Treatment of Natural Killer (NK) Cell activity in unexplained recurrent Spontaneous Abortion

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ABSTRACT

Recurrent spontaneous miscarriage (RSM), defined as ≥2 clinically detectable pregnancy losses before 20 weeks of gestation with the same partner. Many studies have been suggested abnormal natural killer (NK) cell activity can be a high chance division linked with RSM. Aberrant expression of angiogenic cytokines during implantation window in women with RSA is one of the key factors that adversely affect endometrial development, as evidenced by the inadequate expression of various endometrial receptivity markers. This review concerns a brief introduction into the basics of blood vessel development as well as the regulatory mechanisms of this process. This review discussed to work out Intralipid could be consumed as a possibility treatment as a part of RSM. In conclusion Intralipid, bordering on LMWH can overthrow the level of NK cells, which has been accounted for to be estimated angiogenic factors and supportive for increasing RSA results in pregnancy. This review is aimed to produce an evidence-based guidance on treatment of unexplained recurrent spontaneous miscarriage.

Keywords: Recurrent spontaneous abortion, NK cells, Intralipid, LMWH, VEGF-A, VEGF-C, IL-2, IL4I1

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INTRODUCTION

Recurrent spontaneous abortion (RSA), which is defined as two or greater consecutive pregnancy losses in advance than the 20th week of gestation from the ultimate menstrual interval, occurs takes place in kind of 1% to five% of women at reproductive age[1]. Despite the fact that many known reasons of RSA together with anatomic (15%), infectious (1%–2%), hormonal (20%), immunological (20%), and genetic (2%–five%) had been recognized, a big some cases (approximately forty%-50%) do no longer have regarded reasons, and those instances are known as unexplained recurrent spontaneous abortion (URSA)[2]. It's been proposed that URSA belongs to an autoimmune sickness associated with the failure of fetal-maternal immunologic tolerance[3]. Regulatory T cells (Tregs) play a crucial function inside the induction of a privileged tolerant microenvironment at the fetal-maternal interface because the fetus has genetic material from the father and mother the maternal immune method ought to adapt to the presence of overseas cells if a being pregnant is to do[4]This is clearly understood despite the fact that there's now enormous evidence to propose that abnormal immune mechanisms are involved Survival of the allogeneic embryo inside the uterus relies upon on the upkeep of immune tolerance at the maternal-fetal interface[5]. The pregnant uterus is replete with activated maternal immune cells. How this immune tolerance is acquired and maintained has been a subject of excessive research.

The key immune cells that predominantly populate the pregnant uterus are herbal killer (NK) cells[6]. In regular pregnancy, those cells are not killers, but alternatively offer a microenvironment this is pregnancy compatible and supports healthful placentation. In placental mammals, an array of notably orchestrated immune elements to assist a success being pregnant outcome has been integrated. Immune disorders are also recognized as significant contributors to RSA [7]. Peripheral natural killer (NK) cells are an important part of the altered immune repertoire found in RSA[8]. Thus, CD3−CD16+CD56+ cells expressing NK cell-associated molecules include...
CD3–CD16++CD56+ cells, which are cytotoxic and express the killer immunoglobulin-like receptor (KIR) family, as do CD3–CD16+CD56++ cells with lower CD16+ expression, which have a lower cytotoxicity. This includes energetic cooperation between maternal immune cells, specifically NK cells, and trophoblastic cells[9]. This intricate system is required for placentation, immune regulation and to transform the blood delivery to the fetus. For the duration of the beyond decade, diverse forms of maternal immune cells had been thought to be concerned in move-communicate with trophoblasts and in programming immune tolerance.

Tregs have attracted a wonderful deal of attention in promoting implantation and immune tolerance past implantation[10]. However, what has now not been absolutely addressed is how this immune-trophoblastic axis breaks down at some stage in unfavorable being pregnant results, specifically early pregnancy loss, and in response to unscheduled irritation. Extreme research efforts have started to shed mild on the roles of NK cells and Tregs in early pregnancy loss[11], even though an awful lot remains to be unraveled on the way to completely symbolize the mechanisms underlying their negative pastime. An accelerated expertise of host-surroundings interactions that cause the cytotoxic phenotype of those in any other case pregnancy like-minded maternal immune cells is vital for prediction, prevention and remedy of pregnancy maladies, specifically recurrent being the pregnant loss[12].

VASCULAR ENDOTHELIAL DEVELOPMENT FACTOR (VEGF-A) AND (VEGF-C)

On this overview, we discuss applicable information from experimental and human fashions which can give an explanation for the being pregnant disrupting roles of these pivotal sentinel cells on the maternal-fetal interface but, vascular endothelial development factor (VEGF), is a most essential angiogenic cytokine[13]. VEGF regulates proliferation, differentiation, and survival of endothelial cells and complements vascular permeability In vitro and in vivo reports of activated NK cell adhesion to endothelial cells show off that VEGF promotes adhesion[14], whereas everyday fibroblast improvement component inhibits adhesion via the regulation of intracellular and vascular cellular adhesion molecules on tumor vasculature [15]. Whether or not or no longer NK cells unique angiogenic development motives or their receptors has no longer been referred to. We now show that moreover to VEGF-A and VEGF-C[14]. Enormously, uNK cells, gift inside the secretory phase of the cycle, explicit excessive levels of VEGF-C[16]. NK cells specific angiogenic development reasons and advise that uterine NK cells may also just play a fundamental function in endometrial angiogenesis and regeneration[17]. Circulating NK cell numbers and cytotoxicity were extensively investigated in patients with RSA[16], and high ranges of NK cells and cytotoxicity (NK) are viewed to make a contribution to URSA and infertility [18]. Previous studies found that an immemorative NK had an optimistic predictive rate of seventy-one % for a subsequent miscarriage in recurrent miscarriage patients, as while positioned subsequent with 20 % in women with a low NK [19]. Fig-1

Immune-centered remedy alternatives concentrated on RSA women with a high NK have been noted to be powerful for enhancing pregnancy outcomes in this populous [20]. treatment with Heparin in women with recurrent pregnancy loss and NK cell or NKT-like cellular increase has been advised as a chance-loose and priceless healing approach that’s related to high premiums of scientific pregnancy and are living births [21]. Moreover, as a blood product, immunoglobulin has some crucial side-consequences paying homage to transfusion-transmitted illnesses.

INTERLEUKIN-2 (IL-2) AND 4 INDUCED PROTEIN 1 (IL-4I1)

Successful pregnancy depends on the presence of trophoblast growth-promoting cytokines[22]. One subclass of human NK cells (CD56(bright)) constitutively expresses the high-affinity interleukin 2 (IL-2) receptor and produces immunoregulatory cytokines[23]. CD56(bright) NK cells are present in human lymph nodes and that endogenous T cell-derived IL-2, acting through the NK high-affinity IL-2 receptor, co-stimulates CD56(bright) NK cells to secrete IFN-gamma[24].

Thus, adaptive immunoregulators influence innate cytokine production, which in turn may influence the developing antigen-specific immune response. The previous study has shown an active interaction between innate and adaptive human lymphocytes and emphasize the importance of studying interactions between immune components to understand the immune response as a whole[25].

Human Th17 cells as a new subset of helper T cells have been attentive, a producer pro-inflammatory cytokines. However, it is still unknown how Th17 cells effect on pregnancy outcome.[26] IL-2 and interleukin-4 (IL-4), an L-amino acid oxidase (LAAO) are significant in the differentiation of T helper cells into subtypes T helper type-1 (Th1) and type-2 (Th2). Th17 cells exhibition reduced IL-2 production due to IL-4- induced gene 1 (IL4I1) up-regulation[27]. Similarly, IL-4-induced immune deviation as antigen-specific therapy for inflammatory autoimmune disease[28]. Additionally, IL-2 induces the production of IFN-gamma and the ability of IL-2 to induce pro-inflammatory cytokine[29][30]. Elevated maternal serum
levels of interleukin-2 soluble receptor-alpha (IL-2 sRalpha), tumour necrosis factor-alpha (TNF-alpha) and interferon-gamma (IFN-gamma) have been associated with pregnancy loss. [31]

**INTRALIPID AND HEPALRIN**

Intralipid, a 20% intravenous fat emulsion it is again and again used as a supply of fats and calories for patients requiring parental vitamin, has been discovered to modulate immune feature and help pregnancy effects in sufferers with RSA[32][33]. Despite the fact that, few reports were conducted associated with this topic, and as a result, more potential, randomized reports are should overview whether or not or not Intralipid® has the equal effects on NK try and can be used as an opportunity remedy. Our look at aimed to investigate whether or not or not Intralipid® has the equal effects on NK cell cytotoxicity[20].

Even as, we additionally goal to test whether or not or not reducing NK cell toxicity can beautify being pregnant outcomes with recognize to intralipids and immunoglobulins, respectively. Of be aware, Intralipid and NK cells had been reported to have opposite outcomes on trophoblast invasion. Thus, we moreover attempted to deal with adjustments within the in presence of NK cells to mimic the in vivo conditions. Some study has shown that Intralipid have effects in vitro on Th1/Th2 balance and have different modulating effects on the immune response. [34]. It is reported that in the presence of Intralipid, suggesting an interference with the binding of IL-2 to its receptor on these cells. It is conceivable that administration of Intralipid to preterm infants may interfere with the binding of IL-2 to the specific receptors on their activated lymphocytes, with a possible subsequent suppression of their immune response[35].

In recurrent spontaneous abortion the combination of low-molecular-weight heparin (LMWH) and low-dose aspirin is considered to be the treatment of choice[36]. Therapy of RSA is may eliminating the increased thrombotic state, LMWH appear to have additional qualities in preventing adverse pregnancy outcome[37] by their anti-inflammatory and proangiogenic properties and restored VEGF[38] secretion and MMP activity[39]. Aspirin, LMWH are ineffective to pre- vent pregnancy losses and the combination of these drugs have been used in patients with recurrent miscarriage and recurrent implantation failure[40]. Apart from its anti-thrombotic effects, which consist to prevent the endothelial cells adhesion to the endothelium; heparin can bind selectins and integrins and interfere with complement activation. Adaptation of the maternal immune response to accommodate the semi-allogeneic fetus is necessary for pregnancy success, and disturbances in maternal tolerance are implicated in miscarriage[41]. Recently,
attention has focused on the role of uterine NK cells as mediators in successful implantation and placental maturation[42]. It is thought that these cells exert their function through cytokine production rather than cytotoxicity. In fact, the latter function should be inhibited to prevent pregnancy failure. Notwithstanding the fact that spontaneous miscarriage is a typically localized process, it is of great interest to evaluate systemic changes that precede this process, in particular for future diagnostic purposes.

The pathogenesis of URSA remains unclear, and disorders of the immune system may be one of the major risk factors for URSA[43]. NK cells play an important role in human pregnancy, and the systemic regulation of NK cells contributes to reproductive success. Abnormal NK activity has been reported to have association with URSA[44], so therapies aim to decrease NK activity was proved to be useful to treat these patients. In fact, this confirms previous data which showed that high NK cell levels are associated with RSA and subsequent miscarriage[45].

In humans, unlike peripheral blood NK cells of the cytotoxic CD56dim CD16 phenotype, the pregnant uterus (decidua) is populated with specialized uNK cells that are of the non-cytotoxic CD56bright CD16 phenotype[46]. Whereas both cytotoxicity and NK cell numbers increased in the RSA[47]. Recent studies have hypothesized that the signature angiogenic machinery of uNK cells is responsible for their non-cytotoxic behavior during normal pregnancy and the non-cytotoxic phenotype of uNK cells is accomplished through uNK cell produced VEGF-C, VEGF-A, which enhances resistance to lysis of trophoblasts and endothelial cells. [48][49][50][51]

The VEGF-A plays an important role in angiogenesis [52] with many effects including endothelial cell proliferation, migration, increase in vascular permeability and maintenance of vessel fragility[53]. VEGF-C deals with growth of blood vessels and lymphatics.[54] VEGF-A and C binds with high affinity to two related receptor tyrosine kinases expressed on vascular endothelial cells.[55]

Intralipid is effective in the treatment of women experiencing reproductive failure who display elevated NK cell activity and cytokine such as IL-2[56]. Furthermore, Intralipids have been shown to stimulate the reticuloendothelial system and remove “danger signals” that can lead to pregnancy loss. Elevated levels of NK cell cytotoxicity have been linked to recurrent miscarriage and recurrent implantation failure. Whether Intralipid influences trophoblast invasion via stimulating NK PPAR-γ expression or trophoblast PPAR-γ expression is unknown and will be assessed in future studies.[57] In recent study have shown that Intralipid is able to prevent VEGF down-regulation in tissues[58].

CONCLUSION

In women with a history of recurrent spontaneous abortion, low pre-conceptional peripheral NK cell levels are indicative of a subsequent successful pregnancy. Similarly, maternal levels of angiogenic factors and cytokines have played a foremost role in URSA as well as normal pregnant and with subsequent live birth.

LMWH and intralipid can regulate NK cell activity during pregnancy. In recurrent pregnancy loss, the combination of low-molecular-weight heparin and low-dose aspirin is considered to be the treatment of choice. A complementary study demonstrated the beneficial effect of low molecular weight heparin (LMWH) on angiogenesis. Heparins/LMWH and intralipid appear to have qualities in preventing adverse pregnancy outcome by their anti-inflammatory and pro-angiogenic properties and restored VEGF secretion.

This review suggests that these molecules could be used as potential predictive markers of miscarriage in these women presenting with URSA during the first trimester of pregnancy. During pregnancy, URSA women have increased the level of anti-inflammatory and angiogenic factors, associated with the treatment of NK cell cytotoxicity with Intralipid as well as Heparin. Also, there is significantly increase the concentration of pro-inflammatory in women with a history of recurrent spontaneous miscarriage when compared with healthy women. Thus in URSA, diagnostic and therapeutic measures aimed at characterizing and modulating NK cell activity appear promising in the future.

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