Imbalanced Coagulation in the Airway of Type-2 High Asthma with Comorbid Obesity

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Abstract: Asthma is a common, chronic airway inflammatory disease marked by airway hyperresponsiveness, inflammation, and remodeling. Asthma incidence has increased rapidly in the past few decades and recent multicenter analyses have revealed several unique asthma endotypes. Of these, type-2 high asthma with comorbid obesity presents a unique clinical challenge marked by increased resistance to standard therapies and exacerbated disease development. The extrinsic coagulation pathway plays a significant role in both type-2 high asthma and obesity. The type-2 high asthma airway is marked by increased procoagulant potential, which is readily activated following damage to airway tissue. In this review, we summarize the current understanding of the role the extrinsic coagulation pathway plays in the airway of type-2 high asthma with comorbid obesity. We propose that asthma control is worsened in obesity as a result of a systemic and local airway shift towards a procoagulant and anti-fibrinolytic environment. Lastly, we hypothesize bariatric surgery as a treatment for improved asthma management in type-2 high asthma with comorbid obesity, facilitated by normalization of systemic procoagulant and pro-inflammatory mediators. A better understanding of attenuated coagulation parameters in the airway following bariatric surgery will advance our knowledge of biomolecular pathways driving asthma pathobiology in patients with obesity.

Keywords: asthma, obesity, coagulation, bariatric surgery, extrinsic pathway, inflammation

Introduction

Asthma is characterized by airway hyperresponsiveness (AHR), remodeling, and inflammation resulting in moderate-to-severe respiratory symptoms.1 Numerous cluster analyses have identified asthma as a heterogeneous disease consisting of several manifestations, with most patients categorized into two main endotypes: Type-2 high and Type-2 low, with varying phenotypic presentations.1–4 These endotypes are based on a predominance (high) or a scarcity (low) of eosinophilic and type 2 cytokine (interleukin (IL)-13, IL-4, and IL-5)-driven airway inflammation.

Type-2 high asthma further comprises two phenotypes. The first is typically characterized as early-onset (<12 years of age), allergen-sensitized asthma concurrent with elevated sputum eosinophilia (>1%-3%)5 and serum immunoglobulin E (IgE).4–6 The second Type-2 high asthma phenotype is typically characterized as late-onset (>12 years of age), non-allergic, eosinophilic asthma that does not display elevated serum IgE levels.7 Both Type-2 high asthma phenotypes are more responsive to inhaled and oral corticosteroids when compared to the Type-2 low endotype, with the early-onset allergic phenotype relatively the most sensitive to
corticosteroid therapy, and management of the late-onset non-atopic phenotype more reliant on oral corticosteroids.\textsuperscript{7} Type-2 high asthma with comorbid obesity is a significant healthcare problem due to changes in therapeutic success. Compared to lean Type-2 high asthma patients, patients with obesity are more likely to be hospitalized, use oral steroids, be admitted to the intensive care unit (ICU),\textsuperscript{8} and experience greater asthma severity through augmented small airway dysfunction\textsuperscript{9} and bronchial hyperresponsiveness.\textsuperscript{8} These trends strongly suggest that the development of obesity significantly impacts pulmonary function; however, no longitudinal studies have investigated the effect of weight gain on Type-2 high asthma pathophysiology. The increased resistance to corticosteroid therapy and limited alternative therapies for Type-2 high asthma with comorbid obesity\textsuperscript{10,11} places this population at increased risk of poor clinical outcomes.

A key feature present in both obesity and asthma is an imbalance in coagulation. Activation of the coagulation cascade occurs via two mechanisms: the intrinsic and extrinsic pathways. The intrinsic pathway begins with vascular injury, resulting in the exposure of endothelial collagen to plasma, causing activation of factor XII.\textsuperscript{12} The extrinsic pathway also begins as a result of tissue damage; whereby plasma leaks into the extravascular space. There, coagulation factor VII (FVII) complexes with tissue factor (TF), which is presented by various cell types,\textsuperscript{12} activating downstream zymogens, the inactive precursor of an enzyme which lacks catalytic capability. TF is expressed in perivascular cells,\textsuperscript{13} adventitial tissue,\textsuperscript{14} and epithelial cells,\textsuperscript{15} and is in greatest concentration in vital organs such as the brain, lungs, and kidney, allowing for rapid activation of the coagulation cascade following vascular injury.\textsuperscript{15,16}

The two coagulation pathways converge as the common pathway with the activation of factor X (FX), which becomes factor X active (FXa). Downstream processes result in the conversion of prothrombin to thrombin, which cleaves fibrinogen into fibrin fragments, the main component in clot structure. Once the clot has stabilized and tissue repair occurs, fibrinolysis—the breakdown of fibrin clots into fibrin degradation products (FDPs)—is initiated by activation of tissue plasminogen activator (tPA) and urokinase plasminogen activator (uPA). tPA/ uPA cleave the zymogen plasminogen, forming plasmin, the enzyme responsible for fibrinolysis. Figure 1 illustrates extrinsic pathway activation in the airway following injury to the airway epithelium.

In disease states, fluctuation in clotting and fibrinolytic factors results in improper coagulation. In pathologies such as Type-2 high asthma with comorbid obesity, a significant shift in this equilibrium exists, favoring the procoagulant state through increased production of coagulation factors.\textsuperscript{17} Additionally, these pathologies are characterized by increased anti-fibrinolytic mediators, thus prolonging clot time.\textsuperscript{17,18}

While coagulation has been studied separately in obesity and asthma, to date, no studies specifically evaluate how imbalanced coagulation within obesity impacts airway pathology in the Type-2 high asthma phenotype. Given the shift in equilibrium toward clot formation and hypofibrinolysis in both obesity and asthma, it is imperative to better understand coagulation in this setting. These studies would help advance our knowledge of asthma pathophysiology and promote therapeutic development for Type-2 high asthma with comorbid obesity.

Bariatric surgery is a promising therapy for improving asthma with comorbid obesity.\textsuperscript{19–21} Studies evaluating both asthma\textsuperscript{19,20} and coagulation\textsuperscript{22,23} report improvement of coagulant and inflammatory parameters following weight loss through conventional diet and exercise regimens or bariatric surgery, but minimal literature is available regarding the mechanisms by which attenuated coagulation improves Type-2 high asthma symptoms, lung function, airway remodeling, and quality of life for patients within this population.

The goal of this review is to present current knowledge of the extrinsic coagulation pathway and regulatory effects of the coagulation cascade on asthma pathobiology in Type-2 high asthma patients with comorbid obesity. The size and complexity of the coagulation cascades narrows the topic of discussion toward the extrinsic pathway due to its potential therapeutic targets. The intrinsic pathway and numerous autogenic loops in coagulation may also serve as potential targets to regulate asthma exacerbations and other pathologic features, but the breadth of regulators is beyond the scope of this review.

Given the procoagulant and hypo-fibrinolytic environment in both obesity and Type-2 high asthma, the coagulation cascade may serve as a link between the influences of increased adiposity and diminished pulmonary function or AHR in asthma. In an attempt to combat the challenges associated with the development of pharmacologic therapies for improper coagulation, we propose bariatric surgery as a possible therapy, with potential for return to normal coagulation processes and a decrease in
inflammatory and procoagulant mediators following surgery and/or weight loss.

Coagulation Imbalance in Type-2 High Asthma

Pro-Coagulation in Type-2 High Asthma

Type-2 high asthma is a well-characterized endotype, and the airways of patients with Type-2 high asthma show shifts in coagulation profiles resulting in a propensity towards coagulation and with decreased fibrinolysis. Recent evidence supports that the type-2 cytokine, IL-13, increases both the expression and release of TF from human airway epithelial cells, contributing to the procoagulant environment in the airway. Additionally, induced sputum obtained from severe eosinophilic asthma patients exhibits significantly elevated soluble TF compared to sputum from healthy controls. Brims et al noted that this group of severe asthma patients who used inhaled corticosteroids (ICS) had increased expression of plasminogen activator inhibitor-1 (PAI-1)—an anti-fibrinolytic mediator that interferes with activation of plasminogen by inhibiting tPA and uPA—, thrombin activatable fibrinolysis inhibitor (TAFI) which protects fibrin clots from lysis, and fibrin deposition, suggesting that ICS use contributes to airway coagulation and fibrosis in severe eosinophilic asthma.

Furthermore, in an ovalbumin-mouse model of allergic asthma, FVIIa deficiency resulted in improved asthma response seen through decreased lung eosinophilia and AHR. Additionally, a study by Ollivier et al found that TF-FVIIa augmented vascular endothelial growth factor (VEGF) production by human airway fibroblasts, promoting angiogenesis, a hallmark feature of airway remodeling in asthma.

Downstream activation of FX is also predictive of impaired airway physiology. In ovalbumin-allergen challenged mice, FX expression is significantly elevated compared to controls, and mice treated with fondaparinux (FPX), an inhibitor of FX, experience attenuated airway mucin production, AHR, airway thickening, and airway...
fibrosis. Additionally, FXa increases airway fibroblast proliferation and procollagen synthesis, contributing to airway stiffening and reduced lung compliance.

Progressing through the coagulation cascade, further zymogen activation contributes to worsening asthma. Asthma is associated with enhanced thrombin activation, contributing to increased fibrinogen cleavage while also binding with protease-activated receptors (PARs), driving inflammation. Platelet accumulation and fibrin deposition are further known to exacerbate asthma. In asthma, platelet accumulation in the airway contributes to bronchial hyperresponsiveness, bronchoconstriction, airway inflammation, and airway remodeling. Furthermore, FDPs contribute to the inflammatory and infiltration profiles characteristic of asthma. More specifically, the fibrin E fragment increases leukocyte invasion/migration and IL-1β and IL-6 production, while fibrin D dimers augment IL-1β, IL-6, and PAI-1 release by peripheral blood monocytes. During tissue repair, under homeostatic conditions, fibrinolysis facilitates clot breakdown and FDP clearance, potentially decreasing pro-inflammatory cytokine production and immune system activation.

Hypofibrinolysis in Type-2 High Asthma

Augmentation of a procoagulant environment in Type-2 high asthma is also met with hypofibrinolysis. The major direct inhibitor of fibrinolysis is PAI-1, which inhibits both tPA and uPA, prolonging clot presence at the injury site. Type-2 high asthma is associated with increased production of PAI-1, which is produced by numerous cell types in the lung, including macrophages, mast cells, fibroblasts, and bronchial epithelial cells and is strongly associated with impaired lung function. Furthermore, sputum from asthma patients and bronchoalveolar lavage (BAL) fluid from ovalbumin-sensitized rodents both exhibited a marked increase in PAI-1 relative to healthy and saline-treated controls. Administration of aerosolized tPA reduces AHR in ovalbumin-challenged mice, and aerosolized uPA in this same model results in improved AHR and subepithelial fibrosis.

Activated protein C (APC)—a collateral cascade initiated following thrombin activation that decreases the extent of clot formation—is reduced in Type-2 high asthma and is suggestive as a potential therapy. In a randomized human trial, recombinant human APC administration did not alter the procoagulant state in the airway but did decrease neutrophil influx and degranulation while leaving eosinophil influx unaltered. However, inhalation of APC before ovalbumin-challenge in mice resulted in a strong attenuation of allergic inflammation with decreased eosinophil influx, IL-4, IL-5, and IL-13 and IgE levels in BAL fluid, and diminished AHR. These findings suggest that APC plays an anti-inflammatory role through modulation of dendritic cell and neutrophil trafficking in asthma, and these effects are independent of the functions of APC in the coagulation pathway. Therefore, reduced levels of APC in Type 2-high asthma reflect increased inflammation in the airway, and administration of APC inhibits inflammatory cell migration, suggesting a role as an anti-inflammatory therapeutic agent for asthma.

Current asthma therapies target asthma symptoms and exacerbations, but few, if any, specifically address the mediators of the coagulation pathway. This discrepancy may, in part, be due to the complexity of the coagulation cascade and difficulties faced when regulating its mediators. In a study conducted by Peyvandi et al, deficiency in fibrinogen, FX, and FXIII associated strongly with severe bleeding episodes. Additional studies attempted to knockout TF in mice, but this approach resulted in a significant increase in embryonic death due to severe hemorrhagic events and poor vascular development. Therefore, therapeutic strategies attempting to target mediators of the coagulation cascade must carefully consider the detrimental effects of even mildly altering expression and activity of these factors.

Type-2 high asthma with comorbid obesity characteristically involves distal airways and presents as a severe asthma phenotype. Recent findings from a case study of status asthmaticus found extensive fibrin deposition in distal airways of difficult-to-treat steroid-resistant asthma; these findings were also seen in a Type-2 high asthma mouse model conducted by the same group. In addition to increased fibrin deposition, Wagers et al noted an impaired fibrinolytic system within the mouse model. Given that obesity is an important risk factor for increased asthma severity and that impaired coagulation presents in both obesity and Type-2 high asthma alone, Type-2 high asthma patients with comorbid obesity are at elevated risk for rapid disease progression. The interactions of airway mediators with the mediators of the coagulation cascade and their downstream effects on asthma pathobiology are listed in Table 1.
Table 1 Cytokine, Adipokines, and Coagulation Mediators Contributing to Asthma Pathobiology

### Airway Mediators that Influence the Coagulation Pathway

| Mediator | Airway Cellular Source | Impact on Coagulation Mediator Production | Impact on Asthma Pathobiology | References |
|----------|------------------------|-----------------------------------------|-------------------------------|------------|
| IL-1β    | Macrophage             | Restrict APC, platelet hyperactivation, Inhibits tPA | Promotes airway inflammation | [55,78,94] |
| IL-6     | Macrophage             | Increases TF, platelet hyperactivation | Promotes airway inflammation | [55,78]    |
| IL-8     | Macrophage             | Platelet hyperactivation | Promotes airway inflammation | [55,78]    |
| IL-13    | Th2 cells, mast cells, eosinophils, ILC2 | Increases TF | AHR and airway inflammation | [24,116,117] |
| TNF-α    | Macrophage             | Increases FVII, inhibits tPA | AHR                          | [55,70,94] |

### Systemic Mediators that Influence Coagulation and Airway Processes

| Mediator | Tissue/Cellular Source | Impact on Coagulation Mediator Production | Impact on Asthma Pathobiology | References |
|----------|------------------------|-----------------------------------------|-------------------------------|------------|
| CRP      | Adipocytes and macrophage | Increased TF | Airway obstruction and inflammation; IL-1β, TNF-α production in the airway | [94,104] |
| Leptin   | Adipocytes             | Increases TF, PAI-1, platelet activation | AHR and airway inflammation | [55,71,72,76,77] |
| Resistin | Adipocytes             | Increases TF, PAI-1 | Increases airway mucin gene expression | [81,82,118] |
| Adiponectin | Adipocytes          | Reduces TF | Increases wound repair in bronchial epithelial cells; protects against airway inflammation and vascular remodeling | [85,119,120] |
| TF       | Airway epithelial cells, perivascular and adventitial cells, airway neutrophils, peripheral blood mononuclear cells, adipocytes | Complexes with FVII to initiate extrinsic cascade | Increases VEGF production in the airway; angiogenesis; airway remodeling | [13–15,24,27–30,61] |
| FVII/a   | Hepatocytes and adipocytes | Complexes with TF to initiate extrinsic cascade | Increases VEGF production in the airway, angiogenesis; airway remodeling, eosinophilia and AHR | [27–30,60,67,68,70] |
| FX/a     | Hepatocytes             | Inhibited by antithrombin III; fondaparinux | Increases airway mucin production, AHR, peribronchial smooth muscle thickening, airway fibrosis; decreased lung compliance | [31,32,63] |
| Fibrinogen | Hepatocytes            | Hypercoagulation | Increased ICS resistance | [25,40,67–69] |

(Continued)
Coagulation in Obesity

Obesity is typically defined in the literature by body mass index (BMI ≥ 30 kg/m² for adults) or waist circumference (WC ≥ 40 inches for adult males or ≥ 35 inches for adult females). Obesity contributes to chronic low-grade systemic inflammation due to an increase in pro-inflammatory adipokines and cytokines. Pro-inflammatory mediators increased in obesity include IL-1β, IL-6, IL-8, and tumor necrosis factor-alpha (TNF-α), among others. This inflammatory profile associated with obesity is attributed to polarization from anti-inflammatory M2 to pro-inflammatory M1 macrophages within adipose tissue, which results in increased secretion of Type 1 and Type 17 cytokines. The impact of systemic inflammation in obesity is a significant topic of interest due to its contribution to numerous comorbidities such as type II diabetes, stroke, pulmonary embolism, and asthma, among others.

Obesity promotes a systemic and local procoagulant and anti-fibrinolytic environment, which accounts for the increased risk of cardiovascular diseases, as depicted in Figure 2. This shift favoring coagulation with increased adiposity is due, in part, to increased adipose production of TF and PAI-1. These mediators also increase in systemic circulation due to Type 1 and Type 17 cytokine interaction with various organ systems. Mice on high-fat diet, a model of obesity, experience a significant increase in TF expression in numerous vital organs, including the brain, lung, and kidneys, with the largest increase seen in the brain and lungs.

Aside from augmented coagulation and inhibition of fibrinolysis, obesity is also associated with a decrease of the anti-coagulation mediator, anti-thrombin III, which inhibits FX and prothrombin activation. Obesity is characterized by increased APC, despite the persistent procoagulant state, and these levels are attenuated following weight loss.

The mechanism driving the diminished efficacy of APC in obesity may be related to thrombin generation. A majority of coagulation factor synthesis occurs in the liver, and numerous factors increase in obesity, including FVII and fibrinogen. Adipocytes exposed to TNF-α, which is elevated in obesity, also contribute to increased synthesis of FVII which complexes with TF initiating the extrinsic coagulation cascade. Given these findings, the augmented presence of FVII, supplied both by adipocytes and hepatocytes, increases the coagulation potential within obesity.

| Table I (Continued). |
|----------------------|

| Mediator | Airway Cellular Source | Impact on Coagulation Mediator Production | Impact on Asthma Pathobiology | References |
|----------|------------------------|------------------------------------------|-------------------------------|------------|
| Thrombin | N/A                    | Increased fibrinogen cleavage; binds to PARs | Airway inflammation           | [33]       |
| FDPs     | N/A                    | Increased PAI-I                            | Increased airway inflammation; Increased IL-1β, IL-6 production in the airway, leukocyte trafficking | [35–37]   |
| PAI-I    | Adipocytes, airway macrophages, mast cells, fibroblasts, epithelial, endothelial cells | Inhibits tPA and uPA | Increased AHR; impaired lung function and airway fibrosis | [25,36,38,40,42,62,72] |
| PC and APC | Hepatocytes         | Decreases airway thrombin levels | Decreased airway neutrophilia in humans, decreased airway eosinophilia, IL-4, IL-5, IL-13 production and AHR in mice | [46,47,64–66] |
| tPA and uPA | Airway epithelial cells | Inhibited by PAI-1 | Reduced AHR; airway fibrosis | [40,43] |

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A major adipokine elevated in obesity is leptin; a satiety hormone that synapses in the lateral geniculate nucleus of the thalamus and inhibits hunger. Leptin signaling becomes imbalanced during obesity, leading to leptin hyposensitivity, unregulated caloric intake, and adipogenesis. Leptin is a significant contributor to chronic low-grade systemic inflammation and influences mediators within the coagulation pathway, such as TF expression, in neutrophils and peripheral blood mononuclear cells and PAI-1 production by vascular endothelial cells. Leptin also interacts with type-2 cytokines to promote mucin and eotaxin secretion by airway cells and eosinophil chemotaxis in the airway, suggesting a role for leptin in augmenting airway remodeling and inflammation in Type-2 high asthma with comorbid obesity.

Obesity not only contributes to augmented susceptibility to pro-coagulation through increased production of procoagulant and anti-fibrinolytic factors, but also through enhanced platelet hyperreactivity. The effect of leptin on platelet activation is still under debate, but platelets cultured in physiological equivalent levels of leptin, adenosine diphosphate (ADP), and thrombin seen in obesity resulted in increased aggregation, while this did not occur in levels comparable to those observed in lean patients. Additionally, IL-1β, IL-6, and IL-8 were all found to promote hypercoagulability. More specifically, IL-1β impaired anti-coagulation processes restricting activation of PC; IL-6 increased TF, priming for coagulation initiation; and addition of all three cytokines to whole blood produced the most pronounced platelet hyperactivation.

Additional adipokines impacted in obesity include augmented resistin (pro-inflammatory) and decreased adiponectin (anti-inflammatory and atherogenic). The impact of resistin and adiponectin on coagulation factor synthesis is under debate. An in vitro study conducted by Ayer et al found no significant effect of increasing doses of resistin on TF production by peripheral blood mononuclear cells. However, Calabró et al observed augmented TF expression in human coronary artery endothelial cells when treated with resistin. Additionally, resistin gene knockdown in 3T3-L1 adipocytes resulted in decreased...
PAI-1 gene and protein production. A mouse study conducted by Kato et al found no significant difference between platelet count or levels of coagulation parameters between wild type and adiponectin knockout mice. Furthermore, the addition of adiponectin to human platelets had no impact on platelet activation/aggregation. Despite these reports, adiponectin appears to inhibit macrophage TF expression, possibly providing a protective role for the initiation of the extrinsic pathway.

To date, no studies have reported the impact of diet and exercise on coagulation profiles specifically in Type-2 high asthma or in those patients with asthma and comorbid obesity. However, a clinical trial found that a low sodium diet in asthma patients had no impact on systemic inflammation or exercise on coagulation profiles specifically in Type-2 high allergic asthma phenotype. However, this finding is not surprising. Augmented AHR is attributed to increased adiposity through reduced lung and chest wall compliance largely due to compression, which is corrected following weight loss. Despite this, all patients experienced a significant improvement in controlling asthma exacerbations. Lack of change in AHR in patients with high IgE suggests the pre-existing allergic asthma-associated airway pathology is still present following weight loss from bariatric surgery, contributing to the lack of significant effect found by Dixon et al. These findings provide insight into the role that increased adiposity plays in complicating asthma management, which is corrected following weight loss; further studies are needed to better understand the metabolic and cellular signaling mechanisms whereby bariatric surgery impacts features of asthma exacerbations and management.

In addition to improved asthma control, bariatric surgery corrects inflammatory cytokine and adipokine profiles. Circulating levels of leptin, TNF-α, IL-6 and C-reactive protein (CRP), an indirect inhibitor of tPA through augmentation of IL-1β and TNF-α, significantly decrease 3-to-12 months following bariatric surgery, with a concomitant increase seen in circulating adiponectin levels. Serum levels of Type-2 cytokines IL-13 and chitinase-3-like protein 1 (YKL-40) are also shown to decrease following bariatric surgery. Furthermore, the airways following bariatric surgery show a significant decrease in mast cells 12 months post-operative, an important cell type for mediating allergic responses and initiating asthma pathobiology.

The impact of bariatric surgery on parameters of coagulation has also been identified and are shown in Table 2. As previously mentioned, bariatric surgery reduces CRP levels. CRP is known to increase TF expression by monocytes, thereby increasing pro-thrombotic risk. Independent of the impact of CRP on TF production, weight loss alone results in a significant decrease in TF. Serum levels of the co-initiator of the extrinsic coagulation cascade, FVII, also significantly decrease 12 months after bariatric surgery, further reducing the pro-thrombotic potential. Cugno et al further found a significant decrease in serum CRP, PAI-1, and fibrinogen 3 months post-operative, with the lowest levels achieved at 12 months.

The increased circulating fibrinogen in patients with obesity contributes to rapid clot formation through augmented hypercoagulability and increased resistance to
fibrinolysis, which significantly improves following bariatric surgery and weight loss.\textsuperscript{18,23} Furthermore, Stolberg et al found a significant decrease in systemic levels of thrombin, fibrinogen, and PAI-1 antigens following Roux-en-Y gastric bypass (RYGB) at 6- and 24-months post-operative, leading to improved clot lysis.\textsuperscript{22} Similar results were noted by Ay et al who showed significant decreases in PAI-1, TF, and prothrombin fragments 2 years after surgery due to reduced thrombomodulin and thrombin potential.\textsuperscript{64–66} Bariatric surgery also promoted a significant reduction in systemic PAI-1 as early as 3-months post-operative,\textsuperscript{22,65,97,99,100,105} reducing the inhibition of tPA and uPA, leading to improved fibrinolysis.

Indeed, epigenetic mechanisms may provide a link between bariatric surgery and improved coagulation profiles in patients with obesity. DNA methylation presents as a possible molecular response following bariatric surgery and may be involved in reversing some of the detrimental effects of obesity. Compared to lean individuals, adults with obesity exhibit global hypomethylation, with associated increased expression of proinflammatory genes.\textsuperscript{106} A study conducted by Nicoletti et al found that surgically-induced weight loss resulted in decreased methylation of \textit{IL6}, while an increase in \textit{IL6} methylation was observed following weight loss through restricted diet.\textsuperscript{107} These authors further demonstrated that baseline \textit{SERPINE1} (the gene encoding PAI-1) methylation was significantly associated with percentage of weight loss, suggesting that baseline methylation markers of coagulation pathway-associated genes may predict future weight loss in patients.\textsuperscript{107} These results are intriguing and point to the potential of epigenomics studies to offer greater insight into the intersection of obesity and asthma pathobiology.

It should be noted that the attenuated coagulation factors observed following bariatric surgery were seen in systemic circulation. To date, there are no studies investigating how these coagulation parameter profiles change in the airway following bariatric surgery. Despite this knowledge gap, attenuated circulating levels of procoagulant/anti-fibrinolytic mediators are suspected to improve asthma outcomes, which are worsened through an imbalance in the coagulation cascade in Type-2 high asthma patients with comorbid obesity. In asthma, there is an increase in vascularity,\textsuperscript{108,109} vascular permeability,\textsuperscript{110–112} and plasma exudation,\textsuperscript{40,113,114} thereby readily exposing extravascular tissues to the circulating mediators of the coagulation cascade.

**Table 2 The Impact of Bariatric Surgery on Mediator Production**

| Coagulation Mediator | Effect of Bariatric Surgery | Timeframe Following Surgery | Reference |
|----------------------|----------------------------|-----------------------------|-----------|
| TF                   | Decrease                   | 2 years                     | [65]      |
| FVII                 | Decrease                   | 12 months                   | [99]      |
| PAI-I                | Decrease                   | 3, 6, 12, and 24 months     | [22,65,97,99,100] |
| Fibrinogen           | Decrease                   | 3, 6, 12, and 24 months     | [22,99]   |
| Prothrombin          | Decrease                   | 6 and 24 months             | [22,65]   |
| Anti-thrombin III    | Increase                   | 6 and 12 months             | [63,105]  |
| PC/APC               | Decrease                   | 12 months                   | [64–66]   |

**Conclusion**

The increased prevalence of obesity worldwide poses new challenges within health care. More specifically, previously understood and controlled pathologies are being complicated by increased adiposity at an alarming rate, as seen in comorbid obesity and asthma. Impaired coagulation in Type-2 high asthma with comorbid obesity presents as a unique pathologic mechanism; advancing our understanding of this pathway will allow for better therapeutic development for this growing population of patients.

At present, the lack of treatments for Type-2 high asthma with comorbid obesity places these patients at increased risk for hospitalization, ICU admission, poor symptom control, and quality of life.\textsuperscript{8} As presented herein, an imbalance in the extrinsic coagulation cascade resulting from increased adiposity may play a significant role in
impaired management of asthma exacerbations. Increased levels of TF and FVII in the airway and systemic circulation respectively, primes the pathway for rapid activation. Increased presence and synthesis of downstream mediators FX and fibrinogen, among others, not only aids in rapid clot formation within the airway, but also produces collateral effects driving AHR, remodeling, and inflammation. Additionally, augmented production of anti-fibrinolytic mediators prolongs clot presence and impairs responsiveness to ICS. Given the complexity and heterogeneity of asthma and the general influence of obesity on cardio/pulmonary vascular pathobiology, these pathways may be critical in other phenotypic presentations of asthma, including Type-2 low asthma with obesity, an area in need of much greater understanding.

Currently, available asthma therapies do not target mediators within the coagulation cascade. This circumstance is due, in part, to the challenges and complications faced when modulating systemic levels of mediators of this complex pathway. Interestingly, inhaled heparin, an antioxidant and anti-coagulant, has been identified as a novel therapy to enhance asthma management through decreased airway obstruction, improved airway inflammation, and augmented delivery of other inhaled therapies.\(^\text{15}\) Despite heparin’s potential benefit in asthma management, it does not address the underlying shift towards a procoagulant environment in the airway.

Obesity is characterized by a significant pathway modification favoring procoagulant and anti-fibrinolytic mediators, both in the airway and in systemic compartments. The use of concurrent ICS and heparin addresses the symptomatic responses of asthma but cannot address the shift in inflammatory and pro-coagulation profiles. However, bariatric surgery addresses these imbalances on a systemic scale leading to improved responsiveness to therapies.

We hypothesize that bariatric surgery, unlike other pharmacological interventions, provides significant improvements in asthma control as seen through improved AHR, lung compliance, and normalized coagulant and inflammatory parameters. Future research will identify specific metabolic and cellular mechanisms whereby bariatric surgery influences airway coagulation and physiology and reveal potential applications to clinical care. A better understanding of bariatric surgery-induced metabolic and cellular mechanisms regulating mediators of the coagulation pathway, whether those mechanisms be weight loss-dependent or -independent, holds promise for the development of potential therapies to benefit all patients with Type-2 high asthma, not just those with comorbid obesity.

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**Author Contributions**

All authors made substantial contributions to conception and design, acquisition of data, or analysis and interpretation of data; took part in drafting the article or revising it critically for important intellectual content; agreed to submit to the current journal; gave final approval of the version to be published; and agree to be accountable for all aspects of the work.

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