Prosocial behavior relates to the rate and timing of cortical thinning from adolescence to young adulthood

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ARTICLE INFO
Keywords:
Adolescence
Cortical thickness
Development
Longitudinal
Prosocial behavior
Structural MRI

ABSTRACT
Prosocial behavior, or voluntary actions that intentionally benefit others, relate to desirable developmental outcomes such as peer acceptance, while lack of prosocial behavior has been associated with several neurodevelopmental disorders. Mapping the biological foundations of prosociality may thus aid our understanding of both normal and abnormal development, yet how prosociality relates to cortical development is largely unknown. Here, relations between prosociality, as measured by the Strengths and Difficulties Questionnaire (self-report), and changes in thickness across the cortical mantle were examined using mixed-effects models. The sample consisted of 169 healthy individuals (92 females) aged 12–26 with repeated MRI from up to 3 time points, at approximately 3-year intervals (301 scans). In regions associated with social cognition and behavioral control, higher prosociality was associated with greater cortical thinning during early-to-middle adolescence, followed by attenuation of this process during the transition to young adulthood. Comparatively, lower prosociality was related to initially slower thinning, followed by comparatively protracted thinning into the mid-twenties. This study showed that prosocial behavior is associated with regional development of cortical thickness in adolescence and young adulthood. The results suggest that the rate of thinning in these regions, as well as its timing, may be factors related to prosocial behavior.

1. Introduction
Prosociality refers to voluntary actions that intentionally benefit others (Eisenberg et al., 2015). Basic forms of prosociality emerge early in childhood, but prosocial behaviors continue to develop throughout childhood and adolescence and into early adulthood (Eisenberg et al., 2002, 2015; Warneken, 2016). High prosociality has been associated with greater peer acceptance and academic achievement (Caprara et al., 2000), and low prosociality with several developmental disorders, e.g. conduct disorder (CD; Hay et al., 2010). Mapping the brain correlates of prosociality may thus aid our understanding of the neurobiological foundations of both normal and abnormal development.

Theoretical models suggest that prosociality may rely on several neural systems. Firstly, a major source of the increase in prosociality is development of social cognitive skills, e.g. Theory of Mind (ToM; Eisenberg et al., 2015; Kuhnert et al., 2017). Modest, but consistent associations between ToM and prosociality exist and increase in strength with age (Imuta et al., 2016). ToM also predicts prosocial behavior...
prospectively (Caputi et al., 2012; Kuhnert et al., 2017). Secondly, in some situations, prosociality may be evolutionarily adaptive for functioning in social groups (Nowak, 2006). However, excessive sharing and helping can also put a strain on individual’s resources and may thus be maladaptive. The costs of being prosocial must be weighed against the potential benefits, a process requiring value-based decision-making and behavioral control (Steinbeis, 2018). Third, adaptive prosociality may depend on the dynamic interplay of socio-cognitive (e.g. ToM) and socio-affective processes (e.g. empathy, compassion; Preckel et al., 2018; Tusche et al., 2016).

Functional and structural imaging studies support these theoretical models. From childhood to adulthood, functional studies have related prosocial behaviors, e.g. helping and sharing, to activity in regions involved in: (1) social cognition, e.g. medial prefrontal cortex (mPFC), tempo-parietal junction (TPJ), precuneus, superior temporal sulcus (STS; Schreuters et al., 2015; van der Meulen et al., 2016; Van Hoorn et al., 2016; Zanon et al., 2014), (2) behavioral control and decision making, e.g. doro-lateral (dIPFC; Steinbeis et al., 2012; Telzer et al., 2011), and (3) empathy, e.g. anterior cingulate cortex (ACC) and anterior insula (Ai; Hein et al., 2016; Hein et al., 2010). The ACC also contains Von Economo neurons, known to be involved in social cognition (Allman et al., 2005).

Two studies have examined prosociality and brain structure. In children aged 6–9 years, mostly positive associations were observed between prosociality and cortical thickness (CT) in regions involved in (1) social cognition, e.g. anterior temporal lobe, mPFC, cuneus and precuneus, (2) behavioral control, e.g. dIPFC, and (3) empathy, e.g. pars orbitalis (Thijssen et al., 2015; Wildeboer et al., 2017). These relations were predominantly identified in the right hemisphere for both males and females, except for a few regions where these associations were present in males alone (Thijssen et al., 2015). To our knowledge, no studies have examined prosocial behavior and structural change longitudinally across adolescence. As adolescence is a period characterized by changes in social cognition and peer relations, and marked maturation of associated brain structure and function (Foulkes and Blakemore, 2018), such longitudinal studies would arguably be highly informative for understanding the development of prosocial behaviors.

The aim of the current study was to examine how prosociality relates to the longitudinal development of CT in adolescence and young adulthood. Normative cortical development from childhood to adulthood is characterized by regionally variable decreases in CT as derived from MRI (Tannes et al., 2017; Vijayakumar et al., 2016; Wierenga et al., 2014; Zhou et al., 2015). Here, we used the terms cortical thickness (CT) and cortical thinning, but it is in principle possible that the cortex in development is not thinning (further discussion in Supporting Information (S1.1)). Although thinning during development is not uniformly related to adaptive outcomes (Box et al., 2018; Jirsaare et al., 2018; Luby et al., 2018; Román et al., 2018), studies have associated greater cortical thinning in adolescence with reduced neuroticism and aggressive behavior, greater intellectual ability and more adaptive emotion regulation (Ferschmann et al., 2018; Shaw et al., 2006; Vijayakumar et al., 2014a, 2014b, 2014c). Moreover, delayed and slower rates of cortical maturation have been related to symptoms of anxiety and depression in healthy individuals (Ducharme et al., 2013), and ADHD (Shaw et al., 2011). Since the relations between CT trajectories and developmental outcomes may be regionally specific (Fandakova et al., 2017; Vijayakumar et al., 2014a) and since there is some evidence that the rate of CT change is not stable from childhood to adulthood (Zhou et al., 2015), it is possible that a crucial factor in explaining the effects of the rate of cortical thinning may be its timing. Taken together, we therefore hypothesized that 1) higher levels of prosociality would be associated with greater age-expected thinning as many studies linked this developmental pattern to adaptive behaviors, and 2) that the associations would be region-specific. More specifically, given the role of social-cognitive and social-affective processes in prosociality, we also hypothesized that prosociality would be associated with regions implicated in ToM (Blakemore, 2012; Frith and Frith, 2003): mPFC, temporal poles, posterior STS, and TPJ, as well as regions related to empathy, i.e. AI and ACC (Lamm et al., 2011). Moreover, given the crucial role of behavior control in prosociality, we hypothesized that prosociality would be related to maturation of cortical regions associated with behavioral control (Hare et al., 2009; Steinbeis et al., 2014), i.e. dlPFC and ventromedial prefrontal (vmPFC) cortices.

2. Methods

2.1. Participants

The sample was drawn from NeuroCognitive Development (Tannes et al., 2018, 2013; Tannes et al., 2010), an accelerated longitudinal project with three waves, see Supporting Information (S2.1) for details. Table 1 describes the sample; 64 individuals provided data from one time point (TP), 78 from 2 TPs and 27 from 3 TPs.

In total 301 observations were available for individuals who had at least one TP with MRI data of sufficient quality (see 2.4) and self-reported prosocial behavior. For each individual, we included only those TPs where both MRI and prosociality data were available. Table 2 provides an overview of sample characteristics. Refer to Supporting Information for further detail, including attrition analyses (S2.1).

2.2. Prosocial behavior

Prosociality was assessed using the 5-item self-report prosocial subscale of the Strengths and Difficulties Questionnaire (Goodman, 1997). The items assessing different aspects of prosociality (e.g. “I usually share with others”, “I often volunteer to help others”) are rated on a 3-point Likert Scale as “not true”, “somewhat true” and “certainly true”. Mean inter-item correlation for this scale was 0.23, a value within the recommended range (Briggs and Cheek, 1986). See Supporting Information (S2.2.1) for discussion of scale reliability. Mean prosocial score in this sample was 8.38 (SD = 1.41).

2.3. MRI acquisition

The same 12-channel head coil on the same 1.5 T Siemens Avanto scanner (Siemens Medical Solutions, Erlangen, Germany) was used to collect MRI data at all 3 waves. The sequence used for morphological analyses at all three time points was a 3D T1-weighted magnetization prepared rapid gradient echo (MPRAGE), see parameters in Supporting Information (S2.3).

2.4. MRI processing and acquisition

Whole brain segmentation and cortical surface reconstruction was performed using FreeSurfer 6.0 (http://surfer.nmr.mgh.harvard.edu/)

Table 1

| N | Sex | Age Mean (SD) | Age Min-max | Retained from previous waves | Newly recruited |
|---|-----|---------------|-------------|-------------------------------|----------------|
| Wave 1 | 101 | 53/48 | 15.8 (2.35) | 12.0-19.7 | – | 101 |
| Wave 2 | 121 | 61/60 | 16.7 (2.88) | 11.9-21.9 | 61 | 60 |
| Wave 3 | 106 | 58/48 | 20.1 (3.27) | 13.8-26.1 | 106 | 0 |
| Total | 169 | 92/77 | 17.7 (3.43) | 11.9-26.1 | 207 | |

Note: New participants were recruited at wave 1 and wave 2. Column “Retained from previous waves” refers to individuals who participated at a time point 2 (or 3) and also have data from wave 1 and/or 2.
3. Results

3.1. Prosocial behavior development

Results showed main effects of age and sex for prosocial behavior, but no interaction between age and sex (see Supplementary Tables 1 and 2 for model fit). There was an increase in prosocial behavior with age, and females reported themselves as more prosocial than males (Fig. 1).

3.2. Prosocial behavior and cortical thickness development

Analyses testing interactions between prosocial behavior and quadratic age-related cortical development yielded seven significant clusters within the following regions: 1) right posterior middle temporal cortex, 2) left dlPFC, 3) right inferior frontal gyrus (IFG), 4) right (d) mPFC, 5) right intraparietal sulcus, 6) right posterior superior temporal cortex and TPJ, 7) right dorsal (d)ACC. The clusters are shown in Fig. 2 and further details are provided in Supplementary Table 3. Controlling for estimated IQ, parental education, and parental income did not substantially change the results (see Supporting Information (S3.1.2)).

Fig. 1. Developmental trajectory for prosocial behavior. Females are represented in pink and males are represented in blue. The shaded areas correspond to the 95% confidence intervals. Significance testing was performed using likelihood ratio (LRT) test where models were fit by maximum likelihood (ML) estimation method.

2.5. Statistical analyses

2.5.1. Prosocial behavior development

Development of prosociality over time was assessed using the nlmle package (Pinheiro et al., 2016) in R (R Core Team, 2018), in Rstudio (www.rstudio.com). This and subsequent models that were tested in mixed-effects models where subject was entered as a random effect to account for within-individual dependence, i.e. responses from one individual over multiple TPs are likely to be correlated. Mixed-effects models deal with this form of dependency in the data, allow irregularly spaced measurements (Gibbons et al., 2010), and help to preserve power in case of missing data (Matuschek et al., 2017). We began by first identifying the best fitting development model for prosociality by comparing linear, quadratic, and cubic effects of age, as well as the null model. Model testing was performed using the likelihood ratio (LRT) test where models were fit by maximum likelihood (ML) estimation method. We considered both the LRT p-values and Akaike Information Criterion (AIC, (Akaike, 1974)) during model selection, and a more complex model was only chosen if $p < 0.05$ and AIC value indicated better fit (value that was smaller by at least two). In the next step, we investigated whether inclusion of sex improved the “developmental model”, using the same model selection procedure. First, the main effect of sex was added to the model, followed by an interaction of age and sex. Model equations are outlined in Supporting Information (S2.4.1).

2.5.2. Cortical thickness development

Considering the lack of studies on prosociality and CT development, analyses were not limited to a priori selected regions of interests. Instead, whole-brain analyses across the cortical surface were performed to reduce the risk of Type II errors. Background analyses were conducted to assess the developmental trajectory of CT using SurfStat (http://www.math.mcgill.ca/keith/surfstat/), a toolbox created for MATLAB (The MathWorks, Inc., Nathan, MA). Random field theory (RFT) corrections ($p < 0.05$, cluster-defining threshold 0.005) were used to account for multiple comparisons (Worsley et al., 2004). The best-fit model for the developmental trajectory of CT was assessed in three steps whereby a cubic model was tested first, followed by a quadratic model, and a linear model (see (S2.4.2) for equations and (S3.1.1) for results).

2.5.3. Associations between prosocial behavior and cortical thickness development

As the background analyses of CT identified quadratic changes with age (see Supporting Information (S2.4.2 and S3.1.1)), we next tested interactions between prosociality and quadratic age terms using mixed models in SurfStat (see (S2.4.3) for equations). While linear mixed models are a valid approach for rejecting the null hypothesis that the relationship is linear, the quadratic function should not be interpreted literally as the shape of true developmental trajectory (Fjell et al., 2010). While inferring break points of age-trajectories based on quadratic models are heavily influenced by the age-range of the sample, a nonparametric smoothing approach seems to be more robust to such influences (Fjell et al., 2010). Thus, smoothing splines are recommended for interpreting the shapes of developmental trajectories. Therefore, we extracted average CT of significant clusters from the linear mixed model and used generalized additive mixed models (GAMM) with a cubic spline basis in R (Wood, 2006) to visualize developmental trajectories. This methodological choice is further elaborated in Supporting Information (S2.4.3). In a series of post hoc analyses, the model was rerun while controlling for estimated IQ, parental income, and parental education, see Supporting Information (S3.1.2).
To illustrate these relations between prosociality and CT development, participants were divided into two groups depending on their prosocial score throughout the study: stable low or stable high, see Supporting Information (S2.4.3). Cortical development was plotted for each group to estimate the developmental trajectories (Fig. 3). Higher scores on prosocial behavior were associated with greater thinning between early and mid-adolescence, which then slowed to less thinning during the transition to young adulthood. Lower scores on prosocial behavior showed the reverse pattern of slower thinning at younger ages and greater thinning at older ages. This pattern was observed across all the identified significant clusters. Note that the low/high division was made for illustration purposes only and the statistical analyses were run with prosocial behavior as a continuous measure.

To further probe the data, relations between prosociality, CT and age were visualized using contour plots (Fig. 4). The plots indicate that the relations between prosociality and CT were positive for the youngest and...
of the oldest group across all the identified significant clusters; in the younger group high prosocial scores were related to thicker cortex (white/light pink) and lower scores with thinner cortex (pink), the same pattern was found for the oldest group where high prosocial scores were associated with thicker cortex (yellow/light green) and low prosocial scores with thicker cortex (dark green).

4. Discussion

The current study examined the relations between prosociality and longitudinal CT development in adolescents and young adults. Higher self-reported prosociality was associated with greater initial rate of regional thinning during early adolescence, followed by attenuation of this process during the transition to young adulthood in regions involved in social cognition and behavior control. In contrast, lower prosocial scores were related to initially slower thinning, followed by more maturation into the third decade of life. These results suggest that the rate of thinning and possibly its timing, are key factors in prosociality during development.

Existing studies disagree on whether relatively greater thinning in development is adaptive (see e.g. (Bos et al., 2018; Vijayakumar et al., 2014a, 2014b, 2014c)). Similarly, being prosocial is generally considered to be positive, but being too prosocial may be maladaptive (Steinbeis, 2018). In our sample, prosociality was i) negatively related to a number of maladaptive developmental outcomes (see supplementary analyses (S3.2.2)), and ii) associated with greater thinning in early adolescence and more attenuated thinning in young adulthood. Although highly speculative, these results may imply that greater cortical thinning is adaptive in younger ages, but may no longer be optimal in young adulthood. In other words, greater thinning may or may not be optimal depending on age. However, this notion should be scrutinized by future studies. Previous studies have linked interventions promoting prosociality to number of favorable developmental outcomes (Caprara et al., 2014). Although highly speculative, future studies could test a hypothesis that such interventions during this time of plasticity may have lasting impacts through their effect on neural development.

Links between prosociality and longitudinal development of CT were identified in several regions and were in line with our hypotheses. First, anatomical locations previously associated with ToM, including TPJ, pSTS and posterior parts of mPFC, all in the right hemisphere were identified. The precise role of ToM in prosociality is not fully understood. Understanding one’s mental states may directly lead to prosocial behaviors (Eggum et al., 2011), but it is also possible to use this ability for both good (prosocial behaviors) and ill (e.g. antisocial behavior such as manipulation). Therefore, ToM should rather be viewed as an information-gathering tool serving various purposes (Eisenberg et al., 2001). ToM may also be related to prosociality by affecting empathic concern (Van der Graaff et al., 2018), sympathy and moral reasoning (Eisenberg et al., 2001). The role of social approval has also been emphasized (Lane et al., 2010): individuals proficient in perspective taking have the ability to consider how others will react to their behavior, and anticipation of these reactions may motivate prosocial behaviors valued in one’s social environment, e.g. doing the right thing to make others proud of you.

We also found an association between prosociality and cortical development in the right intraparietal sulcus, a part of the so-called mirror neuron system (MNS) in humans. This network is thought to aid understanding of others’ actions and the intentions behind these actions (Cattaneo and Rizzolatti, 2009). Importantly, the MNS has been implicated in a longitudinal link between childhood empathy and prosociality in adolescence, albeit in more frontal components than in the current study (Flournoy et al., 2016). A large meta-analysis (Van Overwalle and Baetens, 2009) of functional neuroimaging studies suggests that the ToM and MNS represent two separate, but complementary neural networks, that both serve to optimize aspects of social processing. Consistently, our findings suggest that both networks may also contribute to prosociality.

Interestingly, these findings were located in the right hemisphere. There is some limited evidence of right hemispheric specialization for social emotions (Benowitz et al., 1983; Blonder et al., 1993), which may partially explain this finding. Additionally, Von Economo cells are more numerous in the right hemisphere (Allman et al., 2005), and studies on ToM have also emphasized this hemisphere (Scholz et al., 2009; Young et al., 2010).

While systems supporting understanding others’ actions, intentions, thoughts and feelings, may be important contributors to prosociality, systems supporting strategic decision-making may also be critical. Pro-social behaviors are vital for living in social groups, an arrangement that has been beneficial for humans throughout evolution, e.g. advantages such as resource sharing or group protection. Groups with larger number of prosocial individuals are more likely to succeed over groups consisting mainly of selfish individuals, and therefore, at least under some

Fig. 4. Contour plots of relations between prosocial behavior, age and cortical thickness. The plots indicate that the relations between prosociality and CT were positive for the youngest and the oldest group across all the identified significant clusters. Such a relationship was not present in the middle of the age range (late adolescence). Each plot contains mean cortical thickness from the clusters where interactions between prosociality and age squared were significant. The green to white color scale indicates low to high cortical thickness values.
circumstances, evolutionary mechanisms may have contributed to the emergence of prosocial behaviors in social groups (see Van Vugt and van Lange, 2006 for review). Not acting in a prosocial manner could cause retaliation and exclusion by group members (Axelrod and Hamilton, 1981; Fehr and Gachter, 2000). On the other hand, acting excessively prosocially can put a strain on one’s resources and may place one in a disadvantage in some situations. Considering these costs and benefits of being prosocial prior to engagement in such acts may thus be adaptive (Steinbeis, 2018). Our results showed that prosociality was associated with CT development in the left dlPFC, but not vmPFC, despite both regions supporting value-based decision-making (Sokol-Hessner et al., 2012). The left dlPFC has previously been identified in behavioral control and implicated in strategic social behavior (Steinbeis et al., 2012), and as a key structure involved in self-control (Hare et al., 2009), a function that impacts decisions to act prosocially (Joosten et al., 2015; Osgood and Muraven, 2015). Additionally, transcranial stimulation of this region has been associated with increased prosociality (Balconi and Canavesio, 2014), and bilateral dlPFC activation has been associated with sharing in the context of social norm compliance (Spitzer et al., 2007). Taken together, prior findings implicate the dlPFC supporting the integration of emotional information in decision making in settings (Balconi and Canavesio, 2014), thus highlighting the potential role of this structure in prosocial behaviors.

In line with our hypotheses, prosociality was also related to CT development of the right dACC, as region that has repeatedly been linked to cognitive control, performance-monitoring, motivation and reward-based decision-making (see Shenhav et al., 2016 for review). The overall expected value of control theory (EVC) integrates these findings and proposes that the dACC is involved in making decisions as to how much control is allocated based on cost/benefit analysis of estimated reward outcome and effort cost (Shenhav et al., 2016). This is in line with the notion that prosocial behaviors depend on thorough situational assessment and cost-benefit analysis. Additionally, this phylogenetically and ontogenetically late developing structure has repeatedly been implicated in empathy (Hein et al., 2016; Hein et al., 2010) and contains Von Economo cells (Allman et al., 2005) that are involved in rapid assessment of complex situations and provide necessary input to regions involved in ToM.

Prior research on samples aged 6–9, have identified predominantly positive associations between prosociality and CT (Thijssen et al., 2015; Wildeboer et al., 2017). Our study differed by focusing on an older age-range, 12–26 years, and by employing a longitudinal design to test for associations with cortical development (i.e. changes in CT over time). Still, some of the same regions in similar medial and lateral frontal cortices were implicated in our study as well as by Thijssen et al. (2015) who used the same measure of prosociality. In line with these earlier findings, we also observed positive associations in the younger age range of our sample (see Fig. 4). Additionally, previous work on symptoms of CD based on data from the first wave of the current project found negative relations between symptoms of CD and CT, with the relation being strongest in the youngest individuals (Walhovd et al., 2012). Prosocial and antisocial behaviors have by some been conceptualized as extreme ends of a continuum (Éron and Huesmann, 1984), although the relation is likely more complex (see Eisenberg et al., 2015 for review). Nonetheless, Waldman et al. (2011) uncovered an overlap between genetic influences on CD and prosociality, and similar areas involved in regulatory processes and social-cognitive processing have been implicated in behavioral studies (see Walhovd et al., 2012). Considering our longitudinal and the cross-sectional results together, although speculative, it is possible that highly prosocial individuals differ from less prosocial individuals by having thicker cortices to begin with in early development, followed by differential pattern of thinning into young adulthood.

While not a primary focus of this study, we found that prosociality showed subtle linear increases from age 12 to young adulthood. This is largely consistent with prior literature that has shown decreases in early adolescence followed by a rebound (Eisenberg et al., 2015). However, we failed to identify any associations between change in prosociality and changes in CT over adolescence, which likely relates to the moderate levels of rank-order stability in prosocial behaviors (see S3.2.1). Females also reported being more prosocial but the rate of increase over time was identical for both sexes. This is in line with existing research, demonstrating less prosociality in males, particularly when self-report measurement is used (Fabes and Eisenberg, 1998). These differences may reflect conceptions of what males and females are supposed to be like, rather than actual behaviors. Additionally, measures of prosocial behavior may include a disproportionate number of items on which girls are more likely to rate themselves as prosocial than boys (see (Fabes and Eisenberg, 1998) for review). However, we note that interactions between cortical development and prosocial behavior were not driven by these sex differences (see Supporting Information (S3.2.6) for further detail).

Future studies should also employ more specific measures of prosociality as different subtypes of prosociality, i.e. helping, sharing and comforting, may depend on different forms of social-cognitive understanding and show unique developmental trajectories (Dunfield, 2014). They may thus also relate to distinct cortical regions and patterns of cortical development. Self-report used in the current study should be supplemented by additional measures of prosociality provided by different observers in order to reduce the impact of social-desirability bias which may have influenced results in this study. Additionally, psychometric properties of the SDQ have not been tested for the entire age-range of the current sample. While this may constitute a possible limitation, the continuity of measures across time points was considered to be more important. Future studies could also compare cortical maturation in highly antisocial individuals to that of highly prosocial individuals to examine whether there is support for a dimensional view of prosocial-antisocial behaviors. Moreover, note that estimates of non-linear development in our study were driven by between-subject differences. Future studies with more time points are needed to model within-subject non-linear trajectories. Finally, future studies should include measures of pubertal maturation, a key factor driving individual differences in brain structural development (Herting and Sowell, 2017). This may provide further insight into the relations between cortical development and prosocial behavior.

In conclusion, the longitudinal nature of the current study offers novel insight into the emergence of the links between prosociality and brain structure, specifically highlighting the moderating role of age; the association changed from positive to negative across early to late adolescence, and subsequently became positive again in young adulthood. These dynamic relations were observed in regions previously associated with social processing and behavioral control. The results emphasize that both the rate and the timing of cortical thinning may be related to prosocial behaviors during development. Given the study’s exploratory nature and limitations, the conclusions are speculative and should be interpreted with caution.

**Funding**

This study was supported by the Research Council of Norway (to AMF, KBW and CKT: 230345, 271839, 288083, 223273)

**Declaration of Competing Interest**

No conflict of interest

**Appendix A. Supplementary data**

Supplementary material related to this article can be found, in the online version, at doi:https://doi.org/10.1016/j.dcn.2019.100734.
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