013), as well as in a longitudinal study over the age of 22 years (Tamnes et al., 2013), whereas hippocampus indicates an inverted U exponential volumetric trajectory from childhood to puberty, peaking at a later age than the basal ganglia (Wierenga et al., 2014). Moreover, an entire...
lifespan study reported that the absolute global GMV follows a complex trajectory with four phases: (1) rapid increase from 0 to 8–10 years, (2) rapid decrease until 40 years, (3) a plateau from 40 to 80 years, and (4) a rapid decrease after 80 years (Coupe et al., 2017). The rapid decrease from 8 to 40 years is similar to the trajectory of MTG sub-regions to some extent. However, this cannot be used as direct evidence to support our results for the following two reasons: (1) trajectory of maturational and aging effects varied considerably over the brain cortex, with an earlier maturation of sensory, motor, and limbic regions relative to association regions (Sowell et al., 2003; Gogtay et al., 2004; Shaw et al., 2008); and (2) the gray matter reductions were markedly greater than the subcortical changes (Sullivan et al., 2011; Tamnes et al., 2013). Anyway, our results of U-shaped trajectories of MTG sub-regions extended their findings and offered detailed information for MTG maturations at the subregional level, which might be helpful to guide interpretation of alterations associated with these regions in many brain disorders.

Functionally, we also identified that RSFC between MTG sub-regions and cortical brain regions showed increased U-shaped trajectories, but RSFC between MTG sub-regions and subcortical regions/cerebellum showed U-shaped way. These results were supported by previous findings of the rewiring and pruning of subcortical–cortical connectivity accompanied by increased cortico-cortical connectivity during development (Supekar et al., 2009), as well as increased correlation strength of subcortical–cortical connectivity but decreased cortico-cortical connectivity during aging (Tomasi and Volkow, 2012). Moreover, these results were also similar to one previous study based on fMRI, which showed inverse U-shaped trajectories between regional nodal properties for cortical regions (e.g., MTG.L, PreCG.L) and age (7–85 years) but U-shaped way for subcortical regions (e.g., HIP.L) (Cao et al., 2014). Interestingly, the maturation ages of RSFC were about 20 years earlier than those of GMV, with about 40 years. On one hand, this aligns well with the finding of age-related U-shaped trajectory of the normalized rich-club coefficient, as well as functional connectivity strength and proportions, with maturation age of approximately 40 years (Cao et al., 2014). On the other hand, our results further supported the suggestion that development and degeneration of structural and functional connectivity is not synchronous (Horn et al., 2014).

To our knowledge, it is the first illustration that LSTM networks can be used to effectively estimate chronological age based on MRI features although it has been widely adopted in other clinical predictions (Wei et al., 2019; Koo et al., 2020). As expected, our trained LSTM model showed relatively higher accuracy in age prediction with a correlation between chronological and predicted age of $r = 0.95$ (MEA = 3.89, $p = 9.13 \times 10^{-64}$). This performance was superior to most previous studies using traditional machine learning methods, such as support vector regression (Wang et al., 2012; Erus et al., 2015; Zhao et al., 2015), least absolute shrinkage and selection operator (Varikuti et al., 2018), kernel regression methods (Franke et al., 2012), ridge regression model (Zhao et al., 2019), and elastic net penalized linear regression model (Khundrakpam et al., 2015; Lewis et al., 2018), among which the best performance was $r = 0.93$. Other deep learning methods using convolutional neural network based on the gray matter (Wang et al., 2019) and raw imaging data (Cole et al., 2017) resulted in slightly worse performances of MAE = 4.45, and MAE = 4.16, respectively, in age predictions. Comparable to our results, a neuroanatomical model, including 231 variables derived from multiple imaging modalities, resulted in a correlation of $r = 0.961$ between chronological and predicted age and a mean prediction error

**FIGURE 4** | Prediction results based on features from MTG sub-regions using LSTM. (A) Features are used to predict an individual’s age in the LSTM network. The scatter plot depicts actual age versus predicted age by all features. (B) Pearson correlation analyses between the actual and predicted ages are performed to evaluate the prediction accuracy. Abbreviations of brain regions are listed in Table 1.
across all ages of just 1.03 years (Brown et al., 2012). However, this study included typically developing individuals between 3 and 20 years rather than across lifespan (7–85 years) and more variables as compared to ours. Moreover, our results indicated that GMV and RSFC of MTG sub-regions could be served as biomarkers to predict individual age. Similar to our results, many previous studies confirmed that the whole-brain GMV (Cole et al., 2017; Varikuti et al., 2018; Wang et al., 2019) and RSFC (Wang et al., 2012), as well as multi-modal MRI features (Franke et al., 2012), were sensitive to age predictions. However, these features were based on the whole-brain level. Our results further extended previous studies by identifying features to particular brain regions at the sub-regional level.

Several limitations should be addressed. First, only GMV and RSFC were examined in the current study, additional future studies using other neuroimaging techniques, such as diffusion tensor imaging, to evaluate the lifespan trajectory of white matter fibers originated from MTG sub-regions are warranted. Second, no behavioral variables were included to measure cognitive and language abilities, which were two important functions associated with MTG (Whitney et al., 2011). Thus, brain behavior associations are missing, weakening the interpretation of our results. Finally, we only included participants from 7 to 85 years of age, investigating earlier developmental changes during infancy and early childhood is also significantly important.

CONCLUSION

We identified specific trajectories for GMV and RSFC of MTG sub-regions across the lifespan. These results not only offered detailed information for MTG maturations and degeneration at the sub-regional level but also might be helpful to guide the interpretation of alterations associated with these regions in many brain disorders. Finally, LSTM networks based on GMV and RSFC of MTG sub-regions can accurately predict chronological age in healthy individuals. This can be achieved using features of particular brain regions, substantially reducing computational complexity to extract features from the whole-brain level.

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DATA AVAILABILITY STATEMENT

Data used in this study are publicly available at the International Neuroimaging Datasharing Initiative (INDI) (http://Icon_1000.projects.nitrc.org/indi/pro/nki.html#LastRelease) from the Nathan Kline Institute (NKI, NY, United States).

ETHICS STATEMENT

The studies involving human participants were reviewed and approved by NKI institutional review board. Written informed consent to participate in this study was provided by the participants’ legal guardian/next of kin.

AUTHOR CONTRIBUTIONS

JX, QH, and HL designed the study. JZ and JC downloaded the data. JX, JZ, JL, and HW analyzed the data. JX wrote the manuscript. All authors reviewed the manuscript.

FUNDING

This work was supported by the National Natural Science Foundation of China (Nos. 62001462 and 62006220), the National key research program (No. 2018YFB1105600), and the Shenzhen Science and Technology Research Program (Nos. JCYJ20180507182441903 and JCYJ20200109114816594).

SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fnagi.2022.799260/full#supplementary-material

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