Antiretroviral Therapy for Mitochondrial Toxicity in HIV-Infected Pregnant Women
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Abstract

Background: The use of antiretroviral therapy (ART) in HIV-infected women is crucial to restore and maintain the immune system and prevent HIV transmission during pregnancy, labor, delivery and breastfeeding. Furthermore, ART reduce the risk of Mother-to-child transmission (MTCT). Therefore, ART has been associated with mitochondrial defences that could induce pre-eclampsia, preterm birth, low birthweight, intrauterine growth restriction (IUGR), stillbirth and sudden infant death.

Objective: To evaluate the effect of antiretroviral therapy on mitochondrial defences in HIV-infected pregnant women.

Methods: We searched eligible studies in MEDLINE, Scopus and WHO Global Index Medicus. Included studies were assessing the effects of antiretroviral therapy on mitochondrial diseases in HIV infected pregnant women and HIV exposed infants. JTL searched eligible studies in different databases and both JLT and JLT critically appraised included studies.

Results: We found five observational studies with low risk of bias. All studies illustrated that ART increased the mean of mitochondrial defences. The results were statistically significant in all studies with P<0.05.

Conclusion: Mitochondrial lesions were very common in HIV infected pregnant women and HIV exposed infants. However, further investigations are needful to strengthen this evidence.

Keywords: Antiretroviral treatment; HIV; Infants; Pregnancy; Mitochondrial defences

Introduction

HIV prevalence in Sub-Saharan Africa is the highest worldwide, with 25.6 million people living with HIV in 2015 [1]; among them, 58% women are HIV infected [2]. MTCT is the highest in the world in Sub-Saharan Africa. The use of antiretroviral therapy (ART) in antenatal and postpartum period is crucial both for preserving maternal health and for PMTCT of HIV [3]. The effectiveness of ART regimens in reducing MTCT of HIV-infection and delaying disease progression has been demonstrated. Then, ART should be offered to all pregnancies as stipulated by the World Health Organization (WHO) [4]. However, systematic reviews have shown that ART in HIV-infected women has been associated with pre-clampsia, preterm birth, low birthweight, intrauterine growth restriction (IUGR), stillbirth and sudden infant death, particularly in Sub-Saharan Africa. Molecular medicine has illustrated that ART may cause severe mitochondrial dysfunction. A recent study ranks ART mitochondrial toxicities as follows (from the most to the less harmful): (ddC). didanosine (ddI)>stavudine (d4T)>Zidovudine (AZT)>lamivudine (3TC)>abacavir = tenofovir (TDF) [5]. In Sub-Saharan Africa, TDF/3TC based regimens are widely prescribed in HIV-infected pregnant women. This regimen could lead to additive effects and cause more mitochondrial dysfunctions. The prevalence of ART associated to toxicities is reported to be more than 47% and 27% for clinical and laboratory manifestations, respectively [6]. The commonest mitochondrial DNA deletion is the most common mtDNA deletion in mitochondrial dysfunction [6]. These negative effects depend on the capacity of the NRTI to inhibit DNA g-polymerase, the only enzyme devoted to the replication of mitochondrial DNA genome, inducing then a decrease in mtDNA copy number and quality, which, in turn, may finally cause mitochondrial defences [7]. As mtDNA encodes for respiratory chain enzyme subunits, a defect in oxidative phosphorylation (OXPHOS) may result because mitochondrial proteins (mitprotein) could be defected [5].

In fact, mitochondrial defences have been associated with increased rates of preterm delivery, stillbirth, IUGR, and sudden infant death [5-8]. Reviewing the literature, ART can induce subclinical transplacental mitochondrial lesions. Therefore, it is unknown whether HIV in pregnancy could injure mitochondria. Mitochondrial defences in a given organism could lead to the development of mitochonidiopathies [9].

Mitochondriopathies are classified as inherited or acquired (derived from toxic substances), with both sharing similar clinical consequences [10]. This scientific paper is focused on acquired mitochondrial defences due to ART affecting both mitochondrial DNA and proteins. The mutations responsible for genetic mitochondrial diseases can be present in both the nuclear or mitochondrial genome [10]. Genetic mitochondrial diseases are already present in 1 in 5000 newborns and 1 in 200 women may carry one of these deleterious mutations [11]. Studies have shown that ART highly increase mitochondrial mutations.

Objectives

To evaluate the effects of antiretroviral therapy on mitochondrial defences in HIV-infected pregnant women and HIV exposed infants.

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Methods

We systematically searched in MEDLINE, Scopus and WHO Global Index Medicus. We included observational studies assessing the effects of antiretroviral therapy on genetic mitochondrial diseases in HIV infected pregnant women and HIV exposed infants.

The criteria for considering included studies for this mini review were HIV pregnant women on ART and HIV exposed newborn. We assessed maternal mtDNA depletion and newborn mtprotein. We included observational studies comparing cases and controls. We considered all types of ART regimen in HIV pregnant women. No other restriction criteria were imposed in terms of inclusion or exclusion studies.

JLT searched eligible studies in different databases. JLT and JLT worked independently to review full-text versions articles. The reviewers independently extracted data for study characteristics, patient characteristics, and outcomes (unadjusted and adjusted associations). Observational studies were assessed with Newcastle-Ottawa Scale. The following domains were used for bias assessment:

1. Is the case definition adequate?
2. Representativeness of the cases
3. Selection of controls
4. Definition of controls
5. Comparability of cases and controls on the basis of the design or analysis
6. Ascertainment of exposure
7. Same method of ascertainment for cases and controls

Results and Discussion

We screened a total number of 217 studies, and included 5 studies; four studies have been conducted in Spain and one in Canada. All the included studies were observational studies among which four cross-sectional and one prospective cohort study. All of studies analyzed the occurrence of maternal mtDNA and mtprotein in infants.

We included 5 observational studies among which four cross sectional studies and one prospective cohort study [5,7,8,12,13] (Table 1). The results were reported narratively. All mitochondrial studies conducting in HIV-pregnancies have described an increased frequency of mtDNA depletions.

The first study [8] reported the mean (SE) depletion of 44.45 ± 3.77% with P<0.001 in HIV-pregnant women and the mean (SE) mt proteins were reduced in HIV exposed infants compared to the control group 48.79 ± 3.41% with P<0.001. The second study [9] found mt proteins were reduced in HIV-infected pregnant women, infants and foetus with respectively: 25.8%, 38.6% and 13.6% (P<0.05). The third study [10] reported the mean (SE) maternal mtDNA of 42.66 ± 5.94% and mt proteins of 12.82 ± 5.73%. The mtDNA mean (SD) in the fourth study [11] was reduced by 39.20% ± 2.78%. The last study [12] revealed that the mean (SE) mtdna/nDNA ratio was reduced by -18.0 ± 6.1. All the results were statistically significant (P<0.05). All included studies were low risk of bias from 6 to 7 on Newcastle-Ottawa Scale [13].

Conclusion

Summary considerations on study quality in general, the studies did not show major problems of selection bias, the majority use the same population (HIV pregnant women and HIV exposed infants) compared to pregnant women HIV negative and HIV non-exposed infants respectively, limiting then problems of comparability. Multiple linear regressions were used in all included studies, and then adjustment for confounding factors was adequate. Ascertainment of outcome has not major biases in relation to the assessment of mtDNA and mtprotein because those outcomes are ascertained through laboratory record. Losses to follow-up would not possibly represent a problem. In most studies, losses to follow-up were not reported and the possible risk of bias is not predictable. All included studies included double-NRTI based regimens. Among them, three studies included AZT contained regimens [5,7,8]. As explained above, AZT based regimens may cause severe mitochondrial deflections implying highly statistically significant results in AZT based regimens studies.

A potential limitation of our search strategy is that only few databases were considered. It is likely to associated publication bias. However, this is a brief research has confirmed the results of previous

| Study ID          | Study design                        | Settings                              | Participants                                        | Interventions               | Outcomes                                           | P-value     |
|-------------------|-------------------------------------|---------------------------------------|----------------------------------------------------|-----------------------------|----------------------------------------------------|-------------|
| Moren et al. [10] | Cross-sectional, exploratory and controlled study. | Hospital St Joan de Deu of Barcelona (Barcelona, Spain) | 35 HIV-infected pregnant women and 17 controls | AZT based regimens and other NRTI regimens | (mtDNA) depletion Mothers (44.45 ± 3.77%) Infants (48.79 ± 3.41%) | P=0.001 P<0.001 |
| Hema’ndez et al. [7] | Single-site, cross-sectional, controlled observational study | Hospital Clinic of Barcelona (Barcelona, Spain) | 27 HIV-infected and treated pregnant women, and 35 uninfected controls | NRTI contained regimens | Newborn mtprotein reduced: Maternal mtprotein: 25.8% Fetal mtprotein: 13.6% | P<0.05 |
| Hernandez et al. [5] | Single-site, controlled observational study | Hospital Clinic of Barcelona (Barcelona, Spain) | Pregnancy and delivery in 27 HIV-infected and treated pregnant women versus 24 uninfected pregnant controls | NRTI based regimen + protease | mtDNA reduced Maternal: 42.66 ± 5.94% Mtprotein reduced: 12.82 ± 5.73% | P<0.01 P<0.01 |
| Hernandez et al. [7] | Single-site, cross-sectional, controlled, and observational study | Hospital Clinic of Barcelona | 24 asymptomatic HIV–infected pregnant women and 32 uninfected pregnant controls | AZT based regimens | mtDNA reduced Maternal: 39.20% ± 2.78% | P=0.001 |
| Money et al. [13] | Prospective longitudinal observational cohort study | Oak Tree Clinic, a provincial referral centre in Vancouver, British Columbia (BC), Canada | (i) HIV+ women (N = 65) using CART during pregnancy (cART started pre-conception or during Pregnancy); (ii) HIV- women (N = 45). | AZT contained regimens and other NRTI based regimens | mtdna/nDNA ratio -18.0 ± 6.1 | P=0.003 |

Table 1: Included studies.
studies conducted on the effects of ART on mitochondrial diseases. Besides, we did not find any study conducted in Sub-Saharan Africa where HIV prevalence is the highest in the world.

In fact, ARTs are indispensable in the treatment and prevention of HIV infection. Until now, the use of ART during pregnancy is considered safe; therefore, there are still many concerns about maternal and foetal outcomes that are not clearly highlighted in clinical practice. Besides, certain ART regimens have more mitochondrial toxicities than others. By the way, specific ART regimens should be used in pregnancy. In addition, the timing of prescribing ART in pregnancy could lower mitochondrial toxicity. Then, mitochondrial toxicity associated to ART inducing newborn and maternal morbidity need further studies, particularly in Sub-Saharan Africa. This could establish whether HIV-pregnancy may be an additional risk for the onset of mitochondrial toxicity, because it is currently unknown.

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