Multimorbidity in Children and Youth Across the Life-course (MY LIFE): protocol of a Canadian prospective study

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Abstract: INTRODUCTION: Multimorbidity, the co-occurrence of a chronic physical condition and mental disorder, affects a substantial number of children and youth and can lead to compromised quality of life, hardship for families, and an increased burden on the healthcare system. We are conducting a study to document the course of mental disorder in children and youth diagnosed with a chronic physical condition; identify predictors of child and youth multimorbidity; examine whether the effects of these predictors are moderated by relevant psychosocial and biological factors; explore potential inflammatory and stress biomarkers that mediate the onset of child and youth multimorbidity; and, assess whether multimorbidity in children and youth alters patterns of mental health service use. METHODS AND ANALYSIS: Multimorbidity in Children and Youth Across the Life-course (MY LIFE) is a prospective study. Two hundred and fifty children and youth aged 2-16 years diagnosed with a chronic physical condition along with one parent will be recruited from the outpatient clinics at a paediatric tertiary care centre. Data will be collected using a multi-informant, multimethod design at four time-points (at recruitment, and at 6, 12 and 24 months postrecruitment). Parents will provide reports for all children/youth. In addition, youth 10 years will self-report. Mental disorder will be assessed using structured interviews. On completion of data collection, participant-reported data will be linked to provincial health records to identify mental health services use. Multilevel analyses (survival, proportional hazard, structural equation modelling) will be used to address MY LIFE objectives. ETHICS AND DISSEMINATION: This study has been approved by the University of Waterloo Human Research Ethics Board and the Hamilton Integrated Research Ethics Board. Findings will be disseminated to key stakeholders using a number of outlets (peer-reviewed publications and conferences, lay informational pamphlets, social media).

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Multimorbidity in Children and Youth Across the Life-course (MY LIFE): protocol of a Canadian prospective study

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ABSTRACT

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Ethics and dissemination This study has been approved by the University of Waterloo Human Research Ethics Board and the Hamilton Integrated Research Ethics Board. Findings will be disseminated to key stakeholders using a number of outlets (peer-reviewed publications and conferences, lay informational pamphlets, social media).

INTRODUCTION

Children and youth with chronic physical conditions face considerable psychosocial and emotional burden due to their health challenges.1 Thus far, research that comprehensively examines child and youth multimorbidity—the co-occurrence of physical and mental disorder—is limited, including research on the mechanisms involved in the onset and outcomes of multimorbidity.

Yet, understanding child and youth multimorbidity is an urgent public health problem. Improvements in healthcare have reduced childhood mortality, but many surviving children grow up into adulthood.2 Furthermore, because most adult mental disorders have their onset in childhood or adolescence,3 the onset of multimorbidity likely typically occurs early in life, with long-term consequences for children and youth, their families, the healthcare system and society. Indeed, cost calculations suggest that a small proportion of children (approximately 5%, often multimorbid) account for nearly 60% of healthcare costs for individuals <18 years of age.4 Taken together, multimorbidity and its sequelae are an important public health concern.5 6

Current research on child and youth multimorbidity suggests that its prevalence is high.7 Population-based studies report prevalence estimates of 20%–30%,8 10 clinical studies ≥50%.11 12 Most extant studies report that multimorbidity has a negative impact on

Strengths and limitations of this study

- Unlike previous studies on child and youth multimorbidity, this is a prospective study that uses a multi-level, multi-method and multi-informant approach.
- Multimorbidity in Children and Youth Across the Life-course aims to determine biological and psychosocial processes by which some children and youth develop multimorbidity whereas others do not; this is essential for establishing effective interventions.
- Findings have the potential to support children/youth and families, improve care and reduce health system costs.
- Recruitment from a single paediatric tertiary care centre might reduce generalisability of findings.
- As with all prospective studies, missing data and attrition may be problematic; however, multiple approaches will be used to minimise sample loss and other biases.
the quality of life of young people (for an exception see Ferro et al. Impact on the family environment appears more mixed, with some studies reporting closer parent–child relationships in those with (vs without) multimorbidity. Studies on family sociodemographic correlates of multimorbidity are rare, with one study reporting no association. Emerging evidence implicates biological mechanisms that link physical and mental disorder, including immunological, inflammatory and stress-related mechanisms. Findings regarding mental health services use among children and youth with multimorbidity are also mixed; clinical studies reporting less use of acute mental health services, whereas population studies reporting more overall and acute mental health services use. Despite this nascent literature on child and youth multimorbidity, many important gaps in knowledge remain. The Multimorbidity in Children and Youth Across the Life-course (MY LIFE) study was developed to address these gaps.

First, studies of clinical samples of children and youth with physical conditions are typically small and cross-sectional. The lack of prospective studies limits our ability to: (1) understand the temporality of effects; (2) document the natural course of multimorbidity over time; (3) calculate incidence rates; (4) generalise findings to the broader population of children and youth with multimorbidity. MY LIFE will recruit a larger sample of children and youth with a variety of physical conditions and follow them and their families prospectively on multiple measurement occasions. This will inform opportunities for prevention/intervention at all levels: primary (preventing onset of mental comorbidity), secondary (reducing symptoms in children and youth with subclinical symptomatology) and tertiary (improving function in children and youth with multimorbidity). With regards to generalisability, few studies (all using population samples) have compared mental disorder across physical conditions. While findings typically show negligible differences across conditions, identifying condition-specific associations may be useful in understanding mechanisms underlying the onset of multimorbidity and informing intervention.

Second, previous studies have typically assessed symptoms of emotional or behavioural problems and not mental disorder, per se. This is relevant given recent evidence that assessment type moderates estimates of association between physical conditions and mental disorder and influences estimates of prevalence in child and youth populations. Furthermore, there is robust evidence that agreement on measures of mental and psychosocial health between parent and child/youth reports are low. Combined, these methodological limitations can bias estimates (via misclassification), which ultimately could misdirect resources for appropriate mental healthcare for children and youth. MY LIFE will implement a multmethod and multi-informant approach to measuring child and youth mental health and related psychosocial outcomes.

Third, little is known about the mechanisms that underlie the development of multimorbidity in children and youth. There has been some work highlighting biological and psychosocial mechanisms independently, but few investigations have sought to combine these approaches to provide a more comprehensive understanding of this phenomenon. One study has demonstrated that maternal depression has direct effects on immunological and stress regulation in mothers and their children, which subsequently influences symptoms of mental disorder in children. However, that study did not examine these mechanisms in the context of children having a physical condition. Recent evidence in this vulnerable population not only shows a confluence of biological and psychosocial pathways linking physical and mental health, but that effects may be different for male and female children. Relatedly, there is robust evidence demonstrating the positive effect of physical activity and exercise on the mental health of, as well as reducing levels of inflammatory markers in, children and youth. MY LIFE will measure physical activity and inflammatory and stress markers not previously assessed in studies of child and youth multimorbidity in an effort to more fully understand its onset and progression. Clarifying the extent to which biomarkers and psychosocial factors, and ultimately physical and mental health, change over time can inform targets for intervention to reduce risk for multimorbidity.

Fourth, use of mental healthcare has increased dramatically for children and youth in recent years, but knowledge as to whether multimorbidity influences service use or help-seeking behaviour is limited. The few studies that exist have reported inconsistent findings: a population study found increased mental healthcare use among youth with a physical condition and bipolar disorder, whereas a clinical study found that youth with multimorbidity were less likely to receive inpatient mental healthcare and had shorter hospital stays. Through data linkage activities with administrative health records, MY LIFE will contribute to the field by identifying how children and youth with multimorbidity access and use mental health services. Understanding the patterns of mental healthcare for children and youth with multimorbidity will provide useful information to support families navigating the health system and the evidence required to implement models of integrated physical and mental healthcare.

Overall, MY LIFE aims to: (1) document the course of mental disorder in children and youth diagnosed with a chronic physical condition; (2) identify predictors of child and youth multimorbidity; (3) examine whether the effects of these predictors are amplified or moderated by relevant psychosocial and biological factors; (4) explore potential inflammatory and stress biomarkers that mediate the onset of child and youth multimorbidity; (5) assess the extent to which child and youth multimorbidity alters patterns of mental health service use.
METHODS

Design
The design of MY LIFE was informed by our pilot study. MY LIFE is a prospective study in which children and youth with chronic physical conditions and their primary caregiving parent/guardian will be recruited from outpatient programmes at a paediatric hospital in Canada. Participants will be followed for 24 months, with assessments at four occasions: recruitment (baseline) and 6, 12 and 24 months. On completion of the in-person data collection, participant-reported data will be linked to provincial health records to document use of mental health services.

Participants
Children and youth aged 2–16 years with one chronic physical condition is the population of interest. Physician-diagnosed physical conditions are those expected to last ≥12 months; and result in ≥1 of the following sequelae—functional limitations, dependencies to compensate for limitations and need for additional healthcare. Children and youth with multiple physical conditions (ie, medically complex) at the time of recruitment will be excluded as these children/youth represent a small subgroup within this population and whose outcomes and healthcare use are established. 47–49 Associations with mental comorbidity in these children and youth is likely different compared with children and youth who suffer from one, physical condition only. Also, children/youth and parents with insufficient English language skills will be excluded, because the selected measures have not been validated in other languages.

Multimorbidity is evenly distributed across age and sex. Thus, we expect to recruit males and females in a 1:1 ratio, evenly distributed across ages. Our age range is broad to increase coverage and generalisability. We will include children and youth who, as a function of their physical condition, have a developmental or learning disability (eg, in certain presentations of epilepsy). For these children/youth, we will rely on parent reports only.

Procedures
Study procedures are outlined in figure 1. Participants will be recruited from the dermatology, endocrinology, gastroenterology, haemophilia, immunology, neurology, respiratory and rheumatology programmes. Research assistants (RAs) will work in tandem with health professionals to help identify eligible families. Clinic nurses will introduce the study to families and invite them to speak with the RA either before or after their medical appointment. The RA will verify eligibility, describe the study and obtain written permission from eligible participants to be contacted a few days later.

The study coordinator will then contact parents to schedule a time for participants to complete the structured interviews and biological sample collections at...
the hospital. Alternatively, the RA may meet with participants at their home to accommodate travel or child care schedules. In the rare case that a family is not able to meet in-person (either at the hospital or in their home), packages will be mailed, followed by telephone calls from the study coordinator with reminders for completion. During study visits at the hospital, the RA will meet with participants in a private research office. Informed written consent (parent and youth aged 16 years) and assent (children and youth aged 7–15 years) will be obtained at the beginning of the first visit.

Data collection for MY LIFE is based on the following a priori consideration that assessments should be far enough apart to allow sufficient time for detectable changes to occur in measures of children and youth, family and clinical characteristics and to not burden participants. Data will be collected every 6 months during the first year given that the period close to diagnosis is a dynamic time in terms of clinical management and family adaptation, and then again 1 year later. The experience of our research team suggests that a 2-year follow-up is adequate for detecting meaningful changes in psychosocial outcomes of children and youth with physical conditions. Data will be collected using independent structured interviews and self-reported questionnaires on a laptop (computer assisted to minimise missing data) for children/youth and parents to ensure privacy. Parent reports will be collected for all participating youth; youth aged ≥10 years will also provide reports. Data collection will take approximately 60–90 min. On completion of each interview, children/youth and parents will both be provided with a $30 gift card and if applicable, a parking voucher.

**Measures**

We will implement a multilevel, multimethod and multi-informant approach to data collection. Variables will be measured at the individual, family and community levels using child/youth and parent-report and linkage with administrative health records obtained for the study period.

**Self-reported measures**

We will be using two measures of child and youth mental health as our primary outcome of interest. The Mini International Neuropsychiatric Interview for Children and Adolescents (MINI-KID) is a clinical diagnostic interview designed to assess the presence of Diagnostic and Statistical Manual of Mental Disorders (DSM)-5 and International Statistical Classification of Diseases and Related Health Problems (ICD)-10 psychiatric disorders in children and youth over the past 6 months. We will be utilising the following modules which represent the most common mental disorders in childhood and adolescence: major depressive episode, separation anxiety disorder, social anxiety disorder, specific phobia, attention-deficit hyperactivity disorder, conduct disorder, oppositional defiant disorder and generalised anxiety disorder. The MINI-KID has demonstrated strong validity and reliability in population and clinical samples of children and youth.

Second, the Ontario Child Health Study Emotional Behavioural Scales (OCHS-EBS) are a set of self-reported checklists measuring symptoms of psychopathology over the past 6 months using a 3-point Likert scale. Items are scored 0, 1, or 2, indicating responses of ‘never or not true’, ‘sometimes or somewhat true’ and ‘often or very true’, respectively. The raw scores are summed to calculate a scale score for each mental disorder (major depressive, generalised anxiety, separation anxiety, social phobia, attention-deficit hyperactivity, oppositional defiant, conduct), as well as total severity score and scores for internalising and externalising disorders. The OCHS-EBS demonstrates robust psychometric properties.

Child/youth and parent mental health and quality of life, child/youth physical and psychosocial functioning, child/youth intelligence, family environment and sociodemographic characteristics will be assessed using psychometrically robust self-reported measures. When applicable, parents will also complete quality of life and mental health measures for a sibling within 3 years of age from the child in the study to understand the broader impact of child and youth multimorbidity on family members. Details of these measures are described in table 1.

**Biological assessments**

To date, the few studies that have assessed markers of systemic inflammation suggest associations with child and youth mental health. We will measure whether systematic inflammatory biomarkers (which tend to be higher in young people with physical conditions) mediate the timing of onset of multimorbidity. We will collect dried blood spots for all children and youth and whole blood for youth ≥10 years at each measurement occasion. Our procedures for collecting dried blood spots by research staff have been adapted from published guidelines. Whole blood samples will only be collected during study interviews in hospital by experienced phlebotomists. Samples will be analysed using high-sensitivity ELISA to quantify levels of C-reactive protein, tumour necrosis factor-α, interleukin-1β and interleukin-6, as these are robust indicators of systemic inflammation.

Cortisol, a glucocorticoid hormone, is released by the hypothalamic-pituitary-adrenal axis in response to stress. Cortisol extracted from saliva, urine and blood provides transient measures of physiological stress, with limited utility for understanding chronic physiological stress in young people with chronic conditions. Hair cortisol concentration has recently been identified as a potentially valuable biomarker, averaging daily fluctuations in circulating cortisol levels. Hair sampling represents a non-invasive method to measure chronic physiological stress and reduces the impact of individual and contextual differences associated with circulating cortisol. Hair will be collected using a standardised
### Table 1  Study questionnaires

| Construct                | Measure and description                                                                 | Informant         |
|--------------------------|----------------------------------------------------------------------------------------|-------------------|
| Child/youth mental health| **Ontario Child Health Study Emotional Behavioural Scales**<sup>55</sup> 52-item symptom checklist providing total, internalising and externalising scores, as well as subscale scores for major depressive disorder, separation anxiety, social phobia, generalised anxiety, attention-deficit hyperactivity, oppositional defiant disorder and conduct disorder | Child/youth and parent |
|                          | **Strengths and Difficulties Questionnaire**<sup>87</sup> 25-item checklist providing total difficulties scores, as well as domains of emotional problems, conduct problems, hyperactivity, peer problems and prosocial | Parent            |
| Child/youth psychosocial health | **KIDSCREEN**<sup>88</sup> 27-item scale of health-related quality of life with domain scores for physical well-being, psychological well-being, autonomy and parent relations, peers and social support and school environment | Child/youth and parent |
|                          | **Self Description Questionnaire**<sup>89</sup> 90 Five items from the General Self-image subscale; adapted from the National Longitudinal Survey of Children and Youth | Child/youth and parent |
|                          | **Self-Perception Profile for Children**<sup>91</sup> 36-item scale assessing self-concept in the domains of scholastic competence, social acceptance, athletic competence, physical appearance, behavioural conduct and global self-worth | Child/youth |
| Child/youth physical health | **WHO Disability Assessment Schedule 2.0**<sup>93</sup> 12-item measure of disability and functioning in the domains of cognition, mobility, self-care, getting along, life activities and participation | Child/youth and parent |
| Child/youth intelligence | **Kaufman Brief Intelligence Test, second edition**<sup>95</sup> Brief measure of verbal and non-verbal intelligence | Child/youth |
| Parent mental health     | **Center for Epidemiologic Studies Depression Scale**<sup>96</sup> 20-item measure of depressive symptoms across four domains depressed affect, positive affect, somatic activity and interpersonal relations | Parent |
|                          | **Generalized Anxiety Disorder**<sup>97</sup> Seven items measuring symptoms of generalised anxiety | Parent |
| Parent quality of life   | **Short Form-36**<sup>98</sup> 36-item scale that assesses eight health domains: physical functioning, bodily pain, role limitations due to physical health problems, role limitations due to personal or emotional problems, emotional well-being, social functioning, energy/fatigue and general health perceptions, and a single item that provides an indication of perceived change in health | Parent |
| Family environment       | **McMaster Family Assessment Device**<sup>99</sup> 100 12 items comprising the General Functioning subscale measures aspects of communication, problem solving, behavioural control, affective responsiveness and involvement and roles | Parent |
|                          | **Parental Stress Scale**<sup>101</sup> 102 18-item scale that includes four domains of parenting: rewards, stressors, loss of control and satisfaction | Parent |
|                          | **Sibling Inventory of Differential Experience**<sup>103</sup> 18-item scale assessing differential treatment by mother and father in relation to closest-aged sibling | Child/youth |
| School environment and peers | **2014 Ontario Child Health Study (OCHS)**<sup>104</sup> 14 items assessing school climate, bullying and participation in extra-curricular activities | Child/youth |
| Mental healthcare        | **Various items obtained from the OCHS and Canadian Community Health Survey** | Parent |
| Sociodemographics        | **Various items obtained from population-based surveys conducted by Statistics Canada** | Parent |

Only youth ≥10 provided self-reported data.

approach by research staff from children/youth and parents by cutting approximately 80 strands (roughly 4 mm diameter) from the posterior vertex of the head, as close to the scalp as possible, the extraction and assaying of cortisol will follow established procedures, which we have also used previously.<sup>34</sup> Variables hypothesised to affect hair cortisol concentration (eg, hair washes, medications, smoking behaviour (including exposure to secondhand smoke)) will be collected via parent report on a separate form.<sup>67</sup> 71 72

While parents are completing study questionnaires, children 2–5 years will complete the Peabody Developmental Motor Scales, second edition (PDMS-2) to assess gross motor development.<sup>73</sup> This measure, which requires 20–30 min to complete, is collected only at study visits conducted at the hospital. We will be using three gross motor subtests—stationary, locomotion and object manipulation—to obtain a gross motor developmental quotient. The PDMS-2 has excellent reliability and is valid for evaluating children/ youth with varying levels of gross motor skills.<sup>74</sup>
Habitual physical activity at each assessment will be monitored using accelerometry, providing the duration, intensity and frequency of movement among children and youth. Following standard practice, we will monitor physical activity over seven consecutive days, reflecting recent levels of physical activity. Activity intensity is analysed according to cut-points developed and validated specifically for this unit (ActiGraph wGT3X-BT). Sitting and standing height and weight will also be collected for children and youth at each assessment.

**Patient and public involvement**

Patients or the public were not involved in the design, or conduct, or reporting of this protocol. The role of participating families with regards to our knowledge translation strategy is described in the Ethics and dissemination section.

**DATA ANALYSIS AND LINKAGE**

**Sample size calculation**

Simulations that altered various parameter estimates were conducted to determine the sample size requirements for this study. Results suggested that a study with 250 children and youth and four measurement occasions would be adequately powered (1−β≥0.80) at α=0.05 to estimate non-linear changes in mental health trajectories (primary study objective), examine relevant mediating and moderating effects and explore subgroup differences (eg, by physical health or mental disorder). Assuming 40% of eligible children/youth or their parents refuse participation after being provided with details of the study, and 20% of initially enrolled children/youth do not complete the baseline assessment, we anticipate approaching approximately 520 families about the study. If appropriate, multiple imputation will be used to account for missing data.

**Data analysis**

Multilevel models will delineate trajectories of psychopathology and psychosocial outcomes over time, survival analysis and proportional hazard models will be used to model risk and predictors of child and youth multimorbidity, and structural equation modelling will identify factors that mediate and moderate outcomes. Multinomial logistic regression will be used to model patterns of mental health service use for children and youth with and without multimorbidity. Additional analyses will be stratified to examine subgroup differences, including by sex, because prevalence of and pathways to mental disorder is not the same for males and females. When possible, we will use age-stratified analysis based on developmental periods to address the challenge of making inferences on heterogeneous samples. The developmental periods will be early childhood (2–6 years), middle childhood/early adolescence (7–12 years) and older adolescence (13–16 years).

**Data linkage**

Study data will be linked using the provincial health numbers of participants with the health data collected by the Ministry of Health to obtain information on mental health service use. Databases to be linked will include the National Ambulatory Care Reporting System, Discharge Abstract Database and Ontario Health Insurance Plan. We will use postal codes to link study data with Canadian census data to measure community-level variables, such as neighbourhood income and geographic proximity to health services.

**DISCUSSION**

Child and youth multimorbidity creates considerable challenges for the individual, family and healthcare system. By focusing on modifiable predictors and associated health and psychosocial outcomes of child and youth multimorbidity, this study will advance knowledge of the aetiology and course of mental disorder in a large vulnerable population and refine our understanding of how child and youth multimorbidity influences interaction with the healthcare system.

We note the following challenges associated with conducting this study. First, the eligibility criteria are purposefully broad—in terms of age and physical health problems—in order to enrol a large enough sample of children and youth to address our objectives. This heterogeneity may limit the applicability of findings to specific subgroups (eg, young children, youth with juvenile arthritis). However, evidence suggests that differences in risk for mental disorder across physical conditions is negligible. In addition, we will conduct age and condition-stratified analyses to adjust for potential residual confounding and identify potential stratum-specific findings.

Second, attrition may be problematic. Because our study includes parents and children/youth, attrition of one member of the dyad may likely result in the other being lost. To mitigate and minimise sample loss, we are oversampling, implementing a strong follow-up strategy which include multiple contacts over time, and using robust statistical methods to compute unbiased estimates in the presence of missing data.

Third, at the time of this proposal, some mental health services were funded by the provincial Ministry of Children, Community and Social Services (MCCSS). These services are not captured in the datasets in which we will link our study data. Our previous experience has shown that linkage with MCCSS is not possible; we will overcome this challenge by directly asking parents about the MCCSS-funded mental health services they access using the methods described in population-based studies conducted by Statistics Canada.

In summary, we expect that findings from MY LIFE will lead to development and implementation of universal and targeted risk-reduction strategies to reduce mental disorder in children and youth with physical conditions and will inform initiatives for the integration of physical and mental healthcare for children and youth—potentially aiding in eliminating stigma surrounding mental health and help-seeking. Ultimately, findings from this
study will improve the quality of life for children and youth and their families, and potentially reduce burdens on the healthcare system.

**ETHICS AND DISSEMINATION**

MY LIFE received ethical approval from the University of Waterloo Human Research Ethics Board (ORE-292183) and the Hamilton Integrated Research Ethics Board (2797). Dissemination of findings will target academic outlets including publication in peer-reviewed journals and presentations at national and international conferences. We will engage health professional and administrative stakeholders with presentations at departmental rounds and divisional meetings to discuss opportunities to improve child and youth health through clinical practice changes informed by findings from MY LIFE.

At the end of the study, participating families will receive a report detailing findings in a coherent, understandable format. Additionally, we will partner with families to co-produce pamphlets containing key messages from this study; what to expect postdiagnosis of a physical condition, supportive services for children and youth and families, and information about the signs and symptoms of mental disorder. Pamphlets will be distributed to clinics and community agencies, as well as websites known for their reach and impact (eg, Mental Health Commission of Canada, Centre for Addiction and Mental Health).

We will also disseminate findings to the mental health community. The principal investigator will give webinars to staff of mental health agencies and their clients which will be recorded and distributed using existing knowledge translation infrastructure. Finally, we will incorporate social media into our dissemination strategy to reach the general community through media releases, summaries on institution websites, and communicate findings via popular outlets (eg, Twitter).

**Twitter** Mark A Ferro @ARCH_Lab and Jan Willem Gorter @Dr_Gorter

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**Contributors** MAF is the principal investigator of MY LIFE, having conceived the study and its design, was responsible for obtaining funding, leading its implementation, and drafting the protocol manuscript. ELL, RJVL, JWG, LS, MB, KG and BT are co-investigators of MY LIFE, having contributed to its design, obtaining funding and editing the manuscript for content. All authors approved the manuscript for publication.

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