Role of Inflammatory Cytokines, Growth Factors and Adipokines in Adipogenesis and Insulin Resistance

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Abstract— Obesity, manifested by increased adiposity, represents a main cause of morbidity in the developed countries, causing increased risk of insulin resistance and type 2 diabetes mellitus. Recruitment of macrophages and activation of innate immunity represent the initial insult, which can be further exacerbated through secretion of chemokines and adipocytokines from activated macrophages and other cells within the adipose tissue. These events can impact adipogenesis, causing dysfunction of the adipose tissue and increased risk of insulin resistance. Various factors mediate adiposity and related insulin resistance including inflammatory and non-inflammatory factors such as pro and anti-inflammatory cytokines, adipokines and growth factors. In this review we will discuss the role of these factors in adipogenesis and development of insulin resistance and type 2 diabetes mellitus in the context of obesity. Understanding the molecular mechanisms that mediate adipogenesis and insulin resistance could help the development of novel therapeutic strategies for individuals at higher risk of insulin resistance and type 2 diabetes mellitus.

Key Words: cytokines; adipokines; growth factors; adipogenesis; IR.

INTRODUCTION

Obesity represents the fourth most frequent cause of morbidity in the developed countries according to the world health organization (WHO) reports [1, 2]. As obesity becomes more prevalent, the global threat of diabetes, particularly type 2 diabetes mellitus (T2DM), is increasing. It is estimated that more than 500 million people are expected to develop T2DM by 2030 [1, 3, 4]. Although several factors contribute to the increased incidents of diabetes, excess body fat [2] and abdominal obesity [5] are thought to constitute the most important risk factors for the development of T2DM. These risk factors have been directly linked to decreased physical activity due to changes in lifestyle, and increased consumption of food containing high fat [2, 5, 6]. At the molecular level, two major factors are associated with obesity‑induced T2DM: insulin resistance (IR) [7] and inflammation [8–10]. People with T2DM usually manifest highly active immune response with increased levels of inflammatory factors in their bodies. In early 1990s, tumor necrosis factor alpha (TNFα) was found to be the major inflammatory factor secreted by the immune cells. Later other cytokines were found to play an important role in suppressing insulin
signaling pathway and producing IR [11]. Additional factors with no direct inflammatory effect were also found to increase the risk of IR and T2DM. These included various growth factors and adipokines [12, 13]. One of the underlying mechanisms linking these factors with IR in obesity is the process of adipogenesis that involves generating fat cells from their precursors [10, 14]. In this review we will discuss the role of inflammatory cytokines, growth factors and adipokines in adipogenesis and development of IR and T2DM in obesity.

Adipogenesis and Adipose Tissue Formation

Adipocytes (fat cells) make up the majority of the adipose tissue, although the latter also contains preadipocytes (fat stem cells), macrophages, fibroblasts, blood cells, and endothelial cells [3, 15–17]. Adipose tissue is classified into three categories based on its morphology and metabolic functions, namely white, brown, and beige adipose tissue [18, 19]. Adipose tissue is present in various bodily compartments, with roughly 80% of total body fat being found under the skin (subcutaneous adipose tissue or SAT) and the remaining 20% around the digestive organs (mesenteric and omental adipose tissue, or OAT) [20]. Adipose tissue’s main function is to store energy in the form of fat (triacylglycerols). However, when the ratio of energy intake exceeds energy expenditure, the number of fat cells (hyperplasia) and/or their size (hypertrophy) rises, leading to obesity (Fig. 1). Adipose tissue also performs a variety of additional functions, including hormone synthesis, adipokines production, and immunological modulation. Metabolism, insulin sensitivity, and immunological function are all influenced by adipose tissue [21].

Several transcription factors regulate adipogenesis throughout late embryonic development and subsequently in adulthood, stimulating the differentiation of mesenchymal stem cells and preadipocytes to produce mature adipocytes [20]. A group of these factors and their effects are described in Fig. 2. Briefly, the process of adipose tissue formation (adipogenesis) involves three distinct phases: The first phase includes the commitment of mesenchymal stem cells (MSCs) into the adipogenic lineage under the influence of adipogenic enhancing signals such as insulin-like growth factor 1 (IGF-1) and insulin itself [22, 23]. This is followed by cell proliferation known as “mitotic clonal expansion phase”. At certain level of expansion, profound changes take place and the cells change from fibroblastic to spherical shape and the preadipocytes start expressing lipogenic genes including peroxisome proliferator-activated receptor-γ (PPARγ), CCAAT/enhancer-binding proteins (C/EBPs) family, CoA carboxylase (ACC) and adipocyte fatty acid binding protein (aP2), triggering adipogenic differentiation and formation of adipocytes [24].

![Fig. 1](image.png)

Fig. 1 Obesity-mediated changes in adipocyte numbers (hyperplasia) and size (hypertrophy).
Relationship Between Adipogenesis and IR

IR is a pathological condition that affects insulin metabolic pathways. Liver, muscle, and fat cells lose their ability to respond to insulin. Obesity, hyperglycaemia, and high blood pressure are among the underlying causes of IR in these tissues. Factors such as lifestyle, smoking, and family history may further increase the risk of IR and associated comorbidities such as diabetes, hypertension and cardiovascular disease [25, 26]. Inflammatory cytokines such as plasminogen activator inhibitor 1, interleukin (IL)-6, TNF-α, monocyte chemoattractant protein-1 (MCP-1), and leptin are signalling molecules generated by immune cells that regulate IR. TNF-α, IL-6, and MCP-1 are obesity linked inflammatory cytokines, particularly abdominal obesity. TNF-α and IL-6 can also trigger IR by inhibiting certain insulin signalling pathways involved in suppressing insulin signal transduction by serine phosphorylation of IRS1 and activation of JAK-STAT signalling pathway, causing a decrease in GLUT4 and IRS1 expression. Moreover, high levels of TNF-α and IL-6 are associated with increased levels of C-reactive protein (CRP), an acute inflammatory marker [27].

Impaired adipogenesis can contribute to the development of IR in target tissues [7]. Some mediators of lipid formation, including protein kinase C (PKC) and ceramides, can be activated by elevated lipid storage. These active lipid molecules enhance lipid accumulation and induce IR in a variety of target organs [28]. When energy intake increases, the storage capacity of the SAT becomes limited. This triggers the deposition of excess fat around internal tissues and organs, including OAT, skeletal muscles, liver, and heart [29]. Excessive lipid storage causes SAT hypertrophy, which leads to adipose tissue malfunction and increased tissue fibrosis [10]. This triggers further inflammatory processes, lipolysis and IR, leading to T2DM (Fig. 3)[10].

Role of Pro-Inflammatory Cytokines in Adipogenesis and IR

Inflammation is an adaptive immune response that is triggered by infection as well as tissue or cell injury or damage [30]. Inflammatory factors such as cytokines, chemokines, and vasoactive amines are activated by tissue resident macrophages and mast cells, which in turn trigger the onset of the inflammatory response [31]. Some inflammatory factors have pro-inflammatory properties, whereas others have anti-inflammatory properties. However some of these factors have both proinflammatory and anti-inflammatory actions [30]. The pro and / or anti-inflammatory effects depend on inflammatory condition/situation. After phagocytosis, resident macrophages secrete proinflammatory cytokines that recruit other immune cells and cause acute inflammation. Pro-inflammatory cytokines enhance inflammation cascade and boost the inflammatory reactions. Some of the known pro-inflammatory cytokines are interleukins (IL-1β,
Pro-inflammatory cytokines have been reported to have both inhibitory and stimulatory characteristics on adipogenesis. Among the proinflammatory cytokines, IL-1β, IL-6, IL-1F6, IL-15, IL-17, IL-18, IL-33, TNF-α, and OSM have been associated negatively with adipogenesis as they impair or reduce adipogenesis. However, other pro-inflammatory cytokines such as IL-7 and IL-34 have been reported to increase adipogenesis (Table 1).

Moreover, all of the listed (Table 1) pro-inflammatory cytokines, except for IL-1F6, IL-15, IL-18 and IL-33, induce IR. IL-15, IL-18, IL-33 have been reported to have protective characteristics against IR, while increasing insulin sensitivity. Whereas IL-1F6 has been reported to have no effect on IR. Table 1 lists pro-inflammatory cytokines expressed in adipose tissue, their effect on adipogenesis and association with IR and T2DM.

Among the pro-inflammatory and immunomodulatory cytokines, IL-6 represents one of the most studied factors associated with impaired adipogenesis and IR. IL-6 levels are higher in obese insulin resistant individuals compared to BMI-matched insulin sensitive counterparts [32]. Elevation in IL-6 levels is an indication of obesity related IR and has been positively associated with hyperplasia of adipose tissue [59]. IL-6 also plays an important role in hepatic IR [60] and as a signaling molecule that inhibits adipogenesis [32–34]. Furthermore, IL-6 can act as an immunomodulator in various diseases such as multiple sclerosis and Covid-19 infection as indicated recently [61, 62]. TNF-α is another important player in obesity-associated adipose tissue dysfunction. The anti-adiogenic properties of TNF-α are mediated by the function of its receptor 1 (TNFRI) [63]. Due to increased levels of mitogen-activated protein kinase 4 (MAP4K4), which is involved in TNF-α signaling pathway, the number of preadipocytes undergoing differentiation in the abdominal subcutaneous tissue is reduced, resulting in hypertrophic fat cells in association with obesity. This suggests an inverse relationship between lipid storage and proinflammatory capacity [35–37]. Moreover, reports have indicated that although the lipid storage capacity remains unrestricted by TNF-α in mature adipocytes, the expression of the insulin signaling intermediates
are downregulated, hindering insulin-mediated glucose uptake [63].

Another pro-inflammatory cytokine that plays an important role in obesity-associated impaired adipogenesis and insulin resistance is IL-1β. Macrophage-derived IL-1β represents an important anti-adipogenic factor that is associated with IR. High concentration of IL-1β can inhibit adipocyte differentiation, however it is not the only component of macrophage-derived conditioned medium that induces the anti-adipogenic activity [38–40]. IL1-β is upregulated in adipose tissue of obese individuals with IR during the development of IR in adipose cells [64]. IL-33, on the other hand, plays an important protective role during the development of adipose tissue associated inflammation in obesity, although obesity decreases the protective activity of IL-33 in adipocytes [41–43]. Hyperinsulinemia and IR were reduced after treatment with IL-33 [65]. The therapeutic administration of IL-33 leads to several anti-obesity benefits, including the reversal of visceral adipose tissue associated inflammation and reduction of IR [65].

Table 1 Pro-inflammatory Cytokines in adipose tissue and their role in adipogenesis and IR

| Pro-inflammatory cytokines | Expression within the adipose tissue | Effect on adipogenesis | Relation to IR and T2DM |
|---------------------------|--------------------------------------|------------------------|------------------------|
| IL-6 [32–34]              | Preadipocytes, monocytes/macrophages | Impairs adipogenesis   | Induces IR             |
| TNFα [35–37]              | Monocytes/macrophage, stromal vascular cells, adipocytes | Impairs/inhibits adipogenesis | Induces IR |
| IL-1β [38–40]             | Monocytes/macrophage, adipocytes      | Impairs adipogenesis   | Induces IR             |
| IL-33 [41–43]             | Adipocytes, preadipocytes, endothelial cells, fibroblast-like reticular cells, necrotic cells, cells under stress | Reduces/impairs adipogenesis | Reduces/protects against IR (increases insulin sensitivity and tolerance) |
| IL-18 [43, 44]            | Stromal vascular cells, macrophages, dendritic cells, epithelial cells, endothelial cells | No reported effect     | Induces/increases insulin sensitivity |
| IL-15 [45–47]             | Adipocytes, stromal vascular cells    | Inhibits/impairs adipogenesis | Induces/improves insulin sensitivity |
| IL-34 [48]                | Adipocytes, stromal vascular cells    | Induces/enhances adipogenesis | Induces IR             |
| IL-7 [49, 50]             | Stromal vascular cells                | Induces/enhances adipogenesis | Induces IR             |
| IL-1F6 & IL-1F8 [51]      | Stromal vascular fraction             | Impairs adipogenesis   | No reported effect     |
| OSM [52–54]               | Stromal vascular fraction, macrophages | Inhibits/impairs adipogenesis | Induces IR             |
| IL-17 [55–57]             | T helper cells, γδ T cells            | Inhibits/impairs adipogenesis | Induces IR             |
| IFN-α [58]                | Fibroblasts, monocytes                | Impairs adipogenesis   | Induces IR             |

Important role early in the inflammatory cascade process [66]. Phosphorylated Akt is increased by IL-18, while phosphorylated P38 MAPK is downregulated. In obesity and diabetes, higher serum IL-18 levels may be a compensatory response to IR [43, 44]. IL-15 is another pro-inflammatory cytokine that directly reduces adipogenesis by upregulating calcineurin [67]. In the absence of IL-15, fat formation in white adipose tissues is reduced, and lipid use is increased by adaptive thermogenesis [45–47]. In addition, IL-15 increases inflammation in adipose tissues, which may contribute to chronic inflammation and obesity-related metabolic syndrome [46]. IL-34 serum concentrations are greatly elevated in obese patients, regardless of their diabetes status. IL-34 levels in the blood are strongly and positively associated with IR-related metabolic parameters [48]. IL-7 is involved in the induction of adipogenesis and IR in response to a high-fat diet [50]. IL-7 modulates adipose tissue mass through a lymphocyte-independent mechanism, while immune cells involved in white adipose tissue inflammation relay its protective role on glucose homeostasis [50].

In mature adipocytes, IL-1 family member 6 (IL-1F6) and IL-1 family member 8 (IL-1F8) can stimulate inflammatory gene expression. IL-1F6 reduces PPARγ expression, which may result in a decreased adipocyte
development, implying that this cytokine has metabolic effects [51]. Recent data has indicated that OSM is produced by immune cells in white adipose tissue and its levels are dramatically increased in obesity and T2DM [52–54]. OSM has a paracrine effect on adipocytes, generating a proinflammatory phenotype in adipose tissue [68]. By modulating C/EBP activity, OSM slows the initiation of terminal differentiation of adipocytes via the Ras/ERK and STAT5 signaling pathways [53].

IL-17 inhibits the expression of several pro-adipogenic transcription factors, such as PPARγ and C/EBPα [55]. Hence, adipogenesis is suppressed by IL-17 due to the combined action of transcription factors that govern adipocyte differentiation [55–57]. Furthermore, reports have suggested that IL-17 acts as a negative regulator of adipogenesis and glucose metabolism, delaying the onset of obesity [56] similar to other pro-inflammatory cytokines such as IL-1β and TNF-α. IFN-α inhibits lipid formation and the expression of adipogenesis-related genes. During early stages of adipogenesis, IFN-α suppresses adipocyte development. Moreover, IFN-α regulates the production of CDK2 and p21 that stops the cell cycle. Furthermore, IFN-α-induced STAT1 phosphorylation inhibits adipocyte development [58].

**Role of Anti-Inflammatory Cytokines in Adipogenesis and IR**

In contrast to pro-inflammatory cytokines, anti-inflammatory cytokines prevent inflammatory reactions and control the pro-inflammatory cytokines responses. They represent immunoregulatory molecules that control the pro-inflammatory cytokine response by acting in concert with specific cytokine inhibitors and soluble cytokine receptors to regulate the human immune response. Table 2 lists some of the anti-inflammatory cytokines, their expression in adipose tissue and effect on adipogenesis and modulating insulin sensitivity. These include IL-1 receptor antagonist (IL-1Ra), IL-4, IL-5, IL-10, IL-11, IL-13, TGF-β.

The anti-inflammatory IL-1Ra operates by suppressing the effects of IL-1. Obese individuals have significantly higher serum levels of IL-1Ra [81] that are associated with increased body mass index (BMI) and IR, and is overexpressed in their white adipose tissues [69, 70]. Insulin sensitivity is reduced by IL-1Ra [69], which causes a muscle-specific decrease in glucose absorption. The link between the anti-inflammatory cytokine IL-4 and T2DM was previously established as IL-4 promotes insulin sensitivity, glucose tolerance, and lipid deposition inhibition in order to regulate glucose and lipid metabolism [71–73]. The majority of regulatory cells in lean adipose tissue maintain tissue homeostasis by excreting type 2 cytokines like IL-4, IL-5, and IL-13, which keeps adipose tissue macrophages in an anti-inflammatory condition [82]. Furthermore, IL-5 deficiency causes impairment of eosinophil buildup in the visceral adipose tissues, leading to increased adiposity and IR [74, 75].

The white adipose tissues of obese insulin resistant individuals exhibit elevated IL-10 secretion by proinflammatory macrophages, which causes a negative effect on insulin sensitivity and fat cell metabolism [76]. IL-10 works on IL-10 receptor alpha (IL-Rα) in adipose tissue [83]. However, unlike its role in mice, IL-10 is suggested to have no effect on human adipocyte activity [76]. Studies in mice have shown that IL-10 inhibits diet-induced

| Anti-inflammatory cytokines | Expression in adipose tissue | Effect on adipogenesis | Relation to IR and T2DM |
|----------------------------|------------------------------|------------------------|------------------------|
| IL-1Ra [69, 70]            | White adipose tissue         | Impairs adipogenesis   | Reduces insulin sensitivity |
| IL-4 [71–73]              | Adipocytes, eosinophils, M2-like reparative macrophages | Inhibits lipid deposition | Promotes insulin sensitivity |
| IL-5 [74, 75]             | Visceral adipose tissue, eosinophils | Deficiency in Visceral fat promotes adiposity | Deficiency in visceral fat promotes IR |
| IL-10 [76, 77]            | Adipocyte progenitors, macrophages, leukocytes | Deficiency in subcutaneous fat promotes browning | Promotes IR in adipocytes and insulin sensitivity in skeletal muscle |
| IL-11 [78]                | Adipocytes                   | Inhibits adipogenesis  | Not documented          |
| IL-13 [79]                | Adipocytes                   | Not documented         | Protects against IR     |
| TGF-β [80]                | Adipocytes                   | Inhibits adipocyte development | Not documented          |
IR by reducing response of macrophages and levels of cytokines in skeletal muscle [84]. Furthermore, energy consumption and adipose thermogenesis are increased in mice missing IL-10. Deletion of IL-10 also protects mice from diet-induced obesity and triggers browning of mouse subcutaneous white adipose tissue. In this model, IL-10 alters chromat shape as well as C/EBP and activating transcription factor (ATF) occupancy [77].

The anti-inflammatory cytokine IL-11 belongs to the gp130 cytokine co-receptor-related family and is actively produced in differentiating cells in response to PGF2α stimulation. PGF2α inhibits adipocyte differentiation via an autocrine negative feedback loop mediated by IL-11, which controls adipogenesis via the STAT1 transcription factor’s crucial actions [78]. The anti-inflammatory cytokine IL-13 is produced in the adipose tissue of obese patients, primarily from adipocytes. The IkB kinase (IKK) stimulates the synthesis of proinflammatory factors in adipocytes, but it also increases the formation of IL-13, which has a specific protective impact by lowering adipose tissue inflammation and IR [79]. Many elements of development, including adipogenesis, are regulated by members of the TGF-β superfamily. TGF-β and activin A inhibit adipogenesis, whereas the rest of the growth factors exhibit positive effects on adipogenesis. On the other hand, FGF21 and TGF-β induce insulin sensitivity while the rest promote IR.

### Role of Growth Factors in Adipogenesis and IR

Growth factors are biologically active molecules secreted in the body, which can affect cell growth and promote mitosis. They can cause altered gene expression by affecting various signal transduction pathways [85]. Several growth factors that are either protein (over 50 amino acid residues) or peptides (2–50 amino acid residues) exhibit high affinity for specific receptors on the cell surface. The target receptor cell surface are mainly plasma membrane-bound proteins that show tyrosine kinase activity. Example of growth factors include granulocyte–macrophage colony-stimulating factor (GM-CSF), vascular endothelial growth factor (VEGF), epidermal growth factor (EGF) and its receptor (EGFR), and platelet-derived growth factor (PDGF). Furthermore, some hormones that affect the cell growth such as estrogen and progesterone are considered as growth factors. Recent literature has shown that growth factors are essential for various physiological function such as wound healing and cancer among others [86]. Table 3 summarizes some growth factors that are expressed in adipose tissues and their impact on adipogenesis and relation to IR and T2DM.

Among the listed (Table 3) growth factors, EGF receptor (EGFR) and TGF-β inhibit adipogenesis, whereas the rest of the growth factors exhibit positive effects on adipogenesis. On the other hand, FGF21 and TGF-β induce insulin sensitivity while the rest promote IR.

The suppression of EGFR activity reduces adipogenesis and Akt phosphorylation in adipose-derived stem cells, but only the action of FGFR-1 decreases adipogenesis and Akt phosphorylation, whereas ErbB2 inhibition has the opposite effect. Furthermore, ErbB2-mediated suppression of adipogenesis in adipose-derived stem cells requires EGFR activation [67], whereas the inhibition of EGFR signaling leads to increased longevity in diabetic nephropathy [68]. Obese individuals have higher VEGF-C and -D levels in their blood, which is linked to poorer lipid metrics. Neutralization of VEGF-C in the subcutaneous adipose tissue during the development of obesity improves metabolic indices and IR in mice. It has been revealed that the lymphangiogenic factors VEGF-C and -D have an unexpected function in the modulation of metabolic syndrome-related adipose tissue inflammation [87]. Increased VEGF-C levels are linked to metabolic degradation and the development of IR. Blocking

### Table 3 Growth factors in adipose tissues and their role in adipogenesis and IR

| Growth Factors | Expression in adipose tissue | Effect on adipogenesis | Relation to IR and T2DM |
|----------------|------------------------------|------------------------|------------------------|
| EGFR (62, 63)  | Subcutaneous adipose tissue | Inhibits adipogenesis  | Induces IR             |
| VEGF-C [87, 88]| Adipose tissue, hepatic lipid | Increases adipogenesis | Induces IR             |
| CTGF [89]      | Preadipocytes                | Increases adipogenesis | Induces IR             |
| IGF-1 [90, 91]| Adipocytes                   | Increases adipogenesis | Improves IR            |
| FGF21 [92]     | Subcutaneous adipose tissue | Induces adipogenesis   | Enhances insulin sensitivity |
| TGF-β [93–95] | White adipose tissue         | Inhibits adipogenesis  | Induces insulin sensitivity |
VEGF-C in obese people may be a good way to prevent the onset of IR [88]. Connective tissue growth factor (CTGF) is found in abundance in preadipocytes and its expression is connected to body fat accumulation, as well as skeletal muscle and hepatic IR, with CTGF positive cells predominantly seen in fibrotic areas. The expression of CTGF in adipose tissue decreases in a stepwise manner as weight reduction progressed. In obese persons, elevated CTGF expression is linked to adipose tissue growth, adipose tissue fibrosis, and multi-organ IR [89].

In the pathophysiology of obesity, the growth hormone (GH)/insulin-like growth factor (IGF) system is linked. This system is engaged in the crosstalk between adipose tissue, liver, and pituitary, and both GH and IGF-I have direct effects on adipocyte proliferation and differentiation. This system appears to play a key role in visceral adiposity, and there is a rationale for targeting it in the treatment of visceral obesity induced by GH deficiency, metabolic syndrome, and lipodystrophies [90]. The increase in IGF-1 and the GH dosage were linked to changes in glucose metabolism following the start of GH therapy. Regardless of pubertal stage, all cases of impaired fasting glycaemia and/or impaired glucose tolerance identified after GH administration are reversible with dietary intervention and do not progress to diabetes mellitus [91]. Fibroblast growth factor 21 (FGF21) promotes the healthy growth of subcutaneous adipose tissue, which increases systemic insulin sensitivity. In insulin-sensitive obese individuals, serum FGF21 levels correlate with the volume of subcutaneous adipose tissue. Circulating FGF21 causes an increase in M2 macrophage polarization and upregulates adiponectin in subcutaneous adipose tissue. In obesity, increased levels of endogenous FGF21 act as a defensive mechanism against systemic IR [92]. Not only does the transforming growth factor-β (TGF-β) signaling pathway play a function in adipogenesis, but it also plays a role in the development process of IR. TGF-β partly reduces adipogenesis via the Smad3-dependent pathway. Smad3 is a complex regulator involved in adipose physiology as well as the etiology of obesity and T2DM, suggesting that it might be utilized to treat obesity and other relevant complications [93–95].

**Role of Adipsokines in Adipogenesis and IR**

The cytokines that are produced by the adipose tissue are called adipokines. Adipokines such as leptin, adiponectin, resistin and chemokine (C–C motif) ligand 2 can affect the insulin function and metabolism of lipids and glucose. Adipokines also have influence on the secretion of some hormones and chemokines. Adipose tissue expansion can lead to imbalance of adipokines, and this imbalance can lead to IR, metabolic syndrome, T2DM and cardiovascular disease. However, each adipokine has a different effect on the obesity and development of IR [96]. Table 4 summarizes adipokines in adipose tissues and their roles in adipogenesis and IR.

Among the listed adipokines, MCPIP1 and progranulin induce IR, whereas the remaining adipokines (Table 4) were shown to increase insulin sensitivity. Moreover, only MCPIP1 was shown to impair adipogenesis whereas the other listed adipokines exhibit enhancing effects on adipogenesis. Leptin is an adipokine produced by white adipose tissue in proportion to the size of fat depots. Leptin reduces body fat by suppressing appetite and raising energy expenditure. Leptin has an indirect effect on metabolism by altering sympathetic nervous

| Adipokines          | Expression in adipose tissue                                      | Effect on adipogenesis                   | Relation to IR and T2DM               |
|---------------------|-------------------------------------------------------------------|-----------------------------------------|---------------------------------------|
| Leptin [97]         | Adipocytes                                                       | Suppresses adipocytes proliferation     | Induces insulin sensitivity           |
| Omentin [98]        | Stromal vascular fraction (SVF) of visceral adipose tissue        | Induce adipogenesis                     | Regulates insulin sensitivity         |
| Adiponectin [99]    | Adipose tissue and endothelial cells                              | Enhances adipogenesis                   | Insulin sensitizing effect            |
| Vaspin [100]        | Visceral & subcutaneous adipose tissue                            | Enhances adipogenesis                   | Insulin sensitizing effect            |
| Apelin [101, 102]   | Adipose tissues and endothelial cells                             | Enhances adipogenesis                   | Increase insulin sensitivity          |
| MCPIP1 [103]        | Immune cells (macrophages/monocytes)                             | Impair adipogenesis                     | Induces IR                            |
| Progranulin [104, 105] | Adipose tissue, epithelial cells                                | Not documented                          | High levels correlate with IR         |
system activity or insulin sensitivity [97]. It affects adiposity by decreasing cell proliferation in white fat cells by generating inhibitory circulatory factors and contributing to sympathetic tone, both of which restrict cell growth. The stromal-vascular portion of visceral adipose tissue produces omentin. In adults and adolescents, obesity lowers omentin serum concentrations and adipose tissue secretion [98]. Although this adipokine is believed to control insulin sensitivity, its clinical significance requires further investigation.

Adiponectin is the most available peptide produced by adipocytes, and its deficit has been linked to

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**Fig. 4** Role of cytokines, growth factors and adipokines in adipogenesis and insulin resistance. The graph shows factors affecting adipogenesis and insulin resistance either in an opposite A or a similar B direction.
obesity-related disorders such as IR, T2DM, and cardiovascular disease. Apart from adipocytes, this adipokine can be produced by a variety of cells, including skeletal and cardiac myocytes, as well as endothelial cells. Adiponectin’s activities are mediated by adiponectin receptors AdipoR1 and AdipoR2. Adiponectin was suggested to protect against IR, diabetes, and atherosclerosis [99]. Vaspin (serpinA12) expression is positively linked with BMI and insulin sensitivity, and it improves glucose tolerance in vivo, suggesting a compensatory function in response to reduced insulin signaling in obesity [100].

Apelin is an adipocyte-produced hormone that plays an important role in energy metabolism. Through the PI3K/Akt and AMPK signaling pathways, apelin-APJ signaling promotes brown adipocyte development by boosting the production of brown adipogenic and thermogenic transcriptional factors. TNF-α suppression of brown adipogenesis is relieved by apelin. Adipocytes’ baseline activity is also boosted by apelin. Apelin is able to increase the brown-like characteristics in white adipocytes. The brown adipogenic and browning effects of apelin suggest a potential therapeutic route to combat obesity and related metabolic disorders. Apelin improves not only brown adipocyte differentiation and metabolic activity, but also white adipocyte browning. Apelin-APJ signaling increases browning of adipose tissue [101, 102].

Monocyte chemoattractant protein-1 induced protein-1 (MCPIP1) is an RNase that reduces the stability of transcripts that code for inflammatory proteins. MCPIP1 also plays a function in the control of adipogenesis in vitro by lowering the expression of critical transcription factors such as C/EBPβ. Recent studies have shown that MCPIP1 is an essential adipogenesis and adipocyte metabolism regulator [103]. The levels of circulating progranulin were related to BMI, HbA1c, IL-6, and TG in distinct manners. Recent data has indicated that T2DM patients and obese individuals have higher plasma progranulin levels, which is associated with glycolipid metabolism, chronic inflammation, and IR [104, 105]. Progranulin gene expression is elevated during adipocyte development and controlled by multiple inflammatory and metabolic stimuli in a gender, location, and cell-specific dependent manners.

CONCLUSION

This article reviews some of the most studied factors related to obesity-associated inflammatory response and insulin resistance, which play a critical role in the metabolic consequences of obesity. Various cells within the adipose tissue can secrete specific cytokines, adipokines and growth factors that cause dysfunction of the adipose tissue and impairment of insulin signalling (Fig. 4). The inflammatory environment associated with obesity triggers various inflammatory cascades of adipose tissues through activating specific kinases, which mediate these events. These endocrine and paracrine inflammatory cues within the adipose tissue can further trigger a systemic insulin resistant, inflammatory, and metabolic dyslipidaemia, causing increased risk of T2DM. Understanding these molecular events associated with reduced adipogenesis in insulin resistant obese individuals could help identifying novel therapeutic targets for individuals at higher risk of IR and T2DM.

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AUTHORS’ CONTRIBUTIONS

All authors have contributed to reviewing literature, writing the manuscript, reviewing and approving the final version of the manuscript.

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AVAILABILITY OF DATA AND MATERIALS

Not applicable.

DECLARATIONS

Ethics Approval and Consent to Participate Not applicable.

Competing Interests The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

Consent for Publication Not applicable.
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