The Effect of HIV/HPV on the Structure of the Placental Villi

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Abstract

A placenta obtained of patient with co-infection HPV/HIV antiretroviral therapy and serology negative for another illness was analysed with the purpose of demonstrate the effect of this infection on the structure of the placental villi. The features of patient, the diagnostic, methods used, exam of material with H&E stain and sampling was done according morphological criterion of the literature as written in a previous article. Villi with interrumped syncytium are seen communicating the stromal region with the intervillous space. A group of them are associated by fibrinoid deposition, cells of the cellular island lack of cytoplasm, severe changes are observed in stem villi, many placental ramifications are immature, zones of calcification are observed and numerous remainders of villi are noted. The presence of this viral combination has provoked a strong attack to the structure of the villi which put in danger the normal interchange of gases and nutrients in the maternal-fetal intervillous space.

Keywords: Co-infection HPV/HIV; Histopathological changes; Placental villi

Abbreviations: LIF: Leukemia Inhibitory Factor.

Introduction

The placenta acquires viral infections proceedings from the mother [1]. Extensive microscopic infarction, distal villous hypoplasia, increased syncytial knots, increased perivillous fibrin, stem villous thrombosis, avascular villi, villous stromal kariorrhesis and villitis of unknown etiology has been found in placentas from pregnancies in HIV-infected women [2].

In the cases where placental pathology findings are found among HIV-infectected and uninfected women the choioamnionitis, deciduitis, Hofbauer cell hyperplasia, villitis, villous stromal edema and intravillous hemorrhage are more prevalent in HIV-infected women [3].

Studies suggest that most perinatal immunodeficiency virus (HIV) type 1 infections occur at or near birth [4]. Others indicate that most transmission occurs in utero through transplacental transmission [5]. It is known that placental inflammation increases the risk of vertical HIV transmission [6].

If the placenta is damaged and the blood from the infected mother transfers into the blood circulation of the foetus as occurs in choioamnionitis there is increased HIV transmission risk. Might be that maternal infected cells travel across the placenta and from here the virus gets to the stromal vessels of the placental villi reaching the foetus. D’Costa, et al. found a non-inflammatory lesion in the pathology of the placenta in HIV infection as cytotrophoblastic hyperplasia and the placental disc did not show any significant decrease in dimensions [7].
Patterson et al presented data to suggest that Leukemia inhibitory factor (LIF) inhibits HIV-1 replication and is upregulated in the placentas of nontransmitters women. LIF, a member of the IL-6 cytokine family has their receptor (LIFR) in the trophoblast and some women that express lower quantities of LIF increase the risk of mother to infant transmission [8].

It would seem that HIV-1 transmission from an infected mother to her infant is the result of a complex interplay between virus and host in which genotypes of virus, mother and infant impact the risk of transmission [9].

Hofbauer cells sequester HIV-1 in intracellular compartments that can be accessed by HIV-1 specific antibodies in the stromal region; activity that may be overridden when a strong association between maternal human cytomegalovirus viremia promotes inflammation, chronic villitis and trophoblast damage provoking no placental protection, facilitating in utero transmission of HIV-1 [10].

Many years ago it was described that the few studies which have reported the histopathology of placentas from HIV seropositive women have failed to demonstrate either villositis or a consistent microscopic lesion [11].

By other hands HPV was associated with chorioamnionitis, lymphohistiocytic villitis, sclerosing villitis, villitis of unknown etiology, perivillous fibrin embracing disintegrated syncytiotrophoblast, fibrosis, vascular obliteration, aggregated villi and syncytium loss [12].

HPV16/18 was detected in 24.4% of placentas from term deliveries by PCR [13]. In a HPV family study this virus was found in mother, neonate and umbilical cord as HPV16, 6, 83 and 39 [14]. Transplacental transmission of HPV was considered when type-specific HPV concordance was found between the mother, the placenta and the newborn or the mother and cord blood in a study where allowed the identification of genotypes 6/11,16,18,31,33,42,52 and 58 [15].

To date, approximately 200 different genotypes of HPVs have been indentified; categorized in the high-risk(HPV16,18,31) and the low-risk types(HPV6,11).Infection with low-risk HPVs usually results in benign epithelial warts, while with the high-risk HPVs can lead to cervical cancer. Infection with HPV may result in death of trophoblasts, malfunction or malignancy and these changes can disrupt the integrity of the trophoblast and cause spontaneous abortions or preterm delivery. HPV infections are more abundant in human HIV-positive patients [16].

Material and Methods

Placenta was obtained of patient of low socioeconomic recourse with co-infection HPV/HIV, at 38 weeks, who had antiretroviral therapy by human acquired immunodeficiency syndrome using protocol of the group of clinical essay of pediatrics AIDS [17]. PACTG 076 according to sigla in English, indicating 3’Azido 3’deoxythymidine (AZT) as Connor, et al. during the third trimester of pregnancy [18].

The features of patients, their diagnostic, methods used, sampling, examination of placenta infected and control, were described according to morphological criterion of the literature as has been written in preliminar article.

Results

Placental villi were seen with interrupted syncytium suffering necrosis and being very thin. A great deal of them are absent of syncytium and the stromal region is related with the intervillous space (Figure 1). Villi that are very near and have lost part of syncytium tend to cellular fusion by means of fibrinoid deposition (Figure 2).

Numerous cells of Cellular Island were observed with cytoplasmic lysis (Figure 3).

**Figure 1:** The interruption of the syncytium is communicating stromal region with the intervillous space in some villi.400x.H&E.

Figure 2: A group of placental villi are associate by fibrinoid deposition.100x.H&E.

Figure 3: Cells of a cellular island have lost their cytoplasm.400x.H&E.

Figure 4: Severe degenerative changes are observed in the vessels of the two stem villi.100x.H&E.

Figure 5: Placental ramification present feature of immaturity (arrow).100x.H&E.

Figure 6: Mature intermediate villi have lost syncytium and lack of terminal villi.100x.H&E.

Figure 7: Distrophic calcification is noted in this zone.100x.H&E.
The stems villi were noted with changes in the tunica intima and media of blood vessels. Damaged endothelium and separate muscle cells were observed (Figure 4). Immature ramifications of placental villi persist with frequency in the observations which not correspond at 38 weeks of pregnancy (Figure 5). Short mature intermediate villi without terminal villi were found with degenerative changes at level of the syncytium (Figure 6). Regions of calcified placental villi can be observed (Figure 7). With frequency debris of stem villi and intermediate villi appeared in the intervillous space (Figure 8).

**Discussion**

The interruption of the syncytial membrane in presence of HIV/HPV can impede the pregnancy provoking death fetal Replication of HPV is increased in trophoblast cells of HIV patients without inducing an immune response [16]. Oncogenetic human papillomavirus cause squamous intraepithelial lesions and cancer of the uterine cervix in human immunodeficiency virus seropositive women [22].

Similar lesions are also observed in syncytium of placental villi under the influence of HIV. This virus can enhance the risk of HPV infection, promote their latent infection and favour their persistence [23]. We don't know for example, if HPV16 viral load is provoking the same effect that induces in human uterine cervix associate to cervical dysplasia. However, we in a preliminary study have found stem villi with fibrotic stroma and numerous fibrotic villi that lost their vessels which could to correspond with a process similar to a fibrodysplasia. The stromal tissue of the placental villi is substituted by fibrous tissue [21].

The loss of a region of syncytium in two villi or more that are very near provably produces in them develops of cytotrophoblasts that secrets matrix-type fibrinoid, which represent the glue that guarantees adhesiveness, the re-epithelialization of damaged villous surface by the viruses and the develops of a new villi [19].

The infection by HIV has a cytolitic effect and the infection by HPV has effect of vacuolization of the cytoplasm as vacuolants agents, both viruses have provoked the cytoplasmic lysis observed in cell island [21].

The changes observed in stem villi have contributed with the fibrous tissue noted which were not seen in control placenta. The stem villous thrombosis that occurs in these cases of infection by HIV associate to avascular villi has been inhibiting the normal develop of the placental villi showing an immature tree at term [2].

Zones of calclification have indicated the presences of severe degenerative changes provoked by the activity of HIV/HPV producing a strong destructive effect on the placenta.

Debris of placental villi which were seen in the intervillous space are expression of the increased viral attack.

The local spread of HPV from the genital tract may result in placental infection that induces cell death of the trophoblast, placental dysfunction, preeclampsia and spontaneous preterm delivery [22].

There is evidence to show that HPV types 11, 18 and 31 can replicate in trophoblast in vivo [23].

The impaired immunity in HIV positive patients will lead possibly to increased HPV infection contributing with an expansion of lesions in the structure of the placental villi as has been suggested in the uterine cervix [24].

HIV related immunodeficiency alters the relative carcinogenicity of HPV types. A lower fraction of invasive cervical cancer is caused by HPV 16 but the attributable fraction for HPV 18 is concomitantly higher [25].

We have not found these carcinogenetic effects in the placental villi.

Although has been presented that in seven placentas, n=7/12, had a high carcinogenic risk of types 16, 18, 52 and 58 [15].

**Figure 8:** Reminders of placental villi shows the destructive effect of these viruses.100x.H&E.
Infected maternal blood macrophages transmit the infection to placental trophoblasts. Transcytosis, endocytosis pathway, is the entry mechanism of these viruses to the placental villi. E5, E6, E7 oncoproteins of HPV are viroporins that generate pores in the syncytium promoting the apoptosis process. This could explain the syncytiotial interruption observed [26].

This higher aggressiveness of the HPV in patients with HIV due to progressive immunosuppression has been reflected in increased damage to the structure of the villi.

The zidovudine treatment could also affect the structure of the villi since this has effect on the damage to the mitochondria of the cells [18].

In conclusion, the presence of this viral combination has provoked a higher attack to the structure of the villi which put in danger the normal interchange of gases and nutrients in the maternal-fetal intervillous space.

References


