Celecoxib versus placebo as an adjunct to treatment-as-usual in children and youth with obsessive–compulsive disorder: protocol for a single-site randomised quadruple-blind phase II study

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ABSTRACT

Background  Cyclooxygenase (COX) enzymes oxidise arachidonic acid to prostaglandins, which modulate neuronal function and inflammation in the central nervous system. Consensus guidelines suggest non-steroidal anti-inflammatory drugs as a possible adjunctive approach in adults with obsessive–compulsive disorder (OCD) and in children with acute-onset OCD subtypes. However, there is limited evidence to support this approach. The primary objective of this study is to determine the efficacy of the COX-2-selective inhibitor celecoxib as an adjunct to treatment-as-usual in children and youth with moderate-to-severe OCD. The safety of this intervention including adverse events will also be systematically assessed.

Methods  The Adjunctive CElexib in childhood-onset OCD (ACE-OCD) study is a single-centre randomised, quadruple-blind, placebo-controlled superiority trial with two parallel groups: celecoxib 100 mg twice daily and placebo. Treatments will be added to participants’ routine clinical care, which will not change over the course of the study. Target recruitment is 80 participants ages 7–18 with no recent treatment changes. The primary outcome is OCD severity after 12 weeks of treatment, measured by clinician-administered Children’s Yale–Brown Obsessive Compulsive Scale (CY-BOCS). Secondary outcomes include CY-BOCS score after 6 weeks; difference in the proportion of participants achieving a clinically meaningful response or remission; mean clinical global impression of severity and improvement after 6 and 12 weeks; and proportion of participants reporting adverse events possibly or probably related to the study intervention. The primary analyses, carried out according to intention-to-treat principles, will compare the celecoxib to placebo group on each outcome of interest, adjusting for baseline scores using analysis of covariance or logistic regression. Participants will be offered a 12-week open-label celecoxib extension and will be invited to participate in an ancillary study for biomarker analyses.

Strengths and limitations of this study

This study is the first randomised, placebo-controlled trial to evaluate the efficacy and safety of adjunctive non-steroidal anti-inflammatory drug therapy in childhood-onset obsessive–compulsive disorder and does not restrict participants to a diagnosis of paediatric acute-onset neuropsychiatric syndrome or paediatric autoimmune neuropsychiatric disorder associated with streptococcal infections.

Study visits may occur virtually and screening blood work can be completed in participants’ local communities, increasing accessibility for participants.

Participants will have the option to consent to an ancillary study involving biosample collection for correlative biology; this will provide preliminary longitudinal data allowing measurement of associations between inflammatory biomarkers and clinical phenotype.

This study incorporates assessment of participants’ and parents’ perspectives on participation, including their experience of virtual visits, to inform future studies of psychopharmacologic interventions in this population.

While heterogeneity of usual therapy may limit power to detect differences between arms, this represents a more pragmatic approach than contemporaneous initiation of a selective serotonin reuptake inhibitor as described in preliminary studies in adults.

INTRODUCTION

Obsessive–compulsive disorder (OCD) is a common neuropsychiatric condition identified by the World Health Organisation as one of the leading causes of worldwide medical disability.1 It affects 1%–3% of the population presentations to multiple stakeholders including patients, parents and healthcare providers.

Trial registration number NCT04673578.
COX-inhibitors have suggested modest symptom improvement with celecoxib in the treatment of depression\(^{16}\) and first-line options for patients with enduring symptoms.

A large body of work suggests an association between infection and an abrupt, early-onset form of OCD, termed paediatric autoimmune neuropsychiatric disorder associated with streptococcal infections (PANDAS),\(^{2,3}\) as well as paediatric acute neuropsychiatric syndrome (PANS).\(^{4}\) Recent epidemiological data suggest that recurrent episodes of infection and inflammation are associated with the development of multiple mental disorders in children,\(^{7}\) including ‘classic’ OCD.\(^{8}\) Moreover, patients with autoimmune disorders have higher rates of comorbid OCD compared with the general population.\(^{9,10}\) A recent cohort study based on Swedish National Register data suggested increased rates of multiple autoimmune diseases among patients with OCD and their first-degree relatives\(^{11}\); we have also described higher-than-expected rates of immune-related conditions in individuals with CO-OCD.\(^{12}\) Positron emission tomography imaging in adults with OCD has demonstrated increased volume of translocator protein-18 distribution in cortico-striato-thalamo-cortical circuits, implicating widespread microglial activation.\(^{13}\) It is unclear whether changes in cellular and soluble inflammatory markers represent underlying aetiology, a consequence of disease progression or associated epiphenomena.

Non-steroidal anti-inflammatory drugs (NSAIDs) inhibit cyclooxygenase (COX) enzymes, which catalyse the metabolism of arachidonic acids to prostanooids. COX-2 and its products play an important physiological role in synaptic plasticity and long-term potentiation and may also contribute to neuropsychology by enhancing glutamate excitotoxicity, promoting neuronal cell death and oxidising endogenous cannabinoids.\(^{14,15}\) Recent meta-analyses suggest a potential role of adjunctive COX-2 inhibitors in the treatment of depression\(^{16}\) and first-episode schizophrenia,\(^{17,18}\) with additional small studies suggesting possible benefit in neurodevelopmental conditions including autism spectrum disorder.\(^{19,20}\) Behavioural effects of COX inhibition may relate not only to anti-inflammatory activity but also to direct effects on neuronal function and synaptic transmission. Consensus guidelines on the use of anti-inflammatory therapy in children with PANDAS suggest NSAIDs as first-line options for patients with mild impairment.\(^{21}\) However, a recent systematic review of treatment for PANS/PANDAS found insufficient evidence to support this practice.\(^{22}\) In adults with OCD, three small randomised controlled trials (RCTs) have suggested modest symptom improvement with celecoxib as an adjunct to fluoxetine,\(^{23}\) fluvoxamine,\(^{24}\) or other selective serotonin reuptake inhibitors.\(^{25}\) This raises the possibility that COX-2 inhibition may be effective in a general OCD population.\(^{26}\) However, no controlled studies to date have tested the effects of COX inhibitors in CO-OCD. This study will determine the efficacy of the COX-2-selective inhibitor celecoxib as an adjunct to treatment-as-usual in children and youth with moderate-to-severe OCD. There is no cost to participate in the study.

METHODS AND DESIGN

Study design

The ACE-OCD trial is a randomised, quadruple-blind, placebo-controlled, single-site study comparing a 12-week course of twice daily celecoxib with placebo as an adjunct to treatment-as-usual in children and youth with moderate-to-severe OCD between the ages of 7 and 18. The protocol was drafted in accordance with the Standard Protocol Items: Recommendations for Interventional Trials statement.\(^{27}\) All parents/guardians participating in the study will give electronically-documented informed consent; child and youth participants will provide informed assent or consent.

Study setting

This is a single-site study based at the British Columbia Children’s Hospital (BCCH) Provincial OCD Program in Vancouver, BC, Canada. Study visits will be conducted virtually, using electronic-consent and survey platforms through Research Electronic Data Capture (REDCap) and Zoom, an online videoconference platform that complies with the Personal Information Protection and Electronic Documents Act and the Personal Health Protection Act. Participants will continue to receive treatment-as-usual from their regular healthcare providers, which will not change as a result of participation in this study.

Patient selection

Participants will be recruited from BCCH and based on self-referral through community paediatrics, psychiatry, and psychology practices. Participants may be receiving concurrent pharmacotherapy or psychotherapy according to their routine clinical care, constituting ‘treatment-as-usual’ as long as there have been no changes in the preceding 4 weeks and during the study period. They must have a previous diagnosis of OCD. Participants with PANS or PANDAS who also meet diagnostic criteria for OCD are eligible to participate. Refer to table 1 for full inclusion and exclusion criteria.

Allocation and randomisation

Participants will be randomly assigned to either placebo or celecoxib with a 1:1 allocation as per a computer-generated randomisation schedule stratified by baseline CY-BOCS score (16–23 versus ≥24) using permuted blocks of random sizes of 2 and 4. Specific information regarding the allocation sequence will be stored in a separate document with access restricted to the study’s statistician, the research pharmacist and a research assistant.
Table 1  Inclusion and exclusion criteria

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<th>Criterion</th>
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| **Inclusion** | 1. Age 7–18 years  
2. Resident of British Columbia, Canada  
3. Diagnostic and Statistical Manual of Mental Disorders (DSM-5) diagnosis of OCD based on (1) history of prior clinician assessment and (2) standardised interview  
4. Children's Yale-Brown Obsessive Compulsive Scale score ≥16 (moderate to severe)  
5. Able to take medication twice daily in capsule form (in whole form or sprinkled contents)  
6. Negative pregnancy test (either serum or urine) in participants with childbearing potential  
7. Use of highly effective and/or double barrier contraception, or abstinence, in participants with childbearing potential |
| **Exclusion** | 1. Lifetime diagnosis of autism spectrum disorder, bipolar disorder, psychotic disorder, substance-use disorder, intellectual disability, significant head injury causing loss of consciousness, renal disease, liver disease, gastrointestinal bleeding, peptic ulcer disease, inflammatory bowel disease, severe or uncontrolled asthma, bleeding disorders, heart disease, heart failure or hypertension  
2. Current major depressive episode, acute psychosis, active substance use, suicidality or restriction of fluid intake  
3. Pregnant or breastfeeding during the study period  
4. Active infection or antibiotic treatment at baseline  
5. Allergy to celecoxib, sulfonamide compounds or NSAIDs, including aspirin  
6. Current or previous regular use of immune-modulating therapies for treatment of OCD symptoms, at an effective anti-inflammatory dose (including NSAIDs, corticosteroids, and biologics)  
7. Use of NSAIDs at any dose by a frequency ≥2 times per month during the 2 months prior to randomisation  
8. Current use of intravenous or oral corticosteroids  
9. Concurrent use of CYP2C9 inhibitors fluconazole, amiodarone, oxandrolone or methotrexate; CYP2C9 inducers including rifampin and phenobarbital; or any other drug that may interact with celecoxib and, in the opinion of study physicians, represents a potential safety risk  
10. Poor CYP2C9 metaboliser (ie, CYP2C9*3/*3 genotype) based on clinical suspicion or previous genotyping  
11. Abnormality identified on baseline serology including leucocytosis, leucopaenia, thrombocytopenia, anaemia, abnormal renal function (creatinine >1.5 × upper limit of normal) or abnormal liver function (alanine aminotransferase, alkaline phosphatase, or aspartate aminotransferase >1.5 × upper limit of normal)  
12. New medication started in the 4 weeks prior to baseline, or change in dose in the 2 weeks prior to baseline  
13. Changes in CBT or other psychotherapy in the 2 weeks prior to baseline (ie, change in regular frequency, modality or care provider)  
14. Notable other treatment changes during the study period (either pharmacotherapy or psychotherapy)  
15. No regular physician (family doctor or specialist) providing usual medical care  
16. Participant/parent unable to provide informed consent or assent or participate in self-care, adverse event reporting or follow-up assessments  
17. Inability to have blood pressure measured within 2 months prior to enrolment (either on-site at BCCH or by a primary care provider)  
18. Intention of pregnancy in participants with childbearing potential |

BCCH, British Columbia Children's Hospital; CBT, cognitive–behavioural therapy; NSAIDs, non-steroidal anti-inflammatory drugs; OCD, obsessive–compulsive disorder.

(RA) not involved in the study. The block sizes will not be disclosed to trial implementers.

**Blinding**

Trial participants, investigators, care providers and outcome assessors will be blinded to treatment allocation. Placebo capsules will be identical in appearance to celecoxib capsules. Unique randomisation codes will be used for each participant to avoid inadvertent loss of blinding for all participants in the event that one is unblinded. Data analysis and manuscript writing will be performed after unblinding once data have been cleaned for primary and secondary endpoints and adverse events (AEs). Participants will be provided with an option to be contacted and informed of their allocation at that time.

Emergency unblinding will occur only in exceptional circumstances when required to maintain participant safety—that is, when knowledge of the actual treatment is essential for further management. The blind will be maintained as far as possible and will not be disclosed to other study personnel unless required for patient management. Unblinding will not be a reason for study drug discontinuation.

**Sample size calculation**

The sample size of 80 participants (40 per arm) was estimated on the basis of the primary hypothesis. If we assume a power to detect a minimally clinically significant between-group difference in CY-BOCS scores of 2.5 with an SD of 5 (equivalent to a Cohen’s d effect size of 0.5 and
roughly based on two existing studies of adjunctive celecoxib in adults, a correlation of 0.5 between baseline and final CY-BOCS score, and a sample size of 40 participants per arm, we will have power of 80% to detect a between-group difference using a directional, one-tailed alpha (celecoxib-placebo) using analysis of covariance. Missing follow-up data due to attrition will be imputed (as described in detail in the Statistical Analysis section below). Our recruitment target is similar to pilot studies of adjunctive celecoxib in other psychiatric disorders.

Interventions

Eligible patients will be randomised to a 12-week course of either celecoxib (generic form) or placebo containing microcrystalline cellulose. Participants receiving celecoxib with weight between 10 and 25 kg, inclusive, will receive 50 mg two times per day (2–5 mg/kg per dose); those >25 kg will receive 100 mg two times per day as per Food and Drug Administration (FDA)-approved paediatric dosing in children (maximum 4 mg/kg per dose). The placebo capsule is effectively indistinguishable from that of the drug. Participants will be instructed to take the capsule with food to reduce the risk of gastrointestinal side effects. Those unable to swallow a capsule may sprinkle the contents on moist food, given similar pharmacokinetics compared with an intact capsule. Adherence will be documented by capsule count and adherence questionnaires. Weekly adherence reminders will be provided by email or text. Participants will also be asked to maintain an electronic diary documenting the first dose, missed doses, AEs and changes to the usual way they take the capsule.

Participant schedule and follow-up

Prior to their first study visit, parents/guardians of participants who provide informed consent will complete a full eligibility screening questionnaire followed by a diagnostic interview that includes the Mini International Neuropsychiatric Interview for Children and Adolescents, a short structured interview that covers a broad range of Diagnostic and Statistical Manual of Mental Disorders (DSM-5) psychiatric diagnoses in children and adolescents. Participants and their parents who are eligible to proceed to the first study visit will complete a demographic/medical questionnaire and Participant Perspective Questionnaire (PPQ) via REDCap prior to the first study visit.

Study visits will proceed according to the flow chart in figure 1. Measures completed by a study physician at the first visit include the CY-BOCS, Clinical Global Impression (CGI) scales, review of diagnostic criteria for PANS/PANDAS, tic disorders and restricted food intake, clinician treatment expectancy and clinician experience of remote study visits. Participants who continue to meet eligibility criteria after study visit 1 will be provided with a requisition for monitoring blood work if not already completed (complete blood count, creatinine, aspartate aminotransferase, alanine aminotransferase, electrolytes, pregnancy test). Participants will have the option to consent to participation in an ancillary study for biosample collection (blood, saliva, buccal swab and stool) for future analyses of inflammatory markers.

For participants who remain eligible for randomisation, the BCCH Pharmacy will dispense the study drug or placebo according to an allocation sequence provided to them by the team’s statistician at a dose based on the patient’s weight. For visits 2-4, parents/participants will again complete a REDCap survey prior to each visit, including adherence and AE questionnaires. Participants will be provided with a requisition for blood work to be completed following visit 3. Participants with ongoing symptoms (CY-BOCS >8) at visit 3 will have the option to continue with a 12-week open-label extension with celecoxib, with a follow-up visit and monitoring blood work at 24 weeks.

Outcome parameters and statistical analyses

Primary outcome

The primary outcome is OCD severity as measured by total CY-BOCS score after 12 weeks in the celecoxib compared with placebo arm, adjusted for baseline OCD severity. This is a more powerful statistical approach in comparison to analysis of change scores.

Secondary outcomes

Secondary outcomes include the following: (1) OCD severity after 6 weeks of treatment in the celecoxib compared with placebo arm, adjusted for baseline OCD severity; (2) difference in the proportion of participants achieving a clinically meaningful response (defined as a 25% reduction in the CY-BOCS score or CGI-I of 1 or 2 based on previous meta-analyses) after 6 and 12 weeks of treatment in the celecoxib compared with placebo arm; (3) difference in the proportion of participants achieving clinical remission (CY-BOCS ≤14) after 6 and 12 weeks of treatment in the celecoxib compared with placebo arm; (4) mean CGI of severity (CGI-S) after 6 and 12 weeks in the celecoxib compared with placebo arm, adjusted for baseline OCD severity; (5) mean CGI of improvement (CGI-I) after 6 and 12 weeks in the celecoxib compared with placebo arm, adjusted for baseline OCD severity; and (6) difference between celecoxib and placebo arms in the proportion of participants reporting AEs that are possibly, probably or definitely related to the study intervention. Definitions of response and remission are applied as described previously to allow for cross-study comparability.

Exploratory outcomes

Exploratory analyses will include determination of the associations among age, sex, race/ethnicity, body mass index (BMI) percentile, treatment at baseline, severity at baseline, presence/severity of PANS/PANDAS symptoms or tics at any time point based on clinician assessment, medical/psychiatric comorbidities, time since

Open access
Recruitment from BCCH or self-referral from community; no change to usual care

↓

Informed consent; phone screening; diagnostic interview

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Participant/parent questionnaires Set 1:
- PPQ including PGI and treatment expectancy
- Medical/demographic questionnaires

Study visit #1

Clinician Measures Set 1: CY-BOCS, CGI, PANS/PANDAS and tic assessments, treatment expectancy

Randomization
(WEEK 0)

Celecoxib
n=40

Placebo
n=40

Participant/parent questionnaires Set 2:
- PPQ; adherence and adverse events

Study visit #2
Week 6

Clinician Measures Set 2: CY-BOCS, CGI, PANS/PANDAS and tic assessments, treatment expectancy

Participant/parent questionnaires Set 3:
- PPQ; adherence and adverse events

Study visit #3
Week 12

Clinician Measures Set 3: CY-BOCS, CGI, PANS/PANDAS and tic assessments

Optional open-label celecoxib extension

Study visit #4
Week 24

Baseline bloodwork:
- CBC, Cr, electrolytes, liver enzymes, pregnancy test; optional ancillary study

Baseline measurements:
- Height, weight, blood pressure

Follow-up bloodwork:
- CBC, Cr, electrolytes, liver enzymes; pregnancy test; optional ancillary study

**Figure 1** Flow diagram of study visits and assessments. *a*MIND-Kid diagnostic interview administered by phone with the participant and parent present. *b*Screening and study visits may be conducted virtually according to patient preference and current COVID-19 restrictions. *c*Height, weight and blood pressure will be determined either on-site or by a participant's regular care provider. *d*Participants will inform study staff of the date and time of their first dose. Weekly reminders regarding adherence and completion of the participant e-diary as required will be sent via email, phone or text according to participant preference and consent. CY-BOCS, Children's Yale-Brown Obsessive Compulsive Scale; BCCH, British Columbia Children's Hospital; CGI, Clinical Global Impression; PANDAS, paediatric autoimmune neuropsychiatric disorder associated with streptococcal infections; PANS, paediatric acute-onset neuropsychiatric syndrome; PGI, Patient/Parent Global Impression; PPQ, Participant Perspective Questionnaire.
The CGI-I typically but not always tracks with CGI-S and has been used to define treatment response in treatment trials of paediatric OCD.

**Participant Perspective Questionnaire**
This is a study-specific questionnaire to be completed online by the participant and parent in conjunction with each study visit. Included measures are listed in table 2.

**PANDAS/PANS scale**
Included in the PPQ, this rating scale assesses severity and change in PANS/PANDAS symptoms and is a parent-report form based on criteria proposed by the PANS Consortium and described previously. This measure has also been used to capture PANS exacerbation, as it asks the rater whether each current symptom had been possibly worse (1 point), dramatically worse (2 points), new (3 points) or better/same (0 points) within the past week. The maximum score possible is 54.

**Clinician assessment**
In addition to assessment of OCD severity, PANS/PANDAS diagnosis and review of tic symptoms/severity will be conducted by the clinician at all study visits. This will include CGI measures for tics and for food intake restriction (a PANS criterion). Clinicians will also complete several questions related to treatment expectancy and their experience of virtual study visits.

**Adverse events**
AEs will be systematically assessed at study visits 2-4 using a questionnaire adaptation of the Safety Monitoring
Uniform Research Form (SMURF). The SMURF is an AE-elicitation tool specifically aimed at paediatric populations, developed by the NIMH-funded Research Units on Paediatric Psychopharmacology. A checklist will also be included in participant electronic diaries to allow for standardisation of reporting and to facilitate recall when completing the AE Questionnaire prior to the visit.

Adherence
Medication adherence questionnaires will be completed on REDCap by participants prior to visits 2-4 and will be reviewed with the family by the RA and study physician. This will consist of two questions regarding the frequency with which participants have taken all doses or missed one dose, with the response rated on a visual analogue scale. An open-ended question will be included regarding the reason for any missed doses. Adherence will also be assessed by capsule count at the end of the study.

Safety monitoring and interim analysis
This study will be reviewed by a data safety and monitoring board (DSMB). An interim analysis of recruitment rates and AEs will be conducted after the first 10 patients and the first year of recruitment. Unblinding of data will occur only at the request of the DSMB by a statistician not directly involved in the conduct of the study. No interim efficacy analysis is planned. Individual participants may be directly involved in the conduct of the study. No interim analysis of recruitment rates and AEs will be conducted after the first 10 patients or the first year of recruitment. Unblinding of data will occur only at the request of the DSMB by a statistician not directly involved in the conduct of the study. No interim efficacy analysis is planned. Individual participants may be asked to leave the study for their own safety, in which case a protocol deviation and analysis plan will be completed.

Statistical analyses
Analyses will be carried out according to the intention-to-treat principle such that participants will be analysed according to the group to which they were randomised regardless of adherence. Descriptive statistics will be conducted on baseline variables to evaluate the characteristics of the total sample and subsamples in each treatment condition. The primary analyses will be conducted on two sets of data. First, the analyses will be conducted on complete case data, which is defined as the set of subjects without missing data on the variables included in the particular statistical model. Second, missing data will be multiply imputed using the multivariate imputation by chained equations approach, which is appropriate when data are missing at random or are missing completely at random. The imputation method for all variables will be semi-parametric predictive mean matching, which restricts imputations to the observed values in the data set. Common diagnostics, including visual inspection of trace plots and examination of R-hat values, will be used to ensure the validity of the imputation procedure. The imputation model will include baseline demographic and clinical characteristics used to form subgroups for exploratory analyses, treatment condition, baseline scores on outcomes measures, as well as observed follow-up scores on the outcomes of interest. The imputation model will create forty imputed data sets, on which statistical analyses will be performed. Statistical estimates will be pooled over the 40 imputed data sets using the Barnard-Rubin procedure to estimate pooled SEs and df.

**Primary analysis**
The primary analyses will compare the celecoxib to the placebo group on the outcome of interest at weeks 6 and 12, adjusting for baseline scores on the outcome, using analysis of covariance (ANCOVA) for continuous outcomes and logistic regression for categorical outcomes. Additionally, baseline CY-BOCS will be included as a covariate in all analyses, even when CY-BOCS is not the outcome variable. ANCOVA produces unbiased treatment effect estimates and less variance in the treatment effect as compared with the commonly-used linear mixed model, resulting in superior statistical power. All continuous outcomes believed to be generated from a Gaussian distribution will be analysed using this approach.

The primary contrast for this study will be the between-group difference (celecoxib vs placebo) in CY-BOCS score at 12 weeks, adjusted for baseline CY-BOCS score, using multiply imputed data and complete case data. The estimated between-group difference using the multiply imputed data will be considered the primary estimate; the estimated between-group difference using complete case data will be considered secondary. The statistical significance threshold for this analysis will be set at a one-sided alpha=0.05, to test whether the celecoxib group has lower adjusted 12-week CY-BOCS scores compared with the placebo group. For this analysis, we will report the between-group point estimate, 95% CI, and p value to three decimal places. A p value less than 0.001 will be reported as p<0.001. Additional analyses of between-group differences in secondary outcomes and in symptom severity at the midpoint assessment will be considered descriptive and will be described using point estimates and 95% CIs.

**Secondary analysis**
Secondary analyses will include analysis of the proportion of patients in each group who achieve a 25% reduction in CY-BOCS score from baseline or CGI-I of 1 or 2 (treatment response) and who achieve a CY-BOCS score ≤14 (remission). Logistic regression will analyse between-group differences in this binary outcome (achieved ≥25% reduction vs did not), adjusting for baselineCY-BOCS score. Similarly, logistic regression will examine group differences in a binary AE variable (experienced at least one AE vs did not), also adjusting for baseline CY-BOCS score. Association between OCD symptoms, Patient/Parent Global Impression (PGI), and treatment expectancy will be estimated using linear regression modelling with treatment group, age, sex, BMI percentile, race/ethnicity, PANS/PANDAS status and tic status as covariates.
Rationale for use of a COX-2-selective versus COX-1-selective inhibitor

While all NSAIDs appear to have anti-inflammatory, antipyretic, and analgesic properties attributable to prostaglandin inhibition, they vary with respect to COX selectivity\(^2\) and may have neuroprotective effects not directly related to their classic anti-inflammatory activity.\(^3\)\(^4\) In the CNS, modulation of glutamate, serotonin, norepinephrine and endocannabinoid signalling has been primarily demonstrated with COX-2 rather than COX-1 inhibitors.\(^4\)\(^5\)\(^6\)\(^7\) Other than a negative RCT of naproxen in geriatric depression\(^8\) and a study of adjuvant aspirin in schizophrenia,\(^12\) few RCTs have evaluated non-selective NSAIDs in primary psychiatric disorders. Given the significance of different COX isofoms and their unknown relative ‘potencies’ in the CNS, careful attention must be given to selection and evaluation of specific NSAIDs to better understand their neurobiology and clinical efficacy. This study uses celecoxib rather than naproxen given evidence of benefit in adults with OCD and preclinical data pointing to modulation of serotonin and glutamate. Celecoxib is also associated with fewer gastrointestinal side effects in adults.

Rationale for dosing regimen

The US FDA has approved the use of celecoxib in the paediatric population for the management of juvenile idiopathic arthritis (JIA)\(^15\) and it is available in the US to children from ages two and up based on a non-inferiority study comparing celecoxib with naproxen.\(^31\) A follow-up registry study from routine clinical practice included 274 children on NSAIDs and found that AEs were similar for non-selective NSAIDs and celecoxib, and that no serious AEs were attributed to NSAID use over a mean duration of treatment of 11–13 months.\(^52\) The dosages used were within the range of those tested in children with JIA over 12 weeks (3–6 mg/kg two times per day).\(^51\) To avoid exceeding plasma levels associated with the 6 mg/kg suspension, the FDA-approved capsule dosing will be used in this study.

Strengths and limitations of this study

While an RCT of naproxen in PANDAS is currently recruiting (NCT04015596), this study has broader inclusion criteria based on emerging evidence for inflammatory dysregulation in ‘classic’ OCD and existing data in adults. The pragmatic approach of adding celecoxib to treatment-as-usual is a potential strength reflecting typical use in clinical practice. Because of this, our study population is likely to be more heterogeneous than that of existing adult studies. It is difficult to predict to what extent and in which direction selection bias will affect the representativeness of the study population, as in our clinical experience families often consider anti-inflammatory therapy at all stages and severities of the disorder. A subset of children may benefit from immune-modulating therapies, but there are no validated strategies for identifying these individuals. This study incorporates

**DISCUSSION**

This study will be the first to assess the efficacy of celecoxib in paediatric OCD. Multiple lines of evidence suggest behavioural effects of COX inhibition, which may relate not only to anti-inflammatory activity but also to direct effects on neuronal function and synaptic transmission. While clinical phenotyping will identify children meeting criteria for PANS/PANDAS, this work will also bring much-needed attention to a heterogeneous population of patients with OCD and may inform future trials of immune-modulating therapies. Participant perspectives on treatment expectancy, outcomes, and trial participation will be used to inform the design of future studies in this population.

**ETHICS AND DISSEMINATION**

**Data collection and confidentiality**

All data are handled confidentially and the information in the datasets for analyses is non-identifiable.

**Ethics**

A no objection letter has been received from Health Canada. This study has been approved by the University of British Columbia/Children’s and Women’s Health Centre of British Columbia Research Ethics Board.

**Withdrawal**

Patients will be informed of their right to withdraw from the study without explanation at any time.

**Dissemination plan**

The findings will be disseminated in peer-reviewed academic journals and presentations to multiple stakeholders including patients, parents and healthcare providers.

Other analysis

Data collected during the 12-week extension period will be reported in a descriptive fashion, for example, number of observations, percentages, means and SDs.

**Patient and public involvement**

The research question addressed by the study has been informed by discussions with families interested in trialling NSAID therapy and the current lack of evidence base to inform treatment recommendations. Feedback from families has been incorporated into trial design, including addition of an open-label phase. Procedures for recruitment, assessment, BioBank sample collection, outcome assessments, follow-up and results dissemination are common to other studies in the BCCH Provincial OCD Program that have provided both patients and families with an opportunity for input. Because this trial is unique in incorporating virtual/remote study visits for a pharmacological intervention within a paediatric psychiatric population in BC, participants’ perspectives on their participation may provide critical information relevant to the design of future studies.
biosample collection preintervention and postintervention, allowing not only for safety monitoring but also for future analyses of pro-inflammatory markers. Given the paucity of data from interventional trials examining longitudinal markers of inflammation and treatment response in paediatric OCD, this will generate much-needed preliminary data to inform further studies of immune-related biomarkers. Due to funding limitations, these samples will be allocated for future analyses.

This study incorporates questionnaires aimed at better understanding participants’ experiences of virtual study visits, which is a novel format for psychopharmaceutical trials at our centre in the context of the COVID-19 pandemic and will increase equitable access to opportunities for research participation. We expect that these data will inform the design of future studies incorporating remote research visits and clinical care.

CONCLUSIONS
NSAIDs are common in clinical practice and referenced in both adult and paediatric treatment guidelines for OCD, but no controlled studies have evaluated the effects of COX inhibitors in CO-OCD. This study will be the first to assess the efficacy and safety of adjunctive celecoxib in this population and will inform clinical management of children and youth with OCD.

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Contributors CW-R drafted the initial protocol under the supervision of SES, who revised for significant content. JRB created the statistical analysis plan. MM, SB, DE and LT provided clinical input into study design and monitoring. AA, ZN, BL and CL drafted subsections of the initial protocol and facilitated research ethics board submission. All authors revised the protocol and approved of the final version to be submitted.

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Competing interests The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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