PHASE III TRIAL OF ANTIBIOTICS TO REDUCE
CHORIOAMNIONITIS–RELATED PERINATAL HIV TRANSMISSION

A Study of the HIVNET Group

Sponsored by:

Division of AIDS, US National Institute of Allergy and Infectious Diseases
US National Institute of Child Health and Human Development
US National Institute on Drug Abuse
US National Institute of Mental Health
US National Institutes of Health

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HIVNET 024

Phase III trial of antibiotics to reduce chorioamnionitis–related perinatal HIV transmission

A Study of the HIVNET Group

Sponsored by:

Division of AIDS, US National Institute of Allergy and Infectious Diseases
US National Institute of Child Health and Human Development
US National Institute on Drug Abuse
US National Institute of Mental Health
US National Institutes of Health

I, the Site Investigator of Record, agree to conduct this study in full accordance with the provisions of this protocol and will comply with all requirements regarding the obligations of clinical investigators as fully outlined in the Statement of Investigator (Form FDA 1572), which I have also signed. I agree to maintain all study documentation for a minimum of two years after FDA clearance or until Division of AIDS, NIAID/NIH and pharmaceutical co-sponsor advise that it is no longer necessary. Publication of the results of this study will be governed by DAIDS policies. Any presentation, abstract, or manuscript will be made available by the investigators to DAIDS, NICHD, and the pharmaceutical co-sponsors for review prior to submission.

I have read and understand the information in the Investigator's Brochure and Package Inserts, including the potential risks and side effects of the product under investigation and will ensure that all associates, colleagues, and employees assisting in the conduct of the study are informed about the obligations incurred by their contribution to the study.

__________________________________       __________________________________
Signature of Site Investigator of Record    Date
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Blantyre, Malawi

Lilongwe Central Hospital and participating antenatal clinics
Lilongwe, Malawi

University Teaching Hospital and participating antenatal clinics
Lusaka, Zambia

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Dar es Salaam, Tanzania
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<td>adverse experience</td>
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<td>zidovudine</td>
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<td>BPM</td>
<td>beats per minute</td>
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<td>Clinical Research Products Management Center</td>
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<td>Data and Safety Monitoring Board</td>
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<td>Family Health International</td>
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<td>gonorrhea culture</td>
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<td>human immunodeficiency virus</td>
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<td>institutional review board</td>
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<td>international units</td>
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<td>LCR</td>
<td>ligase chain reaction</td>
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<td>MCT</td>
<td>maternal-child transmission</td>
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<td>National Institute of Allergy and Infectious Diseases</td>
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<td>NICHD</td>
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<td>NVP</td>
<td>Nevirapine</td>
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<td>Pharmaceutical Affairs Branch, DAIDS, NIAID</td>
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<td>Preparation for AIDS Vaccine Evaluation</td>
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<td>polymerase chain reaction</td>
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<td>PROM</td>
<td>premature rupture of membranes</td>
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<td>QECH</td>
<td>Queen Elizabeth Central Hospital, Blantyre, Malawi</td>
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<td>RAB</td>
<td>Regulatory Affairs Branch, DAIDS</td>
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<td>RNA</td>
<td>ribonucleic acid</td>
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<td>ROM</td>
<td>rupture of membranes</td>
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<td>RPR</td>
<td>rapid plasma reagin (test for syphilis)</td>
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<td>RT-PCR</td>
<td>reverse transcriptase polymerase chain reaction</td>
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<td>SAE</td>
<td>serious adverse experience</td>
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<td>sexually transmitted disease</td>
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<tr>
<td>TID</td>
<td>three times a day</td>
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<td>TPHA</td>
<td>Treponema pallidum hemagglutination</td>
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<td>USAID</td>
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SCHEMA

**HIVNET 024: PHASE III TRIAL OF ANTIBIOTICS TO REDUCE CHORIOAMNIONITIS-RELATED PERINATAL HIV TRANSMISSION**

**DESIGN:** This will be a randomized, double-blinded, controlled Phase III trial of antibiotics to prevent chorioamnionitis-associated perinatal HIV transmission. Using a simple 2-arm design, half the subjects will receive two courses of antibiotics, with the control subjects receiving two courses of a placebo.

**SAMPLE SIZE:** Using a target of reduction in HIV transmission from 11% to 7.5% with 90% power, the sample size will be 3120 HIV-positive pregnant women. In order to prevent stigmatization and to investigate the impact of antibiotics on HIV-negative women in this setting, 600 additional HIV-negative women will be enrolled and treated indistinguishably from those who are HIV-positive.

**POPULATION:** Women will be recruited from hospitals and antenatal clinics. After obtaining informed consent, HIV seropositive and seronegative women will be enrolled, followed during pregnancy, and requested to deliver in the hospital or clinics. Women and infants will attend postnatal follow-up visits in designated clinics.

**REGIMEN:** At 20 – 24 weeks, women randomized to receive antibiotics will receive metronidazole 250 mg three times a day (TID) and erythromycin 250 mg orally TID for 7 days. Women randomized to the control group will receive identically appearing placebos. With the onset of contractions and/or premature rupture of membranes (PROM), study participants will initiate a second oral course of antibiotics consisting of metronidazole 250 mg and ampicillin 500 mg or placebo every 4 hours, continuing after delivery TID until the course is completed. All HIV-infected women and their neonates will be offered the HIVNET 012 Nevirapine (NVP) regimen. If the mother accepts the Nevirapine for herself and her baby, she will be given one oral dose of 200 mg NVP at the 26-30 week antenatal visit to be taken at onset of labor, and her baby will receive one oral dose of 2mg/kg NVP suspension at 72 hours post-birth or discharge, whichever occurs earlier. If the mother refuses Nevirapine or is uninfected, she will receive a matched placebo at the 26-30 week visit to preserve participant confidentiality.

**OBJECTIVE:** To determine if low-cost antibiotic treatment given twice during pregnancy (total cost less than $5.00) aimed at reducing chronic and acute chorioamnionitis will reduce perinatal HIV transmission.
1.0 INTRODUCTION

On February 18, 2003, the Division of AIDS (DAIDS) Data Safety and Monitoring Board (DSMB) reviewed outcome data from nearly 1000 infants with data complete through 2 to 8 weeks. They concluded that, while statistical evidence neither established benefit nor harm, the evidence did rule out targeted levels of benefit in the use of antibiotics to reduce maternal-to-child transmission (MTCT) of HIV. The DSMB therefore recommended that HIVNET 024 stop recruitment. The study sponsor (DAIDS) concurred with this recommendation. Consequently, accrual to HIVNET 024 and administration of the initial course of antibiotics/placebos was halted on February 21, 2003. As of March 5th, dispensing of the labor course of study antibiotics/placebos was halted, all active participants were to be informed of the results of the interim analysis, and sites began retrieving study antibiotics/placebos under the guidance of the Pharmaceutical Affairs Branch (PAB) of DAIDS. In addition, the protocol team leadership agreed that the duration of follow-up of mother/infant pairs should be shortened to 3 months post-delivery rather than 12 months post delivery as all safety and efficacy endpoints will have been met by that time.

The DSMB emphasized that Nevirapine should continue to be dispensed to HIV-infected pregnant women already enrolled in the study as originally intended and described in the protocol (because of its proven efficacy in reducing MTCT of HIV, and because it was not available to HIV-infected pregnant women at the time that the study commenced). The NVP was NOT included in the protocol as part of the study question. However, because it has been provided in the context of this trial and is not licensed in the United States for the prevention of MTCT of HIV, it must be considered an “investigational product” under an IND for this protocol and undergo the safety follow-up and reporting as dictated in the DAIDS reporting manual of serious adverse effects. It is worth noting that, since the start of HIVNET 024 in 2001, the Ministries of Health in all of the participating countries (Malawi, Tanzania and Zambia) have approved the use of NVP for the prevention of mother-to-child transmission of HIV.

This version of the protocol includes amendments to those sections of the protocol that are affected by the recommendations and decisions of the DSMB, DAIDS and the protocol team leadership. The section that has undergone the most revisions is “4.0 Clinical and Laboratory Evaluations.” While no new laboratory assessments have been added, shortening the duration of follow up has necessitated creating a final visit schema that can be applied to all HIV-infected women and their babies who return for follow up at or after the 3 month visit.

1.1 Background

Obstetric risk factors for HIV maternal-child transmission (MCT) include preterm birth, prolonged rupture of the membranes, and both clinical and histologic chorioamnionitis. Each of these risk factors for HIV MCT may operate through a common pathway in which HIV-infected maternal white cells enter the amniotic
fluid following bacterial infections of the placental membranes and result in both “histologic chorioamnionitis” and perinatal HIV transmission.

The relationship between chorioamnionitis and preterm birth has been investigated for a number of years.\(^1\)\(^-\)\(^10\) Findings include the observation that a substantial number of preterm births are associated with and likely caused by chorioamnionitis, with this sub-clinical infection preceding labor or spontaneous rupture of membranes. The earlier the gestational age at onset of spontaneous labor, the more likely it is caused by chorioamnionitis, with histologic chorioamnionitis found in more than 80% of the spontaneous preterm births less than 30 weeks gestation. The organisms associated with this intrauterine infection in every population studied to date are of relatively low virulence and include Ureaplasma, Mycoplasma, Bacteroides, Mobiluncus, and Gardnerella.\(^1\),\(^4\)

Chorioamnionitis prior to delivery is asymptomatic, does not result in fever, chills or abdominal pain, and is not easy to diagnose. The location of the infection, in the chorioidecidual space, is not amenable to culture. Treatment trials have therefore focused on suggestive evidence of this infection and included preterm contractions and the presence of bacterial vaginosis (BV). BV, an overgrowth of Ureaplasma, Mycoplasma, Bacteroides, Mobiluncus, and Gardnerella in the vagina, is associated with and probably a marker for chorioamnionitis. Treatment trials targeting these markers of infection in which metronidazole 250mg TID given orally for a week or more with or without ampicillin or erythromycin have often shown benefit in reducing prematurity, while treatment trials with other antibiotics, or metronidazole given in other dosing regimens, have not been effective. A randomized trial of antibiotics (metronidazole/erythromycin) in women at risk for spontaneous preterm birth who also had BV resulted in a substantial reduction in the rate of preterm birth in that population.\(^7\) Other studies have confirmed this result. In a mass antibiotic treatment trial of STDs in the Rakai district of Uganda using metronidazole in pregnant women, both low birthweight and preterm birth were significantly reduced (Gray RH, et al, unpublished data).

However, in a recently completed NICHD trial targeting vaginal Trichomonas infection, in which metronidazole was given as two 2gm oral doses 48 hours apart, and then repeated 4 weeks later, preterm birth increased – 19% versus 12% in the control group (p<0.001).\(^11\) There are several possibilities for the apparent increase in preterm birth with metronidazole in women with Trichomonas. These include: 1) chance, 2) metronidazole acting to stimulate contractions, or 3) the release of inflammatory products from the action of metronidazole on the Trichomonas organism. In all previous studies, metronidazole use has not been associated with an increase in preterm birth. Therefore, a direct labor-enhancing effect is unlikely. In the U.S., metronidazole is currently the treatment of choice for women with symptomatic Trichomonas, and to date, no one has suggested an increase in preterm birth in women with symptomatic Trichomonas infection treated with metronidazole. In a small study, its use in women with Trichomonas
was not associated with an increase in preterm birth.\textsuperscript{8} This trial used metronidazole and erythromycin as is proposed in the 024 trial, not in doses used in the NICHD \textit{Trichomonas} trial. The addition of erythromycin may also be significant, in that erythromycin is a macrolide antibiotic and has important anti-inflammatory properties in addition to its antibacterial properties. Therefore, even if a release of inflammatory products was the cause of the increase in preterm birth in the NICHD study, the addition of a macrolide antibiotic and the absence of bolus dosing make the 024 approach different. The protocol team believes that the risk of increasing prematurity by the use of the antibiotics in individual women is low, since the bulk of evidence suggests a reduction in preterm birth rather than an increase. Additionally, the populations are different, and as stated earlier, the dosing regimen proposed in this study is different. However, the team remains concerned about a possible interaction between metronidazole use and \textit{Trichomonas} treatment leading to preterm birth, and will monitor outcomes closely for any such relationship.

A better predictor of spontaneous preterm birth than BV is fetal fibronectin (FFN), a placental membrane derived protein which when found in the vagina after 20 weeks gestation is the most potent predictor (odds ratio = 60) of spontaneous preterm birth yet described.\textsuperscript{12} Fetal fibronectin, when found in the vagina at 20 to 24 weeks gestational age, is a strong predictor of neonatal sepsis (odds ratio = 6) and a very potent predictor of clinical chorioamnionitis (odds ratio of 20), and is nearly always associated with histologic chorioamnionitis at delivery.\textsuperscript{13} For reasons of both cost and practicality, it is not expected that fetal fibronectin testing will be used routinely in sub-Saharan Africa. However, if a positive fetal fibronectin result is strongly associated with HIV MCT, our understanding of the mechanism will be greatly increased.

The relationship between BV, preterm birth, histologic chorioamnionitis and perinatal transmission of HIV has been consistently demonstrated. Preliminary results from Malawi show a 2-fold increase (28\% vs. 14\%) in HIV MCT in BV positive vs. BV negative women.\textsuperscript{14,15} Perinatal HIV transmission is more common in preterm infants, and there is now evidence that sub-clinical chorioamnionitis is a substantial risk factor for MCT.\textsuperscript{16,17} For this study, the primary hypothesis is that early and appropriate treatment of sub-clinical chorioamnionitis prior to the onset of spontaneous preterm labor, and/or antibiotic treatment during labor to prevent PROM-associated-chorioamnionitis will reduce the risk of perinatal HIV transmission.\textsuperscript{18}

In this study, a short-course of Nevirapine will be offered to all HIV-infected women and their neonates. Among the non-nucleoside reverse transcriptase inhibitors, Nevirapine is a non-nucleoside benzodiazepine derivative that is a potent inhibitor of HIV-1 replication with an \textit{IC}_{50} of 10 ng/ml, excellent oral absorption and bioavailability, with a high therapeutic index. Nevirapine is highly lipophilic and widely distributed throughout the body, and has been shown to penetrate cell-free HIV-1 and inactivate virion-associated reverse transcriptase in
Nevirapine prophylaxis has been shown to prevent infection in chimpanzees challenged with HIV-1.\(^{19}\) It has been used in adults, children, infants, and as a single dose in the first week of life in neonates.\(^{20-22}\) The drug is well tolerated. Nevirapine is primarily metabolized by the liver. It is an inducer of hepatic cytochrome p450 metabolic enzymes, resulting in an “autoinduction” phenomenon characterized by an approximately 1.5 to 2 fold increase in apparent oral clearance and decrease in terminal half-life after 2-4 weeks of daily dosing in older children and adults. Because of liver metabolism, the half life of Nevirapine in neonates is anticipated to be prolonged compared to older children; the half life of a single dose of Nevirapine administered at age 48-72 hours in neonates was 36.8 hours (range 27.3-49.5 hours).\(^{22}\) The most frequently reported adverse experiences related to Nevirapine are rash, fever, nausea, headache and abnormal liver function tests. The major clinical toxicity is rash, which can rarely be severe or life threatening (Stevens-Johnson Syndrome, occurring in 0.5\% of 1752 adults patients exposed to Nevirapine). Rashes are usually mild to moderate maculopapular erythematous cutaneous eruptions with or without pruritus and located on the trunk, face and extremities. The majority of severe rashes occur in the first 28 days of treatment. No rash toxicity was observed in the PACTG 250, in which infants received a single dose at 48-72 hours of age.\(^{22}\)

Results from HIVNET 012, *A Phase IIB randomized, controlled trial to evaluate the safety, tolerance, and HIV vertical transmission rates associated with short course Nevirapine (NVP) vs. short course zidovudine (ZDV) in HIV infected pregnant women and their infants in Uganda*, were recently published.\(^{23}\) An intrapartum/postpartum regimen of a 200 mg oral dose of NVP given to the mother at the onset of labor and a 2 mg/kg dose given to the infant within 72 hours of life reduced the risk of perinatal transmission among breastfeeding women by 48\% at 14-16 weeks compared to an intrapartum/postpartum regimen of AZT (600 mg, then 300 mg every 3 hours during labor to mother, and 4mg/kg twice daily for one week to the infant). No serious drug-related toxicities were observed in the 626 women and babies who received either drug. In particular, the incidence of rash was similar between both treatment groups, and there were no cases of Stevens-Johnson syndrome. These data demonstrate the safety of a 2-dose NVP regimen, and the reduction in transmission with this NVP regimen persists to 14-16 weeks. This protocol is designed to include a single dose of NVP to the HIV-infected mothers and their infants, as in the HIVNET 012 regimen.

### 1.2 Rationale

This intervention to reduce HIV MCT is chosen for evaluation because:

1. It has biologic plausibility for reducing HIV MCT.
2. It is very low in cost compared to most antiretroviral therapies (i.e., a full course of metronidazole in host countries is $0.60, erythromycin $1.50, and ampicillin $2.50. The two courses of antibiotic treatment will be approximately $4.60). As recommended in a recent meeting sponsored by the Pediatric AIDS Foundation and Emory University, “there is a need to evaluate simple and feasible...
interventions other than antiretroviral agents for their efficacy in diminishing HIV MCT in developing nations."

3. If, ultimately, antiretroviral agents are used only near delivery, it is not likely that they will reduce the portion of HIV MCT occurring earlier in pregnancy. Antibiotic treatment as described in this protocol has biologic plausibility for reducing antepartum as well as peripartum HIV MCT and for achieving additional reductions in HIV MCT over and above that achieved with short course antiretroviral therapy alone, if that treatment is available.

The specific antibiotics to be used prenatally include metronidazole and erythromycin. The rationale for choosing these two antibiotics includes: 1) the organisms discussed above are generally sensitive to one or both of these antibiotics; 2) only metronidazole and/or metronidazole and erythromycin have been shown to be associated with a reduction in spontaneous preterm birth; 3) metronidazole has been used during pregnancy in the United States and elsewhere for a number of years and is safe, especially in the second trimester. Erythromycin and ampicillin are also safe during pregnancy.

An initial treatment course of metronidazole and erythromycin will be given orally three times per day for seven days at 20-24 weeks gestation. This course was chosen for the following reasons: 1) in the second trimester women are being treated for an intrauterine infection and not solely bacterial vaginosis, and 2) the two trials in which a reduction in prematurity has been achieved used a prolonged course of antibiotics (one week or more) in the second trimester.

In labor and with ruptured membranes, since Group B Streptococcus is often the offending organism and rapid antibiotic entrance into the amniotic fluid is an important issue, metronidazole 250 mg in combination with ampicillin 500 mg will be administered orally every 4 hours to prevent the acute chorioamnionitis associated with membrane rupture. Both of these drugs achieve satisfactory serum and amniotic fluid levels within 1-2 hours after oral administration.

Organisms associated with both BV and chorioamnionitis have been similar whether studied in the U.S., Europe, Asia or South Africa, and have not differed substantially between black and white women in the U.S, Great Britain or South Africa. A randomized trial in South Africa showed a reduction in preterm birth in women in early preterm labor with intact membranes who used metronidazole and ampicillin. Therefore, although most studies dealing with infection and preterm birth have been performed elsewhere, it is very likely that they are applicable to sub-Saharan Africa. Even if this study only demonstrates that the antibiotic strategy employed in this protocol reduces preterm births in sub-Saharan Africa, this result in itself will be of major importance to the women of these countries.

Currently, there is little consensus in the United States and other countries about the value of routinely screening for infection and treating pregnant women with
antibiotics either prior to or during labor to prevent preterm birth. For example, while there is some evidence that treating women who had a previous preterm birth who were BV positive reduced preterm birth, there are several recent reports which suggest no benefit when large populations of women are screened and treated. In fact, recently analyzed data from a large-scale NIH trial suggested no reduction in preterm birth using this approach. In view of these findings, no major U.S. organization has yet made a recommendation for widespread screening for and treatment of bacterial vaginosis during pregnancy. Since this practice is not routine in the U.S. and in fact is under study in a number of U.S. locations, the investigators believe it is also appropriate to study antibiotic use during pregnancy using placebo controls in an African setting.

If antibiotic treatment aimed at reducing chorioamnionitis significantly reduces transmission, the study results as well as information regarding the cost of implementing the intervention will be provided to appropriate Ministry of Health officials and other policy makers.

2.0 STUDY OBJECTIVES

While the primary study objective was answered during the interim data analysis, the protocol team will continue to pursue answers to the remaining objectives with the data collected.

2.1 Primary

To determine if antibiotic treatment aimed at reducing chronic and acute chorioamnionitis will reduce HIV MCT.

2.2 Secondary

- To determine if antibiotic treatment reduces preterm birth and maternal/infant morbidity and mortality
- To determine if antibiotic treatment reduces HIV MCT in women with BV or a positive FFN.
- To determine if the reduction in HIV MCT potentially achieved by antibiotics is associated with a reduction in histologic chorioamnionitis, preterm birth, or vaginal viral load.
- In HIV-negative women, to determine if the antibiotic regimen employed in this study is associated with an improvement in birthweight and other perinatal outcomes.
- To determine the correlation between vaginal pH, the whiff test, a non-specific vaginal discharge and clue cells on a wet prep with a Gram stain diagnosis of BV, and to determine the sensitivity, specificity, positive and
negative predictive values of these tests for diagnosing BV and predicting HIV MCT.

The secondary endpoints include preterm birth and infant and maternal morbidity and mortality. In developing countries, most preterm infants die. Even in term infants, mortality is often associated with infections acquired during childbirth. There are substantial data emerging from many sources that preterm birth is associated with a chronic but asymptomatic chorioamnionitis, and that chorioamnionitis at delivery is associated with neonatal infection, morbidity and mortality. It is, therefore, reasonable to hypothesize that antibiotic prophylaxis may reduce preterm birth and its associated morbidity and mortality, as well as the infectious morbidity and mortality associated with chorioamnionitis acquired near delivery. Maternal morbidity and mortality may also be reduced. This study will allow determination in both HIV positive and HIV negative women of whether perinatal morbidity and mortality may be reduced with prophylactic antibiotics.

The investigators are also interested in determining whether or not bacterial vaginosis and fetal fibronectin predict HIV MCT, and if women with these findings are more or less responsive to antibiotics for both the primary and secondary outcomes. The investigators expect the prevalence of BV in this population to be 30-50%. The prevalence of women testing positive for fetal fibronectin is expected to be 6-8%. Both of these “vaginal findings” have been linked to chorioamnionitis. Therefore, there is reason to believe that their presence may be related to both the primary and secondary outcomes. In this study, if antibiotics work to reduce HIV MCT, the cause of the reduction will be of interest. By obtaining the above information, examining the placenta for evidence of histologic chorioamnionitis in the treated women and controls, measuring vaginal viral load and determining if the antibiotics reduced the HIV MCT associated with preterm birth, a better understanding of the relationship between HIV MCT and antibiotics will be achieved.

Finally, although there is preliminary evidence that the presence of BV may be associated with HIV MCT, even this conclusion is uncertain. Furthermore, since BV may be diagnosed in many different ways, and it is not clear which, if any, of the many diagnostic tests predict HIV MCT, correlating this information with the various outcomes may provide useful information about BV and HIV MCT.

3.0 STUDY DESIGN

The original study design is described below. All active participants in HIVNET 024 must be reconsented to participate in the follow-up of this protocol. Sample informed consents for the follow-up are in the appendices; there is a separate consent for those women who were HIV infected at enrollment and those women who were HIV uninfected at enrollment. The original target accrual was 3120 HIV-infected and 600
HIV-uninfected women. At the time that accrual was halted, the trial was approximately 70% enrolled.

This is a randomized, double blinded, controlled phase III clinical trial of antibiotics. Study participants will be randomized to receive either antibiotic treatment or placebo. All HIV-infected participants will be offered the HIVNET 012 Nevirapine regimen. Follow-up will continue until 12 months after the last woman delivers or until sufficient events accrue to provide adequate power for the primary endpoints.

A total of 600 HIV-negative women will be enrolled to conceal the HIV status and avoid stigmatization of participating women at sites where this is deemed appropriate and necessary. This sample of HIV-negative women will also provide valuable information on the secondary objectives (see Section 3.2). Determination of whether the use of this regimen is associated with an improvement in birthweight, a reduction in preterm birth, and a reduction in maternal chorioamnionitis in HIV-negative women, as well as whether adherence to the treatment regimen and tolerance of the antibiotics are similar in the two cohorts, is of interest. Data from the HIV-positive and negative women will not be combined in the analysis.

3.1 Selection and Enrollment of Subjects

The inclusion and exclusion criteria refer to all prospective participants (both HIV positive and negative women with the exception of their HIV status).

3.1.1 Inclusion Criteria

• HIV positive at enrollment;
• 20-24 weeks gestation;
• Willing to give informed consent (for HIV testing and for enrollment into the study);
• Willing to take antibiotic treatment as scheduled;
• Planning to deliver at one of the study sites;
• Willing to come back for follow-up visits for one year post-partum

3.1.2 Exclusion Criteria

• Have taken antibiotics, other than treatment for syphilis or gonorrhea, within the last two weeks;
• Are allergic to penicillin, ampicillin, erythromycin, or metronidazole;
• Have known major illnesses likely to influence pregnancy outcome including diabetes, severe renal or heart disease, or active tuberculosis, prior to randomization;
• Have known major obstetric problems such as placenta previa, ruptured membranes or multiple pregnancy prior to randomization;
• Have known central nervous system diseases, including seizures;
• Are taking anticoagulant drugs.

3.1.3 Enrollment Procedures
Women of unknown HIV status will be recruited at 16-23 weeks gestation from the antenatal clinics. Informed consent for HIV counseling and testing and screening for the study will take place. Women will return to the clinic in one or two weeks to receive their test results. Women who are eligible for the study (those known to be HIV-infected and those who have undergone the screening process) will be scheduled to return for an enrollment visit at 20-24 weeks gestation; these women will be encouraged to bring their baby’s father to the enrollment visit. At the enrollment visit, the study will be explained to the woman and her partner (if available) and informed consent will be obtained. After informed consent is obtained, the woman will be randomized. Randomization will employ permuted block algorithms with randomized block sizes. Randomization of HIV-positive women will be stratified only by site. A separate randomization stratification will occur for HIV-negative women (see section 3.2 below).

3.2 Co-enrollment Guidelines
A sample of 600 HIV-negative women will be included in the study. These women will be randomized using a separate stratified randomization scheme and followed during pregnancy and postpartum (with their babies) in a manner indistinguishable from HIV-positive mother-infant dyads. Inclusion of these HIV-negative women will allow avoidance of stigmatization associated with inclusion of only HIV-positive women from the antenatal clinics at sites where this has been identified as an issue. Furthermore, with a sample of 600 HIV-uninfected women, the effect of antibiotic treatment on some of the secondary outcomes may be evaluable. Establishing a beneficial impact of the proposed treatments among HIV-negative women may justify recommending antibiotics to all women irrespective of their HIV status. There is also potential to study the effect of the antibiotics on early infant morbidity and mortality. Therefore, generalizability of the study findings will be possible.

Enrollment of the 600 HIV-negative women will take place at an appropriate ratio to the HIV-infected participants to insure steady enrollment of negatives over the course of the study. If the selected participant refuses or is not eligible, the HIV-negative woman immediately succeeding her will be approached for participation in the study.

4.0 CLINICAL AND LABORATORY EVALUATIONS

The shortened duration of follow-up has not increased the number or type of clinical or laboratory evaluations to be collected from participants. All evaluations following enrollment through 3 months are unchanged with the exception that:

- no study antibiotics/placebos are to be dispensed and
- the 3-month visit (or the 6-, 9-, or 12-month visit for those mother/infant pairs already beyond the 3-month visit) has become a final visit.
In those rare cases that study antibiotics/placebos were not retrieved from the participant, the site is obligated to complete all documentation relevant to maternal study antibiotics/placebos adherence and to assess for adverse experiences after the first dose and for 2 weeks following the last dose.

In sections 4.1 through 4.3 are the expected clinical and laboratory evaluations for the HIV-infected and HIV-uninfected cohorts and their infants for pregnancy and until three months after delivery/birth; for HIV-infected and HIV-uninfected women and their infants who return for a final follow-up visit at 3, 6, or 9 months after delivery/birth; and for HIV-infected and HIV-uninfected women and their infants who return for a final follow-up visit at 12 months after delivery/birth. Sections 4.4 through 4.8 are from the original protocol and are included for comparison and to provide necessary background information.

Appropriate post-test counseling, as well as infant feeding counseling, should be provided to mothers of all HIV-exposed infants who undergo diagnostic testing for HIV infection. All infants identified as HIV-infected should be referred for specialized care, if available. Mothers of breastfeeding infants with negative HIV diagnostic test results should be counseled about the continued risk of MTCT of HIV through breast milk, and should be cautioned that a negative test result while the infant is breastfeeding does not guarantee that the infant will remain HIV-uninfected.

### 4.1 Maternal and Infant Evaluations through 3 months

All active participants will be consented for continued follow-up in this protocol. Non-laboratory and laboratory evaluations to be collected during pregnancy and through 3 months following birth are described in Table 1.

**Table 1: Non-laboratory and laboratory evaluations from pregnancy through 3 months following birth**

<table>
<thead>
<tr>
<th>Non-Laboratory Data</th>
<th>Labs</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Follow-up (26 – 30 Wks)</strong></td>
<td><strong>For all participants:</strong></td>
</tr>
<tr>
<td>• Interval Medical History, including history of receipt of antibiotics and other medications</td>
<td>• BV Status (Gram Stain, whiff test and pH)</td>
</tr>
<tr>
<td>• Assessment of adherence</td>
<td>• Wet mount for clue cells</td>
</tr>
<tr>
<td>• Dispensing of NVP</td>
<td>• Candida, Trichomonas (2 wet mount slides)</td>
</tr>
<tr>
<td>• Assessment of adverse experiences</td>
<td>• FFN Status (ELISA)</td>
</tr>
<tr>
<td><strong>Follow-up (36 Wks)</strong></td>
<td><strong>For HIV-infected cohort only:</strong></td>
</tr>
<tr>
<td>• Interval Medical/Obstetric History, including history of receipt of antibiotics and other medications</td>
<td>• Cervical swab for HIV viral load</td>
</tr>
<tr>
<td><strong>At Delivery</strong></td>
<td><strong>For HIV-infected cohort only:</strong></td>
</tr>
<tr>
<td>• Length of Labor</td>
<td>• Blood for CBC and CD4 Analysis</td>
</tr>
</tbody>
</table>

For all participants:
## Non-Laboratory Data

- Length of ROM
- Clinical Chorioamnionitis
- Obstetric Interventions
- Infant Weight/Gestational Age
- Antibiotic Usage
- Adverse experiences

## Labs

- Placenta, Membranes and Cord for Histology

### Between birth and 48 hours

<table>
<thead>
<tr>
<th>For infants born to HIV-infected cohort only:</th>
</tr>
</thead>
<tbody>
<tr>
<td>Blood draw for</td>
</tr>
<tr>
<td>DBS for HIV RNA PCR</td>
</tr>
<tr>
<td>CBC</td>
</tr>
</tbody>
</table>

### Follow-up (4-6 wks)

<table>
<thead>
<tr>
<th>For infants born to HIV-infected cohort only:</th>
</tr>
</thead>
<tbody>
<tr>
<td>Blood draw for</td>
</tr>
<tr>
<td>ALT</td>
</tr>
<tr>
<td>CBC</td>
</tr>
</tbody>
</table>

### Final Visit (3 months)

<table>
<thead>
<tr>
<th>For infants born to HIV-uninfected women or infants already confirmed as HIV-infected:</th>
</tr>
</thead>
<tbody>
<tr>
<td>No Blood Draw</td>
</tr>
</tbody>
</table>

### Post-test Counseling Visit (3 months + 4 to 6 weeks)

<table>
<thead>
<tr>
<th>For infants with negative HIV RNA PCR result at 3 months or infants confirmed as HIV-infected:</th>
</tr>
</thead>
<tbody>
<tr>
<td>No blood draw</td>
</tr>
</tbody>
</table>

### Post-test Counseling Visit (3 months + 10 to 12 weeks)

<table>
<thead>
<tr>
<th>For HIV-exposed infants with positive RNA PCR result at 3 months:</th>
</tr>
</thead>
<tbody>
<tr>
<td>Blood draw for</td>
</tr>
<tr>
<td>DBS for HIV RNA PCR</td>
</tr>
</tbody>
</table>

### For infants confirmed as HIV-infected:

<table>
<thead>
<tr>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>No blood draw</td>
</tr>
</tbody>
</table>
4.2 Maternal and Infant Final Visit(s): 3, 6 and 9 months

This section pertains to those mother/infant pairs who return for their scheduled follow-up visit at 3, 6, or 9 months following delivery. The majority of these participants will be terminated at this visit:

- all mothers (HIV-infected and HIV-uninfected),
- all infants with confirmed HIV infection (2 consecutive positive RNA PCR tests drawn on different days), and
- all infants born to HIV-uninfected women.

No laboratories evaluations will be collected from this group although some non-laboratory evaluations will be done.

The remaining active participants are the HIV-exposed infants of unknown or unconfirmed HIV status (no or 1 positive HIV RNA PCR). For these infants, a DBS will be collected at this final visit, but the mother/infant pair will be scheduled to return for post-test counseling 4 to 6 weeks later. If the DBS is negative, that infant will be discontinued (effective the date of the last blood draw). If the DBS is positive and is a confirmatory sample, that HIV-infected infant will be discontinued (effective the date of the last blood draw). If the DBS is positive and a confirmatory sample is necessary, a DBS should be drawn and the mother/infant pair scheduled for post-test counseling 4 to 6 weeks later. At that visit, if the DBS is positive, and it is the second consecutive positive result, that HIV-infected infant will be discontinued (effective the date of the last blood draw). See Table 2 for details.

Table 2: Non-laboratory and laboratory evaluations for final follow-up visit(s)
4.3 Maternal and Infant Final Visit(s): 12 months

This section pertains to those mother/infant pairs who return for their scheduled follow-up visit at 12 months following delivery. The majority of these participants will be terminated at this visit:

- all mothers (HIV-infected and HIV-uninfected),
- all infants with confirmed HIV infection (2 consecutive positive RNA PCR tests drawn on different days), and
- all infants born to HIV-uninfected women.

No laboratories evaluations will be collected from this group although some non-laboratory evaluations will be done including assessments of infant viability, maternal health and determination of feeding strategy.

The remaining active participants are the HIV-exposed infants of unknown or unconfirmed HIV infection status (no or 1 positive HIV RNA PCR). Because levels of maternal antibodies to HIV in an HIV-exposed child begin to disappear between 9 to 18 months of age, an antibody test is to be done to screen out those infants who have seroreverted. Blood should be obtained for both an antibody test and a DBS. Any infant who has a negative antibody test is considered to be HIV-uninfected at the time of the visit. For infants who have a positive antibody test, the DBS should be sent to UNC for an HIV RNA PCR assay. At the post-test counseling visit (1-2 weeks after the 12 month visit), follow-up may be discontinued for those infants with a negative ELISA (effective on the date when
Those infants who had a positive ELISA will have blood obtained for a second DBS, and a post-test counseling session will be scheduled in 8 weeks. By the time of that post-test counseling visit, the site should have results of both of the HIV RNA PCR assays that were obtained and will be able to counsel the infant’s mother about the status of the infant at the time of the final visits. If both tests were negative, the infant will be considered to be uninfected at the time of the final visit. If both tests are positive, the infant is confirmed to be HIV-infected. In both cases, follow-up should be discontinued for this infant (effective on the date when blood was last obtained) and referred for appropriate follow-up. If the results are equivocal, blood should be obtained for a final HIV RNA PCR assay, and a post-test counseling session should be scheduled.

Table 3: Non-laboratory and laboratory evaluations for final visit(s): 12 months

<table>
<thead>
<tr>
<th>Non-Laboratory Data</th>
<th>Labs</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Final Visit (12 months)</strong></td>
<td>For HIV-exposed infants of indeterminate status (1 or no positive HIV RNA):</td>
</tr>
<tr>
<td>• Infant Viability and Health</td>
<td>Blood draw for</td>
</tr>
<tr>
<td>• Infant feeding modality</td>
<td>• ELISA</td>
</tr>
<tr>
<td>• Schedule for post-test-counseling</td>
<td>• DBS for HIV RNA PCR</td>
</tr>
<tr>
<td><strong>Post-test Counseling Visit</strong></td>
<td>For infants with negative ELISA result:</td>
</tr>
<tr>
<td>(12 months + 1 to 2 weeks)</td>
<td>No blood draw</td>
</tr>
<tr>
<td>• Post-test counseling</td>
<td></td>
</tr>
<tr>
<td>• <strong>Discontinue follow up</strong></td>
<td></td>
</tr>
<tr>
<td>• Post-test counseling</td>
<td>For HIV-exposed infants with positive ELISA:</td>
</tr>
<tr>
<td>• Schedule for post-test counseling</td>
<td>Blood draw for</td>
</tr>
<tr>
<td></td>
<td>• DBS for HIV RNA PCR</td>
</tr>
</tbody>
</table>

| **Post-test Counseling Visit** | NO BLOOD DRAW FOR: |
| (12 months + 9 to 10 weeks) | Infants with 2 negative RNA PCRs |
| • Post-test counseling | Infants with 2 positive RNA PCRs |
| • **Discontinue follow up if HIV infection status of the infant has been determined** | For infants with equivocal final RNA results: |
| • Schedule for post-test counseling if test results are equivocal | • DBS for HIV RNA PCR |

The following sections -- 4.4 through 4.7 -- are essentially the instructions for evaluations in the protocol as originally designed. There have been minor modifications made to section 4.4 (“Pre-entry Evaluations”) to clarify that women of known HIV status would be exempt from the 024 screening process; to section 4.5 (“Evaluations During Pregnancy and Labor”) to clarify methods for determining gestational age; and to the title of section 4.5.1 (“Adherence to Study Antibiotics/Placebos”) to clarify that, because this is a blinded study, strategies for determining
adherence applied to both the study antibiotics and to the placebos. All of these changes are highlighted.

4.4 Pre-entry Evaluations

Appropriate pretest HIV counseling by trained HIV counselors will be provided to all women attending the antenatal clinic on the day of screening. Women who had not already undergone VCT through another mechanism and who agree to be HIV tested will sign a screening informed consent form. Screening for syphilis is part of routine antenatal care in this study, and will be performed at the Screening or Enrollment visit, according to site-specific procedures. For those tested, HIV and syphilis test results will be available within one week from the initial antenatal visit, and in general, women will be seen for a repeat counseling visit within two weeks of testing. All women who underwent pre-test counseling and testing will receive post-test counseling upon return to the clinic. HIV-positive women and a sample of HIV-negative women will be requested to participate in the study; an enrollment visit will be scheduled at 20-24 weeks gestation and women will be encouraged to bring the baby’s father.

HIV testing will be performed according to site-specific procedures. All positive test results will be confirmed with an additional test.

Syphilis testing: All women will have blood tested using RPR for screening and TPHA for confirmation. Women with a reactive syphilis test will receive treatment at no cost. Testing will be performed at the Screening or Enrollment visit, according to site-specific procedures.

4.5 Evaluations During Pregnancy and Labor

At the enrollment visit the study procedures will be explained to the women and their babies’ fathers, including the schedule of the follow-up visits. Women who agree to participate in the trial will sign a separate enrollment consent form; signature of the baby’s father will also be requested. HIV-infected women who enroll in the study will be offered the HIVNET 012 Nevirapine regimen. Women who choose to accept the Nevirapine will sign a separate signature line on the study enrollment consent; signature of the baby’s father will also be requested.

Structured questionnaires will be used to collect the non-laboratory data described in the table below (Table 4). In addition, at the first antenatal visit (20-24 weeks) socio-demographic information including age, parity, maternal and paternal education, residence, socioeconomic status, marital status, and number of sexual partners will be collected on all women. The enrollment questionnaire will include symptoms of clinical AIDS (for HIV-positive women) and sexually transmitted diseases. A clinical examination form will also be completed. Table 1 summarizes these activities.

Gestational age will be determined based on patient recall of the last menstrual period, corroborated with uterine size. If uterine size is not consistent (within 2
weeks) with recall, uterine size will define the obstetric gestational age. Ultrasound examinations are rarely performed at these sites; therefore estimating gestational age by this criterion will not be routinely available. However, in the event that an ultrasound examination is performed, the gestational age from that assessment should be used. In addition, at delivery, the babies will have a modified Ballard examination performed to allow for both an obstetric and pediatric estimation of gestational age. The Ballard must be done within 48 hours of delivery to be considered valid. If the two are consistent within 2 weeks, the obstetric gestational age will be used; if not, the final study gestational age will be based on the Ballard examination. Staff at both locations will be trained to determine both the obstetric and pediatric gestational ages. However, this algorithm will result in a single project gestational age which will be used whenever a project gestational age is required as part of the protocol or as an outcome.

Table 4. Non-laboratory and laboratory evaluations during pregnancy and labor.

<table>
<thead>
<tr>
<th>Non-Laboratory Data</th>
<th>Labs</th>
</tr>
</thead>
<tbody>
<tr>
<td>Enrollment (20-24 wks):</td>
<td>Syphilis RPR (if not performed at Screening)</td>
</tr>
<tr>
<td>Maternal Demographics</td>
<td>BV Status (Gram Stain, whiff test and pH)</td>
</tr>
<tr>
<td>Obstetric History</td>
<td>Wet mount for clue cells</td>
</tr>
<tr>
<td>Medical History</td>
<td>Candida, Trichomonas (2 wet mount slides)</td>
</tr>
<tr>
<td>Sexual History</td>
<td>FFN Status (EIA) *</td>
</tr>
<tr>
<td>Gestational Age</td>
<td>Blood for CBC and CD4 Analysis</td>
</tr>
<tr>
<td>Antibiotic Usage</td>
<td>Plasma Viral Load*</td>
</tr>
<tr>
<td>Other Medications</td>
<td>Cervical swab for HIV Load*</td>
</tr>
<tr>
<td></td>
<td>Cervical swab for Gonorrhea Culture and Chlamydia</td>
</tr>
<tr>
<td>Follow-up (26 – 30 Wks)</td>
<td>BV Status (Gram Stain, whiff test and pH)</td>
</tr>
<tr>
<td>Intervening Medical History</td>
<td>Wet mount for clue cells</td>
</tr>
<tr>
<td>Antibiotic Usage</td>
<td>Candida, Trichomonas (2 wet mount slides)</td>
</tr>
<tr>
<td>Other Medications</td>
<td>FFN Status (EIA) *</td>
</tr>
<tr>
<td>Pill Counts/Adherence/Adverse experiences</td>
<td>Cervical swab for HIV Load*</td>
</tr>
<tr>
<td></td>
<td></td>
</tr>
<tr>
<td>Follow-up (36 Wks)</td>
<td>Blood for CBC and CD4 Analysis</td>
</tr>
<tr>
<td>Intervening Medical/OB History</td>
<td></td>
</tr>
<tr>
<td>Antibiotic Usage</td>
<td></td>
</tr>
<tr>
<td>Other Medications</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
</tr>
<tr>
<td>At Delivery</td>
<td>Placenta, Membranes and Cord for Histology</td>
</tr>
<tr>
<td>Length of Labor</td>
<td></td>
</tr>
<tr>
<td>Length of ROM</td>
<td></td>
</tr>
<tr>
<td>Clinical Chorioamnionitis</td>
<td></td>
</tr>
<tr>
<td>Obstetric Interventions</td>
<td></td>
</tr>
<tr>
<td>Infant Weight/Gestational Age</td>
<td></td>
</tr>
</tbody>
</table>
### 4.5.1 Adherence to Study Antibiotics/Placebos

The prenatal antibiotics, metronidazole and erythromycin, and the peripartum antibiotics, metronidazole and ampicillin, and placebos will be packaged in blister packs to be given to each woman at appropriate visits. The initial dose of prenatal antibiotics or placebo will be taken at the enrollment visit under direct observation, and the participant will be asked to bring the blister packs with her to her next visit at 26-30 weeks gestation. Each woman will be given the peripartum antibiotics or placebo at the 26-30 week visit. She will take the antibiotics or placebo at onset of labor and throughout delivery and after delivery until the course is completed. If a woman vomits within one hour of dosing, she will be redosed. She will be asked to bring the blister packs with her to her next visit at 4-6 weeks post-partum. Adherence will be assessed by the number of pills missing from the blister packs at the follow-up visits. If the blister packs are not returned, reported use will be noted.

Erythromycin use is associated with various types of gastrointestinal complaints (but very few serious complications) and continued use is less than optimal for this reason. In an effort to enhance appropriate use of the drug, at randomization, each woman will be informed of the potential for mild but harmless gastrointestinal side-effects and will be encouraged to continue using the medications. If these side effects become intolerable, rather than discontinuing, the woman will be encouraged to reduce the dose, but take the medication over a longer time-period. The amount of each drug taken by the participants will be monitored.

Rarely, patients taking metronidazole and/or erythromycin have been reported to develop a peripheral neuropathy, seizures, or other CNS symptoms. For this reason, patients with central nervous system diseases, including seizures, will be excluded from participation, and any patient developing a neuropathy or other CNS sign or seizures during treatment will have the erythromycin and metronidazole or their placebos discontinued.

Since metronidazole may potentiate the action of certain anticoagulants, women taking any anticoagulant drug will also be excluded from participation.

### 4.5.2 Other STDs
Except for the intervention of study antibiotics (or placebo) and the addition of a vitamin/mineral supplement and the HIVNET 012 NVP regimen (to HIV-infected women and their infants), the women will receive regular prenatal care. At the study sites, women are generally screened for syphilis, and at times for gonorrhea. If reactive for syphilis or positive for gonorrhea, women are generally treated with a penicillin-like drug. Treatment of syphilis or gonorrhea with a penicillin-like drug should have little influence on the development of chorioamnionitis since the organisms involved, including *Gardnerella, Bacteroides, Mycoplasma and Ureaplasma*, are not generally susceptible to the penicillin-like antibiotics. For this reason, women who receive penicillin-like antibiotics or were treated for syphilis or gonorrhea within two weeks prior to randomization will not be excluded. Bacterial vaginosis is not usually screened for or treated in these settings, but if a woman complains of a symptomatic discharge, she will be appropriately treated. *Candida, trichomonas and chlamydia* are not routinely screened for, but occasionally are treated if symptomatic. In any case, for each woman, at the time of enrollment, during chart review and again at delivery, the study physicians will discern which antibiotics, if any, were used during the course of pregnancy and these will be recorded. It is important to nest this clinical trial within the usual health care environment to assess its utility within the actual local medical practice. Other than the improved screening and treatment for syphilis and provision of multivitamin/mineral supplements and Nevirapine, the usual standard of care at the study sites will be provided.

4.5.3 Evaluation of clinical and laboratory events

This study will investigate if BV and FFN are predictors of perinatal HIV transmission, and also if treatment of HIV-positive women with or without BV and/or FFN with the proposed antibiotic regimen results in a reduction in perinatal HIV transmission. Diagnosis of BV (or FFN positivity) will not influence the conduct of the treatment in this study. The diagnosis of BV will be determined based on a Gram stain (Nugent criteria) of vaginal fluid taken at the randomization visit. Since BV will not be used to direct therapy, the slides for Gram stain for BV will be evaluated in batch independent in time from the conduct of the trial. FFN testing is being done to answer a question related to the mechanism of HIV MCT; if MCT is more common in FFN-positive women, this would strongly suggest that chronic chorioamnionitis plays a significant role in this process.

The Gram staining method to diagnose BV requires expertise and training. Therefore, simple inexpensive methods to assess BV are desirable. Clinical criteria to diagnose BV have been developed (three of the following four criteria: vaginal pH >4.5, homogenous vaginal discharge, presence of ≥20% clue cells, and a positive amine test), and are highly correlated with the Gram staining scoring method. In a recent study in
Malawi, BV based on these clinical criteria was shown to be associated with HIV perinatal transmission\textsuperscript{15} and increased acquisition of HIV.\textsuperscript{14}

Vaginal pH will be measured using pH paper on a vaginal swab obtained from lateral and posterior fornicees. An amine (or whiff) test will be performed by mixing a few drops of 10\% potassium hydroxide with vaginal fluid. These tests are simple and can be routinely performed in the hospital laboratories at the study sites. The presence of clue cells will be evaluated by mixing vaginal fluid with a drop of normal saline on a slide and examining under high power magnification. The presence of vaginal discharge will be determined by a speculum-aided pelvic exam conducted by trained nurses. The implications for low cost screening to identify candidate women for therapy in areas of the world with less resources are obvious; hence this nested evaluation of the sensitivity, specificity and predictive value of clinical criteria to diagnose BV is an important subproject within this clinical trial.

FFN is measured using an EIA assay from vaginal fluid collected using a Dacron swab, which is placed in buffer and frozen for subsequent analysis. Adeza Biochemical, Sunnyvale, CA, has provided the investigators the EIA kits necessary to determine levels of fetal fibronectin and will make kits available for this study. If FFN identifies a group at substantially increased risk for HIV MCT, this will enhance knowledge about the mechanism of MCT and it may allow future investigators the opportunity to target transmission in women at high risk for MCT.

At enrollment and approximately 4 weeks later, specimens will be collected to diagnose BV, and two wet mount slides will be prepared and examined for detection of candida and trichomonas. A swab in buffer for FFN and a cervical swab for GC and vaginal viral load evaluations will be collected. At enrollment, a cervical swab will also be collected for chlamydia EIA. Analysis of the Gram stains will be conducted for all women. FFN samples will be stored to be run in batch when feasible.

Blood samples will be collected for CBC from each woman at enrollment and the 36-week visit. The blood for HIV-infected women only will be stored for later viral load analysis at enrollment and tested for CD4 counts at both time points.

### 4.6 Evaluations at Delivery

At delivery, clinical chorioamnionitis will be defined by the presence of a temperature $>$38°C during labor in association with any of the following: 1) fetal tachycardia $>$160 BPM, 2) significant abdominal tenderness, 3) a purulent cervical discharge, or 4) a foul smelling infant or placenta.

After delivery, the placenta will be brought to the laboratory in a sealed plastic bucket containing 500 ml 10\% formol saline and weighed. The membranes will be
rolled in a standard fashion, and after fixation, two sections at least 2 cm apart will be evaluated. Two sections of the cord, one close to the placenta and the other near the fetus will be obtained, as will two representative sections of the placenta. The specimens will be fixed in 10% neutral buffered formalin, processed for routine paraffin embedding, embedded, cut at 5 µm micron thickness and stained by routine hematoxylin-easin method.

Slides will be examined by the designated pathologist without knowledge of outcome or maternal HIV status. Histologic chorioamnionitis will be based on qualitative assessment of the membranes, cord and placenta basalis, looking for mononuclear and polymorphonuclear white cells using a standard protocol. A grading system which divides the histologic chorioamnion evaluation into normal and mild, moderate and severe chorioamnionitis will be used. It will be noted if the predominant infiltrate is mononuclear or polymorphonuclear so that the chorioamnionitis may be divided into chronic or acute. The presence of findings suggestive of placental malarial infiltration will also be noted.

Following delivery, mothers and infants will be evaluated for evidence of disease and adverse experiences to medications. Placental tissue will be stored so that examination of the placenta by in situ and immunohistochemistry using a novel and highly sensitive method utilizing tyramide-fluorophors may be conducted. This method can simultaneously detect HIV RNA, HIV DNA, HIV proteins and cellular proteins and help delineate the role of chorioamnionitis and HIV expression. A section of the placenta will also be kept to examine for malaria parasites and histologic changes associated with malaria. Malaria is endemic in the study sites and is a known risk factor for preterm birth and low birthweight.

As described in the protocol, the investigators plan to obtain biologic specimens that have the potential to answer questions related to mechanisms of transmission, but are not crucial to answer the primary question about efficacy of antibiotics for reducing HIV MCT. These include the testing for BV; FFN; viral load in blood and vaginal fluids; and evidence of malaria in the placenta. If, during the initial enrollment period, it appears that obtaining any of these specimens will reduce the overall ability to carry out this study, the collection of these specimens will be deleted from the protocol.

4.7 Post-Delivery Evaluations (Table 5)
At each postnatal visit infants will have a blood draw by heelstick or venous puncture according to site-specific procedures. Within 48 hours and at 4-6 weeks, the specimens from infants born to HIV-infected women will be tested for HIV RNA using PCR. Positive specimens will be confirmed by a second RNA PCR on a separate specimen no later than the next scheduled visit and sooner if possible. Infants who test positive on initial PCR testing but die or are lost to follow-up prior to a confirmation test on a separate specimen will be considered HIV-infected in the study analysis. Infants who are positive using the definition of an infected infant (section 7.51) will have reached the primary endpoint. Because a
second primary endpoint includes being alive and free of disease at 1 year of age, infants of HIV-positive women who were not previously classified as HIV-infected will be screened for HIV by ELISA at 12 months of age. All who are non-reactive will be considered free of HIV. Any reactive tests will be confirmed with an HIV RNA PCR. Infants who test positive at 12 months will have another PCR for HIV RNA performed and those positive using definitions in section 7.51 will be considered positive for this endpoint. If the recommendations are still current at the time study infants are born, each infant diagnosed as HIV-positive will be provided with prophylactic Bactrim or an equivalent medication during the course of the study. All infants will have a CBC performed on the blood specimens collected within 48 hours, 4-6 weeks, 3 months, and 12 months. Specimens from infants born to HIV-uninfected women will be stored for potential future approved studies.

To monitor the safety of Nevirapine, infants who receive Nevirapine will have an ALT at the 4-6 week visit. If the results from the ALT are a grade 1 or 2 abnormality, the test will be repeated at the 3-month visit. If the results are a grade 3 or 4 abnormality, the mother will be requested to return with her baby within one week of the test result for a repeat test. The ALT will be performed at the on-site laboratory.

Following delivery (within 48 hours), at 4-6 weeks, 3 months, 6 months, 9 months and one year of age, questions related to infant health, maternal health and breastfeeding practices will be asked. At each post-delivery visit, a clinical assessment form will be completed for the mother and the newborn. These activities are summarized in Table 5.

Because there is concern that some antibiotic treatment may lead to an increase in neonatal sepsis with more virulent organisms, occurrence of neonatal sepsis prior to discharge from the hospital or at the 4-6 weeks evaluation will be monitored. The NIAID Vaccine and Prevention DSMB will evaluate mortality and severe morbidity in the treatment and placebo groups at 3-month intervals to ensure that the treatment is not causing harm.

Breastfeeding is nearly universal in all sites. The investigators are aware of the risk associated with HIV transmission to the baby if breastfed. Currently, breastfeeding is encouraged in the host countries during the entire infancy since alternative safe measures are not generally available. However, women will be adequately counseled and the risks and benefits of breastfeeding will be explained thoroughly at each visit to help women make an informed decision. During the entire study period the research team will be guided by the advice and recommendations adopted in-country.

It is understood that some infants will likely become HIV infected due to breastfeeding prior to evaluation at 4-6 weeks. There is no method available to
allow distinction between infection occurring during the perinatal/peripartum period and secondary infection due to early breastfeeding.

Table 5. Non-laboratory and laboratory evaluations during follow-up.

<table>
<thead>
<tr>
<th>Non-Laboratory Data</th>
<th>Labs</th>
</tr>
</thead>
<tbody>
<tr>
<td>Within 48 hours; 4 – 6 wks; 3, 6, and 9 months; 1 year</td>
<td>Within 48 hours, 4-6 weeks</td>
</tr>
<tr>
<td>Infant Viability and Health, Maternal Health Breast vs. Formula Feeding</td>
<td>Blood draw for Infant RNA PCR for HIV&lt;sup&gt;A&lt;/sup&gt;</td>
</tr>
<tr>
<td></td>
<td>Within 48 hours, 4-6 weeks, 3, 12 months</td>
</tr>
<tr>
<td></td>
<td>Blood draw for CBC</td>
</tr>
<tr>
<td></td>
<td>4-6 weeks</td>
</tr>
<tr>
<td></td>
<td>Blood draw for ALT&lt;sup&gt;*&lt;/sup&gt;</td>
</tr>
<tr>
<td></td>
<td>3, 6, 9 months</td>
</tr>
<tr>
<td></td>
<td>Blood draw for storage</td>
</tr>
<tr>
<td></td>
<td>12 months</td>
</tr>
<tr>
<td></td>
<td>Blood draw for Infant ELISA and DBS. If the ELISA is reactive, the DBS sample will be tested for HIV RNA PCR.</td>
</tr>
</tbody>
</table>

<sup>A</sup> To be run on infants of HIV-positive women. Positive specimens will be confirmed by a second RNA PCR on a separate specimen no later than the next scheduled visit and sooner if possible.

<sup>*</sup>In infants receiving Nevirapine. If results indicate a grade 1 or 2 abnormality, the test will be repeated at the 3 month visit. If the results indicate a grade 3 or 4 abnormality, the test will be repeated within one week of the test result.

5.0 DATA COLLECTION AND MONITORING AND ADVERSE EXPERIENCE REPORTING

5.1 Records to Be Kept
Case Report Forms (CRF) will be provided for each subject. Subjects will not be identified by name on any study documents. Subjects will be identified by an identification number unique for each participant; this will be provided by the HIVNET Statistical Center upon randomization. At screening, all women will be provided a temporary screening ID to track clinical and laboratory results. The use of the screening ID will be discontinued upon randomization.

All data on the CRF must be legibly recorded in black ink. A correction should be made by striking through the incorrect entry with a single line and entering the correct information adjacent to it. The correction must be initialed and dated by the investigator or a designated, qualified individual.

5.2 Data Management and Documentation Guidelines
Instructions concerning the recording of study data on case report forms, or the entry of data in the computerized database, will be provided by the HIVNET Statistical Center, who will coordinate data management for the trial.

5.3 Regional Monitoring
5.3.1 Site visits by FHI study monitors will be made in accordance with HIVNET policy to monitor the quality of data collected in the research
records, the accuracy of the data entered in the database, and compliance with all regulatory requirements associated with clinical trials.

5.3.2 Site visits will be made at the study sites at regular intervals.

5.3.3 The investigators will make study documents (e.g., consent forms, drug distribution forms, case report forms) and pertinent hospital or clinic records readily available for inspection by the FHI site monitor, the FDA, Boehringer-Ingelheim staff and pertinent NIH staff for confirmation of the study data.

5.4 Adverse Experience Reporting
An adverse event is defined as any health related reaction, effect, toxicity or abnormal laboratory result that a patient experiences during the course of a study irrespective of relationship to study treatment.

Serious adverse experiences should be reported according to the procedures described in the Study Specific Manual of Operations. Events that occur after exposure to the investigational product (Nevirapine) and that meet the criteria described in the DAIDS SAE Reporting Manual should be reported to DAIDS (ROC) and to the Statistical Center. Events related to the other study products should be reported to the Statistical Center only.

6.0 STUDY TREATMENT

6.1 Drug Regimens, Administration and Duration

As per the recommendations of the DSMB following the interim analysis, no study antibiotics/placebos are to be dispensed. Any study antibiotics/placebos that have been dispensed are to be retrieved. The remainder of this section (6.1.1) describes the original design of the protocol with respect to the study antibiotics/placebos.

6.1.1 Antibiotics/placebos

At 20–24 weeks, the women randomized to the treatment arm will receive metronidazole 250 mg TID and erythromycin 250 mg TID for 7 days. Women randomized to the control group will receive identically appearing placebos.

With the onset of labor and/or rupture of membranes, rapid antibiotic entrance into the amniotic fluid is important. Participants will be given metronidazole 250 mg and ampicillin 500 mg (or placebo) at the 26-30 week visit and will be instructed to take one of each pill at onset of labor. They will be asked to come to the hospital immediately after onset of labor, at which time the study staff will assume administration of the study antibiotics. These drugs were chosen to prevent the acute chorioamnionitis associated with membrane rupture. Both of these drugs
achieve satisfactory serum and amniotic fluid levels within 1-2 hours after oral administration. Since the secondary outcomes include maternal infection and mortality, each woman will be asked to continue using the medications after delivery, three times per day, until the course is completed. Women randomized to the control group will receive identically appearing placebos.

6.1.2 Multivitamins
All women in both treatment and control arms will receive a standard vitamin/mineral preparation daily from enrollment in the study until delivery. This preparation will include 30 mg iron, 400 mcg folic acid, 5000 IU Vitamin A, 400 IU Vitamin D, 30 IU Vitamin E, 50 mg Vitamin C, 2 mg Vitamin B₁, 3 mg Vitamin B₂, 3 mg Vitamin B₆, 5 mcg Vitamin B₁₂, 20 mg Niacin, 250 mg Calcium, 150 mcg Iodine, 100 mg Magnesium, and 15 mg Zinc.

6.1.3 Nevirapine
All HIV-infected women and their neonates will be offered the HIVNET 012 Nevirapine (NVP) regimen. If the mother accepts the Nevirapine for her and her baby, she will be given one oral dose of 200 mg NVP at the 26-30 week antenatal visit to be taken at onset of labor. If the mother does not deliver within 48 hours of taking the Nevirapine dose, she will be redosed immediately, or at onset of active labor in the case of false labor.

If the mother delivers her baby more than one hour and less than 48 hours after taking Nevirapine, her baby will receive one oral dose of 2mg/kg NVP suspension at 72 hours post-birth or discharge, whichever occurs earlier. If the mother delivers her baby less than one hour or more than 48 hours after taking Nevirapine, the infant will be dosed as soon as possible. If the infant vomits within one hour of receiving Nevirapine, the infant will be redosed once.

If the mother delivers somewhere other than the study hospital, she will be asked to return to the study clinic within seven days of delivery to allow infant dosing of Nevirapine. Infants who are delivered elsewhere and do not present at the clinic within seven days will not be dosed.

All HIV-uninfected women and HIV-infected women at sites enrolling HIV-uninfected women who refuse Nevirapine will be given a matched placebo at the 26-30 week visit to maintain participant confidentiality. Sites enrolling only HIV-infected women will not dispense Nevirapine placebo tablets to women who refuse Nevirapine.

6.2 Drug Acquisition
The active antibiotics were acquired through the Division of AIDS contract with SRI. The same contractor will make the placebos.

The multivitamins were manufactured by Tishcon Corporation.
The Nevirapine and Nevirapine placebo will be donated by Boehringer Ingelheim Pharmaceuticals.

6.3 **Drug Supply, Distribution and Pharmacy**

PAB will coordinate the packaging and shipping of the product through existing subcontracts with the Clinical Research Product Management Center (CRPMC). The antibiotics/placebos will be packaged in blister packs. The multivitamins will be packaged in bottles of 100. The Nevirapine and Nevirapine placebo will be packaged in bulk for distribution by the study pharmacist.

The protocol pharmacist is required to maintain complete records of all study agents received from the CRPMC of the Division of AIDS and subsequently dispensed.

6.4 **Concomitant Medications**

Any concomitant medication, if deemed medically necessary, will be permitted for either the mother or neonate while on-study; the administration of concomitant medication will be recorded on CRFs. Because of the reported interaction between metronidazole and alcohol, all women in the study will be asked to refrain from alcohol use during the trial, as reflected in the informed consent form.

6.5 **Toxicity Management**

Allergic reactions are possible with any drug. Severe anaphylactic types of reactions have been reported with penicillin use (1/50,000). Gastrointestinal side effects including nausea and vomiting and diarrhea have been reported with erythromycin use. If the toxicity appears to be a significant allergic reaction, both antibiotics will be stopped immediately. However, if the problem appears to be mild gastrointestinal discomfort associated with erythromycin use, the metronidazole will be continued at full dose. If necessary, the daily erythromycin dose may be reduced, but will not be discontinued unless the woman clearly cannot tolerate the medication. If there is nausea and vomiting or diarrhea, the women will be asked to decrease only the erythromycin, first to 2 tablets per day, and then, if necessary, to one per day until the symptoms abate. If a participant experiences wheezing or rash compatible with study drug reaction while on study drugs, the study drugs will be stopped and the participant will be asked to report to the study clinic immediately.

Extended Nevirapine use has been associated with the occurrence of skin rashes. Nevirapine-related rashes are not expected in this protocol due to the abbreviated course of Nevirapine that will be administered. This protocol will employ the short-course Nevirapine regimen used in HIVNET 012, in which the occurrence of skin rashes was the same in the Nevirapine and zidovudine arms. Despite the fact that Nevirapine-related rashes are not expected, occurrence of skin rashes temporally related to Nevirapine use will be recorded and reported as adverse experiences. The severity of skin rashes will be determined based on the
Supplemental Toxicity Table for Grading Severity of Cutaneous/Skin Rash/Dermatitis Adverse Experiences located in Appendix V. Mothers will be instructed to return to the clinic if they or their baby experiences a rash. In addition to monitoring for skin rash, all infants who receive Nevirapine will have an ALT at the 4-6 week visit. If the results are a grade 1 or 2 abnormality, the test will be repeated at the 3 month visit. If the results are a grade 3 or 4 abnormality, the mother will be requested to return with her baby within one week of the test result for a repeat test.

All mothers will agree to deliver at the study hospital when they enroll in the trial. Despite this fact, some mothers may deliver elsewhere. Mothers will be requested to report to the clinic with their infants within 7 days after delivery if they do not deliver at the clinic. This will aid in toxicity management if any side effects are observed, as well as allow for infant dosing of Nevirapine. As in the HIVNET 012 protocol, infants presented at the clinic within 7 days after birth will be given the infant Nevirapine dose; if the mother presents with the infant after 7 days, no infant dose of Nevirapine will be administered.

6.6 Criteria for Treatment Discontinuation
Study participants may be discontinued from treatment for the following reasons:
- The subject refuses further treatment and/or follow-up evaluations.
- The investigator determines that further participation would be detrimental to the subject's health or well-being.
- The subject experiences a severe allergic reaction to the study drugs.

6.7 Other Infections
A number of questions have arisen about the use of other treatments, especially the use of antibacterial agents that may obscure the results of this study. No appropriate treatment will be withheld because a woman is enrolled in this study. Therefore, if a woman has a symptomatic vaginal infection that would normally be treated at the study site, appropriate treatment will be given. If it is standard treatment to use antibiotics in the face of ruptured membranes for more than 24 hours, or with maternal fever, they will be utilized. If newborns are normally treated with antibiotics following maternal chorioamnionitis they will be so treated. The individual treatments received by study subjects and the prevailing clinical practices that may influence MCT will be documented, but no attempt will be made to restrict what is considered routine medical care.

If during study drug administration a woman has a condition requiring antibiotic use, such as a urinary tract infection, chorioamnionitis or ruptured membranes of more than 24 hours, the woman will be treated with routine antibiotics in addition to the study drugs. Because the active arm of the study uses a relatively low dose of antibiotics, additional use of antibiotics should not place the woman at increased risk. The alternative option, unblinding the woman and providing antibiotics if she is on the placebo, would result in significant risk to bias the
study. However, if the physician, because of severe allergic reaction or any other cause, feels it is necessary or in the patient’s best interest to unblind the study products, the nature of the study drugs will be made available to the attending physician. The list will be kept by the in-country pharmacist, who will be the only person able to break the code.

7.0 STATISTICAL CONSIDERATIONS

7.1 General Design Issues
This is a randomized, double blinded, controlled Phase III clinical trial of antibiotics to reduce chorioamnionitis-associated perinatal HIV transmission. The interventions will be administered to pregnant women during the second trimester of pregnancy and at onset of labor in order to interrupt vertical transmission of HIV.

7.2 Endpoints

7.2.1 Primary endpoints
- Infant HIV infection as determined by a positive RNA PCR on dried blood spots on filter paper taken at 4-6 weeks\(^{33}\)
- Composite of infant HIV infection and mortality at one year of age

7.2.2 Secondary endpoints
- Infant HIV infection at 24-48 hours and 12 months
- Rates of clinical and histologic chorioamnionitis
- Rates of neonatal/infant morbidity and mortality
- Rates of maternal morbidity and mortality
- Safety/tolerance of antibiotic treatment given to pregnant women

7.2.3 Definition of gestational age is found in section 4.2. Preterm birth will be defined as <37 weeks, but evaluation of preterm birth at earlier gestational ages defined as <35 and <32 weeks will also be done. The definition of histologic chorioamnionitis is found in section 4.3. Infant mortality will be divided into neonatal, defined as <28 days, and post-neonatal, defined as 28 to 365 days. Evaluation of neonatal morbidity will include suspected or documented sepsis (based on physician examination or positive culture), pneumonia (based on physician examination or x-ray), and suspected or documented respiratory distress syndrome (based on examination or x-ray). The length of stay on initial hospitalization will be documented. Seizures and the presence of signs and symptoms of meningitis will be noted. Post-discharge medical problems such as seizures, pneumonia, meningitis, severe diarrhea, and all hospital readmissions and diagnoses will be noted. At each visit the infant will be weighed and the height and head circumference measured so that deviations from normal growth (<10\(^{th}\) percentile for an African standard)
may be determined. Medications used, such as any antibiotic treatment will be recorded.

Data on the mothers’ health during the initial hospitalization and following discharge will also be collected. Length of hospital stay and readmissions will be noted as will the use of any antibiotics for treatment of infections. Maternal diagnoses including chorioamnionitis, post-partum endometritis, wound infections, urinary tract infections, pneumonias, etc., will be recorded. Any adverse experiences to the medications as well as the adherence to the medications provided will also be tracked.

7.3 Randomization and Blinding Procedures
Participants will be randomized at 20-24 weeks gestation in a double-blinded fashion to receive either the active agent (metronidazole/erythromycin/ampicillin) or matched placebo. The randomization will be designed by the HIVNET Statistical Center and employ permuted block algorithms with varying block size, blocked within study site to ensure that balance between assignments is maintained within each study site. Study drug will be packaged according to the randomization and sent to the study site. Randomization will be performed on site by assigning study drug to participants in sequential order. These procedures will be coordinated with the HIVNET Statistical Center and detailed in the study specific Manual of Operations. HIV-positive and HIV-negative subjects will be randomized in separate strata.

7.4 Sample Size and Accrual
The rate of HIV-1 transmission at 4-6 weeks is the primary endpoint for this study. The control arm 4-6 weeks transmission probability is estimated to be 11%. This estimate is based on the recent HIVNET 012 results using the Nevirapine regimen involving a single dose to the mother at onset of labor and the single dose to the infant soon after delivery. For a one-sided 0.025-level chi-square statistic to provide 90% power to detect a reduction in vertical transmission from 11% to 7.5%, 2870 mother/infant pairs would be needed in the trial. However, allowing for up to 3% loss-to-follow-up rates during pregnancy, and 5% missing information post-delivery, the majority of this due to early mortality, the trial will require 3120 mother/infant pairs. The investigators will work very hard to achieve the proposed 3% loss to follow-up. However, if that rate is not achieved, sample sizes will be recalculated so that they reflect the actual lost-to-follow-up rates. A 3% lost-to-follow-up rate was achieved in Malawi in the micronutrient study (Taha, personal communication).

7.5 Monitoring and Analysis
An intent-to-treat analysis will be conducted using the entire sample from all sites. The incidence of HIV-1 transmission and mortality rates will be determined among infants in each study arm. Binary endpoints will be compared between treatment arms using contingency tables or, if time dependent (death/drop-out), Kaplan Meier curves. Adjustment for potential confounders will be done through logistic regression
or proportional hazards models. Single continuous outcomes (viral load, CD4 counts, etc.) will be compared with t-tests or non-parametric analysis. If needed, variable transformations such as logarithms will be used. Adjustment for confounders will be made with linear models. Repeated measures such as symptoms and anthropometric indices will be evaluated using generalized estimation equations.

An initial safety analysis will be performed after the first three months of accrual. Additional analyses of safety and formal interim analyses of efficacy will then be performed at approximately six-month intervals during the remainder of the projected 18-24 month study duration. Recommendations for early termination of positive or negative results will be guided by the symmetric group sequential O’Brien-Fleming Boundary. The O’Brien-Fleming design allows for early termination if extreme initial results are seen, while essentially allowing employment of the standard single stage one-sided 0.025-level test statistic at the final analysis if it is reached. This enables one to maintain the power of the single-stage design in the presence of interim monitoring without having to increase the maximum sample size. The Lan-Demets implementation of the O’Brien-Fleming use function will be employed to define proper significance levels at the time of each formal interim analysis by the NIAID Vaccine and Prevention DSMB.

Recommendations regarding trial continuation and modification of study conduct will be based on safety as well as efficacy considerations. In the assessment of safety measures, particular attention will be given to monitoring the incidence of preterm birth, motivated in part by recent data showing a relationship between asymptomatic trichomonas, treatment with metronidazole, and increased risk for preterm birth.\(^{11}\)

Timing of transmission will be estimated by assuming that if the infant HIV PCR test was positive within 48 hours of birth, that transmission likely occurred prenatally, while if it was negative within 48 hours of birth but positive at 4-6 weeks, that MCT occurred in the intrapartum period. The investigators are aware that some of the infants who turn HIV-positive between discharge and 4-6 weeks will do so because of breast feeding, but at this time have no way to determine the proportion of the converters due to breast feeding vs. perinatal transmission.

In addition to viral load and CD4 counts, several variables will be evaluated for their relationship to MCT (and child health) in each group including BV and FFN status, the presence of STDs, histologic chorioamnionitis, length of PROM and labor, obstetric interventions, and gestational age and/or preterm birth.

7.5.1 Definition of HIV Infected/Uninfected Infant for HIVNET Perinatal Protocols

Following is the standard HIVNET definition adopted by the HIVNET Perinatal Working Group; if this definition changes before implementation of this protocol, adoption of these alterations will be considered so that a
standard definition is used for all HIVNET perinatal transmission protocols:

An infant less than 15 months of age will be considered to be infected with HIV if two separate peripheral blood specimens from different days are drawn and each specimen is positive by at least one of the following assays: HIV-1 culture, HIV-1 DNA PCR, HIV RNA RT-PCR. At least one of these tests will be done in a laboratory that is approved to perform the assay for protocol testing. A positive result will be confirmed no later than the next scheduled visit and sooner if possible.

The Organon-Teknika NucliSens assay will be used for PCR on dried blood spots from the Malawi and Zambia sites. The Roche 1.5 assay will be used on the samples from the Tanzanian site.

Infants greater than 15 months of age who are reactive for HIV-1 antibody by two different EIAs or HIV-1 Western Blot will be considered to be HIV-1 infected.

A non-breastfed infant born to an HIV-infected mother will be considered to be uninfected with HIV when two separate peripheral blood specimens are drawn on different days and both are negative either for HIV DNA, HIV RNA, and/or HIV culture. All of these tests will be performed in a laboratory that is approved to perform the assay for protocol testing. Specimens will be drawn at least 4 weeks apart and will be drawn when the infant is four weeks of age or older and has been off antiretrovirals for at least two weeks. At least one specimen will be drawn when the infant is greater than 8 weeks of age. Infants who are >9 months of age will be considered to be uninfected if they are nonreactive by EIA for HIV-1 antibody.

A breastfed infant born to an HIV-infected mother will be considered to be HIV uninfected if after 8 weeks from the time of weaning two separate peripheral blood specimens are drawn on different days and both are negative either for HIV DNA, HIV RNA and/or HIV culture. All of these tests will be performed in a laboratory that is approved to perform the assay for protocol testing. Specimens will be drawn at least 4 weeks apart and after the infant has been off antiretrovirals for at least two weeks.

In cases where there is one positive specimen for HIV DNA, HIV RNA, or HIV culture, the infant will be considered uninfected only if the last two subsequent specimens are negative by the same initially positive marker and one other marker.
8.0 HUMAN SUBJECTS

8.1 Institutional Review Board (IRB) Review and Informed Consent
This protocol and the informed consent documents (Appendix II) and any subsequent modifications will be reviewed and approved by the Institutional Review Boards or Ethics Committees responsible for oversight of the study. The mother must give written informed consent for herself and her baby's participation in the study. According to Federal regulations, for studies in fetuses in utero when the purpose of the study is to meet the health needs of the fetus, and the fetus will be placed at minimum risk necessary to meet these needs, or the risk to the fetus is minimal and the purpose of the study is to gain important biomedical knowledge that cannot be obtained by other means, the father's written informed consent is also required, unless his identity or whereabouts cannot reasonably be determined, or he is not reasonably available, or the pregnancy resulted from rape. The informed consents will describe the purpose of the study, the procedures to be followed, and the risks and benefits of participation. A copy of the consent form will be given to the mother (and father, if applicable).

8.2 Subject Confidentiality
All laboratory specimens, evaluation forms, reports, and other records will be identified only by a coded number to maintain subject confidentiality. All records will be kept in a locked file cabinet. All computer entry and networking programs will bers. Clinical information will not be released without written permission of the subject, except as necessary for monitoring by the HPTN or the NIAID.

8.3 Study Discontinuation
With proper justification, the study may be discontinued at any time by the HPTN Scientific Steering Group or the NIAID.

9.0 PUBLICATION OF RESEARCH FINDINGS
Publication of the results of this trial will be governed by DAIDS and HPTN policies. Any presentation, abstract, or manuscript will be made available for review by the HPTN Manuscript and Presentation Review Committee prior to submission.

10.0 BIOHAZARD CONTAINMENT
As the transmission of HIV and other blood-borne pathogens can occur through contact with contaminated needles, blood, and blood products, appropriate blood and secretion precautions will be employed by all personnel in the drawing of blood and shipping and handling of all specimens for this study, as currently recommended by the Centers for Disease Control and Prevention.
11.0 REFERENCES

APPENDICES

I. SCHEDULE OF EVALUATIONS

II. SAMPLE INFORMED CONSENTS

III. ADVERSE EVENT REPORTING REQUIREMENTS

IV. DIVISION OF AIDS TOXICITY TABLES FOR GRADING SEVERITY OF ADULT ADVERSE EXPERIENCES

V. DIVISION OF AIDS TOXICITY TABLES FOR GRADING SEVERITY OF PEDIATRIC ADVERSE EXPERIENCES

VI. NEVIRAPINE TOXICITY: SUPPLEMENTAL TOXICITY TABLE FOR GRADING SEVERITY OF CUTANEOUS/SKIN RASH/DERMATITIS ADVERSE EXPERIENCES
APPENDIX I SCHEDULE OF EVALUATIONS

A. Evaluations for HIV-infected and HIV-uninfected women from pregnancy through 3 months

<table>
<thead>
<tr>
<th>Maternal Evaluations</th>
<th>26-30 weeks</th>
<th>36 weeks</th>
<th>Labor and Delivery</th>
<th>4-6 weeks</th>
<th>3 months</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Laboratory:</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>BV status (Gram Stain, clue cells, whiff test, and pH)</td>
<td>X</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Presence of Candida, Trichomonas (2 wet mounts)</td>
<td>X</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>FFN status (ELISA)</td>
<td>X</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>HIV-infected cohort only:</strong> Blood for CBC, CD4</td>
<td>X</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>HIV-infected cohort only:</strong> Cervical swab for HIV load</td>
<td>X</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Placenta, membranes and cord for histology</td>
<td>X</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Non-Laboratory</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Obstetric/Medical History</td>
<td>X</td>
<td>X</td>
<td></td>
<td>X</td>
<td></td>
</tr>
<tr>
<td>Sexual History</td>
<td>X</td>
<td></td>
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<td></td>
<td></td>
</tr>
<tr>
<td>Gestational Age</td>
<td>X</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Concomitant medication usage</td>
<td>X</td>
<td>X</td>
<td></td>
<td>X</td>
<td></td>
</tr>
<tr>
<td>Assess for Adverse experiences</td>
<td>X</td>
<td></td>
<td></td>
<td>X</td>
<td></td>
</tr>
<tr>
<td>Length of labor/ROM</td>
<td>X</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Clinical chorioamnionitis</td>
<td>X</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Obstetric interventions</td>
<td>X</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Terminate</td>
<td>X</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
B. Evaluations for HIV-exposed and HIV-unexposed infants from birth through 3 months

<table>
<thead>
<tr>
<th>Neonate Evaluations</th>
<th>Visit 4 Birth - 48 hours</th>
<th>Visit 5 4-6 weeks</th>
<th>Visit 6 3 months</th>
<th>Post-Test Counseling Visit 3 months + 4 to 6 weeks</th>
<th>Post-test Counseling Visit 3 months + 8 to 12 weeks</th>
</tr>
</thead>
<tbody>
<tr>
<td>Laboratory: TO BE OBTAINED ON THE HIV-EXPOSED COHORT ONLY</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Blood for CBC</td>
<td>X</td>
<td>X</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>DBS for HIV RNA PCR</td>
<td>X</td>
<td>X</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>If already confirmed HIV-infected (i.e., 2 positive tests), do not draw DBS; terminate.</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>If confirmation of previous single positive result needed, collect DBS and schedule for post-test counseling.</td>
<td></td>
<td></td>
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</tr>
<tr>
<td>If all previous DBS samples have been negative, collect DBS and schedule for post-test counseling.</td>
<td></td>
<td></td>
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</tr>
<tr>
<td>If result of Visit 6 confirms HIV status, do not draw DBS; terminate.</td>
<td></td>
<td></td>
<td></td>
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</tr>
<tr>
<td>If result of Visit 6 is negative, no blood draw; terminate.</td>
<td></td>
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</tr>
<tr>
<td>If result of previous visit confirms HIV status, no blood draw; terminate.</td>
<td></td>
<td></td>
<td></td>
<td></td>
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</tr>
<tr>
<td>Blood for ALT</td>
<td>X</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Non-Laboratory: TO BE OBTAINED ON ALL INFANTS</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Weight</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td></td>
</tr>
<tr>
<td>Length, Head Circumference</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Gestational Age</td>
<td>X</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>General Health</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td></td>
<td>X</td>
</tr>
<tr>
<td>Assess for AEs</td>
<td>X</td>
<td>X</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Concomitant meds</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td></td>
<td>X</td>
</tr>
<tr>
<td>Post-test counseling</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

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C. Evaluations for women/infant pairs who return for final follow up at 3, 6 or 9 months post delivery

<table>
<thead>
<tr>
<th>Evaluations</th>
<th>Visit N (Visit 6, 7, or 8)</th>
<th>Post-Test Counseling</th>
<th>Post-Test Counseling</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Visit N + 4 to 6 weeks (HIV-indeterminate infants only)</td>
<td>Visit N + 8 to 12 weeks (HIV-indeterminate infants only)</td>
</tr>
<tr>
<td>Maternal:</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Discontinuation</td>
<td>X</td>
<td></td>
<td></td>
</tr>
<tr>
<td>All Infants:</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Weight</td>
<td>X</td>
<td>X</td>
<td>X</td>
</tr>
<tr>
<td>General Health</td>
<td>X</td>
<td>X</td>
<td>X</td>
</tr>
<tr>
<td>Concomitant meds</td>
<td>X</td>
<td>X</td>
<td>X</td>
</tr>
<tr>
<td>HIV-Indeterminate Infants ONLY</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Post-test counseling</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>DBS</td>
<td>If already confirmed HIV-infected (i.e., 2 positive tests), do not draw DBS; terminate.</td>
<td>If result of Visit N confirms HIV status, do not draw DBS; terminate.</td>
<td>If result of last visit confirms HIV status, no blood draw; terminate.</td>
</tr>
<tr>
<td></td>
<td>If confirmation of previous single positive result needed, collect DBS and schedule for Visit post-test counseling.</td>
<td>If result of Visit N is negative, no blood draw; terminate.</td>
<td>If result of Visit N is positive and confirmation is necessary, obtain DBS and schedule for post-test counseling</td>
</tr>
<tr>
<td></td>
<td>If all previous DBS studies have been negative, collect DBS and schedule for post-test counseling.</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
### D. Evaluations for women/infant pairs who return for final follow up at 12 months post delivery

<table>
<thead>
<tr>
<th>Evaluations</th>
<th>Visit 9 12 months</th>
<th>Post-Test Counseling 12 months + 1-2 weeks (HIV-indeterminate infants only)</th>
<th>Post-Test Counseling 12 months + 9-10 weeks (HIV-indeterminate infants only)</th>
<th>Post-Test Counseling 12 months + 13 to 16 weeks (HIV-indeterminate infants only)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Maternal: Terminations</td>
<td>X</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>All Infants: Weight</td>
<td>X</td>
<td>X</td>
<td></td>
<td>X</td>
</tr>
<tr>
<td>General Health</td>
<td>X</td>
<td>X</td>
<td></td>
<td>X</td>
</tr>
<tr>
<td>Concomitant meds</td>
<td>X</td>
<td>X</td>
<td></td>
<td>X</td>
</tr>
<tr>
<td>HIV-exposed infants ONLY Post-test counseling</td>
<td>X</td>
<td>X</td>
<td></td>
<td>X</td>
</tr>
<tr>
<td>ELISA, DBS</td>
<td>If already confirmed HIV-infected (i.e., 2 positive tests), do not draw DBS or ELISA; terminate. If of unknown HIV status with no previous positive HIV results, obtain ELISA and DBS. Schedule for post-test counseling. Note: • If ELISA negative, do not send DBS to UNC • If ELISA positive, send DBS to UNC</td>
<td>If ELISA negative, do not draw blood; terminate. If ELISA positive, draw DBS and schedule for post-test counseling.</td>
<td>If both RNA PCRs are positive, HIV infection is confirmed. No blood draw; terminate. If both RNA PCRs are negative, status at final visit is uninfected. No blood draw; terminate.</td>
<td>If result of last visit confirms HIV status, no blood draw; terminate.</td>
</tr>
</tbody>
</table>
### E: ORIGINAL SCHEDULE OF EVALUATIONS

<table>
<thead>
<tr>
<th>Maternal Evaluations</th>
<th>Pre-entry/Screening</th>
<th>Enrollment (20-24 weeks)</th>
<th>26-30 weeks</th>
<th>36 weeks</th>
<th>Labor and Delivery</th>
<th>Infant Follow-up visits (4-6 weeks; 3, 6, 9 12 months)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Laboratory:</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>HIV test</td>
<td>X</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Syphilis test (RPR for screening, TPHA for confirmation), at screening or enrollment</td>
<td>X</td>
<td>X</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>BV status (Gram Stain, clue cells, whiff test, and pH)</td>
<td>X</td>
<td>X</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Presence of Candida, Trichomonas (2 wet mounts)</td>
<td>X</td>
<td>X</td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>PFF status (ELISA)</td>
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<td>X</td>
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<td></td>
<td></td>
<td></td>
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<tr>
<td>Blood for CBC, CD4; Plasma for Viral Load</td>
<td>X</td>
<td>X</td>
<td>X³</td>
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<tr>
<td>Cervical swab for HIV load</td>
<td>X</td>
<td>X</td>
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<tr>
<td>Cervical swab for Gonorrhea Culture and Chlamydia</td>
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<tr>
<td>Placenta, membranes and cord for histology</td>
<td>X</td>
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<tr>
<td><strong>Non-Laboratory</strong></td>
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<tr>
<td>Informed consent</td>
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<td>Sexual History</td>
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<td>X</td>
<td>X</td>
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<tr>
<td>Antibiotic/other medication usage</td>
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<td>X</td>
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<tr>
<td>Pill counts/adherence</td>
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<td>X (4-6 weeks only)</td>
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<td>Adverse experiences</td>
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<td>X (4-6 weeks only)</td>
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<tr>
<td>Length of labor/ROM</td>
<td>X</td>
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<tr>
<td>Clinical chorioamnionitis</td>
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<tr>
<td>Obstetric interventions</td>
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<tr>
<td>General Health</td>
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<tr>
<td>Breastfeeding practices</td>
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¹ Specimens to be stored for later analysis
² In HIV-positive women only
³ No viral load is to be obtained at this visit
<table>
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<tr>
<th>Neonate Evaluations</th>
<th>Birth - 48 hours</th>
<th>4-6 weeks</th>
<th>3 months</th>
<th>6 months</th>
<th>9 months</th>
<th>12 months</th>
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<td>Blood for CBC</td>
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<td>Blood for HIV RNA PCR(^2)</td>
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<td>X(^4)</td>
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</tbody>
</table>

\(^1\) Only specimens from infants born to HIV positive women will be tested.
\(^2\) A positive test will be confirmed by a RNA PCR on a separate specimen no later than the next scheduled visit and sooner if possible. 12 month sample to be run only if HIV ELISA is reactive. If positive, will need to be confirmed with a separate sample as soon as possible.
\(^3\) In infants receiving Nevirapine. If the ALT result is a grade 1 or 2 abnormality, the test will be repeated at the 3 month visit. If the result is a grade 3 or 4 abnormality, the test will be repeated within one week of the test result.
APPENDIX II.  SAMPLE INFORMED CONSENTS

1.0 FOLLOW-UP INFORMED CONSENT FOR PREGNANT HIV-INFECTED PARTICIPANTS

2.0 FOLLOW-UP INFORMED CONSENT FOR PREGNANT HIV-UNINFECTED PARTICIPANTS

3.0 FOLLOW-UP INFORMED CONSENT FOR POSTPARTUM HIV-INFECTED PARTICIPANTS

4.0 FOLLOW-UP INFORMED CONSENT FOR POSTPARTUM HIV-UNINFECTED PARTICIPANTS

5.0 ORIGINAL SCREENING INFORMED CONSENT

6.0 ORIGINAL ENROLLMENT INFORMED CONSENT FOR SITES ENROLLING HIV-INFECTED AND HIV-UNINFECTED PARTICIPANTS

7.0 ORIGINAL ENROLLMENT INFORMED CONSENT FOR SITES ENROLLING HIV-INFECTED PARTICIPANTS ONLY
INTRODUCTION:
You are being asked to continue in the research study named above. When you first agreed to participate in the study, we said that we would tell you about any new information. As you know, this study used three drugs approved to fight infections in the uterus in the hopes that these three medications would reduce the chance that an HIV-infected mother would transmit the HIV to her baby. We now know that these three medications DO NOT help to reduce the chance that an HIV-infected mother would transmit the HIV virus to her baby. We also know that these medications are safe and are still effective for use in treating other illnesses.
We have answered the primary research question, and the study is continuing. However, the sponsors and researchers have decided to shorten the duration of follow-up so that, for each participant, the study will end earlier than what had previously been explained to you. Before you agree to participate in the remainder of the study, we would like to explain what the changes will mean for you.

This informed consent document gives you information about the study, which will be discussed with you. Once you understand the study, and if you agree to take part, you and your baby’s father, if available, will be asked to sign this consent or make your mark in front of someone. You will be given a copy to keep.

Please note that:

a. Your continued participation in this research is entirely voluntary;
b. You may decide not to complete or to withdraw from the study at any time without losing the benefits of your standard medical care.

PURPOSE OF THE STUDY:

The main purpose of this research study was to see if antibiotic drugs given to treat an infection of the uterus during pregnancy can reduce the chances of HIV, the virus that causes AIDS, being passed from an HIV-positive mother to her baby. There were other purposes, as well. As you know, antibiotic drugs are drugs used to fight infections in the body. This study will also see if these drugs improve the birth weight of the baby or prevent delivering the baby too early in an HIV positive or an HIV negative woman. The three drugs used in this study to treat infection of the uterus are metronidazole, erythromycin and ampicillin. These drugs have been approved for this use. Many women have used these drugs. The antibiotic drugs were not given to treat HIV infection. As you have been told, the study results showed that the antibiotics did not reduce the chance that an HIV-infected mother would transmit the virus to her baby. Therefore, the antibiotics are no longer being provided in this study.

Some drugs called antiretrovirals have been shown to reduce the chances of HIV being passed from an HIV-infected mother to her baby when given to mothers for several weeks during pregnancy. AZT is one of these drugs, and is approved for this use in the United States. A study in Uganda has shown that a drug called Nevirapine can help reduce the chance (by about half) of a mother with HIV from passing the HIV virus to her baby, when given as a one-time dose to the mother during labor/delivery and to the baby soon after birth. Although Nevirapine has been approved as a treatment for HIV in adults and children in the US, the use of Nevirapine to prevent passing of HIV from an infected mother to her child is still considered experimental in the United States. However, because of its proven safety and effectiveness, Nevirapine has been approved by the Ministry of Health in your country to help reduce the transmission of HIV from an infected mother to her baby. Since you have HIV, you and your baby have already been given or will be offered Nevirapine. Because you have been given Nevirapine as part of this study, you and your infant should continue to be followed until your infant is at least 3 months old.

A total of approximately 2500 women and their infants have participated in the study at this site and the other sites. Now that this study has changed, you and your baby will be followed up for 3 months following delivery or until the next follow up visit if your baby is older than three months.

PROCEDURES:
If all of your questions are answered and you agree to continue with follow up, you will be asked to sign this informed consent. Your follow-up schedule of visits will depend on whether you are still pregnant and/or your baby is less than 3 months old or whether your baby is 3 months of age or older. The main differences are that the changes in the study follow-up will reduce the number of study visits that you were originally asked to complete and will also reduce the number of times that blood will be obtained from your baby. As was explained in the previous informed consent, your baby will be tested for HIV and you will be asked to return for the results. If your baby tests positive for HIV at any of these times, we will need to obtain another sample of blood to make certain that the first result was correct.

If you are still pregnant and/or if your baby is less than 3 months old: At each prenatal visit you will be asked about your health and any medications you have taken recently. You will be asked questions about your sexual history. Some people may be embarrassed by these questions. You may choose not to answer any of the questions if you wish. Between 26 and 30 weeks of gestation, you will have a pelvic exam (an examination of your vagina) and will be tested for sexually transmitted diseases. You will have about 2 teaspoons of blood drawn for tests. You will be given vitamins to take by mouth every day until delivery.

If study antibiotics/placebos were given to you, you will be asked to return them.

At about 36 weeks of gestation, you will be asked about your health and any medications that you have taken since the last visit. You will have about 2 teaspoons of blood drawn for tests.

You will return to the hospital to deliver your baby, and you will be asked about medications you have taken since your last visit. After delivery, your placenta and cord will be tested to see if you had an infection during delivery. Your baby will be weighed and examined. A few drops of blood will be collected from your baby’s heel for tests. Your baby’s blood will be tested for HIV. The results of your baby’s HIV test will be given to you.

Since you have HIV, you and your baby will be offered Nevirapine. If you choose to accept the Nevirapine for you and your baby, at the second visit – when you are between 26 and 30 weeks gestation -- you will be given two pills to take by mouth at the beginning of labor. If you do not give birth to your baby within 48 hours of taking your Nevirapine dose, you will be given another Nevirapine dose to take. Your baby will receive one dose of Nevirapine by mouth in a syrup after birth. If you do not want you or your baby to take Nevirapine, you do not have to accept it. At the end of this form you and your baby’s father, if available, will sign or make your mark to indicate whether you want Nevirapine for you and your baby.

You and your baby will return to this clinic when your baby is between 4 to 6 weeks of age and at 3 months of age. At each visit you will be asked questions about your health, your baby’s health, and your breastfeeding practices. At each of these visits, your baby will have a small spoonful (1 ml) of blood drawn with a needle and will be tested for HIV. The results of your baby’s HIV test will be given to you. Because we cannot be certain that an infant is HIV infected until we have two positive tests from two different days, you may be asked to return for a follow-up visit to confirm your infant’s HIV status. You may return to this clinic at any time during the study to see a study nurse or doctor if you or your baby is sick.

If something unexpected causes you to deliver somewhere besides this hospital, you will still be asked to continue study visits so that the research staff can gather more information which may be
useful and important for future subjects. Ideally you should return to the clinic with 48 to 72 hours so that your infant can receive the Nevirapine. However, we can give the Nevirapine to your infant as long as you return within 7 days after the birth of your baby.

Some of your blood and placenta and some of your baby’s blood will be stored for later tests for this study. Some of these samples will be stored for future approved studies. These future studies may be related to HIV, preterm birth, or causes of infant death. Your name and your baby’s name will not be linked to any of these samples. They will be identified by a code to protect your privacy.

All pills given to you should be kept at room temperature away from heat and light.

If your infant is 3 months or older: If your baby is 3 months of age or older, and we don’t already know that your baby is infected with HIV, your baby will have a small spoonful (1 ml) of blood drawn with a needle and will be tested for HIV. You will be given an appointment to return for the results of your baby’s HIV test. If your baby’s test is positive, we will need to draw another small sample with a needle to test for HIV and will make an appointment for you to return for your baby’s test results.

RISKS and/or DISCOMFORTS:
Since you and your baby will take only one dose of Nevirapine, the risk of having side effects to the medication is unlikely. There have been many research studies of NVP to prevent the transmission of HIV from a mother to her baby; there were no significant side effects reported in any of them. Moreover, NVP has been dispensed to many mother and infants under the guidance of many Ministries of Health of many countries with no reports of significant side effects.

However, Nevirapine may cause some side effects when taken as a treatment on a daily basis. These are listed below. This list includes the more serious or common side effects:
• Rash, which has rarely been severe enough to require hospitalization and has been fatal
• Fever
• Headache
• Upset stomach (nausea)
• Swelling of the liver, which may rarely lead to severe and life-threatening liver damage, and very rarely fatal liver failure.

If you have questions about these or other side effects, ask your study doctor or nurse. At the time you and your baby are discharged from the hospital, you will receive instructions on what to do if you see any rash on you or your baby. It is very important that you tell the study doctor or nurse right away about any rash.

Some women experience mild discomfort during a pelvic exam. Some people experience discomfort when blood is drawn and may feel dizzy or even faint. You or your baby may have a bruise or swelling where the needle goes into to draw the blood.

POTENTIAL BENEFITS:
By continuing to take part in this research study, you will receive vitamins to take during pregnancy that may help you and your baby. Being in this study may reduce the chance of having your baby too soon and may help the overall health of your baby, but no guarantee can be made. Since you have HIV, you and your baby will be offered Nevirapine. Taking Nevirapine may reduce the
chances of your baby being infected with HIV, but no guarantee can be made. You may receive no benefit from this study. However, knowledge gained from this study may help others in the future.

NEW FINDINGS:
You will be told of any new information learned during the remaining study period that might cause you to change your mind about staying in the study.

REASONS WHY YOU MAY BE WITHDRAWN FROM THE STUDY WITHOUT YOUR CONSENT:
You may be removed from the study without your consent for the following reasons:

a. the study doctor decides that continuing in the study would be harmful to you;
b. the study is cancelled by the United States National Institutes of Health (NIH);
c. the study is cancelled by the United States Food and Drug Administration;
d. other administrative reasons;
e. your baby’s father objects to your baby being in the research study.

ALTERNATIVES TO PARTICIPATION:
There are programs in [your country] that can help to prevent a pregnant woman from passing HIV to her baby. If you would prefer not to continue with this study and are still pregnant, you can be referred to one of these programs.

If you do not participate in this study or withdraw from this study, you and your baby will receive standard medical care at this clinic/hospital and you and your baby will not have to undergo any further procedures as a part of this study.

COSTS TO YOU:
There is no cost to you for continuing to participate in the study. The study visits, multivitamins and Nevirapine are free.

CONFIDENTIALITY:
Your research records will be confidential to the extent permitted by local and United States law. You will be identified by a code, and personal information from your records will not be released without your written permission. You will not be personally identified in any publication about this study.

Your records may be reviewed by the National Institutes of Health, the U.S. agency that sponsors this research; the U.S. Food and Drug administration, the agency that reviews research; the pharmaceutical sponsors of this research and the study monitors.

RESEARCH-RELATED INJURY:
If you are injured as a result of participation in this study, the study clinic will give you immediate necessary treatment for your injuries. The cost of this treatment will not be charged to you. You will then be told where you may receive additional treatment for injuries. There is no compensation provided for research-related injuries.

NOTE: You are not giving up any of your legal rights by signing this informed consent document.

PERSONS TO CONTACT FOR PROBLEMS OR QUESTIONS:
If you ever have questions about this study or in case you are injured as a result of participation in this research study, you should contact (name of local investigator or study clinician) at (telephone number or address). If you ever have questions about your rights as a research subject you may call (name and title of IRB member) at (telephone number or address).
PHASE III TRIAL OF ANTIBIOTICS TO REDUCE CHORIOAMNIONITIS-RELATED PERINATAL HIV TRANSMISSION, HIVNET 024, Version 5.0

SIGNATURE PAGE

Client Name: ___________________________

Literate? Yes: __________________________ No: __________________________

CONSENT TO CONTINUE WITH THE STUDY

If you have read the informed consent or had it read and explained to you and understand the information, and you voluntarily agree to follow up with this study, please sign your name or make your mark below.

Volunteer’s name __________________________ Volunteer’s signature __________________________ Date __________________________

Person obtaining consent __________________________ Signature __________________________ Date __________________________

Witness to consent __________________________ Signature of witness __________________________ Date __________________________

If reasonably available:

Father’s name __________________________ Father’s signature __________________________ Date __________________________

CONSENT TO TAKE NEVIRAPINE (PREGNANT PARTICIPANTS ONLY)

If you have read the informed consent or had it read and explained to you and understand the information, and you want to receive Nevirapine for you and your baby, please sign or make your mark below.

Volunteer’s name __________________________ Volunteer’s signature __________________________ Date __________________________

Person obtaining consent __________________________ Signature __________________________ Date __________________________

Witness to consent __________________________ Signature of witness __________________________ Date __________________________

If reasonably available:

Father’s name __________________________ Father’s signature __________________________ Date __________________________
2.0 FOLLOW-UP INFORMED CONSENT FOR HIV-UNINFECTED PARTICIPANTS

Sample Informed Consent
for trials sponsored by the
Division of AIDS, NIAID, NIH, NICHD

REMINDER TO RESEARCH SITES: DO NOT USE PREAMBLE IN LOCAL CONSENTS

NOTE FROM OPRR (OFFICE FOR PROTECTION FROM RESEARCH RISKS) TO SITES ENROLLING SUBJECTS IN THIS STUDY:

PLEASE NOTE THAT THIS SAMPLE LANGUAGE DOES NOT PREEMPT OR REPLACE LOCAL IRB REVIEW AND APPROVAL. INVESTIGATORS ARE REQUIRED TO PROVIDE THE LOCAL IRB WITH A COPY OF THIS SAMPLE CONSENT ALONG WITH THE LANGUAGE INTENDED FOR LOCAL USE. LOCAL IRBS ARE REQUIRED TO WEIGH THE UNIQUE RISKS, CONSTRAINTS, AND POPULATION CONSIDERATIONS AS A CONDITION OF ANY APPROVAL. ANY DELETION OF SUBSTANTIVE CHANGE OR INFORMATION CONCERNING RISKS OR ALTERNATIVE TREATMENTS MUST BE JUSTIFIED BY THE INVESTIGATOR, APPROVED BY THE LOCAL IRB AND NOTED IN THE IRB MINUTES. JUSTIFICATION AND IRB APPROVAL OF SUCH CHANGES MUST BE FORWARDED TO FHI FOR ANY IMC-SPONSORED TRIAL. SPONSOR-APPROVED CHANGES IN AN IMC-PROTOCOL MUST BE APPROVED BY THE LOCAL IRB BEFORE USE UNLESS INTENDED FOR THE ELIMINATION OF APPARENT IMMEDIATE HAZARD. NEW INFORMATION SHALL BE SHARED WITH EXISTING SUBJECTS AT NEXT ENCOUNTER, WITH ALL NEW SUBJECTS PRIOR TO INVOLVEMENT, OR AS THE LOCAL IRB MAY OTHERWISE ADDITIONALLY REQUIRE.

PHASE III TRIAL OF ANTIBIOTICS TO REDUCE CHORIOAMNIONITIS-RELATED PERINATAL HIV TRANSMISSION, HIVNET 024, Version 5.0

FOLLOW-UP INFORMED CONSENT FORM

US Investigator In-country co-investigator
Contact information Contact information

INTRODUCTION:
You are being asked to continue in the research study named above. When you first agreed to participate in the study, we said that we would tell you about any new information. As you know, this study used three drugs approved to fight infections in the uterus in the hopes that these three medications would reduce the chance that an HIV-infected mother would transmit the HIV to her baby. We now know that these three medications DO NOT help to reduce the chance that an HIV-infected mother would transmit the HIV virus to her baby. We also know that these medications are safe and are still effective for use in treating other illnesses.
We have answered the primary research question, and the study is continuing. However, the sponsors and researchers have decided to shorten the duration of follow-up so that, for each participant, the study will end earlier than what had previously been explained to you. Before you agree to participate in the remainder of the study, we would like to explain what the changes will mean for you.

This informed consent document gives you information about the study, which will be discussed with you. Once you understand the study, and if you agree to take part, you and your baby’s father, if available, will be asked to sign this consent or make your mark in front of someone. You will be given a copy to keep.

Please note that:

a. Your continued participation in this research is entirely voluntary;
b. You may decide not to take part or to withdraw from the study at any time without losing the benefits of your standard medical care.

PURPOSE OF THE STUDY:
The main purpose of this research study was to see if antibiotic drugs given to treat an infection of the uterus during pregnancy can reduce the chances of HIV, the virus that causes AIDS, being passed from an HIV-positive mother to her baby. There were other purposes, as well. As you know, antibiotic drugs are drugs used to fight infections in the body. This study will also see if these drugs improve the birth weight of the baby or prevent delivering the baby too early in an HIV positive or an HIV negative woman. The three drugs used in this study to treat infection of the uterus are metronidazole, erythromycin and ampicillin. These drugs have been approved for this use. Many women have used these drugs. The antibiotic drugs were not given to treat HIV infection. As you have been told, the study results showed that the antibiotics did not reduce the chance that an HIV-infected mother would transmit the virus to her baby. Therefore, the antibiotics are no longer being provided in this study.

A total of approximately 2500 women and their infants have participated in the study at this site and the other sites. Now that this study has changed, you and your baby will be followed up for 3 months following delivery or until the next follow up visit if your baby is older than three months.

PROCEDURES:
If all of your questions are answered and you agree to continue with follow up, you will be asked to sign this informed consent. Your follow-up schedule of visits will depend on whether you are still pregnant and/or your baby is less than 3 months old or whether your baby is 3 months of age or older. The main differences are that the changes in the study follow-up will reduce the number of study visits that you were originally asked to complete and your baby will not have to have any blood drawn as part of the study.

If you are still pregnant and/or if your baby is less than 3 months old: At each prenatal visit you will be asked about your health and any medications you have taken recently. You will be asked questions about your sexual history. Some people may be embarrassed by these questions. You may choose not to answer any of the questions if you wish. Between 26 and 30 weeks of gestation, you will have a pelvic exam (an examination of your vagina) and will be tested for sexually transmitted diseases. You will have about 2 teaspoons of blood drawn for tests. You will be given vitamins to take by mouth every day until delivery.
If study antibiotics/placebos were given to you, you will be asked to return them.

At about 36 weeks of gestation, you will be asked about your health and any medications that you have taken since the last visit. You will have about 2 teaspoons of blood drawn for tests.

You will return to the hospital to deliver your baby, and you will be asked about medications you have taken since your last visit. After delivery, your placenta and cord will be tested to see if you had an infection during delivery. Your baby will be weighed and examined.

You and your baby will return to this clinic when your baby is between 4 to 6 weeks of age and at 3 months of age. At each visit you will be asked questions about your health, your baby’s health, and your breastfeeding practices. You may return to this clinic at any time during the study to see a study nurse or doctor if you or your baby is sick.

If something unexpected causes you to deliver somewhere besides this hospital, you will still be asked to continue study visits so that the research staff can gather more information which may be useful and important for future subjects.

Some of your blood and placenta will be stored for later tests for this study. Some of these samples will be stored for future approved studies. These future studies may be related to HIV, preterm birth, or causes of infant death. Your name and your baby’s name will not be linked to any of these samples. They will be identified by a code to protect your privacy.

All pills given to you should be kept at room temperature away from heat and light.

If your infant is 3 months or older: We will ask you questions about your breastfeeding practices and examine your baby.

**RISKS and/or DISCOMFORTS:**

Some women experience mild discomfort during a pelvic exam. Some people experience discomfort when blood is drawn and may feel dizzy or even faint. You may have a bruise or swelling where the needle goes into to draw the blood.

**POTENTIAL BENEFITS:**

By taking part in this research study, you will receive vitamins to take during pregnancy that may help you and your baby. Being in this study may reduce the chance of having your baby too soon and may help the overall health of your baby, but no guarantee can be made. You may receive no benefit from this study. However, knowledge gained from this study may help others in the future.

**NEW FINDINGS:**

You will be told of any new information learned during the course of the study that might cause you to change your mind about staying in the study.

**REASONS WHY YOU MAY BE WITHDRAWN FROM THE STUDY WITHOUT YOUR CONSENT:**

You may be removed from the study without your consent for the following reasons:

a. the study doctor decides that continuing in the study would be harmful to you;

b. the study is cancelled by the United States National Institutes of Health (NIH);
c. the study is cancelled by the United States Food and Drug Administration;
d. other administrative reasons;
e. your baby’s father objects to your baby being in the research study.

**ALTERNATIVES TO PARTICIPATION:**
If you do not participate in this study follow-up or withdraw from this study, you and your baby will receive standard medical care at this clinic/hospital and you and your baby will not have to undergo any further procedures as a part of this study.

**COSTS TO YOU:**
There is no cost to you for participating in the study. All of your drugs and study visits are free.

**CONFIDENTIALITY:**
Your research records will be confidential to the extent permitted by local and United States law. You will be identified by a code, and personal information from your records will not be released without your written permission. You will not be personally identified in any publication about this study.

Your records may be reviewed by the National Institutes of Health, the U.S. agency that sponsors this research; the U.S. Food and Drug administration, the agency that reviews research; the pharmaceutical sponsors of this research and the study monitors.

**RESEARCH-RELATED INJURY:**
If you are injured as a result of participation in this study, the study clinic will give you immediate necessary treatment for your injuries. The cost of this treatment will not be charged to you. You will then be told where you may receive additional treatment for injuries. There is no compensation provided for research-related injuries.

**NOTE:** You are not giving up any of your legal rights by signing this informed consent document.

**PERSONS TO CONTACT FOR PROBLEMS OR QUESTIONS:**
If you ever have questions about this study or in case you are injured as a result of participation in this research study, you should contact *(name of local investigator or study clinician)* at *(telephone number or address)*. If you ever have questions about your rights as a research subject you may call *(name and title of IRB member)* at *(telephone number or address)*.
PHASE III TRIAL OF ANTIBIOTICS TO REDUCE CHORIOAMNIONITIS-RELATED PERINATAL HIV TRANSMISSION, HIVNET 024, Version 5.0

SIGNATURE PAGE

Client Name:________________________________________

Literate? Yes: ____________________ No: ______________________

CONSENT TO CONTINUE WITH THE STUDY

If you have read the informed consent or had it read and explained to you and understand the information, and you voluntarily agree to follow up with this study, please sign your name or make your mark below:

____________________  ______________________  _____________________
Volunteer’s name     Volunteer’s signature     Date

____________________  ______________________  _____________________
Person obtaining consent Signature     Date

____________________  ______________________  _____________________
Witness to consent    Signature of witness     Date

If reasonably available:

____________________  ______________________  _____________________
Father’s name        Father’s signature     Date
3.0 FOLLOW-UP INFORMED CONSENT FOR POSTPARTUM HIV-INFECTED PARTICIPANTS

Sample Informed Consent for trials sponsored by the Division of AIDS, NIAID, NIH, NICHD

REMINDER TO RESEARCH SITES: DO NOT USE PREAMBLE IN LOCAL CONSENTS

NOTE FROM OPRR (OFFICE FOR PROTECTION FROM RESEARCH RISKS) TO SITES ENROLLING SUBJECTS IN THIS STUDY:

PLEASE NOTE THAT THIS SAMPLE LANGUAGE DOES NOT PREEMPT OR REPLACE LOCAL IRB REVIEW AND APPROVAL. INVESTIGATORS ARE REQUIRED TO PROVIDE THE LOCAL IRB WITH A COPY OF THIS SAMPLE CONSENT ALONG WITH THE LANGUAGE INTENDED FOR LOCAL USE. LOCAL IRBS ARE REQUIRED TO WEIGH THE UNIQUE RISKS, CONSTRAINTS, AND POPULATION CONSIDERATIONS AS A CONDITION OF ANY APPROVAL. ANY DELETION OF SUBSTANTIVE CHANGE OR INFORMATION CONCERNING RISKS OR ALTERNATIVE TREATMENTS MUST BE JUSTIFIED BY THE INVESTIGATOR, APPROVED BY THE LOCAL IRB AND NOTED IN THE IRB MINUTES. JUSTIFICATION AND IRB APPROVAL OF SUCH CHANGES MUST BE FORWARDED TO FHI FOR ANY IMC-SPONSORED TRIAL. SPONSOR-APPROVED CHANGES IN AN IMC-PROTOCOL MUST BE APPROVED BY THE LOCAL IRB BEFORE USE UNLESS INTENDED FOR THE ELIMINATION OF APPARENT IMMEDIATE HAZARD. NEW INFORMATION SHALL BE SHARED WITH EXISTING SUBJECTS AT NEXT ENCOUNTER, WITH ALL NEW SUBJECTS PRIOR TO INVOLVEMENT, OR AS THE LOCAL IRB MAY OTHERWISE ADDITIONALLY REQUIRE.

PHASE III TRIAL OF ANTIBIOTICS TO REDUCE CHORIOAMNIONITIS-RELATED PERINATAL HIV TRANSMISSION, HIVNET 024, Version 5.0

FOLLOW-UP INFORMED CONSENT FORM

<table>
<thead>
<tr>
<th>US Investigator</th>
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<td>Contact information</td>
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INTRODUCTION:
You are being asked to continue in the research study named above. When you first agreed to participate in the study, we said that we would tell you about any new information. As you know, this study used three drugs approved to fight infections in the uterus in the hopes that these three medications would reduce the chance that an HIV-infected mother would transmit the HIV to her baby. We now know that these three medications DO NOT help to reduce the chance that an HIV-infected mother would transmit the HIV virus to her baby. We also know that these medications are safe and are still effective for use in treating other illnesses.
We have answered the primary research question, and the study is continuing. However, the sponsors and researchers have decided to shorten the duration of follow-up so that, for each participant, the study will end earlier than what had previously been explained to you. Before you agree to participate in the remainder of the study, we would like to explain what the changes will mean for you.

This informed consent document gives you information about the study, which will be discussed with you. Once you understand the study, and if you agree to take part, you and your baby’s father, if available, will be asked to sign this consent or make your mark in front of someone. You will be given a copy to keep.

Please note that:

a. Your continued participation in this research is entirely voluntary;

b. You may decide not to complete or to withdraw from the study at any time without losing the benefits of your standard medical care.

PURPOSE OF THE STUDY:
The main purpose of this research study was to see if antibiotic drugs given to treat an infection of the uterus during pregnancy can reduce the chances of HIV, the virus that causes AIDS, being passed from an HIV-positive mother to her baby. There were other purposes, as well. As you know, antibiotic drugs are drugs used to fight infections in the body. This study will also see if these drugs improve the birth weight of the baby or prevent delivering the baby too early in an HIV positive or an HIV negative woman. The three drugs used in this study to treat infection of the uterus are metronidazole, erythromycin and ampicillin. These drugs have been approved for this use. Many women have used these drugs. The antibiotic drugs were not given to treat HIV infection. As you have been told, the study results showed that the antibiotics did not reduce the chance that an HIV-infected mother would transmit the virus to her baby. Therefore, the antibiotics are no longer being provided in this study.

Some drugs called antiretrovirals have been shown to reduce the chances of HIV being passed from an HIV-infected mother to her baby when given to mothers for several weeks during pregnancy. AZT is one of these drugs, and is approved for this use in the United States. A study in Uganda has shown that a drug called Nevirapine can help reduce the chance (by about half) of a mother with HIV from passing the HIV virus to her baby, when given as a one-time dose to the mother during labor/delivery and to the baby soon after birth. Although Nevirapine has been approved as a treatment for HIV in adults and children in the US, the use of Nevirapine to prevent passing of HIV from an infected mother to her child is still considered experimental in the United States. However, because of its proven safety and effectiveness, Nevirapine has been approved by the Ministry of Health in your country to help reduce the transmission of HIV from an infected mother to her baby. Since you have HIV, you and your baby have already been given. Because you have been given Nevirapine as part of this study, you and your infant should continue to be followed until your infant is at least 3 months old.

A total of approximately 2500 women and their infants have participated in the study at this site and the other sites. Now that this study has changed, you and your baby will be followed up for 3 months following delivery or until the next follow up visit if your baby is older than three months.

PROCEDURES:
If all of your questions are answered and you agree to continue with follow up, you will be asked to sign this informed consent. Your follow-up schedule of visits will depend on whether your baby is less than 3 months old or whether your baby is 3 months of age or older. The main differences are that the changes in the study follow-up will reduce the number of study visits that you were originally asked to complete and will also reduce the number of times that blood will be obtained from your baby. As was explained in the previous informed consent, your baby will be tested for HIV and you will be asked to return for the results. If your baby tests positive for HIV at any of these times, we will need to obtain another sample of blood to make certain that the first result was correct.

If your infant is less than 3 months old: You and your baby will return to this clinic when your baby is between 4 to 6 weeks of age and at 3 months of age. At each visit you will be asked questions about your health, your baby’s health, and your breastfeeding practices. At each of these visits, your baby will have a small spoonful (1 ml) of blood drawn with a needle and will be tested for HIV. The results of your baby’s HIV test will be given to you. Because we cannot be certain that an infant is HIV infected until we have two positive tests from two different days, you may be asked to return for a follow-up visit to confirm your infant’s HIV status. You may return to this clinic at any time during the study to see a study nurse or doctor if you or your baby is sick.

If your infant is 3 months or older: If your baby is 3 months of age or older, and we don’t already know that your baby is infected with HIV, your baby will have a small spoonful (1 ml) of blood drawn with a needle and will be tested for HIV. You will be given an appointment to return for the results of your baby’s HIV test. If your baby’s test is positive, we will need to draw another small sample with a needle to test for HIV and confirm the result of the first test. We will make an appointment for you to return for your baby’s test results.

Whatever the age of your infant: Some of your blood and placenta and some of your baby’s blood have been and will be stored for later tests for this study. Some of these samples will be stored for future approved studies. These future studies may be related to HIV, preterm birth, or causes of infant death. Your name and your baby’s name will not be linked to any of these samples. They will be identified by a code to protect your privacy.

**RISKS and/or DISCOMFORTS:**

Your baby may have a bruise or swelling where the needle goes in to draw the blood.

**POTENTIAL BENEFITS:**

Taking Nevirapine may have helped to reduce the chances of your baby being infected with HIV, but no guarantee can be made. You may receive no benefit from this study. However, knowledge gained from this study may help others in the future.

**NEW FINDINGS:**

You will be told of any new information learned during the remaining study period that might cause you to change your mind about staying in the study.

**REASONS WHY YOU MAY BE WITHDRAWN FROM THE STUDY WITHOUT YOUR CONSENT:**

You may be removed from the study without your consent for the following reasons:

a. the study doctor decides that continuing in the study would be harmful to you;
b. the study is cancelled by the United States National Institutes of Health (NIH);
c. the study is cancelled by the United States Food and Drug Administration;
d. other administrative reasons;
e. your baby’s father objects to your baby being in the research study.

**ALTERNATIVES TO PARTICIPATION:**
If you do not participate in this study or withdraw from this study, you and your baby will receive standard medical care at this clinic/hospital and you and your baby will not have to undergo any further procedures as a part of this study.

Before you decide to take part in this study, your study clinician will give you information about the risks and any potential benefits of participating in this research study.

**COSTS TO YOU:**
There is no cost to you for continuing to participate in the study. The study visits, are free.

**CONFIDENTIALITY:**
Your research records will be confidential to the extent permitted by local and United States law. You will be identified by a code, and personal information from your records will not be released without your written permission. You will not be personally identified in any publication about this study.

Your records may be reviewed by the National Institutes of Health, the U.S. agency that sponsors this research; the U.S. Food and Drug administration, the agency that reviews research; the pharmaceutical sponsors of this research and the study monitors.

**RESEARCH-RELATED INJURY:**
If you are injured as a result of participation in this study, the study clinic will give you immediate necessary treatment for your injuries. The cost of this treatment will not be charged to you. You will then be told where you may receive additional treatment for injuries. There is no compensation provided for research-related injuries.

**NOTE:** You are not giving up any of your legal rights by signing this informed consent document.

**PERSONS TO CONTACT FOR PROBLEMS OR QUESTIONS:**
If you ever have questions about this study or in case you are injured as a result of participation in this research study, you should contact (name of local investigator or study clinician) at (telephone number or address). If you ever have questions about your rights as a research subject you may call (name and title of IRB member) at (telephone number or address).
SIGNATURE PAGE

Client Name:___________________________

Literate? Yes: __________________
No: ________________________________

CONSENT TO CONTINUE WITH THE STUDY

If you have read the informed consent or had it read and explained to you and understand the information, and you voluntarily agree to follow up with this study, please sign your name or make your mark below.

Volunteer’s name ___________________________ Volunteer’s signature ___________________________
Date ___________________________

Person obtaining consent ___________________________ Signature ___________________________
Date ___________________________

Witness to consent ___________________________ Signature of witness ___________________________
Date ___________________________

If reasonably available:

Father’s name ___________________________ Father’s signature ___________________________
Date ___________________________
4.0 FOLLOW-UP INFORMED CONSENT FOR POSTPARTUM HIV-UNINFECTED PARTICIPANTS

REMINDER TO RESEARCH SITES: DO NOT USE PREAMBLE IN LOCAL CONSENTS

NOTE FROM OPRR (OFFICE FOR PROTECTION FROM RESEARCH RISKS) TO SITES ENROLLING SUBJECTS IN THIS STUDY:

PLEASE NOTE THAT THIS SAMPLE LANGUAGE DOES NOT PREEMPT OR REPLACE LOCAL IRB REVIEW AND APPROVAL. INVESTIGATORS ARE REQUIRED TO PROVIDE THE LOCAL IRB WITH A COPY OF THIS SAMPLE CONSENT ALONG WITH THE LANGUAGE INTENDED FOR LOCAL USE. LOCAL IRBS ARE REQUIRED TO WEIGH THE UNIQUE RISKS, CONSTRAINTS, AND POPULATION CONSIDERATIONS AS A CONDITION OF ANY APPROVAL. ANY DELETION OF SUBSTANTIVE CHANGE OR INFORMATION CONCERNING RISKS OR ALTERNATIVE TREATMENTS MUST BE JUSTIFIED BY THE INVESTIGATOR, APPROVED BY THE LOCAL IRB AND NOTED IN THE IRB MINUTES. JUSTIFICATION AND IRB APPROVAL OF SUCH CHANGES MUST BE FORWARDED TO FHI FOR ANY IMC-SPONSORED TRIAL. SPONSOR-APPROVED CHANGES IN AN IMC-PROTOCOL MUST BE APPROVED BY THE LOCAL IRB BEFORE USE UNLESS INTENDED FOR THE ELIMINATION OF APPARENT IMMEDIATE HAZARD. NEW INFORMATION SHALL BE SHARED WITH EXISTING SUBJECTS AT NEXT ENCOUNTER, WITH ALL NEW SUBJECTS PRIOR TO INVOLVEMENT, OR AS THE LOCAL IRB MAY OTHERWISE ADDITIONALLY REQUIRE.

PHASE III TRIAL OF ANTIBIOTICS TO REDUCE CHORIOAMNIONITIS-RELATED PERINATAL HIV TRANSMISSION, HIVNET 024, Version 5.0

FOLLOW-UP INFORMED CONSENT FORM

INTRODUCTION:
You are being asked to continue in the research study named above. When you first agreed to participate in the study, we said that we would tell you about any new information. As you know, this study used three drugs approved to fight infections in the uterus in the hopes that these three medications would reduce the chance that an HIV-infected mother would transmit the HIV to her baby. We now know that these three medications DO NOT help to reduce the chance that an HIV-infected mother would transmit the HIV virus to her baby. We also know that these medications are safe and are still effective for use in treating other illnesses.
We have answered the primary research question, and the study is continuing. However, the
sponsors and researchers have decided to shorten the duration of follow-up so that, for each
participant, the study will end earlier than what had previously been explained to you. Before you
agree to participate in the remainder of the study, we would like to explain what the changes will
mean for you.

This informed consent document gives you information about the study, which will be discussed
with you. Once you understand the study, and if you agree to take part, you and your baby’s father,
if available, will be asked to sign this consent or make your mark in front of someone. You will be
given a copy to keep.

Please note that:

a. Your continued participation in this research is entirely voluntary;
b. You may decide not to take part or to withdraw from the study at any time without losing the
   benefits of your standard medical care.

**PURPOSE OF THE STUDY:**
The main purpose of this research study was to see if antibiotic drugs given to treat an infection of
the uterus during pregnancy can reduce the chances of HIV, the virus that causes AIDS, being
passed from an HIV-positive mother to her baby. There were other purposes, as well. As you
know, antibiotic drugs are drugs used to fight infections in the body. This study will also see if
these drugs improve the birth weight of the baby or prevent delivering the baby too early in an HIV
positive or an HIV negative woman. The three drugs used in this study to treat infection of the
uterus are metronidazole, erythromycin and ampicillin. These drugs have been approved for this
use. Many women have used these drugs. The antibiotic drugs were not given to treat HIV
infection. As you have been told, the study results showed that the antibiotics did not reduce the
chance that an HIV-infected mother would transmit the virus to her baby. Therefore, the antibiotics
are no longer being provided in this study.

A total of approximately 2500 women and their infants have participated in the study at this site and
the other sites. Now that this study has changed, you and your baby will be followed up for 3
months following delivery or until the next follow up visit if your baby is older than three months.

**PROCEDURES:**
If all of your questions are answered and you agree to continue with follow up, you will be asked to
sign this informed consent. Your follow-up schedule of visits will depend on whether your baby is
less than 3 months old or whether your baby is 3 months of age or older. The main differences are
that the changes in the study follow-up will reduce the number of study visits that you were
originally asked to complete. In addition, no blood will be collected from your infant.

If your infant is less than 3 months old: You and your baby will return to this clinic when your
baby is between 4 to 6 weeks of age and at 3 months of age. At each visit you will be asked
questions about your health, your baby’s health, and your breastfeeding practices. You may return
to this clinic at any time during the study to see a study nurse or doctor if you or your baby is sick.

Some of your blood and placenta have been and will be stored for later tests for this study. Some of
these samples will be stored for future approved studies. These future studies may be related to
HIV, preterm birth, or causes of infant death. Your name and your baby’s name will not be linked
to any of these samples. They will be identified by a code to protect your privacy.
If your infant is 3 months or older: We will ask you questions about your breastfeeding practices and examine your baby. We will not collect any blood samples from your baby.

RISKS and/or DISCOMFORTS:
There are no risks to continuing with participation in this trial.

POTENTIAL BENEFITS:
You may receive no benefit from this study. However, knowledge gained from this study may help others in the future.

NEW FINDINGS:
You will be told of any new information learned during the course of the study that might cause you to change your mind about staying in the study.

REASONS WHY YOU MAY BE WITHDRAWN FROM THE STUDY WITHOUT YOUR CONSENT:
You may be removed from the study without your consent for the following reasons:

a. the study doctor decides that continuing in the study would be harmful to you;
b. the study is cancelled by the United States National Institutes of Health (NIH);
c. the study is cancelled by the United States Food and Drug Administration;
d. other administrative reasons;
e. your baby’s father objects to your baby being in the research study.

ALTERNATIVES TO PARTICIPATION:
If you do not participate in this study or withdraw from this study, you and your baby will receive standard medical care at this clinic/hospital and you and your baby will not have to undergo any further procedures as a part of this study.

COSTS TO YOU:
There is no cost to you for participating in the study. All of your study visits are free.

CONFIDENTIALITY:
Your research records will be confidential to the extent permitted by local and United States law. You will be identified by a code, and personal information from your records will not be released without your written permission. You will not be personally identified in any publication about this study.

Your records may be reviewed by the National Institutes of Health, the U.S. agency that sponsors this research; the U.S. Food and Drug administration, the agency that reviews research; the pharmaceutical sponsors of this research and the study monitors.

RESEARCH-RELATED INJURY:
If you are injured as a result of participation in this study, the study clinic will give you immediate necessary treatment for your injuries. The cost of this treatment will not be charged to you. You will then be told where you may receive additional treatment for injuries. There is no compensation provided for research-related injuries.
NOTE: You are not giving up any of your legal rights by signing this informed consent document.

PERSONS TO CONTACT FOR PROBLEMS OR QUESTIONS:
If you ever have questions about this study or in case you are injured as a result of participation in this research study, you should contact (name of local investigator or study clinician) at (telephone number or address). If you ever have questions about your rights as a research subject you may call (name and title of IRB member) at (telephone number or address).
PHASE III TRIAL OF ANTIBIOTICS TO REDUCE CHORIOAMNIONITIS-RELATED PERINATAL HIV TRANSMISSION, HIVNET 024, Version 5.0

SIGNATURE PAGE

Client Name: ________________________________

Literate? Yes: ____________________ No: ________________________________

CONSENT TO CONTINUE WITH THE STUDY

If you have read the informed consent or had it read and explained to you and understand the information, and you voluntarily agree to follow up with this study, please sign your name or make your mark below.

Volunteer’s name  Volunteer’s signature  Date

Person obtaining consent  Signature  Date

Witness to consent  Signature of witness  Date

If reasonably available:

Father’s name  Father’s signature  Date
5.0 ORIGINAL SCREENING INFORMED CONSENT

Sample Informed Consent for trials sponsored by the Division of AIDS, NIAID, NIH, NICHD

REMINDER TO RESEARCH SITES: DO NOT USE PREAMBLE IN LOCAL CONSENTS

NOTE FROM OPRR (OFFICE FOR PROTECTION FROM RESEARCH RISKS) TO SITES ENROLLING SUBJECTS IN THIS STUDY:

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PHASE III TRIAL OF ANTIBIOTICS TO REDUCE CHORIOAMNIONITIS-RELATED PERINATAL HIV TRANSMISSION, HIVNET 024

SCREENING INFORMED CONSENT FORM

INTRODUCTION:

You are being asked to take part in the screening for the research study named above, because you are pregnant. This study will use three drugs approved to fight infection. Before you decide whether or not to take part in the screening for this study, we would like to explain to you the purpose of the study, any risks to you, and what is expected of you.

This informed consent document gives you information about the screening visit part of the study that will be discussed with you. Once you understand the study, and if you agree to take part in the screening, you will be asked to sign this consent or make your mark in front of someone. You will be given a copy to keep.
Please note that:

a. Your participation in this research is entirely voluntary;
b. You may decide not to take part or to withdraw from the study at any time without losing the benefits of your standard medical care.

**PURPOSE OF THE STUDY:**
The purpose of this research study is to see if antibiotic drugs given to treat an infection of the uterus during pregnancy can reduce the chances of HIV, the virus that causes AIDS, being passed from an HIV-positive mother to her baby. Antibiotic drugs are drugs used to fight infections in the body. The study will also see if these drugs improve the birthweight of the baby or prevent delivering the baby too early in an HIV positive or an HIV negative woman. Three drugs will be used in this study. These drugs have been approved for the treatment of infection of the uterus. Many women have used these drugs. Some studies have shown that an infection of the uterus during pregnancy is associated with a mother with HIV passing the virus to her baby. None of these drugs is being given to treat HIV infection.

This research study will enroll about 3700 women at this site and other study sites. Your participation in the study will last about 16 months.

If you are interested in volunteering for this research study, you will need to meet the requirements for enrollment. You are being asked to give your permission to be tested for HIV and to see if you can be in the study.

If you meet the study requirements, you will be given more information about the study in another consent form. If you agree to participate in the study you will be asked to sign or make your mark on that consent form.

**PROCEDURES:**
After you sign this consent form, you will be asked to give a blood sample. About two teaspoons of blood will be needed. Your blood sample will be tested to see if you are infected with HIV or syphilis. Your test results will be available in one week. You will be asked to return to the clinic in one to two weeks to get your test results. Your test results will be explained to you. If you want to be in this research study, you must receive your HIV test results.

If your syphilis test shows that you have syphilis, you will be offered treatment at this clinic. You will not have to pay for this treatment.

You will be asked questions about any previous pregnancies and your medical history. Based on the results of your HIV test and your pregnancy and medical history, you may be asked to be in the study.

If your HIV test result is not positive but it is also not negative, you will not be able to be in this study. This type of test result means that doctors cannot tell whether you are or are not infected with HIV.

**RISKS and/or DISCOMFORTS:**
Some people experience discomfort when blood is drawn and may feel dizzy or even faint. You may have a bruise or swelling where the needle goes into your arm.
If you have HIV, knowing your HIV status may cause you anxiety. If others find out your HIV status, you may have trouble finding or keeping a job and have problems being accepted in your family and community.

**POTENTIAL BENEFITS:**
You may receive no direct benefit from these tests. However, you will receive information about your HIV status.

**COSTS TO YOU:**
There is no cost to you for these tests.

**CONFIDENTIALITY:**
Your research records will be confidential to the extent permitted by law. You will be identified by a code, and personal information from your records will not be released without your written permission. You will not be personally identified in any publication about this study.

Your records may be reviewed by the National Institutes of Health, the U.S. agency that sponsors this research; the U.S. Food and Drug administration, the agency that reviews research; the pharmaceutical sponsors of this research and the study monitors.

**RESEARCH-RELATED INJURY:**
If you are injured as a result of participation in this screening, the study clinic will give you immediate necessary treatment for your injuries. The cost of this treatment will not be charged to you. You will then be told where you may receive additional treatment for injuries. There is no compensation provided for research-related injuries.

**NOTE:** You are not giving up any of your legal rights by signing this informed consent document.

**PERSONS TO CONTACT FOR PROBLEMS OR QUESTIONS:**
If you ever have questions about this screening or in case you are injured as a result of participation in this screening, you should contact (name of local investigator or study clinician) at (telephone number or address). If you ever have questions about your rights as a research subject you may call (name and title of IRB member) at (telephone number or address).
SIGNATURE PAGE:
If you have read the informed consent, or had it read and explained to you, and you understand the information and voluntarily agree to be screened to join this study, please sign your name or make your mark below.

____________________ ______________________ _____________________
Volunteer’s name  Volunteer’s signature  Date

____________________ ______________________ _____________________
Witness’ name  Witness’ signature  Date
Sample Informed Consent for trials sponsored by the Division of AIDS, NIAID, NIH, NICHD

REMINDER TO RESEARCH SITES: DO NOT USE PREAMBLE IN LOCAL CONSENTS

NOTE FROM OPRR (OFFICE FOR PROTECTION FROM RESEARCH RISKS) TO SITES ENROLLING SUBJECTS IN THIS STUDY:

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PHASE III TRIAL OF ANTIBIOTICS TO REDUCE CHORIOAMNIONITIS-RELATED PERINATAL HIV TRANSMISSION, HIVNET 024

ENROLLMENT INFORMED CONSENT FORM

INTRODUCTION:
You are being asked to take part in the research study named above, because you are pregnant. This study will use three drugs approved to fight infection. Before you decide whether or not to take part in this study, we would like to explain to you the purpose of the study, any risks to you, and what is expected of you.

This informed consent document gives you information about the study which will be discussed with you. Once you understand the study, and if you agree to take part, you and your baby’s father,
if available, will be asked to sign this consent or make your mark in front of someone. You will be given a copy to keep.

Please note that:

a. Your participation in this research is entirely voluntary;
b. You may decide not to take part or to withdraw from the study at any time without losing the benefits of your standard medical care.

**PURPOSE OF THE STUDY:**
The purpose of this research study is to see if antibiotic drugs given to treat an infection of the uterus during pregnancy can reduce the chances of HIV, the virus that causes AIDS, being passed from an HIV-positive mother to her baby. Antibiotic drugs are drugs used to fight infections in the body. The study will also see if these drugs improve the birthweight of the baby or prevent delivering the baby too early in an HIV positive or an HIV negative woman. The three drugs used in this study to treat infection of the uterus are metronidazole, erythromycin and ampicillin. These drugs have been approved for this use. Many women have used these drugs. Some studies have shown that an infection of the uterus during pregnancy is associated with passing HIV from an HIV positive mother to her baby. None of the antibiotic drugs is being given to treat HIV infection.

Some drugs called antiretrovirals have been shown to reduce the chances of HIV being passed from an HIV-infected mother to her baby when given to mothers for several weeks during pregnancy. AZT is one of these drugs, and is approved for this use in the United States. More recently, a study in Uganda has shown that a drug called Nevirapine can help reduce the chance (by about half) of a mother with HIV from passing the HIV virus to her baby, when given as a one-time dose to the mother during labor/delivery and to the baby soon after birth. Nevirapine has been approved as a treatment for HIV in adults and children in the US. However, the use of Nevirapine to prevent passing of HIV from an infected mother to her child is still considered experimental. Experimental means that it may only be used in a research study in a limited number of people. None of these antiretroviral drugs are widely available in this country. If you have HIV, you and your baby will be offered Nevirapine. You do not have to accept Nevirapine to participate in this study.

This research study will enroll about 3700 women at this site and other study sites. Participation in the study will last about 16 months.

**PROCEDURES:**
After you sign the informed consent and have had a chance to ask questions, you will have three study visits before you go into labor. At this first visit you will be asked about your health and any medications you have taken recently. You will be asked questions about your sexual history. Some people may be embarrassed by these questions. You may choose not to answer any of the questions if you wish. You will have a pelvic exam (an examination of your vagina) and will be tested for sexually transmitted diseases. You will have about 2 teaspoons of blood drawn for tests. If you do not have HIV, some of your blood will be stored for future approved studies. You will be given vitamins to take by mouth every day until delivery.

At this visit you will be assigned by chance (like the tossing of a coin) to take the antibiotic study drugs or placebo. A placebo is a pill that looks just like the study drug but has no medicine in it. One woman of every two will receive antibiotic study drugs. One woman of every two will receive placebo. The study doctors and you will not know if you are taking the antibiotic study drug or placebo. Both groups of women, those assigned the antibiotic study drugs (metronidazole and
erythromycin) and those assigned the placebo will take their pills by mouth three times a day for 7 days. You will be asked to bring your pill packs, even if they are empty, to your next study visit.

At your second visit, you will be asked questions about your health and any medications you have taken since the last visit. You will have a pelvic exam and will be tested for sexually transmitted diseases. Your pill packs given to you at the first visit will be collected from you. You will be given pill packs to take home with you for when you go into labor. If you decide to join this study, you must agree to deliver your baby at the study hospital/clinic.

At your third visit you will be asked about your health and any medications that you have taken since the last visit. You will have about 2 teaspoons of blood drawn for tests.

When you go into labor, you will again begin taking either antibiotic study drugs or placebo as assigned. Women assigned antibiotic study drugs (metronidazole and ampicillin) and those assigned placebo will take pills by mouth every four hours until the baby is delivered. You will return to the hospital to deliver your baby. Your pill packs will be collected from you and you will be asked about medications you have taken since your last visit. After delivery, your placenta and cord will be tested to see if you had an infection during delivery. Your baby will be weighed and examined. A few drops of blood will be collected from your baby’s heel for tests. If you have HIV, your baby’s blood will be tested for HIV. The results of your baby’s HIV test will be given to you.

If you have HIV, you and your baby will be offered Nevirapine. If you choose to accept the Nevirapine for you and your baby, at the second visit you will be given one or two pills to take by mouth at the beginning of labor, in addition to the antibiotic study drugs. If you do not give birth to your baby within 48 hours of taking your Nevirapine dose, you will be given another Nevirapine dose to take. Your baby will receive one dose of Nevirapine by mouth in a syrup after birth. If you do not want you or your baby to take Nevirapine, you do not have to accept it. You and your baby can still be in the study even if you do not take Nevirapine. At the end of this form you and your baby’s father, if available, will sign or make your mark to indicate whether you want Nevirapine for you and your baby.

If you do not want Nevirapine for you and your baby, or if you do not have HIV, you will be given one or two placebo pills at the second visit to take when you go into labor, in addition to the antibiotic study drugs. These pills have no medicine in them. You do not have to accept these pills. You and your baby can still be in the study even if you do not take these pills.

After delivery, you will continue taking the antibiotic study drugs or placebo 3 times a day until you do not have any more drugs. You will bring your pill packs to your next clinic visit, even if they are empty.

You and your baby will return for about 5 study visits during the year after your baby is born. At each visit you will be asked questions about your health, your baby’s health, and your breastfeeding practices. During at least two of the visits your baby will have a heelstick to get a few drops of blood for tests. At the last study visit, your baby will have a small spoonful (1 ml) of blood drawn with a needle. If you have HIV, your baby’s blood will be tested for HIV. The results of your baby’s HIV test will be given to you. You may return to this clinic at any time during the study to see a study nurse or doctor if you or your baby is sick.
If something unexpected causes you to deliver somewhere besides this hospital, you will still be asked to continue study visits so that the research staff can gather more information which may be useful and important for future subjects. You will be asked to come to the hospital within 7 days after the birth of your baby so your baby can get the Nevirapine, if you choose to accept it for your baby.

Some of your blood and placenta and some of your baby’s blood will be stored for later tests for this study. Some of these samples will be stored for future approved studies. These future studies may be related to HIV, preterm birth, or causes of infant death. Your and your baby’s name will not be linked to any of these samples. They will be identified by a code to protect your privacy.

You and your baby will receive all of your standard medical care as part of this study. You should tell your study nurse or doctor when taking any non-study medications or enrolling in other research studies.

All pills given to you should be kept at room temperature away from heat and light. If you miss a dose, take it as soon as possible unless it is almost time for the next dose. If it is almost time for the next dose, then skip the missed dose and go back to your regular dosing schedule. Do not take two doses at once.

**RISKS and/or DISCOMFORTS:**
All three antibiotics have been studied and used extensively in pregnant women. However, we want to let you know of possible but rare problems. Some women are allergic to the antibiotics. Allergic means you have a certain kind of reaction after taking the drug. A very small number of people who take each of these drugs have serious reactions. If you experience any of the following, you should stop taking the pills and contact the study clinic immediately: skin rash; vomiting; severe stomach cramping; tightness of chest; swelling of eyelids, face or lips; wheezing or difficulty breathing.

Rarely, people taking these antibiotics have numbness or tingling in their arms and legs, seizures, and other nervous disorders. If you experience any of these side effects, you should stop taking the pills and contact the study clinic immediately.

Some studies have shown that women who take metronidazole during pregnancy are more likely to deliver their baby too early. Some studies have shown that taking metronidazole has no effect on when a woman delivers her baby. Some studies have shown that women who take metronidazole during pregnancy are less likely to deliver their baby too early.

If the medicine upsets your stomach, take your erythromycin pill two times a day for 2-3 days. If the medicine still upsets your stomach, take your erythromycin pill one time a day for 2-3 days. Continue taking your medicine until it is gone.

Do not drink alcohol while taking the pills. This may make you feel very sick. Wait for at least 3 days after you have stopped taking the pills to drink any alcohol.

Taking Nevirapine may cause some side effects, which are listed below. This list includes the more serious or common side effects that may be related to taking Nevirapine. If you have questions about these or other side effects, ask your study doctor or nurse. Possible side effects of Nevirapine are:
• Rash, which has rarely been severe enough to require hospitalization and has been fatal
• Fever
• Headache
• Upset stomach (nausea)
• Swelling of the liver, which may rarely lead to severe and life-threatening liver damage, and very rarely fatal liver failure.

Since you and your baby will take only one dose of Nevirapine, the risk of having these effects is much less likely. In a study conducted with about 600 babies who received one dose of Nevirapine within 2-3 days after birth, no rashes related to Nevirapine were seen. At the time you and your baby are discharged from the hospital, you will receive instructions on what to do if you see any rash on you or your baby. It is very important that you tell the study doctor or nurse right away about any rash.

Nevirapine has not been given with the antibiotics used in this study. We do not know if there will be an interaction between the Nevirapine and study antibiotics.

There may be some more risks from taking non-study drugs with the study drugs. There may be risks to your baby from taking these drugs, but no risks are known at the moment.

Some women experience mild discomfort during a pelvic exam. Some people experience discomfort when blood is drawn and may feel dizzy or even faint. You may have a bruise or swelling where the needle goes into your arm.

POSSIBLE BENEFITS:
By taking part in this research study, you will receive vitamins to take during pregnancy that may help you and your baby. Being in this study may reduce the chance of having your baby too soon and may help the overall health of your baby, but no guarantee can be made. If you have HIV, taking part in this study may reduce the chance of your baby being infected with HIV, but no guarantee can be made. If you have HIV, you and your baby will be offered Nevirapine. Taking Nevirapine may reduce the chances of your baby being infected with HIV, but no guarantee can be made. You may receive no benefit from this study. However, knowledge gained from this study may help others in the future.

NEW FINDINGS:
You will be told of any new information learned during the course of the study that might cause you to change your mind about staying in the study.

REASONS WHY YOU MAY BE WITHDRAWN FROM THE STUDY WITHOUT YOUR CONSENT:
You may be removed from the study without your consent for the following reasons:

a. the study doctor decides that continuing in the study would be harmful to you;
b. the study is cancelled by the United States National Institutes of Health (NIH);
c. the study is cancelled by the United States Food and Drug Administration;
d. other administrative reasons;
e. your baby’s father objects to your baby being in the research study.
If you need a treatment not allowed on this study or you have a bad reaction to the study drugs, you will stop taking the study drugs, but you and your baby will be requested to continue to have follow-up study visits.

**ALTERNATIVES TO PARTICIPATION:**
There are no drugs widely available to women in this country to help prevent an HIV-positive mother from passing HIV to her baby.

If you do not participate in this study or withdraw from this study, you and your baby will receive standard medical care at this clinic/hospital.

Before you decide to take part in this study, your study clinician will give you information about the risks and any potential benefits of participating in this research study.

**COSTS TO YOU:**
There is no cost to you for participating in the study. All of your drugs and study visits are free.

**CONFIDENTIALITY:**
Your research records will be confidential to the extent permitted by law. You will be identified by a code, and personal information from your records will not be released without your written permission. You will not be personally identified in any publication about this study.

Your records may be reviewed by the National Institutes of Health, the U.S. agency that sponsors this research; the U.S. Food and Drug administration, the agency that reviews research; the pharmaceutical sponsors of this research and the study monitors.

**RESEARCH-RELATED INJURY:**
If you are injured as a result of participation in this study, the study clinic will give you immediate necessary treatment for your injuries. The cost of this treatment will not be charged to you. You will then be told where you may receive additional treatment for injuries. There is no compensation provided for research-related injuries.

**NOTE:** You are not giving up any of your legal rights by signing this informed consent document.

**PERSONS TO CONTACT FOR PROBLEMS OR QUESTIONS:**
If you ever have questions about this study or in case you are injured as a result of participation in this research study, you should contact (name of local investigator or study clinician) at (telephone number or address). If you ever have questions about your rights as a research subject you may call (name and title of IRB member) at (telephone number or address).
SIGNATURE PAGE

CONSENT TO JOIN THE STUDY
If you have read the informed consent or had it read and explained to you and understand the information, and you voluntarily agree to join this study, please sign your name or make your mark below.

____________________  ______________________  _____________________
Volunteer’s name  Volunteer’s signature  Date

____________________  ______________________  _____________________
Witness’ name  Witness’ signature  Date

If reasonably available:

____________________  ______________________  _____________________
Father’s name   Father’s signature  Date

CONSENT TO TAKE NEVIRAPINE (HIV POSITIVE PARTICIPANTS ONLY)
If you have read the informed consent or had it read and explained to you and understand the information, and you want to receive Nevirapine for you and your baby, please sign or make your mark below.

____________________  ______________________  _____________________
Volunteer’s name  Volunteer’s signature  Date

____________________  ______________________  _____________________
Witness’ name  Witness’ signature  Date

If reasonably available:

____________________  ______________________  _____________________
Father’s name   Father’s signature  Date
7.0 ORIGINAL ENROLLMENT INFORMED CONSENT FOR SITES ENROLLING HIV-INFECTED PARTICIPANTS ONLY

Sample Informed Consent
for trials sponsored by the
Division of AIDS, NIAID, NIH, NICHD

REMINDER TO RESEARCH SITES: DO NOT USE PREAMBLE IN LOCAL CONSENTS

NOTE FROM OPRR (OFFICE FOR PROTECTION FROM RESEARCH RISKS) TO SITES ENROLLING SUBJECTS IN THIS STUDY:

PLEASE NOTE THAT THIS SAMPLE LANGUAGE DOES NOT PREEMPT OR REPLACE LOCAL IRB REVIEW AND APPROVAL. INVESTIGATORS ARE REQUIRED TO PROVIDE THE LOCAL IRB WITH A COPY OF THIS SAMPLE CONSENT ALONG WITH THE LANGUAGE INTENDED FOR LOCAL USE. LOCAL IRBS ARE REQUIRED TO WEIGH THE UNIQUE RISKS, CONSTRAINTS, AND POPULATION CONSIDERATIONS AS A CONDITION OF ANY APPROVAL. ANY DELETION OF SUBSTANTIVE CHANGE OR INFORMATION CONCERNING RISKS OR ALTERNATIVE TREATMENTS MUST BE JUSTIFIED BY THE INVESTIGATOR, APPROVED BY THE LOCAL IRB AND NOTED IN THE IRB MINUTES. JUSTIFICATION AND IRB APPROVAL OF SUCH CHANGES MUST BE FORWARDED TO FHI FOR ANY IMC-SPONSORED TRIAL. SPONSOR-APPROVED CHANGES IN AN IMC-PROTOCOL MUST BE APPROVED BY THE LOCAL IRB BEFORE USE UNLESS INTENDED FOR THE ELIMINATION OF APPARENT IMMEDIATE HAZARD. NEW INFORMATION SHALL BE SHARED WITH EXISTING SUBJECTS AT NEXT ENCOUNTER, WITH ALL NEW SUBJECTS PRIOR TO INVOLVEMENT, OR AS THE LOCAL IRB MAY OTHERWISE ADDITIONALLY REQUIRE.

PHASE III TRIAL OF ANTIBIOTICS TO REDUCE CHORIOAMNIONITIS-RELATED PERINATAL HIV TRANSMISSION, HIVNET 024

ENROLLMENT INFORMED CONSENT FORM

US Investigator In-country co-investigator
Contact information Contact information

INTRODUCTION:
You are being asked to take part in the research study named above, because you are pregnant. This study will use three drugs approved to fight infection. Before you decide whether or not to take part in this study, we would like to explain to you the purpose of the study, any risks to you, and what is expected of you.

This informed consent document gives you information about the study which will be discussed with you. Once you understand the study, and if you agree to take part, you and your baby’s father,
if available, will be asked to sign this consent or make your mark in front of someone. You will be given a copy to keep.

Please note that:

a. Your participation in this research is entirely voluntary;
b. You may decide not to take part or to withdraw from the study at any time without losing the benefits of your standard medical care.

**PURPOSE OF THE STUDY:**
The purpose of this research study is to see if antibiotic drugs given to treat an infection of the uterus during pregnancy can reduce the chances of HIV, the virus that causes AIDS, being passed from an HIV-positive mother to her baby. Antibiotic drugs are drugs used to fight infections in the body. The study will also see if these drugs improve the birthweight of the baby or prevent delivering the baby too early in an HIV positive or an HIV negative woman. The three drugs used in this study to treat infection of the uterus are metronidazole, erythromycin and ampicillin. These drugs have been approved for this use. Many women have used these drugs. Some studies have shown that an infection of the uterus during pregnancy is associated with passing HIV from an HIV positive mother to her baby. None of the antibiotic drugs is being given to treat HIV infection.

Some drugs called antiretrovirals have been shown to reduce the chances of HIV being passed from an HIV-infected mother to her baby when given to mothers for several weeks during pregnancy. AZT is one of these drugs, and is approved for this use in the United States. More recently, a study in Uganda has shown that a drug called Nevirapine can help reduce the chance (by about half) of a mother with HIV from passing the HIV virus to her baby, when given as a one-time dose to the mother during labor/delivery and to the baby soon after birth. Nevirapine has been approved as a treatment for HIV in adults and children in the US. However, the use of Nevirapine to prevent passing of HIV from an infected mother to her child is still considered experimental. Experimental means that it may only be used in a research study in a limited number of people. None of these antiretroviral drugs are widely available in this country. If you have HIV, you and your baby will be offered Nevirapine. You do not have to accept Nevirapine to participate in this study.

This research study will enroll about 3700 women at this site and other study sites. Participation in the study will last about 16 months.

**PROCEDURES:**
After you sign the informed consent and have had a chance to ask questions, you will have three study visits before you go into labor. At this first visit you will be asked about your health and any medications you have taken recently. You will be asked questions about your sexual history. Some people may be embarrassed by these questions. You may choose not to answer any of the questions if you wish. You will have a pelvic exam (an examination of your vagina) and will be tested for sexually transmitted diseases. You will have about 2 teaspoons of blood drawn for tests. You will be given vitamins to take by mouth every day until delivery.

At this visit you will be assigned by chance (like the tossing of a coin) to take the antibiotic study drugs or placebo. A placebo is a pill that looks just like the study drug but has no medicine in it. One woman of every two will receive antibiotic study drugs. One woman of every two will receive placebo. The study doctors and you will not know if you are taking the antibiotic study drug or placebo. Both groups of women, those assigned the antibiotic study drugs (metronidazole and
erythromycin) and those assigned the placebo will take their pills by mouth three times a day for 7 days. You will be asked to bring your pill packs, even if they are empty, to your next study visit.

At your second visit, you will be asked questions about your health and any medications you have taken since the last visit. You will have a pelvic exam and will be tested for sexually transmitted diseases. Your pill packs given to you at the first visit will be collected from you. You will be given pill packs to take home with you for when you go into labor. If you decide to join this study, you must agree to deliver your baby at the study hospital/clinic.

At your third visit you will be asked about your health and any medications that you have taken since the last visit. You will have about 2 teaspoons of blood drawn for tests.

When you go into labor, you will again begin taking either antibiotic study drugs or placebo as assigned. Women assigned antibiotic study drugs (metronidazole and ampicillin) and those assigned placebo will take pills by mouth every four hours until the baby is delivered. You will return to the hospital to deliver your baby. Your pill packs will be collected from you and you will be asked about medications you have taken since your last visit. After delivery, your placenta and cord will be tested to see if you had an infection during delivery. Your baby will be weighed and examined. A few drops of blood will be collected from your baby’s heel for tests. Your baby’s blood will be tested for HIV. The results of your baby’s HIV test will be given to you.

You and your baby will be offered Nevirapine. If you choose to accept the Nevirapine for you and your baby, at the second visit you will be given one or two pills to take by mouth at the beginning of labor, in addition to the antibiotic study drugs. If you do not give birth to your baby within 48 hours of taking your Nevirapine dose, you will be given another Nevirapine dose to take. Your baby will receive one dose of Nevirapine by mouth in a syrup after birth. If you do not want you or your baby to take Nevirapine, you do not have to accept it. You and your baby can still be in the study even if you do not take Nevirapine. At the end of this form you and your baby’s father, if available, will sign or make your mark to indicate whether you want Nevirapine for you and your baby.

After delivery, you will continue taking the antibiotic study drugs or placebo 3 times a day until you do not have any more drugs. You will bring your pill packs to your next clinic visit, even if they are empty.

You and your baby will return for about 5 study visits during the year after your baby is born. At each visit you will be asked questions about your health, your baby’s health, and your breastfeeding practices. During at least two of the visits your baby will have a heelstick to get a few drops of blood for tests. At the last study visit, your baby will have a small spoonful (1 ml) of blood drawn with a needle. Your baby’s blood will be tested for HIV. The results of your baby’s HIV test will be given to you. You may return to this clinic at any time during the study to see a study nurse or doctor if you or your baby is sick.

If something unexpected causes you to deliver somewhere besides this hospital, you will still be asked to continue study visits so that the research staff can gather more information which may be useful and important for future subjects. You will be asked to come to the hospital within 7 days after the birth of your baby so your baby can get the Nevirapine, if you choose to accept it for your baby.
Some of your blood and placenta and your baby’s blood will be stored for later tests for this study. These future studies may be related to HIV, preterm birth, or causes of infant death. Your and your baby’s name will not be linked to any of these samples. They will be identified by a code to protect your privacy.

You and your baby will receive all of your standard medical care as part of this study. You should tell your study nurse or doctor when taking any non-study medications or enrolling in other research studies.

All pills given to you should be kept at room temperature away from heat and light. If you miss a dose, take it as soon as possible unless it is almost time for the next dose. If it is almost time for the next dose, then skip the missed dose and go back to your regular dosing schedule. Do not take two doses at once.

**RISKS and/or DISCOMFORTS:**

All three antibiotics have been studied and used extensively in pregnant women. However, we want to let you know of possible but rare problems. Some women are allergic to the antibiotics. Allergic means you have a certain kind of reaction after taking the drug. A very small number of people who take each of these drugs have serious reactions. If you experience any of the following, you should stop taking the pills and contact the study clinic immediately: skin rash; vomiting; severe stomach cramping; tightness of chest; swelling of eyelids, face or lips; wheezing or difficulty breathing.

Rarely, people taking these antibiotics have numbness or tingling in their arms and legs, seizures, and other nervous disorders. If you experience any of these side effects, you should stop taking the pills and contact the study clinic immediately.

Some studies have shown that women who take metronidazole during pregnancy are more likely to deliver their baby too early. Some studies have shown that taking metronidazole has no effect on when a woman delivers her baby. Some studies have shown that women who take metronidazole during pregnancy are less likely to deliver their baby too early.

If the medicine upsets your stomach, take your erythromycin pill two times a day for 2-3 days. If the medicine still upsets your stomach, take your erythromycin pill one time a day for 2-3 days. Continue taking your medicine until it is gone.

Do not drink alcohol while taking the pills. This may make you feel very sick. Wait for at least 3 days after you have stopped taking the pills to drink any alcohol.

Taking Nevirapine may cause some side effects, which are listed below. This list includes the more serious or common side effects that may be related to taking Nevirapine. If you have questions about these or other side effects, ask your study doctor or nurse. Possible side effects of Nevirapine are:

- Rash, which has rarely been severe enough to require hospitalization and has been fatal
- Fever
- Headache
- Upset stomach (nausea)
• Swelling of the liver, which may rarely lead to severe and life-threatening liver damage, and very rarely fatal liver failure. Since you and your baby will take only one dose of Nevirapine, the risk of having these effects is much less likely. In a study conducted with about 600 babies who received one dose of Nevirapine within 2-3 days after birth, no rashes related to Nevirapine were seen. At the time you and your baby are discharged from the hospital, you will receive instructions on what to do if you see any rash on you or your baby. It is very important that you tell the study doctor or nurse right away about any rash.

Nevirapine has not been given with the antibiotics used in this study. We do not know if there will be an interaction between the Nevirapine and study antibiotics.

There may be some more risks from taking non-study drugs with the study drugs. There may be risks to your baby from taking these drugs, but no risks are known at the moment.

Some women experience mild discomfort during a pelvic exam. Some people experience discomfort when blood is drawn and may feel dizzy or even faint. You may have a bruise or swelling where the needle goes into your arm.

POTENTIAL BENEFITS:
By taking part in this research study, you will receive vitamins to take during pregnancy that may help you and your baby. Being in this study may reduce the chance of having your baby too soon and may help the overall health of your baby, but no guarantee can be made. Taking part in this study may reduce the chance of your baby being infected with HIV, but no guarantee can be made. You and your baby will be offered Nevirapine. Taking Nevirapine may reduce the chances of your baby being infected with HIV, but no guarantee can be made. You may receive no benefit from this study. However, knowledge gained from this study may help others in the future.

NEW FINDINGS:
You will be told of any new information learned during the course of the study that might cause you to change your mind about staying in the study.

REASONS WHY YOU MAY BE WITHDRAWN FROM THE STUDY WITHOUT YOUR CONSENT:
You may be removed from the study without your consent for the following reasons:
a. the study doctor decides that continuing in the study would be harmful to you;
b. the study is cancelled by the United States National Institutes of Health (NIH);
c. the study is cancelled by the United States Food and Drug Administration;
d. other administrative reasons;
e. your baby’s father objects to your baby being in the research study.

If you need a treatment not allowed on this study or you have a bad reaction to the study drugs, you will stop taking the study drugs, but you and your baby will be requested to continue to have follow-up study visits.

ALTERNATIVES TO PARTICIPATION:
There are no drugs widely available to women in this country to help prevent an HIV-positive mother from passing HIV to her baby.
If you do not participate in this study or withdraw from this study, you and your baby will receive standard medical care at this clinic/hospital.

Before you decide to take part in this study, your study clinician will give you information about the risks and any potential benefits of participating in this research study.

**COSTS TO YOU:**
There is no cost to you for participating in the study. All of your drugs and study visits are free.

**CONFIDENTIALITY:**
Your research records will be confidential to the extent permitted by law. You will be identified by a code, and personal information from your records will not be released without your written permission. You will not be personally identified in any publication about this study.

Your records may be reviewed by the National Institutes of Health, the U.S. agency that sponsors this research; the U.S. Food and Drug administration, the agency that reviews research; the pharmaceutical sponsors of this research and the study monitors.

**RESEARCH-RELATED INJURY:**
If you are injured as a result of participation in this study, the study clinic will give you immediate necessary treatment for your injuries. The cost of this treatment will not be charged to you. You will then be told where you may receive additional treatment for injuries. There is no compensation provided for research-related injuries.

**NOTE:** You are not giving up any of your legal rights by signing this informed consent document.

**PERSONS TO CONTACT FOR PROBLEMS OR QUESTIONS:**
If you ever have questions about this study or in case you are injured as a result of participation in this research study, you should contact *(name of local investigator or study clinician)* at *(telephone number or address)*. If you ever have questions about your rights as a research subject you may call *(name and title of IRB member)* at *(telephone number or address)*.
SIGNATURE PAGE

CONSENT TO JOIN THE STUDY
If you have read the informed consent or had it read and explained to you and understand the information, and you voluntarily agree to join this study, please sign your name or make your mark below.

____________________  ______________________  _____________________
Volunteer’s name  Volunteer’s signature  Date

____________________  ______________________  _____________________
Witness’ name  Witness’ signature  Date

If reasonably available:

____________________  ______________________  _____________________
Father’s name  Father’s signature  Date

CONSENT TO TAKE NEVIRAPINE
If you have read the informed consent or had it read and explained to you and understand the information, and you want to receive Nevirapine for you and your baby, please sign or make your mark below.

____________________  ______________________  _____________________
Volunteer’s name  Volunteer’s signature  Date

____________________  ______________________  _____________________
Witness’ name  Witness’ signature  Date

If reasonably available:

____________________  ______________________  _____________________
Father’s name  Father’s signature  Date
APPENDIX III. DIVISION OF AIDS TOXICITY TABLE FOR GRADING SEVERITY OF ADULT ADVERSE EXPERIENCES

Guidelines

ABBREVIATIONS: Abbreviations utilized in the Table:

ULN = Upper Limit of Normal  LLN = Lower Limit of Normal
Rx = Therapy  Req = Required
Mod = Moderate  IV = Intravenous
ADL = Activities of Daily Living  Dec = Decreased

ESTIMATING SEVERITY GRADE

For abnormalities NOT found elsewhere in the Toxicity Table use the scale below to estimate grade of severity:

GRADE 1  Mild
Transient or mild discomfort (< 48 hours); no medical intervention/therapy required

GRADE 2  Moderate
Mild to moderate limitation in activity - some assistance may be needed; no or minimal medical intervention/therapy required

GRADE 3  Severe
Marked limitation in activity, some assistance usually required; medical intervention/therapy required, hospitalizations possible

GRADE 4  Life-threatening
Extreme limitation in activity, significant assistance required; significant medical intervention/therapy required, hospitalization or hospice care probable.

SERIOUS OR LIFE-THREATENING AEs

ANY clinical event deemed by the clinician to be serious or life-threatening should be considered a grade 4 AE. Clinical events considered to be serious or life-threatening include, but are not limited to: seizures, coma, tetany, diabetic ketoacidosis, disseminated intravascular coagulation, diffuse petechiae, paralysis, acute psychosis, severe depression.
# TABLE FOR GRADING SEVERITY OF ADULT ADVERSE EXPERIENCES

<table>
<thead>
<tr>
<th>PARAMETER</th>
<th>GRADE 1 MILD</th>
<th>GRADE 2 MODERATE</th>
<th>GRADE 3 SEVERE</th>
<th>GRADE 4 POTENTIALLY LIFE THREATENING</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>HEMATOLOGY</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Hemoglobin</td>
<td>9.5 g/dL – 10.5 g/dL</td>
<td>8.0 g/dL – 9.4 g/dL</td>
<td>7.9 g/dL – 6.5 g/dL</td>
<td>&lt;6.5 g/dL</td>
</tr>
<tr>
<td>Absolute Neutrophil Count</td>
<td>1000 – 1500/mm³</td>
<td>750 – 999/mm³</td>
<td>500 - 749/mm³</td>
<td>&lt;500/mm³</td>
</tr>
<tr>
<td>WBC</td>
<td>&gt;13,000</td>
<td>&gt;15,000</td>
<td>&gt;20,000</td>
<td>&gt;30,000 or &lt;1,000</td>
</tr>
<tr>
<td>Percent Polys + Bands</td>
<td>&gt;80%</td>
<td>90%</td>
<td>≥95%</td>
<td>---------</td>
</tr>
<tr>
<td>Platelets</td>
<td>100,000 – 120,000/mm³</td>
<td>75,000 – 99,999/mm³</td>
<td>50,000 – 74,999/mm³</td>
<td>20,000 – 49,999/mm³</td>
</tr>
<tr>
<td>CD4 Counts</td>
<td>300 – 400/mm³</td>
<td>&lt;300/mm³</td>
<td>&lt;200/mm³</td>
<td>&lt;100/mm³</td>
</tr>
<tr>
<td>Uninfected Infected</td>
<td>&lt;300 or &lt;20%</td>
<td>&lt;200 or &lt;18%</td>
<td>&lt;100 or &lt;15%</td>
<td>&lt;50 or &lt;12%</td>
</tr>
<tr>
<td>Fibrinogen</td>
<td>100-200 mg/dl OR 400-600 mg/dl</td>
<td>&lt;100 mg/dl OR &gt;600mg/dl</td>
<td>&lt;50 mg/dl OR associated with gross bleeding OR associated with disseminated coagulation</td>
<td>---------</td>
</tr>
<tr>
<td>Prothrombin Time (PT)</td>
<td>&gt;1.0 – 1.24 x ULN</td>
<td>&gt;1.25 – 1.49 x ULN</td>
<td>&gt;1.5 – 3.0 x ULN</td>
<td>&gt;3.0 x ULN</td>
</tr>
<tr>
<td>PTT</td>
<td>&gt;1.0 – 1.66 x ULN</td>
<td>&gt;1.66 – 2.33 x ULN</td>
<td>&gt;2.33 – 3.0 x ULN</td>
<td>&gt;3.0 x ULN</td>
</tr>
</tbody>
</table>

<p>| <strong>CHEMISTRIES</strong>               |              |                  |                |                                      |
| CPK                           | ≥4 ULN        | ≥6 ULN           | ≥10 ULN        | ≥20 ULN                              |
| Creatinine                    | &gt;1.0 – 1.5 x ULN | &gt;1.5 – 1.9 x ULN | &gt;2.0 – 6.0 x ULN | &gt;6. x ULN                           |
| SODIUM                        | 130 – 135 meq/L | 123 – 129 meq/L  | 116 – 122 meq/L | &lt;116 meq/L                           |
| Hyponatremia                  | 146 – 150 meq/L | 151 – 157 meq/L  | 158 – 165 meq/L | &gt;165 meq/L                           |</p>
<table>
<thead>
<tr>
<th>PARAMETER</th>
<th>GRADE 1 MILD</th>
<th>GRADE 2 MODERATE</th>
<th>GRADE 3 SEVERE</th>
<th>GRADE 4 POTENTIALLY LIFE THREATENING</th>
</tr>
</thead>
<tbody>
<tr>
<td>POTASSIUM</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Hyperkalemia</td>
<td>5.0 – 5.5 meq/L</td>
<td>5.6 – 6.0 meq/L</td>
<td>6.1 – 6.5 meq/L</td>
<td>&gt;6.6 meq/L</td>
</tr>
<tr>
<td>Hypokalemia</td>
<td>3.2 – 3.4 meq/L</td>
<td>3.0 – 3.1 meq/L</td>
<td>2.5 – 2.9 meq/L</td>
<td>&lt;2.5 meq/L</td>
</tr>
<tr>
<td>PHOSPHATE</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Hypophosphalemia</td>
<td>2.0 – 2.4 mg/dL</td>
<td>1.5 – 1.9 mg/Dl</td>
<td>1.0 – 1.4 mg/dL</td>
<td>&lt;1.0 mg/Dl</td>
</tr>
<tr>
<td>CALCIUM</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Hypocalcemia</td>
<td>7.8 – 8.4 mg/dL</td>
<td>7.0 – 7.7 mg/dL</td>
<td>6.1 – 6.9 mg/dL</td>
<td>&lt;6.1 mg/dL</td>
</tr>
<tr>
<td>Hypercalcemia</td>
<td>10.6 – 11.5 mg/dL</td>
<td>11.6 – 12.5 mg/dL</td>
<td>12.6 – 13.5 mg/dL</td>
<td>&gt;13.5 mg/dL</td>
</tr>
<tr>
<td>MAGNESIUM</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Hypomagnesemia</td>
<td>1.2 – 1.4 meq/L</td>
<td>0.9 – 1.1 meq/L</td>
<td>0.6 – 0.8 meq/L</td>
<td>&lt;0.6 meq/L</td>
</tr>
<tr>
<td>BILIRUBIN</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Hyperbilirubinemia</td>
<td>&gt;1.0 – 1.5 x ULN</td>
<td>&gt;1.5 – 2.5 x ULN</td>
<td>&gt;2.5 – 5 x ULN</td>
<td>&gt;5 x ULN</td>
</tr>
<tr>
<td>GLUCOSE</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Hypoglycemia</td>
<td>55 – 64 mg/dL</td>
<td>40 – 54 mg/dL</td>
<td>30 – 39 mg/dL</td>
<td>&lt;30 mg/dL</td>
</tr>
<tr>
<td>Hyperglycemia</td>
<td>116 – 180 mg/dL</td>
<td>161 – 250 mg/dL</td>
<td>251 – 500 mg/dL</td>
<td>&lt;500 mg/dL</td>
</tr>
<tr>
<td>(nonfasting and no prior diabetes)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Triglycerides</td>
<td>--------------</td>
<td>400 – 750 mg/dL</td>
<td>751 – 1200 mg/dL</td>
<td>&gt;1200 mg/dL</td>
</tr>
<tr>
<td>URIC ACID</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Hyperuricemia</td>
<td>7.5 – 10.0 mg/dL</td>
<td>10.1 – 12.0 mg/dL</td>
<td>12.1 – 15.0 mg/dL</td>
<td>&gt;15.0 mg/dL</td>
</tr>
<tr>
<td>PARAMETER</td>
<td>GRADE 1 MILD</td>
<td>GRADE 2 MODERATE</td>
<td>GRADE 3 SEVERE</td>
<td>GRADE 4 POTENTIALLY LIFE THREATENING</td>
</tr>
<tr>
<td>------------------------------</td>
<td>--------------</td>
<td>------------------</td>
<td>---------------</td>
<td>-------------------------------------</td>
</tr>
<tr>
<td>LIVER TRANSAMINASE (LFTs)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>AST (SGOT)</td>
<td>1.25 – 2.5 x ULN</td>
<td>&gt;2.5 – 5.0 x ULN</td>
<td>&gt;5.0 – 10.0 x ULN</td>
<td>&gt;10.0 x ULN</td>
</tr>
<tr>
<td>ALT (SGPT)</td>
<td>1.25 – 3.0 x ULN</td>
<td>&gt;3.0 – 5.0 x ULN</td>
<td>&gt;5.0 – 10.0 x ULN</td>
<td>&gt;10.0 x ULN</td>
</tr>
<tr>
<td>GGT</td>
<td>1.25 – 2.5 x ULN</td>
<td>&gt;2.5 – 5.0 x ULN</td>
<td>&gt;5.0 – 10.0 x ULN</td>
<td>&gt;10.0 x ULN</td>
</tr>
<tr>
<td>Alk Phos</td>
<td>1.25 – 2.5 x ULN</td>
<td>&gt;2.5 – 5.0 x ULN</td>
<td>&gt;5.0 – 10.0 x ULN</td>
<td>&gt;10.0 x ULN</td>
</tr>
<tr>
<td>PANCREATIC ENZYMES</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Amylase</td>
<td>&gt;1.0 – 1.5 x ULN</td>
<td>&gt;1.5 – 2.0 x ULN</td>
<td>&gt;2.0 – 5.0 x ULN</td>
<td>5.0 x ULN</td>
</tr>
<tr>
<td>Pancreatic amylase</td>
<td>&gt;1.0 – 1.5 x ULN</td>
<td>&gt;1.5 – 2.0 x ULN</td>
<td>&gt;2.0 – 5.0 x ULN</td>
<td>5.0 x ULN</td>
</tr>
<tr>
<td>Lipase</td>
<td>&gt;1.0 – 1.5 x ULN</td>
<td>&gt;1.5 – 2.0 x ULN</td>
<td>&gt;2.0 – 5.0 x ULN</td>
<td>5.0 x ULN</td>
</tr>
<tr>
<td>CARDIOVASCULAR</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cardiac Arrhythmia</td>
<td>--------------</td>
<td>Asymptomatic; transient dysrhythmia, no Rx req</td>
<td>Recurrent/persistent dysrhythmia; symptomatic Rx req</td>
<td>Unstable dysrhythmia; hospitalization and Rx req</td>
</tr>
<tr>
<td>Hypertension</td>
<td>Transient, increase &gt;20 mm Hg diastolic BP, no Rx req</td>
<td>Recurrent; chronic increase &gt;20 mm Hg diastolic BP, Rx req</td>
<td>Acute Rx req; outpatient OR hospitalization possible</td>
<td>Hospitalization req OR end organ damage</td>
</tr>
<tr>
<td>Hypotension</td>
<td>Transient orthostatic hypotension with heart rate increased by &gt;20 beats/min OR decreased by &lt;10 mm Hg systolic BP, no Rx req</td>
<td>Symptoms OR BP decreased by &lt;20 mm Hg systolic, correctable with oral fluid Rx</td>
<td>IV fluid req. OR hospitalization</td>
<td>Mean arterial pressure &lt;60 mm Hg, OR end organ damage, OR shock, vasopressor Rx req</td>
</tr>
<tr>
<td>Pericarditis</td>
<td>Minimal effusion</td>
<td>Mild/mod asymptomatic effusion, no Rx</td>
<td>Symptomatic effusion, pain, EKG changes</td>
<td>Tamponade OR pericardiocentesis OR surgery req</td>
</tr>
<tr>
<td>Hemorrhage, blood loss</td>
<td>---------------</td>
<td>Mildly symptomatic no Rx req</td>
<td>Gross blood loss OR 1-2 units transfused</td>
<td>Massibe blood loss OR &gt;2 units transfused</td>
</tr>
</tbody>
</table>
### TABLE FOR GRADING SEVERITY OF ADULT ADVERSE EXPERIENCES

#### -continued-

<table>
<thead>
<tr>
<th>PARAMETER</th>
<th>GRADE 1</th>
<th>GRADE 2</th>
<th>GRADE 3</th>
<th>GRADE 4</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>MILD</td>
<td>MODERATE</td>
<td>SEVERE</td>
<td>POTENTIALLY</td>
</tr>
<tr>
<td>GASTROINTESTINAL</td>
<td></td>
<td></td>
<td></td>
<td>LIFE THREATENING</td>
</tr>
<tr>
<td>Nausea</td>
<td>Mild OR transient, reasonable intake maintained</td>
<td>Mod discomfort OR intake decreased for &lt;3 days</td>
<td>Severe discomfort OR minimal intake for ≥ 3 days</td>
<td>Hospitalization req</td>
</tr>
<tr>
<td>Vomiting</td>
<td>Mild OR transient; 2-3 episodes per day OR mild vomiting lasting &lt; 1 week</td>
<td>Mod OR persistent 4-5 episodes per day; OR vomiting lasting ≥ 1 week</td>
<td>Severe vomiting of all food/fluids in 24 hrs OR orthostatic hypotension OR IV Rx. req.</td>
<td>Hypotensive shock OR hospitalization req. for IV Rx. req.</td>
</tr>
<tr>
<td>Diarrhea</td>
<td>Mild OR transient; 3-4 loose stools per day OR mild diarrhea lasting &lt; 1 week</td>
<td>Mod OR persistent; 5-10 loose stools per day OR diarrhea lasting ≥ 1 week</td>
<td>&gt; 10 loose stools/day, bloody diarrhea; OR orthostatic hypotension OR electrolyte imbalance, &gt;2 L IV fluid required</td>
<td>Hypotensive shock OR severe electrolyte imbalance</td>
</tr>
<tr>
<td>Oral Discomfort/Dysphagia</td>
<td>Mild discomfort, no difficulty swallowing</td>
<td>Difficulty swallowing but able to eat and drink</td>
<td>Unable to swallow solids</td>
<td>Unable to drink fluids; IV fluids req.</td>
</tr>
<tr>
<td>Constipation</td>
<td></td>
<td></td>
<td></td>
<td>Distention with vomiting OR obstipation</td>
</tr>
<tr>
<td>RESPIRATORY</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cough (for aerosol studies)</td>
<td>Transient, no R</td>
<td>Treatment associated cough; inhaled bronchodilator</td>
<td>Uncontrolled cough; systemic Rx req.</td>
<td>--------------</td>
</tr>
<tr>
<td>Bronchospasm Acute</td>
<td>Transient; no R, FEV1 or peak flow reduced to 70%-80%</td>
<td>R,req; normalizes with bronchodilator; FEV1 or peak flow 50%-60%</td>
<td>No normalization with bronchodilator; FEV1 or peak flow 25% - 49%, retractions</td>
<td>Cyanosis; FEV1 or peak flow &lt;25% OR intubated</td>
</tr>
<tr>
<td>Dyspnea</td>
<td>Dyspnea on exertion</td>
<td>Dyspnea with normal activity</td>
<td>Dyspnea at rest</td>
<td>Dyspnea requiring O₂ therapy</td>
</tr>
</tbody>
</table>
### TABLE FOR GRADING SEVERITY OF ADULT ADVERSE EXPERIENCES

-continued-

<table>
<thead>
<tr>
<th>PARAMETER</th>
<th>GRADE 1 MILD</th>
<th>GRADE 2 MODERATE</th>
<th>GRADE 3 SEVERE</th>
<th>GRADE 4 POTENTIALLY LIFE THREATENING</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>NEUROLOGIC</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Neuro-cerebellar</td>
<td>Slight incoordination OR dysdiadochokinesia</td>
<td>Intention tremor OR dysmetria OR slurred speech OR nystagmus</td>
<td>Ataxia requiring assistance to walk or arm incoordination interfering with ADLs</td>
<td>Unable to stand</td>
</tr>
<tr>
<td>Neuro-psych/mood</td>
<td>--------------</td>
<td>--------------</td>
<td>--------------</td>
<td>-------------------------------------</td>
</tr>
<tr>
<td>Paresthesia (burning, tingling, etc.)</td>
<td>Mild discomfort; no Rx req.</td>
<td>Mod discomfort; non-narcotic analgesia required</td>
<td>Severe discomfort; OR narcotic analgesia req with symptomatic improvement</td>
<td>Acute psychosis req hospitalization; Suicidal ideation</td>
</tr>
<tr>
<td>Neuro-motor</td>
<td>Mild weakness in muscle of feet but able to walk and/or mild increase or decrease in reflexes</td>
<td>Mod weakness in feet (unable to walk on heels and/or toes), mild weakness in hands, still able to do most hand tasks and/or loss of previously present reflex or development of hyperreflexia and/or unable to do deep knee bends due to weakness</td>
<td>Marked distal weakness (unable to dorsiflex toes or foot drop, and mod proximal weakness e.g., in hands interfering with ADLs and/or requiring assistance to walk and/or unable to rise from chair unassisted</td>
<td>Incapacitating; OR not responsive to narcotic analgesia</td>
</tr>
<tr>
<td>Neuro-sensory</td>
<td>Mild impairment (decreased sensation, e.g., vibratory, pinprick, hot/cold in great toes) in focal area or symmetrical distribution</td>
<td>Mod impairment (mod decreased sensation, e.g., vibratory, pinprick, hot/cold to ankles) and/or joint position or mild impairment that is not symmetrical</td>
<td>Severe impairment (decreased or loss of sensation to knees or wrists) or loss of sensation of at least mod degree in multiple different body areas (i.e., upper and lower extremities)</td>
<td>Sensory loss involves limbs and trunk</td>
</tr>
<tr>
<td>PARAMETER</td>
<td>GRADE 1 MILD</td>
<td>GRADE 2 MODERATE</td>
<td>GRADE 3 SEVERE</td>
<td>GRADE 4 POTENTIALLY LIFE THREATENING</td>
</tr>
<tr>
<td>---------------------</td>
<td>--------------</td>
<td>------------------</td>
<td>----------------</td>
<td>-------------------------------------</td>
</tr>
<tr>
<td>MUSCULOSKELETAL</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Arthralgia/Arthritis</td>
<td>Arthralgia</td>
<td>Arthralgia with joint effusion or moderate impairment of activity</td>
<td>Frank arthritis with or without effusion OR resulting in severe impairment of activity</td>
<td></td>
</tr>
<tr>
<td>Myalgia</td>
<td>Myalgia without limitation of activity</td>
<td>Muscle tenderness at other than injection site or with moderate impairment of activity</td>
<td>Frank myonecrosis OR with severe impairment of activity</td>
<td></td>
</tr>
<tr>
<td>CUTANEOUS</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Rash/Dermatitis</td>
<td>Erythema, pruritus</td>
<td>Diffuse maculopapular rash OR dry desquamation</td>
<td>Vesiculation OR moist desquamation OR ulceration</td>
<td>ANY ONE: mucous membrane involvement, suspected Stevens-Johnson (TEN), erythema multiforme, necrosis req surgery, exfoliative dermatitis</td>
</tr>
<tr>
<td>Local Reaction</td>
<td>Erythema OR induration &lt;15 x 15 cm (225 cm²)</td>
<td>Erythema, induration, or Edema &gt;15 x 15 cm (225 cm²)</td>
<td>Ulceration OR super infection OR phlebitis</td>
<td>Necrosis of the skin</td>
</tr>
<tr>
<td>URINALYSIS</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Proteinuria</td>
<td>1+</td>
<td>2 – 3+</td>
<td>4+</td>
<td>Nephrotic syndrome</td>
</tr>
<tr>
<td>Random urine</td>
<td>200 mg-1 g loss/day OR &lt;0.3% OR &lt;3 g/l</td>
<td>1 – 2 g loss/day OR 0.3 – 1.0% OR 3 – 10 g/l</td>
<td>2 – 3.5 g loss/day OR &gt;1.0% OR &gt;10 g/l</td>
<td>Nephrotic syndrome OR &gt;3.5 g loss/day</td>
</tr>
<tr>
<td>24 hour urine</td>
<td></td>
<td></td>
<td></td>
<td>Obstructive OR transfusion req</td>
</tr>
<tr>
<td>Hematuria</td>
<td>Microscopic only ≤10 rbc/hpf</td>
<td>&gt;10 rbc/hpf</td>
<td>Gross, with or without clots OR RBC casts</td>
<td></td>
</tr>
<tr>
<td>PARAMETER</td>
<td>GRADE 1 MILD</td>
<td>GRADE 2 MODERATE</td>
<td>GRADE 3 SEVERE</td>
<td>GRADE 4 POTENTIALLY LIFE THREATENING</td>
</tr>
<tr>
<td>-------------------</td>
<td>--------------</td>
<td>------------------</td>
<td>----------------</td>
<td>-------------------------------------</td>
</tr>
<tr>
<td>MISCELLANEOUS</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Fever</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Oral &gt;12 hours</td>
<td>37.7 – 38.9°C (100.0 – 101.5°F)</td>
<td>39.0 – 39.5°C (101.6 – 102.9°F) or max temp of 103°F</td>
<td>39.6 – 40.5°C (103 – 105°F) or max temp of 103.5°F</td>
<td>&gt;40.5°C (205°F) or max temp &gt;105°F</td>
</tr>
<tr>
<td>Headache</td>
<td>Mild; no Rx req, OR non-narcotic analgesia Rx</td>
<td>Mod; OR responds to initial narcotic Rx</td>
<td>Severe; intractable; OR requiring repeated narcotic Rx</td>
<td>Requiring hospitalization, associated with neurologic, respiratory or cardiovascular abnormalities</td>
</tr>
<tr>
<td>Allergic Reaction</td>
<td>Pruritus without rash at injection site</td>
<td>Localized urticaria at injection site</td>
<td>Generalized urticaria angioedema</td>
<td>Anaphylaxis</td>
</tr>
<tr>
<td>ADL*</td>
<td>Normal activity reduced &lt;48 hours</td>
<td>Normal activity reduced 25-50% &gt;48 hours</td>
<td>Normal activity reduced &gt;50%; cannot work &gt; 48 hours</td>
<td>Unable to care for self</td>
</tr>
<tr>
<td>EYE</td>
<td>Mild pain, visual changes, conjunctival erythema, abnormal slit lamp</td>
<td>Loss of vision, clinically diagnosed uveitis, mod-severe pain, glaucoma</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

* ADL = Activities of Daily Living
### APPENDIX IV. DIVISION OF AIDS TOXICITY TABLE FOR GRADING SEVERITY OF PEDIATRIC ADVERSE EXPERIENCES

**DIVISION OF AIDS (DAIDS) TABLE FOR GRADING SEVERITY OF PEDIATRIC (<3 MONTHS OF AGE) ADVERSE EXPERIENCES**

For Vaccine & Prevention Research Programs

For other findings, the Toxicity Table for children >3 months of age (September 1993) is applicable.

Values are for term newborns

Preterm infants should be judged by a comparison of local normal ranges and the newborn ranges identified here.

<table>
<thead>
<tr>
<th>PARAMETER</th>
<th>GRADE 1</th>
<th>GRADE 2</th>
<th>GRADE 3</th>
<th>GRADE 4</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>HEMATOLOGY</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Hemoglobin 1-7 days old</td>
<td>13.0-14.0</td>
<td>12.0-12.9</td>
<td>&lt;12</td>
<td>Cardiac Failure Secondary to Anemia</td>
</tr>
<tr>
<td>8-21 days old</td>
<td>12.0-13.0</td>
<td>10.0-11.9</td>
<td>&lt;10.0</td>
<td>Cardiac Failure Secondary to Anemia</td>
</tr>
<tr>
<td>22-35 days old</td>
<td>9.5-10.5</td>
<td>8.0-9.4</td>
<td>&lt;8.0</td>
<td>Cardiac Failure Secondary to Anemia</td>
</tr>
<tr>
<td>36-56 days old</td>
<td>8.5-9.4</td>
<td>7.0-8.4</td>
<td>&lt;7.0</td>
<td>Cardiac Failure Secondary to Anemia</td>
</tr>
<tr>
<td>57-90 days old</td>
<td>9.0-9.9</td>
<td>7.0-8.9</td>
<td>&lt;7.0</td>
<td>Cardiac Failure Secondary to Anemia</td>
</tr>
<tr>
<td><strong>Abs Neutrophil Cnt</strong> 1 day old</td>
<td>5000-7000</td>
<td>3000-4999</td>
<td>1500-2999</td>
<td>&lt;1500</td>
</tr>
<tr>
<td>2-7 days old</td>
<td>1750-2500</td>
<td>1250-1749</td>
<td>750-1249</td>
<td>&lt;750</td>
</tr>
<tr>
<td>8-56 days old</td>
<td>1200-1800</td>
<td>900-1199</td>
<td>500-899</td>
<td>&lt;500</td>
</tr>
<tr>
<td>57-90 days old</td>
<td>750-1200</td>
<td>400-749</td>
<td>250-399</td>
<td>&lt;250</td>
</tr>
<tr>
<td><strong>Bilirubin</strong> &lt;7 days old</td>
<td>20-25</td>
<td>26-30</td>
<td>&gt;30</td>
<td></td>
</tr>
<tr>
<td>7-60 days old</td>
<td>1.1-1.9xN</td>
<td>2.0-2.9xN</td>
<td>3.0-7.5xN</td>
<td>&gt;7.5xN</td>
</tr>
<tr>
<td>61-90 days old</td>
<td>1.1-1.9xN</td>
<td>2.0-2.9xN</td>
<td>3.0-7.5xN</td>
<td>&gt;7.5xN</td>
</tr>
<tr>
<td><strong>Creatinine</strong> &lt;7 days old</td>
<td>1.0-1.7</td>
<td>1.8-2.4</td>
<td>2.5-3.0</td>
<td>&gt;3.0</td>
</tr>
<tr>
<td>7-60 days old</td>
<td>0.5-0.9</td>
<td>1.0-1.4</td>
<td>1.5-2.0</td>
<td>&gt;2.0</td>
</tr>
<tr>
<td>61-90 days old</td>
<td>0.6-0.8</td>
<td>0.9-1.1</td>
<td>1.2-1.5</td>
<td>&gt;1.5</td>
</tr>
<tr>
<td><strong>Cr Clearance</strong> &lt;7 days old</td>
<td>35-40</td>
<td>30-34</td>
<td>25.29</td>
<td>&lt;25</td>
</tr>
<tr>
<td>7-60 days old</td>
<td>45-50</td>
<td>40-44</td>
<td>35-39</td>
<td>&lt;35</td>
</tr>
<tr>
<td>61-90 days old</td>
<td>60-75</td>
<td>50-59</td>
<td>35-49</td>
<td>&lt;35</td>
</tr>
<tr>
<td><strong>Low Calcium</strong> &lt;7 days old</td>
<td>6.5-6.9</td>
<td>6.0-6.4</td>
<td>5.5-5.9</td>
<td>&lt;5.5</td>
</tr>
<tr>
<td>7-60 days old</td>
<td>7.6-8.0</td>
<td>7.0-7.5</td>
<td>6.0-6.9</td>
<td>&lt;6.0</td>
</tr>
<tr>
<td>61-90 days old</td>
<td>7.8-8.4</td>
<td>7.0-7.7</td>
<td>6.0-6.9</td>
<td>&lt;6.0</td>
</tr>
<tr>
<td><strong>High Calcium</strong> &lt;7 days old</td>
<td>12.0-12.4</td>
<td>12.5-12.9</td>
<td>13.0-13.5</td>
<td>&gt;13.5</td>
</tr>
<tr>
<td>7-60 days old</td>
<td>10.5-11.2</td>
<td>11.3-11.9</td>
<td>12.0-13.0</td>
<td>&gt;13.0</td>
</tr>
<tr>
<td>61-90 days old</td>
<td>10.5-11.2</td>
<td>11.3-11.9</td>
<td>12.0-12.9</td>
<td>≥13.0</td>
</tr>
</tbody>
</table>
**APPENDIX V: DIVISION OF AIDS (DAIDS) TABLE FOR GRADING SEVERITY OF PEDIATRIC (>3 MONTHS OF AGE) ADVERSE EXPERIENCES**

For

Vaccine & Prevention Research Programs

<table>
<thead>
<tr>
<th>PARAMETER</th>
<th>GRADE 1</th>
<th>GRADE 2</th>
<th>GRADE 3</th>
<th>GRADE 4</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>HEMATOLOGY</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Hemoglobin</td>
<td>9.0-9.9</td>
<td>7.0-8.9</td>
<td>&lt;7.0</td>
<td></td>
</tr>
<tr>
<td>&gt;3 mo to &lt;2 y.o.</td>
<td></td>
<td></td>
<td></td>
<td>Cardiac Failure Secondary to anemia</td>
</tr>
<tr>
<td>Hemoglobin</td>
<td>10-10.9</td>
<td>7.0-9.9</td>
<td>&lt;7.0</td>
<td></td>
</tr>
<tr>
<td>≥2 y.o.</td>
<td></td>
<td></td>
<td></td>
<td>Cardiac Failure Secondary to anemia</td>
</tr>
<tr>
<td>Abs Neutrophil Ct</td>
<td>750-1200</td>
<td>400-749</td>
<td>250-399</td>
<td>&lt;250</td>
</tr>
<tr>
<td>Platelets</td>
<td>50,000-75,000</td>
<td>25,000-49,999</td>
<td>25,000</td>
<td>25,000 or bleeding</td>
</tr>
<tr>
<td>PT</td>
<td>1.1-1.25xN</td>
<td>1.26-1.5xN</td>
<td>1.51-3.0xN</td>
<td>&gt;3xN</td>
</tr>
<tr>
<td>PTT</td>
<td>1.1-1.66xN</td>
<td>1.67-2.33xN</td>
<td>2.34-3.0xN</td>
<td>&gt;3xN</td>
</tr>
<tr>
<td><strong>GASTROINTESTINAL</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Bilirubin</td>
<td>1.1-1.9xN</td>
<td>2.0-2.9xN</td>
<td>3.0-7.5xN</td>
<td>&gt;7.5xN</td>
</tr>
<tr>
<td>AST (SGOT)</td>
<td>1.1-4.9xN</td>
<td>5.0-9.9xN</td>
<td>10.0-15.0xN</td>
<td>&gt;15.0xN</td>
</tr>
<tr>
<td>ALT (SGPT)</td>
<td>1.1-4.9xN</td>
<td>5.0-9.9xN</td>
<td>10.0-15.0xN</td>
<td>&gt;15.0xN</td>
</tr>
<tr>
<td>GGT</td>
<td>1.1-4.9xN</td>
<td>5.0-9.9xN</td>
<td>10.0-15.0xN</td>
<td>&gt;15.0xN</td>
</tr>
<tr>
<td>Pancreatic Amylase</td>
<td>1.1-1.4xN</td>
<td>1.5-1.9xN</td>
<td>2.0-3.0xN</td>
<td>&gt;3.0xN</td>
</tr>
<tr>
<td>Total Amylase + Lipase*</td>
<td>1.1-1.4xN</td>
<td>1.5-2.4xN</td>
<td>2.5-5.0xN</td>
<td>&gt;5.0xN</td>
</tr>
<tr>
<td>Uric Acid</td>
<td>7.5-9.9</td>
<td>10-12.4</td>
<td>12.5-15.0</td>
<td>&gt;15.0 or Gout</td>
</tr>
<tr>
<td>CPK</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Abdominal Pain</td>
<td>Mild</td>
<td>Moderate- No Rx Needed</td>
<td>Moderate Rx Needed</td>
<td>Severe- Hospital and Rx</td>
</tr>
<tr>
<td>Diarrhea</td>
<td>Soft stools</td>
<td>Liquid stools</td>
<td>Liquid Stools and Mild Dehydration Bloody stools</td>
<td>Dehydration requiring IV therapy or Hypotensive Shock</td>
</tr>
<tr>
<td>Constipation</td>
<td>Mild</td>
<td>Moderate</td>
<td>Severe</td>
<td>Distention and Vomiting</td>
</tr>
<tr>
<td>Nausea</td>
<td>Mild</td>
<td>Moderate- Decreased po intake</td>
<td>Severe- Little po intake</td>
<td>Unable to ingest food or fluid for &gt;24 hours</td>
</tr>
<tr>
<td>Vomiting</td>
<td>&lt;1 episode/day</td>
<td>1-3 episodes/day or duration&gt;3d</td>
<td>&gt;3 episodes/day or duration&gt;7d</td>
<td>Intractable Vomiting</td>
</tr>
</tbody>
</table>

*Both amylase and lipase must be elevated to the same grade or higher (i.e. if total amylase is Grade 4, but lipase is only Grade 1, the Toxicity Grade is 1.) In pediatric HIV patients, the most common source of serum amylase is the salivary glands. Salivary amylase elevations are generally not clinically significant. When amylase is released from damaged pancreatic cells, it can be a marker of pancreatitis. In most cases of clinical pancreatitis, lipase will also be elevated. However, lipase is also a non-specific marker. Combined elevation of amylase and lipase (each >5 x normal) often indicates pancreatic disease and requires evaluation. However, in the absence of pancreatic disease, drug can be resumed even at Grade 3 and 4 toxicities.
### DIVISION OF AIDS (DAIDS)

**TABLE FOR GRADING SEVERITY OF PEDIATRIC (>3 MONTHS OF AGE) ADVERSE EXPERIENCES**

*For Vaccine & Prevention Research Programs -continued-

<table>
<thead>
<tr>
<th>PARAMETER</th>
<th>GRADE 1</th>
<th>GRADE 2</th>
<th>GRADE 3</th>
<th>GRADE 4</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>RENAL AND ELECTROLYTES</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>CREATININE</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>2 Months-2 Years</td>
<td>0.6-0.8</td>
<td>0.9-1.1</td>
<td>1.2-1.5</td>
<td>&gt;1.5</td>
</tr>
<tr>
<td>2 Years-Adolescent</td>
<td>0.7-1.0</td>
<td>1.1-1.6</td>
<td>1.7-2.0</td>
<td>&gt;2.0</td>
</tr>
<tr>
<td>Adolescents</td>
<td>1.0-1.7</td>
<td>1.8-2.4</td>
<td>2.5-3.5</td>
<td>&gt;3.5</td>
</tr>
<tr>
<td>Creatinine Clearance</td>
<td>60-75 cc/min/1.73 m²</td>
<td>50-59 cc/min/1.73 m²</td>
<td>35-49 cc/min/1.73 m²</td>
<td>&lt;35 cc/min/1.73 m²</td>
</tr>
<tr>
<td><strong>ELECTROLYTES</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>High Sodium</td>
<td>145-149</td>
<td>150-155</td>
<td>&gt;155 or mental status changes</td>
<td></td>
</tr>
<tr>
<td>Low Sodium</td>
<td>130-135</td>
<td>129-124</td>
<td>&lt;124 or mental status changes</td>
<td></td>
</tr>
<tr>
<td>High Potassium</td>
<td>5.0-5.9</td>
<td>6.0-6.4</td>
<td>6.5-7.0</td>
<td>&gt;7.0 or Cardiac arrhythmias</td>
</tr>
<tr>
<td>Low Potassium</td>
<td>3.0-3.5</td>
<td>25-2.9</td>
<td>2.0-2.4</td>
<td>&lt;2.0</td>
</tr>
<tr>
<td>High Calcium</td>
<td>10.5-11.2</td>
<td>11.3-11.9</td>
<td>12.0-12.9</td>
<td>≥ 13.0</td>
</tr>
<tr>
<td>Low Calcium</td>
<td>7.8-8.4</td>
<td>7.0-7.7</td>
<td>6.0-6.9</td>
<td>&lt;6.0</td>
</tr>
<tr>
<td>Low Magnesium</td>
<td>1.2-1.4</td>
<td>0.9-1.1</td>
<td>0.6-0.8</td>
<td>&lt;0.6 or Cardiac arrhythmias</td>
</tr>
<tr>
<td>Hypoglycemia</td>
<td>55-65</td>
<td>40-54</td>
<td>30-39</td>
<td>&lt;30 or Mental status changes</td>
</tr>
<tr>
<td>Hyperglycemia</td>
<td>116-159</td>
<td>160-249</td>
<td>250-400</td>
<td>&gt;400 or Ketoacidosis</td>
</tr>
<tr>
<td>Proteinuria</td>
<td>Tr-1 +, &lt;150 mg/day</td>
<td>2+ 150-499 mg/day</td>
<td>3+ 500-1000 mg/day</td>
<td>4+ or nephrotic syndrome &gt;1000 mg/day</td>
</tr>
<tr>
<td>Hematuria</td>
<td>Microscopic ≤ 25 cells/hpf</td>
<td>Microscopic ≥ 25 cells/hpf</td>
<td>Gross</td>
<td>Obstruction or Transfusion requirement</td>
</tr>
</tbody>
</table>

**Comments**

Calcium values are corrected for albumin concentration. CrCl values do not apply to infants <2 months old.

<table>
<thead>
<tr>
<th>OTHER</th>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Allergy</td>
<td>Pruritis without Rash</td>
<td>Pruritic Rash</td>
<td>Mild Urticaria</td>
<td>Severe urticaria Anaphylaxis, Angioedema</td>
</tr>
<tr>
<td>Drug Fever (Rectal)</td>
<td>38.5-40</td>
<td>&gt;40</td>
<td>Sustained Fever: &gt;40, &gt;5 days</td>
<td></td>
</tr>
<tr>
<td>Cutaneous</td>
<td>Diffuse maculopapular rash, dry desquamation</td>
<td>Vesiculation, ulcers</td>
<td>Exfoliative dermatitis, Stevens-Johnson or Erythema multiforme, Moist desquamation</td>
<td></td>
</tr>
<tr>
<td>Stomatits</td>
<td>Mild discomfort</td>
<td>Painful; difficulty swallowing, but able to eat and drink</td>
<td>Painful; unable to swallow solids</td>
<td>Painful; requires IV</td>
</tr>
</tbody>
</table>

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### DIVISION OF AIDS (DAIDS)
### TABLE FOR GRADING SEVERITY OF PEDIATRIC (> 3MONTHS OF AGE) ADVERSE EXPERIENCES
For Vaccine & Prevention Research Programs
-continued-

<table>
<thead>
<tr>
<th>SYMPTOM</th>
<th>GRADE 1</th>
<th>GRADE 2</th>
<th>GRADE 3</th>
<th>GRADE 4</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>CENTRAL NERVOUS SYSTEM</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Seizures</td>
<td>None</td>
<td>1 Uncomplicated Sz +/- Temp Elevation</td>
<td>1 Sz/Month for ≥2 consecutive Months Or 3 Sz over 6 Months; No Temp Elevation</td>
<td>&gt;1 Sz/ Month; No Temp elevation; No Decrease in Sz Frequency Despite dose reduction</td>
</tr>
<tr>
<td>Seizures are an ubiquitous symptom of numerous systemic or CNS disturbances; alternative explanations should be vigorously sought and eliminated. Status epilepticus represents a severe end of the seizure spectrum, but should be considered as a single seizure event. The need for chronic or acute anticonvulsant medication should be made on a clinical basis. Seizures as a manifestation of drug toxicity are usually primarily generalized. Focal (partial onset) seizures are suggestive of focal central nervous system pathology and should be appropriately investigated, although they may be a manifestation of drug toxicity. Beware of focal seizures which secondarily generalize; these should be approached diagnostically as partial onset seizures. Children with underlying epileptic conditions who experience persistent breakthrough seizures despite maximal anticonvulsant therapy coincident with beginning the trial medication should be considered Grade 4.</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Headache</td>
<td>≤1/Month &lt;2hrs duration</td>
<td>&gt;1/Month &gt;2 hrs duration Moderate to Severe Responds to non-narcotic analgesia or prophylaxis</td>
<td>&gt;2/Month &gt;2hrs duration Moderate to Severe Responds to narcotic analgesia, or does not respond to prophylaxis</td>
<td>&gt;4/Month; &gt;2hrs Duration; Moderate to Severe; Non-Responsive to narcotic Analgesia; or persistently Recurrent despite prophylaxis No decrease in frequency or Severity despite dose reduction</td>
</tr>
<tr>
<td>Headache is a non-specific symptom, but may be a symptom of CNS/intracranial pathology. Appropriate diagnostic measures should be pursued. Duration refers to the waxing and peak phases, not to the resolution/waning phases of the headache. Mild refers to a grade of headache pain which does not affect function or activity. Moderate to sever refers to a grade of headache which affects function or activity.</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mental Status and Behavior</td>
<td>Changes which do not Affect Function</td>
<td>Changes requiring pharmacologic or other therapy; or mild lethargy, sedation or somnolence which resolves with rest</td>
<td>Changes not improved by drugs or other therapies; or onset of convulsion, memory impairiment, lethargy, sedation, or somnolence which does not respond to rest</td>
<td>Onset of delirium, obtundation, coma, or psychosis, or Grade 3 toxicity which does not respond to dose reduction</td>
</tr>
<tr>
<td>Behavior refers to the development of attention deficits with or without hyperactivity, depression, mania, agitation, sleep disorders, phobias, obsessive-compulsive behaviors, or anxiety. Mental status refers to the level of consciousness, memory function, language and analytical operation, and non-dominant hemisphere functioning. Alternative explanations should be sought.</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Balance and Posture</td>
<td>None</td>
<td>None</td>
<td>Ataxia, dizziness, vertigo, tremor, impaired postural balance</td>
<td>Onset of movement disorder; or Grade 3 toxicity which does not respond to dosage adjustment</td>
</tr>
<tr>
<td>&quot;Ataxia” can be mistakenly diagnosed in the face of central weakness or peripheral neuropathy, which should not be considered a drug toxicity of this category. Movement disorders refer to tardive or other dyskinesias, dystonias, chorea, or ballismus. Alternative explanations should be sought.</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
**DIVISION OF AIDS (DAIDS)**

**TABLE FOR GRADING SEVERITY OF PEDIATRIC (> 3MONTHS OF AGE) ADVERSE EXPERIENCES**

For

Vaccine & Prevention Research Programs

-continued-

<table>
<thead>
<tr>
<th>SYMPTOM</th>
<th>GRADE 1</th>
<th>GRADE 2</th>
<th>GRADE 3</th>
<th>GRADE 4</th>
</tr>
</thead>
<tbody>
<tr>
<td>Visual</td>
<td>None</td>
<td>Blurriness, diplopia, or horizontal nystagmus of &lt;1 hour duration, with spontaneous resolution</td>
<td>≥1 episode of Grade 2 symptoms per week, or an episode of Grade 2 Sx lasting 1hour with spontaneous resolution by 4 hours or vertical nystagmus</td>
<td>Decrease in visual acuity, visual field deficit, or oculogyric crisis, or Grade 3 Sx which persist after dose reduction</td>
</tr>
<tr>
<td>Many of the symptoms in this category can be the result of CNS pathology, or alternatively can be an external (i.e., non-CNS) neuro-ophthalmologic disorder. Appropriate diagnostic investigations should be pursued.</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Myelopathy</td>
<td>None</td>
<td>None</td>
<td>None</td>
<td>Myelopathic/spinal cord symptoms, such as</td>
</tr>
<tr>
<td>HIV can cause spinal cord syndromes rarely in children. Other infectious agents can cause myelopathies as well. Alternative explanations should be sought</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Neuropathy/Lower Motor Neuroneopathy</td>
<td>None</td>
<td>Mild transient Paresthesia only</td>
<td>Persistent or progressive paresthesias, burning sensation in feet, or mild dysesthesia; no weakness; mild to moderate deep tendon reflex changes; no sensory loss</td>
<td>Onset of significant weakness, decrease or loss of DTRs, sensory loss in “stocking glove” distribution, radicular sensory loss, multiple cranial nerve involvement; bladder or bowel dysfunction, fasciculations, respiratory embarrassment from chest wall weakness. Grade 3 symptoms which do not resolve with dose reduction</td>
</tr>
<tr>
<td>Infectious agents other than HIV can precipitate a neuropathy and should be considered, especially CMV. Neuropathies which do not resolve after dose reduction or discontinuation should be pursued for alternative infectious or non-infectious etiologies, since drug-related neuropathies will usually resolve after dose reduction or drug discontinuation. It should be borne in mind that many subjects will worsen for up to one month after drug discontinuation prior to improvement (“coasting”). Abnormalities should be confirmed by nerve conduction studies (NCS) +/- electromyographic studies (EMG).</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Myopathy or Neuromuscular Junction Impairment</td>
<td>Normal or mild (&lt;2xN) CPK elevation</td>
<td>Mild proximal weakness and/or atrophy not affecting gross motor function. Mild myalgias, +/- mild CPK elevation(&lt;2xN)</td>
<td>Proximal muscle weakness and/or atrophy affecting motor function +/-CPK elevation; or severe myalgias with CPK elevation; or severe myalgias with CPK &gt;2 x N; Consider confirmatory EMG and/or muscle bx</td>
<td>Onset of myasthenia-like symptoms (fatiguable weakness with external, variable ophthalmoplegia and/or ptosis), or neuromuscular junction blockade (acute paralysis) symptoms(confirm with EMG); or Grade 3 symptoms which do not resolve on dose adjustment; confirm with muscle bx</td>
</tr>
<tr>
<td>HIV can produce a myopathy, and should be differentiated. Drug-induced myopathy can be accompanied by normal CPK levels. On occasion neuropathic or central weakness can mimic myopathic weakness.</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

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TABLE FOR GRADING SEVERITY OF  
PEDIATRIC (> 3MONTHS OF AGE) ADVERSE EXPERIENCES  
For  
Vaccine & Prevention Research Programs  
-continued-

<table>
<thead>
<tr>
<th>SYMPTOM</th>
<th>GRADE 1</th>
<th>GRADE 2</th>
<th>GRADE 3</th>
<th>GRADE 4</th>
</tr>
</thead>
<tbody>
<tr>
<td>Clinical symptoms not otherwise specified in this table</td>
<td>No therapy; monitor condition</td>
<td>May require minimal intervention and monitoring</td>
<td>Requires medical care and possible hospitalization</td>
<td>Requires active medical intervention, hospitalization, or hospice care</td>
</tr>
<tr>
<td>Laboratory values not otherwise specified in this table</td>
<td>Abnormal, but requiring no immediate intervention; follow</td>
<td>Sufficiently abnormal to require evaluation as to causality and perhaps mild therapeutic intervention, but not of sufficient severity to warrant immediate changes in study drug</td>
<td>Sufficiently severe to require evaluation and treatment, including at least temporary suspension of study drug</td>
<td>Life-threatening severity. Requires immediate evaluation, treatment, and usually hospitalization. Study drug must be stopped immediately and should not be restarted until the abnormality is clearly felt to be caused by some other mechanism that study drug.</td>
</tr>
</tbody>
</table>
APPENDIX VI: NEVIRAPINE TOXICITY: SUPPLEMENTAL TOXICITY TABLE FOR GRADING SEVERITY OF CUTANEOUS/SKIN RASH/DERMATITIS ADVERSE EXPERIENCES

USE THIS APPENDIX FOR GRADING CUTANEOUS/SKIN RASH/DERMATITIS ADVERSE EXPERIENCES

<table>
<thead>
<tr>
<th>GRADE 1</th>
<th>GRADE 2</th>
<th>GRADE 3*</th>
<th>GRADE 4*</th>
</tr>
</thead>
<tbody>
<tr>
<td>CUTANEOUS/SKIN RASH/DERMATITIS</td>
<td>Cutaneous/skin rash/dermatitis</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Erythema, with or without pruritis</td>
<td>A. Diffuse erythematous macular or maculopapular cutaneous eruption or dry desquamation with or without pruritis; OR typical target lesions without blistering, vesicles, or ulcerations in the lesions.</td>
<td>A. Diffuse erythematous macular or maculopapular cutaneous eruption or moist desquamation with or without pruritis together with any of the following constitutional findings considered related to study drug: 1. 5 x ULN AST, ALT or 2x baseline if baseline &gt; ULN. 2. fever, &gt;39°C. 3. blistering and/or vesiculation of cutaneous eruptions. 4. any site of mucosal lesions; OR B. angioedema; OR C. exfoliative dermatitis defined as severe widespread erythema and dry scaling of the skin, with generalized superficial lymphadenopathy, and with other constitutional findings such as fever, weight loss, hypoproteinemia possibly related to study drug; OR D. diffuse rash and serum sickness-like reactions defined as a clinical symptom complex manifested as fever, new lymphadenopathy, edema, myalgia, and/or arthralgia; OR E. diffuse cutaneous eruptions, usually starting on the face, trunk or back, often with prodromal symptoms plus one of the following: 1. cutaneous bullae, sometimes confluent with widespread sheet-like detachment of skin (&lt;10% body surface area), (Nikolski's sign); 2. two or more anatomically distinct sites of mucosal erosion or ulceration not due to another cause.</td>
<td>Diffuse cutaneous eruptions, usually starting on the face, trunk or back, often with prodromal symptoms plus cutaneous bullae with widespread sheet-like detachment of skin (&gt;10% of body surface area), (Nikolski's sign), (SJS/Toxic Epidermal Necrolysis (TEN) overlap syndrome; TEN)</td>
</tr>
<tr>
<td>B. Urticaria</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

*When a Grade 3 or 4 cutaneous/skin rash/dermatitis adverse experience is suspected, a Dermatology consult for photographs and biopsies is required.