Introduction

Recurrent pregnancy loss (RPL) is defined as the occurrence of 2 or 3 consecutive abortions prior to the 20th week of pregnancy [1,2]. The incidence of RPL is about 1–2%, affecting 5% of families who want to have a child [2]. So far, the known pathogenesis of RPL comprises only 50% of the causes, which include immune factors, endocrine factors, genetic factors, infection factors, metabolic abnormalities, anatomic abnormalities, and other unexplained factors [3]. We speculate that RPL results from various interactions of the aforementioned factors. At least 50% of patients who have undergone a complete evaluation remain clueless as to why they are unable to achieve pregnancy [3,4]. A comprehensive understanding of all possible causes of RPL is necessary, and in the current study, various methods have been applied to gather relevant information. This review focuses on the role of immune response, infection factors, growth factors, and vitamin D, and evaluates the application of proteomics and microRNA (miRNA) analyses in RPL pathogenesis.

Etiological factors for recurrent pregnancy loss with the perspective of proteomics

Proteomics is a useful tool for recognizing altered protein expression, as well as proteins involved in disease pathogenesis [5]. In a previous study, we applied 2-dimensional-gel-based proteomic tools and identified, for the first time, RPL-associated proteins in the follicular fluid of RPL patients [6]. Our results indicated that coagulation factors (fibrinogen γ and antithrombin) play an important role in maintaining
normal pregnancy [6]. To conduct a comparative proteomic study, blood samples from normal and RPL patients were analyzed in a later study, and these results suggested that inter-α-trypsin inhibitor heavy chain family member 4 expression is a useful biomarker [7,8]. In a recent study, Pan et al. [5] analyzed placenta samples from normal and RPL patients and found 314 differentially expressed proteins, all playing important roles in the growth of the embryo. Furthermore, a network analysis demonstrated that angiotensinogen affects embryonic development via the rennin-angiotensin system [9]. In addition, mitogen-activated protein kinase 14 (MAPK14) is involved in the MAPK signaling pathway, which in turn mediates embryonic responses and apoptosis [10]. Defects in prothrombin (F2) are a cause of thrombosis (THR) susceptibility, and THR can result in RPL [11]. Other proteins include lamin B1 (involved in the development and nuclear integrity of the mouse) [12], nuclear autoantigenic sperm protein (required for preimplantation development during the blastocyst stage in cattle) [13], and secreted protein acidic and rich in cysteine (SPARC). SPARC is a matricellular glucoprotein that promotes cardiomyocyte differentiation by promoting the expression of bone morphogenetic protein 2 (BMP2), and is an essential auxiliary factor for the up-regulation of NKX2.5 expression in cardiomyocytes, mediated by BMP2 [14].

Parietal endoderm-secreted S100A4 is a calcium-binding protein belonging to the EF-hand family. In vitro studies have shown that it influences early cardiomyocyte differentiation and proliferation by increasing NKX2.5, mef2C, and mhcα expression [15]. RNA polymerase II prolonged by the yeast transcription-export (TREX) complex promotes gene expression by combining RNA processing factor and nuclear RNA output factor [16]. TREX consists of THO subcomplexes containing Hpr1p, Tho2p, Mft1p, and Thp2p proteins [17]. The functional linear homogen of Hpr1p in metazoa is Thoc1, Hpr1p, or p84 [18]. Thioredoxins are small redox proteins that regulate the activities of nuclear factor (NF)-κB and activator protein 1, mediate the antioxidant properties of peroxidase, and participate in early embryo formation [19]. Fibronectin and vitronectin are extracellular matrix protein components that promote the endoderm differentiation of human embryonic stem cells through interaction with integrin α5 and integrin α [20].

**DNA methylation in recurrent pregnancy loss**

Epigenetics and epigenetic research are involved in investigating hereditary gene expression or cell phenotype alterations through certain mechanisms, without changing the DNA sequence [21]. DNA methylation is one of the major epigenetic modifications, and plays important roles in embryonic implantation and development [22]. Abnormal DNA methylation is associated with miscarriage, preeclampsia, abnormal embryonic development, and birth abnormalities [23-26]. Among the 539 differential methylation regions (DMRs) found in RPL patients, p53 and SP transcription factors are recruited in the CAMP-responsive element binding protein 5 (CREB5) DMR by CREB5 hypomethylation, which subsequently increases CREB5 expression [27]. Notably, knock-down of CREB5 results in increased levels of tumor necrosis factor (TNF-α) and decreased levels of interleukin (IL)-10, and enhances the expression of NF-κB and p-NF-κB in monocytes, thereby causing immunosuppression [28]. In addition, CREB5 methylation and expression are regulated by IL-6 levels [29]. CREB5 also plays a key role in RPL pathogenesis. Forkhead Box P3 (FOXP3), specifically expressed in CD4+CD25+ T cells, is a human transcription regulator [30], and high expression levels of the FOXP3 gene are key to the development and function of T-reg cells [31]. Methylation levels of the FOXP3 promoter were reported to be higher in patients with RPL than in the control group, and conversely, FOXP3 protein levels were found to be lower in the RPL group than in the control group. The effect of FOXP3 on the differentiation of T-reg cells is the probable cause of immune tolerance failure and subsequent RPL [32].

**Novel genes and mutations in recurrent pregnancy loss**

Since innumerable genes are involved at every physiological step to ensure successful mammalian reproduction, mutations could be the causative factors of the molecular etiology of RPL [33,34]. Quintero-Ronderos et al. [35] reported that 27 coding variants in 22 genes are potentially related to the phenotype resulting in RPL. The 27 coding variants are associated with biological processes involved in cell adhesion-trophoblast endometrium interaction (TRO, CDH11, and CDH1), coagulation (THBD, F5, and FGA), extracellular matrix
remodeling (MMP10, MMP9, COL6A3, ADAMTS1, and TNC), angiogenesis (FLT1 and EPAS1), cell proliferation, differentiation, migration, apoptosis (LIFR, FGFR2, and BMP7), metabolism (AMN), immunological function modulation (IDO2, CR1, TLR3, and TRAF3IP1), and steroidal nuclear receptor activation (NCOA1) [35]. Nitric oxide is a transient gaseous lipophilic molecule, which has a variety of physiological regulatory functions in reproductive regulation. It plays important roles in oviductal functions, trophoblast invasion, trophoblast differentiation, blastocyst implantation, ovulation, fetal-maternal circulatory system establishment, and uterine blood flow regulation [36,37]. Shin et al. [38] reported that eNOS gene polymorphisms, variable number tandem repeat in intron 4, -786T>C, and 894G>T, are associated with risk of RPL. Azani et al. [39] reported that the eNOS-786C-4a-894G haplotype and -786 T>C polymorphism are associated with high risk of RPL. Through next generation sequencing analysis, Ryu et al. [40] found that the combination of stop-gain polymorphism OR4C16G>A and A-A (TAS2R46 and OR4C16) alleles was strongly associated with RPL prevalence.

**Immune response in recurrent pregnancy loss**

Embryos express paternal antigens that are foreign to the mother and may therefore be viewed as allografts [41]. Hence, the immune response of a pregnant woman plays a critical role in maintaining the pregnancy [42]. Immune responses in RPL are mainly divided into immune suppression and immune tolerance [41,43]. Antigens expressed on the surface of fetal or placental tissues possibly induce alloimmune responses by the mother, along with certain immunologic mechanisms that sustain the continuation of normal pregnancy. Type 1 T helper (Th1) and type 2 T helper (Th2) cells play important roles in immune responses, particularly in immune rejection and tolerance [43,44]. Based on a previous report, the normal range of the Th1/Th2 cell ratio should be less than 10.3 [45]. Moreover, Th2 cell dominance is important for the maintenance of normal pregnancy. Ota et al. [46] found that vitamin D deficiency increases the risk of RPL by promoting cellular immunity and autoimmunity. Vitamin D deficiency increases the expression levels of CD19+ B cells, CD56+ natural killer (NK) cells, Th1/Th2 ratio, NK cell cytotoxicity, and the presence of various autoantibodies in the peripheral blood [46]. Wegmann [47] proposed an immunotrophic theory wherein some cytokines are produced by maternal cells, which recognize the fetal antigens, promote proliferation of trophoblastic cells, and help in sustaining the pregnancy. Li et al. [48] reported that signaling by the T cell immunoglobulin (type I membrane protein) and mucin-containing protein 3 (Tim-3) in natural NK cells has an essential protective role. The number of Tim-3+ pNK cells (peripheral NK cells) transiently increased during the first trimester of pregnancy, as compared with Tim-3− NK cells [48]. Tim-3+ NK cells display immunosuppressive activity during early pregnancy by producing increased levels of anti-inflammatory cytokines, including transforming growth factor (TGF)-β1, IL-10, IL-4, and decreased pro-inflammatory cytokines, including TNF-α [48]. Tim-3 on NK cells is stimulated by its ligand galectin-9, leading to activation of signaling by the kinases (c-Jun N-terminal kinase [JNK] and protein kinase B). The abundance of Tim-3 on the surface of NK cells is enhanced through IL-4-signal transducer and activator of transcription 6 or progesterone signaling. Furthermore, the Tim-3 blockade disrupts immune tolerance and induces pregnancy failure (Fig. 1) [48]. Human amniotic epithelial cells play a...
potent regulatory role in the immune response by reducing TGF-β production and regulating the secretion of interferon (IFN) and IL-17 by T cells [49]. In autoimmune diabetes, it is generally believed that destruction of inflammatory β cells in the pancreas is due to the involvement of autoreactive T cells [50]. The chronic inflammation of pancreatic islets plays a critical role in disease development, and immune responses are the milestones in the mechanism of endothelial activation and atherosclerosis plaque development. This is demonstrated by the correlation between inflammatory marker levels and the risk of circulatory events in patients with diabetes mellitus (DM) [51]. Autoimmune antibodies and metabolic abnormalities are known to trigger RPL [50]. A slowly progressive form of type 2 diabetes, latent autoimmune diabetes in adults (LADA), has a prevalence rate of about 10–15% in patients diagnosed with type 2 diabetes [50]. Small studies have described that abnormal DNA methylation of CD4 T cells [52], T-reg cells [53], and NK cells [54] leads to loss of function of the pancreatic β cells by LADA, and this abnormal glucose metabolism eventually has adverse effects on pregnancy [50]. The glutamic acid decarboxylase (GAD) antibody is an immune marker in the early stages of type 1 DM [55]. During pregnancy, routine examination of the GAD antibody can help to improve glucose metabolism and the pregnancy outcome.

**Cellular roles of tumor necrosis factor-α in recurrent pregnancy loss**

Formerly called cachectin, TNF-α encoded on chromosome 6 is a Th1 proinflammatory cytokine [56]. TNF-α, one of 20 genes for the HLA system, is located on chromosome 6p21.3; it spans approximately 3 kb and has 4 exons. More than 80% of secreted proteins are encoded by the last exon [57]. Macrophages are the primary producers of TNF-α in pregnant women, and TNF-α is involved in the signal transduction pathway in the placenta [58]. TNF-α binds to its homologous receptor TNF-R1 to release slencer of death domains and to form complexes with receptors containing the adaptor proteins TNF receptor associated factor 2, fas-associated protein with death domain, TNF receptor type 1-associated DEATH domain protein, and receptor-interacting protein [59]. These adaptor proteins then recruit additional linchpin pathway-specific enzymes (for example, IκB kinase β and caspase-8) to the TNF-R1 complex, where they become activated and pulse-on downstream events, consequently resulting in apoptosis, JNK, and NF-κB activation [59]. In pregnancy, the immunological system plays critical roles both in the development of complications and in ensuring normal pregnancy development. A pregnancy is successful when the balance of Th1, Th2, Th17 cytokines, and T-reg cells works appropriately [60]. It is reported that TNF-α upregulates the level of programmed death-1 (PD-1) in monocytes, which binds with PD-L to promote the production of IL-10 in monocytes, and relies on IL-10 to inhibit the expansion of CD4 T cells [61]. Along with reduced IL-10 levels, TNF-α is increased throughout pregnancy and has been associated with preterm birth, miscarriages, pre-eclampsia, and fetal losses [62].

TNF-α plays a primary role in the immunological mechanism that regulates placentation and subsequent implantation in pregnancy outcomes. TNF-α is secreted by the innate immune cells and the placental cells. An appropriate balance between Th1 (mainly TNF-α, and Th17) and Th2 cytokines (including IL-10) is essential for achieving a good obstetric outcome. Conversely, several obstetric disorders, especially RPL, may occur due to an increase in Th1-dependent cytokines, particularly TNF-α [43]. TNF-α-targeted therapies therefore constitute a promising approach in improving or curing these disorders [63,64]. Hence, among the new and old immunosuppressive drugs, TNF-α blockers have become an effective tool for the treatment of pregnant women presenting with inflammatory- and immune-mediated diseases. A recent study by Alijotas-Reig et al. [65] revealed neither any maternal nor fetal major adverse reactions (including fetal malformations) to TNF therapy. In fact, the incidence of fetal malformations was 3% lower than expected.

**Effect of colony stimulating factor on recurrent pregnancy loss**

Colony stimulating factor (CSF)-1/M-CSF plays an important role in pregnancy. The expression of CSF-1/M-CSF significantly increases during pregnancy by 2-fold and 1,000-fold in serum and the uterus, respectively [66]. The CSF family members were detected in human cytotrophoblasts and syncytiotrophoblasts of the decidual stromal cells, endometrial glands, and epithelium of the placental and maternal cells, in addition to being located on local NK cells. CSFs are
involved in numerous reproductive functions, including ovulation, embryo implantation, placental growth, and embryo development [67]. During ovulation, CSF-3/G-CSF promotes ovulation through leukocyte attraction and activation [68]. After ovarian hyper-stimulation for in vitro fertilization, the cumulus oophorus cells synthesize and secrete CSF-1/M-CSF, and this synthesis is highly dependent on progesterone. In embryo implantation, a dominant Th-2 environment, suitable for the establishment and maintenance of a local tolerant immune environment, is required for successful implantation. CSF-3/G-CSF-mobilized Th-2 cells enhance IL-10 expression, which subsequently activates a signaling pathway leading to a dominant Th-2 environment [69]. For example, local uterine production of CSF-3/G-CSF may contribute to modulating the cytotoxicity of uterine NK cells by IFNγ and IL-18 production [70]. Uterine dendritic cells (DCs) are critical for activating the T cell response, thereby mediating maternal immune tolerance of the semiallogeneic fetus. Absence of GM-CSF, a known regulator of DCs, may impair the generation of T cell-mediated immune tolerance at the outset of pregnancy. This may contribute to the altered DC profile and dysregulated T cell tolerance evident in infertility and miscarriage [71].

**Effects of sodium-glucose transport proteins on recurrent pregnancy loss**

The cloning of intestinal glucose transporter sodium-glucose transport (SGLT)1 has been executed for over 30 years, and there have been several advances in physiological and medical research. The SLC5 gene codes for cotransporters for glucose and/or fructose (SGLT1, SGLT2, SGLT4, and SGLT5), inositol (sodium-myo-inositol co-transporter 1 [SMIT1] and SMIT2), short chain fatty acids (sodium monocarboxylate transporter 1 [SMCT1] and SMCT2), iodide (NIS), choline (choline transporter 1), and biotin (sodium dependent multivitamin transporter) [72]. The SLC5A1 gene for SGLT1 codes for a 72 kDa protein with 14 transmembrane helices (TM1–13) [73]. In both humans and mice, the SGLT1 transcripts were detected in endometrial cells, especially in humans during the early to mid-luteal phase [74]. The SGLT1 gene and protein were found in mouse endometrial glandular and surface epithelial cells, but not in Sglt1-null mice [75]. Endometrial glycogen, litter size, and pup weight were all lower in the Sglt1-null mice than in wild-type mice. Similarly, in patients with RPL, the SGLT1 gene and protein expression were significantly lower during the implantation window as compared to control patients [76]. It was therefore concluded that SGLT1 deficiency in the human endometrium at implantation may predispose early pregnancy failure, obstetrical complications, and low fetal growth. Taken together, these results indicate that it may not be advisable to prescribe SGLT inhibitors to women of childbearing age.

**Endometritis associated with recurrent pregnancy loss**

Endometritis is subdivided into the categories of acute endometritis and chronic endometritis [77]. Acute endometritis is characterized by neutrophil infiltration into superficial endometrial epithelium, the uterine cavity, and the glandular cavity [77]. One of the pathological features of chronic endometritis is the infiltration of phagocytes into the basal layer of the uterus [77]. Endometritis is the outcome of various endometrial structural changes, resulting in the production of numerous inflammatory cells and inflammation, thereby affecting implantation of the fertilized eggs and eventually leading to abortion [78]. Kitaya and Yasuo [79] reported that 8–28% of patients with RPL were diagnosed with chronic endometritis. However, RPL resulting from chronic endometritis can be treated, and the successful pregnancy rate and live birth rate are very high. The pregnancy rate and live birth rate are 76.3% vs. 20% and 65.8% vs. 6.6% in patients with cured and uncured chronic endometritis, respectively [80].

Metwally et al. [81] reported that obesity or being overweight is an independent factor influencing endometrial protein expression, including haptoglobin, transthyretin, and β-globulin. As an inflammatory marker, haptoglobin is an important component of the extra embryonic matrix, which may play a role in blastocyst implantation [81].

**Endocrine effects on recurrent pregnancy loss**

Polycystic ovary syndrome (PCOS) is a disease accompanied
by elevated plasma androgen levels, affecting approximately 5–10% of women of childbearing age [82]. High levels of androgen result in increased abortion rates [83]. Ishikawa cells were treated with high concentrations of androgen and physiological concentrations of androgen. Eight up-regulated proteins and 10 down-regulated proteins were subsequently identified in the high androgen group. One of these proteins was the cyclin-dependent kinase inhibitor 2a; decreased levels of protein expression led to reduced Ishikawa cell migration, invasion, proliferation, and Jar spheroid attachment. These results indicate that proteins associated with PCOS may result in RPL [84]. Luteal phase deficiency (LPD) refers to the inability to maintain physiological levels of progesterone during pregnancy and to enable normal embryo implantation and growth [85]. Progesterone is secreted in a pulsating manner over a short period of time. It is therefore not possible to design an experiment to compare the level of progesterone in the blood between healthy patients and patients with RPL [86]. However, Meresman et al. [87] found that caspase-3 expression is higher in endometrial biopsy samples from women with both RPL and LPD than from women without LPD. Embryonic trophoblast cells secrete human chorionic gonadotropin (hCG), which maintains the corpus luteum function in early pregnancy and plays a key role in implantation and endometrial receptivity [88]. A retrospective cohort study reported that miscarriage rate and time to pregnancy decreased in the 45 pregnant women who underwent 135 monitored cycles without hCG support, compared to 53 pregnant women who underwent 142 cycles with a single injection of mid-luteal hCG [89].

**Vitamin D associated with recurrent pregnancy loss**

Vitamin D is required to maintain normal human physiological activities. It affects not only bone metabolism, but also regulation of cell proliferation, differentiation, and apoptosis [90]. Vitamin D is expressed in the placenta, decidua, and ovary, and may be involved in the synthesis and secretion of hormones, thereby influencing the pregnancy outcome. Vitamin D receptor is also expressed in endometrial stromal cells and affects embryo implantation. Vitamin D is a newly discovered neuroendocrine regulatory factor affecting pregnancy, which is involved in a low inflammatory response. The fact that vitamin D participates in inflammation and maintains genomic stability in women is of interest [91,92].

1,25(OH)2-vitamin D3 (1,25(OH)2D3), an activated vitamin D, inhibits the number and function of CD4 cells and induces CD8 cell expression. 1,25(OH)2D3 inhibits the proliferation of Th1 cells and suppresses the secretion of IFN-γ, IL-2, and TNF-α. In contrast, 1,25(OH)2D3 induces Th2 cells to secrete cytokines including IL-4, IL-5, IL-6, IL-9, IL-10, and IL-13 [93-95]. In addition, 1,25(OH)2D3 inhibits IL-6 and IL-23 secreted by Th17 cells, and promotes the differentiation and expansion of T-reg cells. Expansion factor Fox3 activates T-reg cells. 1,25(OH)2D3 inhibits the maturation and stimulation of DC cells, which negatively affects the proliferation of B cells, plasma cell differentiation, and immunoglobulin production. Thus, the abnormal immune response of maternal and fetal surfaces subsequently results in RPL (Fig. 2) [96].

**Cellular roles of microRNA in recurrent pregnancy loss**

The noncoding RNA molecules belonging to the miRNA family are composed of 21 to 24 nucleotides. These bind to complementary sequences on target messenger RNA, promote mRNA degradation or inhibit translation, and play a
negative role in gene expression [97,98]. More than 1,500 miRNA target more than 5,000 human genes, which may regulate normal pregnancy. Parveen and Agrawal [99] found that gene polymorphisms of miR-146aC>G (rs2910164), miR-149T>C (rs2292832), miR-196a2T>C (rs11614913), and miR-499A>G (rs3746444) are present in RPL patients. The miR-146a regulates FAS expression by binding to FAS mRNA, subsequently reducing apoptosis [100]. The increase of miR-146a levels regulated by NF-κB promotes the survival of mesenchymal stem cells [100]. AKT and E2F1 genes are targets of miR-149, promoting cell growth and cell cycle progression [101]. Jeon et al. [102] reported that 196a2T>C and miR-499A>G polymorphisms may be involved in RPL.

**Conclusion**

This review describes the pathogenesis of RPL from multiple perspectives, such as proteomics, DNA methylation, gene mutations, the immune response, cytokines, endometritis, endocrine, vitamin D deficiency, and miRNA. Repeated abortions can occur due to a single factor in some RPL patients, while a variety of factors are involved in RPL pathogenesis. Although there are many questions that have yet to be answered, a profound understanding of the known etiological factors for RPL may provide translational implications towards therapeutic strategies.

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**Conflict of interest**

No potential conflict of interest relevant to this article was reported.

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