Of the protocol at the STUDY PROTOCOL CTN 238

A RANDOMIZED CONTROL CLINICAL TRIAL OF MICRONUTRIENT & ANTIOXIDANT SUPPLEMENTATION IN PERSONS WITH UNTREATED HIV INFECTION

The MAINTAIN Study

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i. LAY SUMMARY

Infection with human immunodeficiency virus (HIV) causes decline in immunity or the ability to fight infection and progresses to acquired immunodeficiency disease (AIDS). Anti-HIV drug treatment has improved the prognosis of persons with HIV infection, but is expensive and potentially toxic. Low micronutrient levels occur in the blood even in early stages of HIV infection and increase risk of a poorer prognosis, but the role of micronutrient and antioxidant supplements in medical management of HIV/AIDS is not well defined. The proposed clinical trial aims to assess if supplementation of untreated HIV-infected adults with a micronutrient and antioxidant preparation (109 study participants) can delay disease progression or start of anti-HIV drug treatment compared with supplementation with standard multivitamins (109 participants). Participants will be treated for two years and followed quarterly in clinic for two years, starting anti-HIV drug treatment as needed according to current treatment guidelines. We will study changes in number of immune cells in the blood, tolerance of study medication, and indicators of safety, well being and health. We expect supplementation with study medication to delay disease progression or start of anti-HIV medication. If the findings are positive, the study has implications for health and health care savings.
ii. EXECUTIVE SUMMARY

Background: Antiretroviral therapy (ART) has improved the prognosis of persons with human immunodeficiency virus (HIV) infection, but is expensive and potentially toxic. Micronutrient deficiencies occur even in early stages of HIV infection and increase risk of morbidity, disease progression to acquired immunodeficiency syndrome (AIDS) and mortality, but the role of micronutrient antioxidant supplements in medical management of HIV/AIDS is not clear.

Objective: To determine if supplementation of untreated asymptomatic HIV-infected persons with a broad-spectrum micronutrient and antioxidant preparation will reduce the rate of decline of CD4 T lymphocyte count, or delay emergence of documented CDC-defined AIDS-defining illness, or start of ART compared to 100% recommended daily allowance (RDA) multivitamins and minerals, and is safe.

Study design: A prospective, randomized, controlled, double blind clinical trial of supplementation of 218 untreated asymptomatic HIV-infected adults with a micronutrient and antioxidant preparation or identical appearing RDA multivitamins and minerals for two years, with quarterly follow-up in clinic for assessment of time from baseline to CD4 count <350 mm³, or emergence of documented CDC-defined AIDS-defining illness, or start of ART.

Participants and sample size: 218 participants from clinics in Ontario and other participating centres of the CIHR Canadian HIV Trials Network (CTN).

Study duration: Three years, allowing one year for participant accrual and two years follow-up.

Eligibility criteria: The main eligibility criteria are:

- Asymptomatic HIV-infected adults at least 18 years of age
- CD4+ cells ≥375 and ≤750 cells/mm³
- No previous ART (excluding less than seven days and perinatal transmission prophylaxis)

Study intervention: Oral supplementation with a broad spectrum micronutrient and antioxidant preparation (n=109) or identical appearing RDA multivitamins and minerals (n=109).

Primary outcome: Time from baseline to CD4+ cell count <350 cells/mm³ (confirmed by two measures at least one week apart), or emergence of documented CDC-defined AIDS-defining illness, or start of ART.

Secondary outcomes:

- Non-AIDS related adverse events
- Tolerance of and adherence to study medication
- Time from baseline to CD4+ cell count <350 cells/mm³ (confirmed by two measures at least one week apart)
- Time from baseline to emergence of documented CDC-defined AIDS-defining illness
- Time from baseline to start of ART
- Serial quarterly lymphocyte measures: absolute lymphocyte count (ALC), CD4+, CD8+, and CD3+ cell counts, CD4%, CD8%, CD4:CD8
- Serial quarterly HIV RNA plasma viral load
- Serum chemistries: Glucose, BUN, creatinine, total protein, C-reactive protein, albumin, alkaline phosphatase, ALT, AST, total bilirubin
- Serum micronutrient levels: Carotene (quarterly) and vitamin B₁₂ (quarterly), folate (six monthly) and vitamin D (25-OHD, six monthly)
- Quality of Life measures: MOS HIV, EuroQol, and Health Utilities Index (HUI)

Statistical analysis: Analysis of the primary outcome by intention-to-treat will compare time from baseline to primary outcome. Interim analyses are planned once 100 participants are followed for one year.
iii. STUDY FLOWCHART

218 consenting eligible participants will be enrolled during routine clinic visits and randomized to two groups. Neither research staff nor participants will know assigned treatment.

<table>
<thead>
<tr>
<th>Treatment group</th>
<th>Control group</th>
</tr>
</thead>
<tbody>
<tr>
<td>n=109</td>
<td>n=109</td>
</tr>
<tr>
<td>Will receive daily a micronutrient and antioxidant preparation for two years</td>
<td>Will receive identical appearing standard multivitamins</td>
</tr>
</tbody>
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Participants will be followed in quarterly clinic visits for two years. They will be offered opportunistic infection prophylaxis and begin ART as indicated according to current treatment guidelines.

On reaching primary outcome (CD4+ cells <350/mm³, or CDC-defined AIDS-defining illness, or start of ART), participants in both groups will be offered open label study medication with continued follow up until two years total, but censored from analysis.

Analyses of primary outcome by intention-to-treat of all participants randomized will compare time from baseline to primary outcome between groups. Secondary outcomes relate to indicators of safety, well being and health.
1. BACKGROUND

Low serum levels of micronutrients are common in persons with human immunodeficiency virus (HIV) infection and acquired immunodeficiency syndrome (AIDS), even in early stages of infection, and are associated with morbidity and mortality, disease progression, and mother-to-child viral transmission. Besides low dietary intake, low micronutrient concentration in HIV-infected persons may be due to malabsorption, diarrhea, altered gut barrier function, and altered metabolism, in addition to the depletion produced by increased levels of oxidative stress.

While micronutrient deficiencies are to be expected in poor countries, they also occur in developed countries, where dietary intake by HIV-infected individuals is adequate and unsupervised micronutrient use is common. Despite the association between low micronutrient levels and poor clinical outcomes in HIV-infected adults, clinical trials of micronutrient supplementation show no conclusive evidence of benefit of micronutrients on morbidity and mortality and on mother-to-child viral transmission, although some but not all studies show some clinical and immunological benefit.

Indefinite antiretroviral therapy (ART) prolongs life, but is expensive and toxic.

In a 12-week placebo-controlled trial, micronutrient and antioxidant supplementation with the proposed preparation showed a significant 24% increase in CD4 T lymphocytes in patients on a stable highly active antiretroviral therapy (HAART) regimen, while without antioxidant supplementation immunologic benefit was modest and slower. In laboratory experiments and in clinical studies, the antioxidants acetyl L-carnitine (LAC), N-acetyl-cysteine (NAC) and alpha-lipoic acid have each shown the potential of beneficial effects in HIV infection. In a randomized placebo-controlled clinical trial in 481 ART-untreated adults with HIV infection, supplementation with micronutrients showed a salutary effect on survival, with no accompanying changes in T lymphocyte counts. In a large randomized placebo-controlled factorial clinical trial in ART-untreated pregnant women with HIV infection, supplementation with vitamins B, C, and E but not vitamin A showed an increase in CD4 and CD8 T lymphocyte counts and delay in the progression of HIV disease when compared to placebo.

β-carotene supplementation enhances cellular immunity. In a cross-sectional study in HIV-1 seropositive adults, low serum concentrations of beta-carotene correlated with markers of HIV infection. In a controlled clinical trial in 331 AIDS patients, low serum carotene was associated with mortality and supplementation with natural mixed carotenoids and multivitamins and minerals was associated with improved survival compared with multivitamins and minerals alone. Supplementation with vitamins B, C, and E but not vitamin A may be associated with reduction in viral load. Deficiency of vitamin A is prevalent in HIV-infected persons despite adequate dietary vitamin B6 intake and has been associated with immunological changes. Vitamin B12 deficiency is prevalent in HIV infection and associated with HIV disease progression.

Supplementation with the antioxidant LAC was associated with a decrease in CD4 T lymphocyte apoptosis in both symptomatic and asymptomatic zidovudine and didanosine treated HIV-infected patients. In a study of oral supplementation of HIV-infected neuropathic patients with LAC for 33 months, distal symmetrical polyneuropathy symptoms improved, accompanied by significant nerve fiber regeneration.

Laboratory studies show that NAC is associated with rescuing of peripheral blood lymphocytes from apoptosis, by protection from cyclohexamide and necrosis factor-alpha (TNF). Sulfur is lost in HIV-infected patients as a consequence of peripheral tissue cysteine catabolism and this loss is not ameliorated by HAART. Supplementation of NAC in HIV-infected adults in placebo-controlled randomized clinical trials was associated with a significant increase in...
immunological function\textsuperscript{26,27} and in exposed HIV-negative individuals to increased release of chemokines that may decrease susceptibility to HIV infection\textsuperscript{28}.

Alpha lipoic acid has a dose dependent inhibition of HIV-1 replication in cytopathic effect formation, reverse transcriptase activity and plaque formation on CD4-transformed HeLa-cells and no toxic effect on blood mono-nuclear cells and T-cell lines\textsuperscript{32}. Micronutrient interactions may also be associated with reduced morbidity of infections\textsuperscript{44} and improved response to drugs\textsuperscript{45}.

We propose a clinical trial to assess if micronutrient and antioxidant supplementation in untreated asymptomatic HIV-infected persons will reduce the rate of decline in CD4 T lymphocyte count, or emergence of CDC-defined\textsuperscript{46} AIDS-defining illness, or start of ART versus recommended daily allowance (RDA) multivitamins and minerals. Since multivitamins are commonly used by HIV-infected persons\textsuperscript{4} and it is not our intent to evaluate study medication in the absence of multivitamins, participants in the control arm will receive RDA multivitamins and minerals. In our previous clinical trial of mixed carotenoids supplementation also, both the treatment and control arms received RDA multivitamins and minerals\textsuperscript{38}.

\textit{Safety of the proposed medication}

The proposed medication has been used safely for three months in HIV-infected individuals\textsuperscript{21} and is sold over the counter in the United States. Other longer studies of micronutrient and antioxidant supplementation in comparable dosage in HIV-infected and -uninfected adults show no harm\textsuperscript{38,47}. A meta-analysis of clinical trials shows no harm of antioxidant supplementation, except possible increased mortality with some antioxidants in a secondary analysis\textsuperscript{48}. A possible adverse effect of β-carotene in smokers and those exposed to asbestos\textsuperscript{49}, and with high intake in HIV-infected adults in an observational study\textsuperscript{50} have not been confirmed in large randomized studies. Carotenemia and skin discoloration, both reversible conditions, have not been reported in HIV infection even with beta-carotene doses higher than proposed here. In non-HIV individuals, flushing of the skin that is generally mild and transient occasionally occurs with nicotinic acid (vitamin B\textsubscript{3}) at a daily dose greater than 10 mg. Supplementation of nucleoside reverse transcriptase inhibitor-treated HIV-infected adults with vitamins C and E and NAC was associated with increased fasting glucose levels at 24 weeks, but the study was small and uncontrolled and all the participants either had lipoatrophy or sustained hyperlactatemia\textsuperscript{51}. These and other micronutrients and antioxidants were well tolerated in other studies\textsuperscript{21,38}. There is no evidence of harm of iron supplementation in HCV co-infected HIV-infected adults\textsuperscript{52}. Micronutrient and iron supplementation of coinfected female injection drug users improves anemia without increasing plasma HCV or HIV RNA levels or altering liver enzymes\textsuperscript{53}, and no harm has ever been shown at the proposed iron dosage of 18 mg. Hepatitis coinfection patients are not excluded from the study as they are more likely to be nutrient depleted and oxidative stressed and could be helped by the intervention.

2. HYPOTHESES

\textbf{Primary}: Supplementation of untreated asymptomatic HIV-infected persons with broad spectrum micronutrients and antioxidants will delay decline from baseline in CD4 T lymphocyte count to \textless 350 cells/mm\textsuperscript{3}, or emergence of CDC-defined AIDS-defining illness\textsuperscript{46}, or start of ART compared with 100% RDA multivitamins and minerals.

\textbf{Secondary}: Supplementation of untreated asymptomatic HIV-infected persons with broad spectrum micronutrients and antioxidants will be well tolerated and safe.
3. OBJECTIVES

3.1 Primary Objective
- To evaluate if supplementation of untreated asymptomatic HIV-infected adults with micronutrients and antioxidants will reduce the rate of decline of CD4 T lymphocyte count, or delay emergence of CDC-defined AIDS-defining illness\(^46\), or delay start of ART versus RDA multivitamins and minerals.

3.2 Secondary Objectives
- To evaluate safety and adverse events
- To evaluate if study medication is well tolerated
- To evaluate effect of study medication on T lymphocyte immunophenotype subsets
- To evaluate effect of study medication on emergence of CDC-defined AIDS defining illness
- To evaluate if study medication can delay initiation of ART
- To evaluate effect of study medication on plasma HIV RNA viral load
- To evaluate effect of study medication on health related quality of life

4. OVERVIEW OF STUDY DESIGN

This will be a prospective randomized, controlled, double blind, multi-centre clinical trial of supplementation of 218 untreated asymptomatic HIV-infected consenting, eligible participants with a micronutrient and antioxidant preparation or 100% RDA multivitamins and minerals to assess time from baseline to CD4 T lymphocyte count <350 cells/mm\(^3\), or emergence of CDC-defined AIDS-defining illness\(^46\), or start of ART. Patients will be approached during routine clinic visits, and consenting participants screened. Eligible participants will be randomized to one of two equal study groups to receive the study or control medications for two years and be followed during quarterly clinic visits for two years. Participants will be offered opportunistic infection prophylaxis. They will begin ART as and when indicated according to current treatment guidelines (2008)\(^54\), with indicated changes to the ART regimen permitted. The primary outcome measure is time from baseline to CD4 T lymphocyte count <350 cells/mm\(^3\), or emergence of documented CDC-defined AIDS-defining illness\(^46\), or start of ART. Secondary outcomes include tolerance of study medication and indicators of safety, well being, and health. Analyses of the primary outcome will be by intention-to-treat of all participants randomized to the intervention, comparing outcome measures between study groups. Once participants in either group reach outcome (CD4 T lymphocyte count <350 cells/mm\(^3\), or emergence of CDC-defined AIDS-defining illness\(^46\), or start of ART), they will be offered open label study medication with continued quarterly follow up, but censored from analysis.

5. PARTICIPANTS

Study participants will be from CIHR Canadian HIV Trials Network (CTN) centres in Ontario and across Canada.

5.1 Inclusion Criteria
The participant must:
- be an asymptomatic HIV infected adult
- at least 18 years of age
• have CD4+ cells ≥375 and ≤750 cells/mm³
• have received no ART (excluding less than seven days and perinatal transmission prophylaxis)
• if a woman of child bearing potential, have a negative pregnancy test within two weeks prior to randomization and agree to practice barrier method of birth control during the study
• be willing and able to sign informed consent and to comply with the study protocol

5.2 Exclusion Criteria
The participant must not:
• have HIV-2 infection alone
• have known allergy or intolerance to any study medication ingredient
• be pregnant
• have active treatment for an acute opportunistic infection or malignancy
• have ALT greater than 3 x normal range
• have known cirrhosis of the liver
• have serum creatinine >133 µmol/L
• abuse alcohol and recreational drugs
• be taking micronutrient (except vitamin D*) or natural health product supplements within 30 days of randomization

6. STUDY DURATION

We expect the study to last three years, with accrual of participants requiring approximately 12 months and two years follow up of the last enrolled participant.

7. STUDY METHODS

7.1 Enrollment and Screening
Participants will be enrolled in the study during routine clinic visits. Research staff will do a preliminary screening for eligibility, explain the clinical trial to the potential participant, ask them to read and sign an information sheet and consent form, and screen for eligibility to participate in the study.

7.2 Randomization
Each participant will receive a unique Participant Identification (ID) number. A permuted blocked randomization method, stratified by centre and CD4+ cell count balanced above and below 500 cells/mm³, will be used to allocate participants to study group. An independent biostatistician from the Ottawa Methods Centre will generate the randomization scheme. The randomization process will consist of a computer-generated random listing of the treatment allocations stratified by centre and with CD4+ cell count balanced equally above and below 500 cells/mm³ in variable permuted blocks of 2 and 4. The randomization scheme output will be given to The Ottawa Hospital (TOH) Pharmacy in an unblinded fashion. Personnel at TOH Pharmacy will prepare the medication kits and assign a participant ID number to each kit, and will send labeled kits to sites. The system will have backup in the form of a statistician and designate at the coordinating centre. Only the study statistician and designate and designated pharmacist(s) at the coordinating centre will have knowledge of treatment allocation. Sites will access the password protected MAINTAIN Study Web Randomization site to obtain the Participant ID number after screening.
Procedure for users (sites) to obtain Participant ID number from MAINTAIN Randomization Web site

After screening the participant for eligibility and obtaining informed consent, the site research nurse (user) will assign the participant a screening number. The user will login to the password protected Web system, and the system will ask a series of questions. The system will take the user to their specific site, and will ask for the participant screening number. The system will check to ensure the screening number is not used already. Users will then be asked if the participant’s CD4+ cell count is below 500 cells/mm³ or if it is 500 or greater. The system will then assign blinded randomization number (Participant ID number) for that participant, and the user will print results and log off the Web system. The participant will receive the pharmacy kit matching their participant ID number.

Methods for protecting against other sources of bias

In order to minimize selection and ascertainment biases, physicians, nurses, investigators, and research staff will be blinded to the randomization schemes and treatments administered. The trial statistician will designate another statistician to prepare all randomization schemes and interim analyses. Only a designated research pharmacist(s) at the coordinating centre will be aware of the treatment allocation for individual participants. The research pharmacist(s) will not have contact with the study teams or participants and will be expressly forbidden to discuss individual treatment allocation with the clinical care teams. To help ensure an effective masking of treatments, we will test whether physicians, research nurses, and participants can physically distinguish the treatment medication from control RDA multivitamins and minerals. This will be conducted before the start of the trial. At the end of the follow-up period, we will ask investigators, outcome assessors, and participants their hunches as to what treatment they received.

7.3 Safety Considerations

Adverse events attributable to the intervention and laboratory parameters of safety will be assessed and monitored by the Data Safety and Monitoring Board (DSMB). If there is concern for safety as a result of the study, or due to new information regarding the study formulation or its components, the Study Steering Committee will make a decision regarding dissemination of the information to the participants. If new information regarding the study or the study medication or its components raises concerns for the safety of the participants, the Study Steering Committee will defer the decision to CTN’s Safety and Efficacy Review Committee (SERC), an autonomous body with authority to discontinue the study. The DSMB and all study committees will include a community representative.

7.4 Outcome Measures

Primary

- Time from baseline to CD4+ cell count <350 cells/mm³ (confirmed by two measures at least one week apart), or emergence of documented CDC-defined AIDS-defining illness⁴⁶, or start of ART

To accommodate biological and systematic variation and confirm the CD4+ cell count measure, each participant will have two measures of CD4+ cell count at least one week apart at baseline (and up to six weeks apart), at outcome (CD4+ cell count <350 cells/mm³), and at end of study (Week 96 or premature withdrawal). CD4+ cell counts will be done in clinical laboratories.
participating in Canadian CD4+ standardization program with the National HIV Laboratory (Ottawa).

**Secondary**
- Non-AIDS related adverse events
- Tolerance of and adherence to study medication
- Time from baseline to CD4+ cell count <350 cells/mm³ (confirmed by two measures at least one week apart)
- Time from baseline to emergence of documented CDC-defined AIDS-defining illness⁴⁶*
- Time from baseline to start of ART
- Serial quarterly lymphocyte measures: absolute lymphocyte count (ALC), CD4+, CD8+, and CD3+ cell counts, CD4%, CD8%, CD4:CD8
- Serial quarterly HIV RNA plasma viral load
- Serum chemistries: Glucose, BUN, creatinine, total protein, C-reactive protein, albumin, alkaline phosphatase, ALT, AST, total bilirubin
- Serum micronutrient levels: Carotene (quarterly), vitamin B₁₂ (quarterly), folate (six monthly) and vitamin D (25-OHD, six monthly)
- Quality of Life measures: MOS HIV, EuroQol, and Health Utilities Index (HUI)
  *See Appendix A (oral thrush is not considered an AIDS-defining illness)

Study outcomes will be measured by procedures at study visits, described in section 7.9 and Table 1 (Section 11).

**Efficacy** will be measured by assessment of the primary outcome of time from baseline to reduction of CD4+ cell count below 350 cells/mm³, or emergence of AIDS-defining illness, or start of ART. Secondarily, efficacy will be measured by assessing each component of the primary outcome, T lymphocyte immunophenotype subsets, plasma viral load, micronutrient levels, and validated quality of life questionnaires. Micronutrient levels measurements will assess baseline deficiency and its correction with treatment. **Adherence** will be measured by pill count, the validated HIV Adherence Treatment Scale (HATS) questionnaire⁵⁵, and serial measurement of serum carotene, which is a measure of intestinal absorption. **Safety** will be measured by adverse events, urinalysis, hematology and serum chemistries.

### 7.5 Disease Progression Requirements
If the participant has experienced any new or recurrent documented HIV-related or CDC-defined AIDS-defining illness⁴⁶ (see Appendix A) since the last visit, the Investigator/study coordinator is required to complete the HIV Disease Progression section of the case report forms (CRF). The HIV-related and AIDS-defining illnesses need not be reported in the Adverse Event section.

### 7.6 Evaluating and Reporting Adverse Events
#### 7.6.1. Adverse Events
The Investigator or designate must complete the Adverse Event Record Form if the participant has experienced a new event or change in existing medical condition or symptom that is non-HIV related, and is either observed or volunteered, serious or non-serious. The reporting of adverse events will be elicited by non-leading questions. The Investigator will designate and document each adverse event as being mild, moderate or severe in nature, and estimate the possible relationship of the event to study medication. The Investigator must record a detailed description of the event with
onset and resolution dates and times, an indication of its severity, and any action taken as a consequence in the CRF. Appropriate procedures to investigate any adverse event may be initiated at the discretion of the Investigator. All adverse events must be followed until they are fully explained or until they have resolved or stabilized.

7.6.2 Serious or Unexpected Adverse Event Reporting
The Investigator shall make an accurate and adequate special report to the coordinating centre (OHRI), and to any Institutional Review Board/Research Ethics Board (IRB/REB) that has reviewed and is continuing to review the investigation, on any serious adverse events. All serious and/or significant unexpected adverse events that occur during the study or post-therapy period, regardless of treatment group or relationship to medication, MUST BE REPORTED WITHIN 24 HOURS BY TELEPHONE TO THE COORDINATING CENTRE. The coordinating centre must receive an initial report of all serious adverse events as soon as practically possible and a written report within 72 hours. All reports of death or life-threatening adverse events must be reported as soon as possible and no later than seven working days to the aforementioned parties.

7.7 Withdrawal from the Study
Participants may withdraw from the study at any time and for any reason, however, they will be urged to stay on their assigned medications in the absence of illness. The coordinating centre (OHRI) or the Investigator may also withdraw the participant at their discretion, or in the event of intercurrent illness, intolerance to study formulation, adverse events, pregnancy, protocol violation, or administrative reasons. All participants discontinued due to an adverse event must be followed until the event resolves, or becomes chronic or stable. Appropriate medical care must be provided until signs and symptoms have remitted, stabilized, or until abnormal laboratory findings have returned to acceptable or pre-study limits. Unnecessary withdrawal of participants should be avoided. If discontinued due to pregnancy, fetal outcome must be monitored, because of unknown potential risk to the fetus.

7.8 Concomitant Medications
While participating in the study and within 30 days of start of the study, the participant may not take another micronutrient (except vitamin D*) or natural health product, and alcohol intake will be limited to three drinks per day or about 50 gm. Illicit and recreational drugs must not be abused. The participant will be offered opportunistic illness prophylaxis. Antiretroviral therapy will be started and changes made as and when indicated according to current treatment guidelines (2008)\textsuperscript{54}. *Daily vitamin D intake must not exceed 2000 IU (Health Canada guidelines).

7.9 Study Visits
Participants will come for study visits as described below. See Section 11 for Study Schedule. Laboratory tests will be conducted at or through the laboratory of each centre. Results of blood tests are acceptable within 30 days of the visit. Each participant will have two measures of CD4\(^+\) cell count at least one week apart at baseline (and up to six weeks apart), at outcome (CD4\(^+\) cell count <350 cells/mm\(^3\)) and at end of study (Week 96 or premature withdrawal). For CD4\(^+\) cell counts, the clinical laboratories must be participating in Canadian CD4\(^+\) cell standardization program with the National HIV Laboratory (Ottawa).

Pre-treatment evaluation (Screening Visit, Week -4 to -2)
The participant will be assessed for study eligibility by matching the results of the following procedures to eligibility criteria:
- Informed consent and demographics
Medical history including previously diagnosed medical conditions, CDC-defined HIV-related or AIDS-defining illness, opportunistic infections, and previous and current medications (including any prior ART and their start and stop dates) and alcohol, illicit and recreational drug, and natural health product use

Complete physical examination, including vital signs and signs and symptoms including of HIV disease

Height and weight, for calculation of BMI

HIV serology, if result not documented, send for testing

HIV RNA (PCR)

Lymphocyte measures: absolute lymphocyte count (ALC), CD4+, CD3+, and CD8+ cell counts, CD4%, CD8%, CD4:CD8

Hematology: white blood cell (WBC) count with differential, red blood cell (RBC) count, hemogoblin, hematocrit, MCV, ESR, platelet count

Coagulation test (INR)

Serum chemistry: Glucose, blood urea nitrogen (BUN), creatinine, total protein, C-reactive protein, albumin, alkaline phosphatase, alanine transaminase (ALT), aspartate transaminase (AST), total bilirubin

Pregnancy test: A urine pregnancy test will be performed at screening for all females able to bear a child. A β-HCG blood test will be performed for re-test of positive or equivocal urine result

Urinalysis; microscopic examination will be performed only if indicated

Baseline Visit (Week 0)

The following procedures will be done at this visit:

Height and weight, for calculation of BMI

Update medical history (see screening visit) and confirm eligibility

Subjective complaints or symptoms

Medications, including over-the-counter and ART, and alcohol, illicit and recreational drug and natural health product use

Targeted physical examination, including for objective findings, e.g., Weight change, fever, diarrhea, oral hairy leukoplakia, seborrheic dermatitis, oral candidiasis, lymphadenopathy, hepatosplenomegaly, and other documented HIV-related or CDC-defined AIDS-defining illness (see Appendix A)

Adverse events (not related to study medication)

Questionnaires (MOS-HIV, EuroQol, and Health Utilities Index (HUI))

Hepatitis serology (HBV DNA or HBsAg) and (HCV RNA or HCV Ab); if results not documented, send for testing

Haematology, consisting of WBC count with differential, RBC count, hemogoblin, hematocrit, MCV, ESR, platelet count

Serum chemistry: Glucose, BUN, creatinine, total protein, C-reactive protein, albumin, alkaline phosphatase, ALT, AST, total bilirubin

Serum micronutrient levels: carotene, vitamin B₁₂, vitamin D (25-OHD) and folate

Coagulation test (INR)

Lymphocyte measures: ALC, CD4+, CD3+, and CD8+ cell counts, CD4%, CD8%, CD4:CD8. CD4+ cell count confirmed by two measures at least one week and up to six weeks apart

Plasma HIV RNA (PCR)

Urinalysis; microscopic examination will be performed only if indicated
• Pregnancy test: A urine pregnancy test will be performed at baseline for all females able to bear a child. A β-HCG blood test will be performed for re-test of positive or equivocal urine result. Subsequently, a pregnancy test will be done only as needed (delayed menses)
• Dispense study medication

Follow up Quarterly Visits (Weeks 12, 24, 36, 48, 60, 72, 84, and 96)

Follow up visits may be scheduled within six days of the actual visit date

• Subjective complaints or symptoms
• Targeted physical examination including for objective findings, e.g., Weight change, fever, diarrhea, oral hairy leukoplakia, seborrheic dermatitis, oral candidiasis, lymphadenopathy, hepatosplenomegaly, and other documented HIV-related or CDC-defined AIDS-defining illness (see Appendix A). A complete physical examination must be done at end of study (Week 96 or premature withdrawal)
• Adverse events and their considered relationship to the study medication (if applicable)
• All adverse events and HIV/AIDS events to be followed until resolution or become stable
• Questionnaires (MOS-HIV, EuroQol, and Health Utilities Index (HUI))
• Weight, for calculation of BMI
• Residual pill count and HATS adherence questionnaire to assess adherence to study medication
• Medications, including over-the-counter and ART, and alcohol, illicit and recreational drug, and natural health product use
• Haematology, consisting of WBC count with differential, RBC count, hemoglobin, hematocrit, MCV, ESR, platelet count
• Serum chemistry: Glucose, BUN, creatinine, total protein, C-reactive protein, albumin, alkaline phosphatase, ALT, AST, total bilirubin
• Serum micronutrient levels: carotene (quarterly), vitamin B₁₂ (quarterly), vitamin D (25-OHD, six monthly) and folate (six monthly)
• Coagulation test (INR)
• Lymphocyte measures: ALC, CD4+, CD3+, and CD8+ cell counts, CD4%, CD8%, CD4:CD8. CD4+ cell count confirmed by two measures at least one week apart at outcome (CD4+ cell count <350 cells/mm³) and end of study (Week 96 or premature withdrawal)
• Plasma HIV RNA (PCR)
• Urinalysis; microscopic examination will be performed only if indicated
• Dispense study medication up to Week 84 visit

Study termination visit (two to six weeks after 96-week visit or premature withdrawal from the study)
All the procedures of a quarterly visit will be done.

7.10 Sample Size and Statistical Considerations
7.10.1 Sample Size Summary
Estimated time to CD4 count <350 cells/mm³ or start of ART was based on previous research⁵⁶ and expert opinion. In the UK study, 45% of participants had either a CD4+ count below 350/mm³ or started ART at 2 years and 60% met the outcome at 3 years⁵⁶. In addition to the CD4+ count <350/mm³ and start of ART estimates based on more than 25,000 patients, we will measure all observed CDC-defined AIDS-defining illness. As such, we have estimated that the control median time from baseline to CD4+ count <350 cells/mm³, or emergence of documented CDC-defined
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The MAINTAIN Study

AIDS-defining illness, or start of ART to be 50% at 2 years. Based on our previous trial, and expert opinion and consensus, we hypothesized that treatment with micronutrients and antioxidants would decrease the median to 32% (18% absolute change in median at 2 years). A two-sided log rank test with an overall sample size of 218 participants (of which 109 are in the treatment group and 109 in the control group) achieves 80% power at a 0.05 significance level to detect a difference of 18% between the proportions surviving in the control and treatment groups after 2 years. Participants will be enrolled during an accrual period of one year and 50% of the enrollment will be complete when 50% of the accrual time has passed. We have also accounted for a modest non-compliance rate of 15% over the two years for those participants allocated to active treatment medication and 20% for those allocated to control. These estimates are based on consensus of opinion among investigators and the observations seen in our previous trials. See Appendix B for detailed calculation.

7.10.2 Statistical Analysis
Principal analysis of primary outcome measure: All statistical analyses will be based on an intention-to-treat approach. All participants enrolled will be analyzed according to the intervention to which they were allocated, whether they received it or not. The primary outcome (CD4 count <350 cells/mm³, or emergence of CDC-defined AIDS-defining illness, or start of ART) will be analyzed using a two-sided log-rank test. We will also conduct Cox-proportional hazards models to adjust for important prognostic risk factors and/or clinically important baseline imbalances. Participants who reach the primary outcome of confirmed CD4 T lymphocyte count <350 cells/mm³, or emergence of CDC-defined AIDS-defining illness, or start of ART will be censored.

Analysis of secondary outcome measure: Continuous outcomes including T cell measures (ALC, CD4+, CD4%, CD3+, CD8+ count, CD8%, CD4:CD8 ratio, HIV RNA plasma viral load), serum chemistry measures (such as ALT, creatinine, C-reactive protein, glucose, carotene, folate and vitamins B₁₂ and D) will be analyzed using either parametric (independent t-test) or non-parametric procedures (Wilcoxon Rank Sum) followed by generalized linear regression models that adjust for important risk factors. Mean differences in Quality of Life measures between groups will be compared with a two-sample t-test if values are normally distributed. Alternatively, we will use a Mann Whitney test. Adverse events (not related to AIDS) and tolerance of medication will be compared using unadjusted chi-square tests.

Co-interventions, adherence and losses to follow-up: A number of secondary analyses will be performed to better understand the influence of co-interventions, adherence and losses to follow-up on the robustness of the intention-to-treat analysis. The general approaches will include: (a) an analysis of primary and secondary outcomes including only participants who completed the study as per protocol; (b) inclusion of co-interventions in all multivariate analyses; and (c) a best-case, worse-case scenario for losses to follow-up if they exceed 5% in either group.

Interim analyses are planned once 100 participants have been followed for one year.

8. STUDY MEDICATION FORMULATION

8.1 Overview
The micronutrient antioxidant preparation (K-PAX Ultra®) and identical appearing control medication will be supplied by K-PAX Inc. (USA). See Appendix C for composition.
Compassionate provision of study medication
If study results are positive, for one year after the study ends the manufacturer will make available study medication free to control arm participants and at discounted prices to all participants. Provision of the medication and discount will depend on obtaining regulatory approval from Health Canada, as K-PAX is not sold in Canada.

8.2 Dosage and Regimen
8.2.1 Treatment allocation
Treatment group (Group A): Will receive micronutrients and antioxidants.
Control group (Group B): Will receive identical appearing RDA multivitamins and minerals.

8.2.2 Dosage: The treatment and control medications will be taken as one packet (8 capsules) twice a day with meals. Because of the presence of calcium, iron and zinc in the study medication, any other medication must be taken at least two hours before or after taking it. Participants may initiate the intervention at half dose (one packet of 8 capsules once a day) and increase to full dose after one week.

8.2.3 Duration of the intervention
The intervention will last for two years. On reaching primary outcome (CD4+ cell count <350 cells/mm³, or emergence of documented CDC-defined AIDS-defining illness, or start of ART), the participant may be switched to labeled (unblinded) study medication, irrespective of the assigned Group. The participant will continue to be followed for two years total, but will be censored from analysis.

9. ADMINISTRATIVE CONSIDERATIONS

9.1 Confidentiality
All information pertaining to the study, including pre-clinical data, product monograph, protocols, CRFs, and verbal and written information, will be kept strictly confidential and confined to the clinical personnel involved in conducting the study. This information may be relayed in confidence to the REB/IRB or appropriate regulatory authorities, except if required by law. CRFs and all other documents containing confidential participant information leaving the Investigator’s site will not identify the participant by name, but only by a code.

9.2 Ethics Review and Informed Consent
The study protocol and informed consent form were developed with the help of a Community Advisory Committee. Each participating site will obtain approval to conduct this study from their REB/IRB. Each site may make minor modifications to the sample Participant Information Sheet and Informed Consent Form according to the recommendations of their Ethical Review Committee. A copy of the protocol, including protocol amendments, all versions of informed consent forms, information to be completed by participants such as survey instruments or questionnaires must be reviewed and approved by the REB/IRB of each participating centre before implementation. The investigator will notify the REB/IRB of violations from the protocol and serious adverse events. The Investigator will provide a copy of the signed informed consent to the participant, and an original shall be maintained in the participant's medical record. This study will be conducted in accordance with the Therapeutic Products Directorate’s Good Clinical Practice (GCP) guidelines
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(www.hc-sc.gc.ca/dhp-mps/prodpharma/appli-demande/guide-l/i/ch/efficac/e6_e.html) and Health Canada Food & Drug Regulations, Part C Division 5: http://laws.justice.gc.ca/en/F-27/C.R.C.-c.870/index.html and in accordance with the current Declaration of Helsinki (http://www.wma.net/e/policy/b3.htm) and Code of Ethical Conduct for Research Involving Humans (http://www.ethics.ubc.ca/code/july97/). Appropriate reports on the progress of this study will be made by the Principal Investigator to the REB/IRB and the sponsor in accordance with applicable government regulations. Upon completion of the study and finalization of the study report, the Investigator must provide a summary to the REB/IRB.

10. QUALITY ASSURANCE

10.1 Responsibilities of the Institution
The monitoring of the study at each participating site will be in accordance with ICH Harmonized Tripartite Guidelines for Good Clinical Practice and Conduct of Clinical Investigations (Drugs Directorate, Health Canada) (www.hc-sc.gc.ca/dhp-mps/prodpharma/appli-demande/guide-l/i/ch/efficac/e6_e.html). Quality control procedures will be implemented beginning with the data entry system and data quality control checks that will be run on the database generated. Any missing data or data anomalies will be communicated to the site(s) for clarification/resolution.

10.2 Responsibilities of the Investigator(s)
The Investigator undertakes to perform the study in accordance with GCP.

10.3 Responsibilities of the Manufacturer
There will be quality assurance of the study medication with frequent and uninterrupted re-supply. Contents of study medication have been shown to be stable (they were at or above label claim) at one year post-production and will be assayed every six months by the manufacturer.
### 11. STUDY SCHEDULE

#### Table 1: Study Schedule

<table>
<thead>
<tr>
<th>PROCEDURES</th>
<th>Screen Weeks –4 to -2</th>
<th>Baseline Week 0</th>
<th>Weeks 12, 24, 36, 48, 60, 72, 84, and 96 Weeks and Study Termination Visit*</th>
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<tbody>
<tr>
<td>Informed consent and demographics</td>
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<tr>
<td>Medical history**</td>
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<td>Physical examination***</td>
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<td>Subjective complaints or symptoms</td>
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</tr>
<tr>
<td>Adverse events</td>
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<td></td>
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</tr>
<tr>
<td>Resolution/follow up of adverse event</td>
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<tr>
<td>HIV related or CDC-defined AIDS defining illness</td>
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</tr>
<tr>
<td>Resolution/follow up of HIV or AIDS illness</td>
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</tr>
<tr>
<td>Weight</td>
<td>x</td>
<td>x</td>
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<tr>
<td>Medications taken†</td>
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<td>x</td>
</tr>
<tr>
<td>Serum chemistry††</td>
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<td>x</td>
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</tr>
<tr>
<td>Serum micronutrient levels†††</td>
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<tr>
<td>Hematology‡</td>
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<tr>
<td>Coagulation tests (INR)</td>
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<tr>
<td>Plasma HIV RNA</td>
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<td></td>
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</tr>
<tr>
<td>Lymphocyte measures‡‡</td>
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<tr>
<td>HIV serology#</td>
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</tr>
<tr>
<td>Hepatitis serology¶</td>
<td>x</td>
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<td></td>
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<tr>
<td>Quality of life questionnaires</td>
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<td></td>
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<tr>
<td>Residual pill count</td>
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<tr>
<td>HATS Adherence questionnaire</td>
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</tr>
<tr>
<td>Dispense study medications</td>
<td>x</td>
<td>x (up to Week 84)</td>
<td></td>
</tr>
<tr>
<td>Urinalysis (microscopy only if indicated)</td>
<td>x</td>
<td>x</td>
<td></td>
</tr>
<tr>
<td>Pregnancy test§</td>
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</tr>
</tbody>
</table>

Test results are acceptable if results of testing within one month of visit are available

*Follow up quarterly visits may be scheduled within six days of visit date. Study termination visit will be 2-6 weeks after the Week 96 visit or premature withdrawal from the study

**Medical history including previously diagnosed medical conditions, previous opportunistic infections, and previous and current medications, including ART, alcohol, illicit and recreational drug, and natural health product use

***Complete physical examination at screening, end of study (Week 96 or premature withdrawal); targeted physical exam at other visits

†Medications including ART, alcohol, illicit and recreational drug, and natural health product use

††Glucose, blood urea nitrogen (BUN), creatinine, total protein, C-reactive protein, albumin, alkaline phosphatase, alanine transaminase (ALT), aspartate transaminase (AST), total bilirubin

†††Serum carotene (quarterly), vitamin B12 (quarterly), folate (six monthly), and vitamin D (25-OHD, six monthly)

‡White blood cell (WBC) count with differential, red blood cell count (RBC), hemoglobin, hematocrit, MCV, ESR, platelet count

‡‡Absolute lymphocyte count (ALC), CD4+, CD3+, and CD8+ cell counts, CD4%, CD8%, CD4:CD8 ratio. Each participant will have two measures of CD4+ cells at least one week apart to confirm each of baseline (and up to six weeks apart), outcome (<350 cells/mm³) and end of study (96 weeks or premature withdrawal) values

#Test, if result not previously documented

¶(HBV DNA or HBsAg) and (HCV RNA or HCV Ab); test, if result not previously documented

MOS-HIV, EuroQol, and Health Utilities Index (HUI)

§If urine pregnancy test is positive, a serum β-HCG test will be performed. After the screening and baseline visits, a pregnancy test will be done only as needed (e.g., if menses are delayed)
REFERENCES


42. Tang AM, Graham NM, Chandra RK, Saah AJ. Low serum vitamin B-12 concentrations are associated with faster human immunodeficiency virus type 1 (HIV-1) disease progression. *J Nutr* 1997;127(2):345-51.


56. HIV Diagnosis at CD4 Count Above 500 Cells/mm³ and Progression to Below 350 Cells/mm³ Without Antiretroviral Therapy. *J Acquir Immune Defic Syndr* 2007;46(3):257-8.

APPENDIX A

List of Events and Conditions Considered Manifestations of HIV Infection

1. Candidiasis:
   - *Bronchi
   - *Esophagus
   - *Lungs
   - Oropharyngeal (Thrush)
   - *Trachea
   - Vulvovaginal (persistent, frequent, or poorly responsive to therapy)
   - Other Candidiasis

2. Cytomegalovirus:
   - *Retinitis (with loss of vision)
   - *Cytomegalovirus Disease (other than liver, spleen or nodes)

3. Herpes Simplex Virus
   - *Bronchitis
   - *Esophagitis
   - *Pneumonitis
   - *Chronic Ulcer(s) (>1 month in duration)
   - Other Sites

4. Lymphoma (HIV-Related)
   - *Burkitt's (or equivalent term)
   - *Immunoablative (or equivalent term)
   - *Primary of brain

5. Mycobacterial Disease:
   - *Mycobacterium avium complex
   - *Mycobacterium kansasii, disseminated or extra pulmonary
   - *Mycobacterium tuberculosis, any site (pulmonary or extrapulmonary)
   - *Other Mycobacterial species, disseminated or extrapulmonary

6. Other
   - *Cervical cancer, invasive
   - Cervical cancer, in situ
   - Cervical dysplasia
   - *Coccidioidomycosis, disseminated or extra pulmonary
   - *Cryptococcosis, extra pulmonary
   - *Cryptosporidiosis, chronic intestinal (>1 month's duration)
   - *Encephalopathy HIV-related
   - Hairy leukoplakia, oral
   - Herpes zoster
   - *Histoplasmosis, disseminated or extrapulmonary
   - Isosporiasis
   - Kaposi's sarcoma
   - Listeriosis
   - *Pneumocystis carinii pneumonia
   - Pneumonia, recurrent
   - *Progressive multifocal leukoencephalopathy
   - Salmonella septicemia, recurrent
   - *Toxoplasmosis of the brain
   - *Wasting syndrome due to HIV

*AIDS-defining event as described by CDC Surveillance Case Definition of 1993
APPENDIX B
Details of Sample Size Calculation

Two-Sample Log-Rank Power Analysis

**Event:** CD4 count <350 cells/mm³ or emergence of documented CDC-defined AIDS-defining illness or start of ART

Null Hypothesis: Median K-PAX Event Rate at 2 years = Median Control Event Rate at 2 years

Alternative Hypothesis: Median K-PAX Event Rate at 2 years <> Median Control Event Rate at 2 years

<table>
<thead>
<tr>
<th>Power</th>
<th>Sample Size</th>
<th>Event rate K-PAX</th>
<th>Event rate Control</th>
<th>Accrual Time</th>
<th>Follow-up Time</th>
<th>Non-compliance K-PAX</th>
<th>Non-compliance Control</th>
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<tbody>
<tr>
<td>80%</td>
<td>101</td>
<td>50%</td>
<td>24%</td>
<td>1</td>
<td>2</td>
<td>15%</td>
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<td>80%</td>
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<td>20%</td>
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<tr>
<td><strong>80%</strong></td>
<td><strong>218</strong></td>
<td><strong>50%</strong></td>
<td><strong>32%</strong></td>
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<td><strong>2</strong></td>
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<tr>
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<td>276</td>
<td>50%</td>
<td>34%</td>
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<td>2</td>
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<td>363</td>
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<td>36%</td>
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<td>2</td>
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<td>20%</td>
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<tr>
<td>80%</td>
<td>496</td>
<td>50%</td>
<td>38%</td>
<td>1</td>
<td>2</td>
<td>15%</td>
<td>20%</td>
</tr>
</tbody>
</table>

Base Time: 2 years

Proportion loss to follow up in K-PAX during the base time: 0.15000
Proportion loss to follow up in control during the base time: 0.20000
Percent of accrual time until 50% enrollment is reached: 50.00%

**Report Definitions**

Power is the probability of rejecting a false null hypothesis. Power should be close to one. Alpha is the probability of rejecting a true null hypothesis. It should be small. Beta is the probability of accepting a false null hypothesis. It should be small.

**Summary**

A two-sided log rank test with an overall sample size of 218 subjects (of which 109 are in control group 1 and 109 are in the treatment group) achieves 80% power at a 0.05 significance level to detect a difference of 18% between the proportions achieving the outcome in the control and treatment groups after 2 years. Participants will be enrolled during an accrual period of 1 year and 50% of the enrollment will be complete when 50% of the accrual time has past. We have also accounted for a modest non-compliance rate of approximately 15% over the two years for those participants allocated to active treatment medication and 20% for those allocated to control.
## Composition of Medications (Manufacturer: K-PAX (USA))

<table>
<thead>
<tr>
<th>High Potency Antioxidant Formula (K-PAX Ultra&lt;sup&gt;®&lt;/sup&gt;)</th>
<th>Low Dose Multivitamins (equivalent to Centrum&lt;sup&gt;®&lt;/sup&gt;)</th>
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</thead>
<tbody>
<tr>
<td></td>
<td><strong>Daily Dosage</strong></td>
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<tr>
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<td><strong>Vitamins</strong></td>
</tr>
<tr>
<td>Vitamin A (100% b carotene) 20,000 IU 400</td>
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<td>Vitamin C 2,000 mg 3000</td>
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<td>Vitamin E (a tocopherol) 200 IU 667</td>
<td>Vitamin D3 400 IU 100</td>
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<td>Vitamin B-1 60 mg 4000</td>
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<td>Vitamin B12 2.5 mg 41666</td>
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<td>Bioflavinoid Complex 300 mg</td>
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<td>Mixed tocopherols 200 IU</td>
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<td>Choline (Citrate) 60 mg</td>
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<td>Inositol 60 mg</td>
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</tr>
<tr>
<td>Glutamic Acid HCL 400 mg</td>
<td></td>
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<tr>
<td><strong>Potent Antioxidants</strong></td>
<td></td>
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<tr>
<td>Alpha Lipoic Acid 400 mg</td>
<td></td>
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<tr>
<td>Acetyl L-Carnitine 1000 mg</td>
<td></td>
</tr>
<tr>
<td>N-Acetyl Cysteine 1200 mg</td>
<td></td>
</tr>
</tbody>
</table>

*Divided in 16 capsules, taken as one packet of 8 capsules twice daily
†Recommended daily allowance