Sex hormone has become a "hot topic" to evaluate the hormonal therapeutic potential in severe asthma. Th17 cell is one of the main influencing factors involved in the pathogenesis of severe asthma, hence also called as kernel of severe asthma, and Th17 subtype of non-T2 asthma is less responsive (resistance) to inhaled corticosteroid (ICS), so severe in nature. Methyl-CpG binding domain protein 2 (MBD2) is overexpressed and regulates the Th17 differentiation, showing the possibility of therapeutic target in treating Th17 mediated severe asthma. Sex hormone fluctuates at the different physiobiological conditions of the human body and affects the asthma pathobiology showing its role in asthma prevalence, severity, remission, and therapy. This review briefly overviews the sex hormones, their influence in asthma at the different physiobiological conditions of human body, and MBD2 severe asthma connection with the possible therapeutic potential of sex steroids in MBD2 mediated Th17 predominant severe asthma. Male sex hormone tends to show a beneficial effect and possibly downregulates the expression of Th17 cells via regulating MBD2 through a mechanism distinct from corticosteroid treatment and guides us towards discovery of new therapeutic agent, reduces the asthma-related complications, and promotes long-term survival by lowering the risk of therapy-resistant issues of old age severe asthma.

1. Introduction

Asthma is a complex, chronic, heterogenous inflammatory disease, with diverse endotypes and phenotypes, characterized by airway inflammation and airway hyper-responsiveness (AHR), relieved spontaneously or by medications [1]. The T2 (T2 high) asthma is early-onset allergic, and late-onset nonallergic eosinophilic asthma, sensitive to ICS, and Th2 biomarkers are commonly used in diagnosis [2]. T2 low (non T2) asthma, also known as noneosinophilic asthma, is still obscure and is late-onset neutrophilic, paucigranulocytic, or mixed, and inflammation is driven through varieties of asthma-related inflammatory cells and may be associated with airway smooth muscle or neural dysfunction as well as comorbidities occasionally [3, 4]. T2 low asthma is generally less responsive to inhaled corticosteroid, hence also known as severe asthma.

Th17 cells and their cytokines are considered as the main influencing factors in the pathogenesis of severe asthma. That is why Th17 is also entitled as a kernel of severe asthma. Pro-inflammatory cytokines, oxidative stress, neuronal and hormonal responses, and epigenetic regulation are also involved in Th17 cell-mediated severe asthma, but the underlying mechanisms are still unknown. Figure 1 shows factors associated with sex hormones affecting severe asthma.

The methyl-CpG-binding domain (MBD) family proteins determine the transcriptional state of the epigenome, can interpret DNA methylation in epigenetics [5, 6], and are equally crucial for emerging roles in immunity [5]. MBD2 binds to the target gene’s promoter region, induces posttranscriptional histone modification, changes the chromatin structure, and ultimately regulates the expression of target genes, and this is believed to be an essential critical mediator in asthma epigenetics mechanism.
It is a well-known fact that the loss of MBD2 inclines towards Th2 cellular polarization [7] and deficiency in Th17 differentiation. Our initial study in patients showed increased expression of MBD2 in Th17 mediated severe asthma from peripheral blood samples. We also found the increased MBD2 expression after stimulus differentiation in splenic CD4+ T-cells in our animal model, showing involvement of MBD2 in immunological pathogenesis of Th17 mediated neutrophilic severe asthma and differentiation of CD4+ T-cells.

Sex steroids are essential for sex differentiation and reproduction. Testosterone (TES) and other androgens, such as dihydrotestosterone (DHT), have broad immunoregulatory effects that suppress immune responses [8]. Estrogen also regulates innate immune cells and the signaling pathways [9]. The natural physiobiological conditions like menstrual cycle, pregnancy, menopause, and oral contraceptives pills (OCP), hormone replacement therapy (HRT), and epigenetics [10, 11].

Estrogen and progesterone are involved in the differentiation of Th17 cells. IL-17A production with IL-17A mediated inflammation and asthma severity. The interaction and association of sex hormones with internal and external factors are complex. However, understanding this complexity is beneficial for understanding the asthma pathobiology and generating a patient approach therapy.

2. Sex Hormone Steroidogenesis

Cholesterol is the precursor of all steroids, including sex hormones that primarily come from cholesterol ester, uptake by plasma proteins lipoprotein receptors, scavenger receptor class B member 1 (SR-B1) [17, 18]. The side-chain cleavage enzyme p450scc, an essential enzyme to synthesis all steroid hormones, converts the cholesterol into pregnenolone [19]. Pregnenolone is a progestogen and can be metabolized to progesterone by 3β-hydroxysteroid dehydrogenase (3β-HSD), and P450c17 transforms this progestogen to 17α-hydroxyprogrenenolone and dehydroepiandrosterone (an androgen, DHEA). Hydroxilation of DHEA to androstenedione and androstenediol is mediated by 3β-hydroxysteroid dehydrogenase 2 (3β-HSD2) and 17β-hydroxysteroid dehydrogenase 3 (17β-HSD3), respectively [19]. TES is the transformed product of androstenedione and androstenediol...
Figure 2: Sex hormones steroidogenesis. Cholesterol is the precursor of all steroids, including sex hormones, uptaken by plasma proteins lipoprotein receptors scavenger receptor class B member 1 (SR-B1) [17, 18]. P450scc, the side-chain cleavage enzyme, converts the cholesterol into pregnenolone [19]. Pregnenolone is metabolized to progesterone by 3β-hydroxysteroid dehydrogenase (3β-HSD), and P450c17 transforms progestogen to 17β-hydroxypregnenolone and dehydroepiandrosterone (an androgen, DHEA). Hydroxylation of DHEA to androstenedione and androstenediol is mediated by 3β-hydroxysteroid dehydrogenase 2 (3β-HSD2) and 17β-hydroxysteroid dehydrogease 3 (17β-HSD3), respectively [19]. Androstenedione and androstenedion biotransferred to TES by 3β-HSD2 and 17β-HSD3, respectively. 5α-dihydrotestosterone (5α-DHT) [20] and 5β-dihydrotestosterone (5β-DHT) [21] are the reducing TES product by 5α-reductase type 1 or 2 and 5β-reductase, respectively [20, 21]. P450 aromatase (P450aro) bioconverts androstenedione to estrone and TES to 17β-estradiol [19]. Estrone is also bioconverted to 17β-estradiol by 17β-HSD3. TES: testosterone.

3. Early Childhood, Sex Hormones, and Asthma

During childhood, boys experience more frequent asthma symptoms with higher asthma prevalence than girls (ratio of 2:1 before the age of 5 years) [22, 23]. This ratio further increases to 4-fold until the age of 14, and the gender switch occurs to 1:2 as the plasma TES level increases in boys [24–27] (Table 1 shows asthma and sex hormone cycles in human life).

(Note: in boys, asthma prevalence in childhood is higher than girls, and there is a gender switch to females from puberty to adulthood as the male sex hormone and airway caliber is believed to be involved because old aged male tends to have increased asthma episodes than previous stage as a result of decreased level of male sex hormones. In contrast, female tends to have increased asthma prevalence from puberty onset to adulthood mainly due to increased level of female sex hormone. This tendency is reversed during old age mainly due to decreased levels of female sex hormone resulting in a minimal gap between old age male female asthma ratio than the previous stage showing the beneficial role of androgen in old age severe asthma. ↑: minimal increase; ↑↑: high increase; ↑↑↑: highest increase; ↓: minimal decrease; ↓↓: high decrease; ↑↑↓: decrease than previous stage and in decreasing status)

Besides sex hormones, some other factors are also influential in the pathogenesis of childhood asthma. Dysanapsis refers to the differences in airways’ growth to lung, and boys are the sufferer of this mechanism as they have smaller airway diameters compared to lung volumes than the girls [28], and females have lower specific lung resistance than males. This phenomenon also makes boys more likely to have asthma episodes than the girls [29]. Obesity is

<table>
<thead>
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<th>Table 1: Asthma prevalence and sex hormone cycle in male and female.</th>
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<td>Childhood</td>
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<td>Male asthma</td>
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associated with childhood asthma in girls but not in boys [30, 31], and obesity is also associated with hormonal imbalances.

Why do female children experience lower asthma prevalence than the male? Is it anatomical cause or gender differences? Is there any protective effect of female sex hormones for girls or natural selection? Is there any immunologic mechanism that is driving allergic childhood asthma more towards boys? Though the mechanism of age and gender-related asthma differences with hormonal influences is still not precise, we believe the increased prevalence of asthma in male child and shift of prevalence to adult female is strongly associated with the role of sex hormones.

4. Menarche, Menstrual Cycle, Sex Hormones, and Asthma

Menarche is the first menstrual cycle that shows the beginning of the onset of reproducing capacity, development of secondary sexual characteristics [32], and beginning in physiological changes in hormonal cycle. The median age of menarche has been decreasing from the last century and is less than 13 years nowadays [33]. Early menarche fluctuates the hormones, increases the asthma incidence [34], and is also associated with increased risk of breast cancer, cardiovascular disease, obesity, and type 2 diabetes [35, 36].

Multiparous women and girls with a history of early menarche have higher exposure to estrogen, increased cumulative exposure to sex hormones, and are at risk of asthma and severe asthma [37, 38]. The twofold increased asthma prevalence in girls with an early history of menarche is possibly due to the role of sex hormones [37, 39, 40]. Low TES levels and premenstrual and menstrual asthma aggravation are correlated in some asthmatic females [41]. Increased adulthood asthma and lower lung function are associated with early menarche, supporting the influence of hormones in women’s respiratory health status [42].

Similarly, menstrual cycle worsening of asthma symptoms is linked with the fluctuations of serum levels of estradiol and progesterone [43]. Nevertheless, there is a decrease of provocative concentration of methacholine (PC20) by more than 50% in adult women with stable well-controlled asthma during the menstrual cycle, and the highest decrease occurs in the luteal phase at peak estrogen-progesterone levels [44]. The abnormal β2 adrenoceptor regulation is attributed to the cyclic changes in PC20 premenstrual asthma [44, 45].

Two to sevenfold higher serum estradiol and progesterone levels and lower forced expiratory volume in one second (FEV1) and forced vital capacity (FVC) are observed in women with and without asthma during the luteal phase of the menstrual cycle [44, 46]. That is why lower FEV1 and more respiratory symptoms are observed in the premenstrual and menstrual period asthma with an increased level of airway hyperresponsiveness and probably increased medical attendance [44, 47]. Irregular menstruation is also associated with lower lung function [43], and lung function predicts respiratory health. High fractional exhaled nitric oxide (FeNO) correlates with eosinophilic inflammation and increases in women with premenstrual asthma symptoms [48].

Together, these studies indicate the role of circulating estrogen and androgen levels as a reason for more prevalent asthma in females after menarche and during the menstrual and perimenstrual period, but the underlying mechanisms are still unclear.

5. Puberty, Sex Hormones, and Asthma

Puberty is a dynamic process of physical changes and sexual maturation regulated by hormones. In boys, adrenal glands produce weaker androgens, and testes increase the production of TES, whereas in girls, the estrogen production is increased from ovaries leading to thelarche, menarche, and adrenal glands that produce androgens such as androstenedione and DHEA leading to pubarche [49].

There is a gender switch in asthma prevalence at puberty from male to female as asthma is more prevalent and severe in female [50, 51]. Fluctuation and surges of endogenous sex hormones are believed to be involved in the gender switch of asthma [52]. The Childhood Asthma Management Program (CAMP) study tracked the progression of asthma and average asthma symptom score in puberty with Tanner scores. At the age of 10 years, the average asthma symptoms increased with the increasing score but started to decline in boys. The score continually increased with the increasing ages in girls [53].

The problem of dysanapsis seems to be corrected with the high levels of TES during puberty, as this hormone induces a more pronounced airway caliber in boys than girls and gives protection from asthma. This may be the reason behind the more prevalent asthma in old age males than women as the hormone level decreases in aged male [28, 54]. In boys, the increase in the dose of PC20 to decrease FEV1 to 20% shows the pubertal improvement in airway responsiveness, but not observed in girls [55], and the PC20 was further increased with sexual maturation in boys [55]. Hormonal fluctuations and correction of anatomical problems are the main influencing factors in the prevalence and gender switch of asthma at puberty.

6. Pregnancy, Sex Hormones, and Asthma

Asthma is a common, sometimes fatal medical condition that has a variable impact on pregnancy. Generally, 33% of pregnant women experience worsening asthma symptoms with bronchial hyperresponsiveness, 33% improve, and the remaining one-third have unchanged asthma status [56]. Controlling asthma and maintaining asthma control are vital for reducing fetal and maternal risks linked with asthma during pregnancy. Because poorly controlled asthma and severe asthma exacerbations are associated with increased risk newborn comorbidities like higher rates of prematurity, intrauterine growth retardation, respiratory complications, and hyperbilirubinemia [57], preeclampsia and gestational diabetes are seen on the maternal side. But interestingly, about 7.2% of pregnant women with improved asthma status present with asthma exacerbations [56].
Pregnancy with more severe asthma phenotypes is likely to exacerbate and worsen asthma symptoms frequently [58].

There is a linear increase in asthma prevalence in multiparous women with the increasing order of birth [38], and multiparous women and girls with early menarche are exposed to higher estrogen levels with significant accumulative doses of sex hormones exposure. They tend to have a higher risk of severe asthma [37, 38].

The sex of the fetus is considered a risk factor for worsening asthma symptoms during pregnancy. A female fetus has been linked with worsening maternal asthma symptoms and related to hormonal changes in pregnancy, with the speculation of discharge of fetal sex hormone in the maternal blood resulting in asthma exacerbations, increase in corticosteroid use, and even hospitalization independent of maternal age, smoking status, and body mass index (BMI) [59–62]. Interestingly, female newborns with reduced birthweights were remarkably seen from asthmatic mothers but not in male newborns postulating gender-specific effect [63] and probably male hormonal effect. However, a larger Canadian study did not confirm this finding [64]. The male fetus and increased level of TES might be protective against maternal asthma [62].

It is concluded that pregnancy-induced physiological processes may confound the hormone effect, but mainly hormonal fluctuations either from maternal or fetal side possibly involved in pregnancy-related asthma.

7. Adulthood, Middle Age, Sex Hormones, and Asthma

Adulthood is a state where humans are believed to be fully grown and matured with complete physical and intellectual maturity around the completion of second decade. Defining middle age is arbitrary, but it is generally believed to be between 40 and 60 years. This is the age where people start to experience physiological, psychological, and hormonal changes and a somewhat gradual decline in physical activities.

In this age group, asthma is more prevalent in women until the age of 45 years and is also associated with higher mortality [65, 66] and beyond that increased in men suggesting the age-gender interaction [54], with possible hormonal influence. Similarly, women have a higher incidence of non-atopic asthma during the reproductive period, but no gender difference is noted in the incidence of atopic asthma [67, 68].

In females, the transition of lower to higher risk and prevalence of asthma from childhood to adulthood is associated with hormonal transitional fluctuations during normal physiobiological stages like puberty, menarche, menstruation, and menopause [50, 69]. Similarly, the male sex hormone is believed to be involved in the opposite asthma status in males and, overall, suggest sex hormones might have a role in modulating pathways associated with asthma pathogenesis.

8. Menopause, Sex Hormones, and Asthma

Menopause is a natural physiobiological process that heralds the penultimate menstrual cycle, generally occurs at late 40s or early 50s, and the decrement of sex hormones usually occurs from this phase. The perimenopausal period is the transition phase from where the asthma in females changes from high incidence occurring in adulthood to low incidence, and less severe from menopause onwards, and is believed to be related to fluctuations in female sex hormones. Age, asthma duration, obesity, HRT, and allostatic load (AL) are the confounding factors that affect the independent effect between hormonal fluctuation and asthma status in menopause.

In postmenopausal women, age-adjusted relative risk of asthma may drop compared with premenopausal women [70, 71], and minimal postmenopausal difference of asthma prevalence is observed between men and women compared to adulthood [50]. The age-adjusted lower relative risk of asthma incidence is noted in postmenopausal women who never received HRT [70], and interestingly, the protective effect of menopause was reversed by estrogen HRT [70].

Overall, the reduced asthma prevalence and severity in postmenopausal women compared to men postulates that asthma improves after menopause, and menopause is the phase of reduced levels of sex hormones that probably have an impact on asthma status changes.

9. Old Age, Sex Hormones, and Asthma

The age approaching or exceeding life expectancy is old age, and aging is the process of growing old. The cumulative “wear and tear” on the body due to regular and/or exaggerated response of body physiological systems to environmental challenges is known as AL [72]. It can result in premature ageing [73] and a proper mechanism to understand mortality and morbidity in asthma. AL as early aging is present in the mildest asthma group not receiving ICS [74]. Another aging change associated with asthma is autophagy, a biological process of intracellular degradation associated with conditions like asthma, chronic obstructive pulmonary disease (COPD), cystic fibrosis, pulmonary hypertension, cancer, infection, and even in normal aging. Plasma autophagy marker (LC3B) increases in asthmatics, a more pronounced increase of LC3B seen with severe asthmatics with inverse FEV1 association and direct association with increased age. Eosinophilic persistent asthma has a close correlation with accelerated aging [75]. The immune system and various structural changes occur on aging asthmatics like increased chest wall rigidity, less elastic recoil, and reduced respiratory muscle power that ultimately furnish the declined lung function [76–78] and confound the hormonal impact of asthma.

A noticeable decline in serum TES level is noted with increasing age in an old-aged man [79, 80], resulting in more pronounced and severe asthma prevalence. Low androgen level in childhood and their deficiency in aging males postulate the significant androgen role in childhood and old age asthma [80]. Interestingly, circulating androgens are more related to metabolic risk factors than medical comorbidities in males from young age to middle age to old age, but female sex hormone estrogen is not associated with metabolic risk traits. That is why the association of sex hormones with age, metabolic factors, and medical comorbidities is responsible
for alteration in health-related factors [81] and even asthma transition with ages.

Conceivably, low TES in older men possibly exacerbate asthma indicates the association of declining level of sex hormone. It might be used as a potential asthma therapy in old-aged males and possibly even in old-aged females.

10. Obesity, Hormonal OCP, Neuronal Factors, Sex Hormones, and Asthma

10.1. Obesity, Sex Hormones, and Asthma. Obesity is associated with higher asthma risks, especially in girls and women. Leptin, a key player in body weight regulation, regulates TH1 responses, and the production of asthma-related proinflammatory mediators is associated with obese females in asthma [82]. Regardless of physical fitness, higher asthma prevalence and asthma-related morbidity are seen in girls but not in boys [83], and about five to seven times higher asthma prevalence are seen in obese female above 11 years of age than the normal BMI female [84, 85]. Increasing BMI has also direct association with asthma severity [65]. In contrast, it was interesting that low BMI smoker women are at higher risk of asthma than the high BMI nonsmoker women [86].

Although obese-asthma phenotype gender dimorphism is supported by many reports [55], it is still unclear whether this is caused by gender-specific factors confounded with sex hormones. However, we strongly believe in the role of sex hormones in obesity-mediated asthma.

10.2. Hormonal OCP, Sex Hormones, and Asthma. The administration of exogenous hormones and the development of asthma have been explored in many studies. No significant association is found between OCP and the development of asthma [71]. However, there is a slight significant increase in asthma risk with HRT in women. Recently, it has been found that hormonal contraceptives are beneficial for reproductive-age women and may reduce the risk of severe asthma exacerbations [87]. However, the biological basis of how the hormones influence the asthma outcome needs to be evaluated.

Hormonal contraceptives are beneficial in premenstrual asthma and associated with a lower risk of current asthma and fewer symptoms, (OR 0.68; 95% CI, 0.47–0.98) and (OR 0.68; 95% CI, 0.47–0.98), respectively [37, 88]. The luteal phase sex hormones repression by OCP attenuates the cyclical change in airway hyperresponsiveness as seen through changes in PC 20% to drop FEV1 and PEF [89].

Risk of asthma and wheeze with shortness of breath is increased in menstruating women (OR 1.42; 95% CI, 1.09–1.86) and (OR 1.27; 95% CI, 1.02–1.60) using OCP but not in the women without OCP [38, 90]. However, the risk was not seen in menstruating underweight women [26]. Similarly, OCP is associated with increased wheezing risk in asthmatic women [90, 91] and decreased wheezing in some studies [37, 92]. In 2021, the role of HRT in perimenopausal and postmenopausal women with asthma has been explored, and increased risk of asthma exacerbations in perimenopausal and postmenopausal women is associated with previous history of long-term HRT of at least more than two years but not current [86].

Evidently, hormonal OCP or HRT plays crucial role in asthma pathogenesis and prevalence and influences the outcome. However, the mechanism of how HRT work needs to be evaluated thoroughly.

10.3. Neuronal Factors, Sex Hormones, and Asthma. Women with attention deficit hyperactivity disorder (ADHD) reported to have higher prevalence of asthma than men with ADHD which is related to higher prevalence of smoking and obesity [93] and might have a confounding hormonal role. Studies have shown that, during the decline of estrogen and progesterone, i.e., weeks before menstruation, ADHD symptoms may worsen [94] and improve during elevated estrogen and progesterone, i.e., during pregnancy [95]. However, in another study during the menstrual cycle, enhanced ADHD symptoms were associated with reduced estradiol (E2) levels and increased progesterone or TES levels on the following day [96]. That is the reason while treating the asthmatic female with ADHD, and it is advised to inquire about regularity of the menstrual cycle, hormonal profile, hormone therapy, and other physiobiological stages.

Together, the confounding association between ADHD, obesity, smoking, menstrual cycle, pregnancy, and gender with sex hormones in asthma needs to be thoroughly evaluated.

11. Sex Hormones Functional Activities in Asthma

Sex hormones are linked with asthma, but the effect and the progression, remission, or protection mechanism of asthma are complex and primarily associated with the hormonal fluctuation [97], mainly through regulating hormonal receptors that act differently. Male gets beneficial protective effect of sex hormones after puberty, and at the same time, girls before puberty and female after menopause have the beneficial effect of sex hormones in asthma. The cyclic changes of hormones are believed to be linked with asthma exacerbations in females [98]. Below, we evaluate the functional status of sex hormones in asthma (Figure 3 for the functional role of sex hormones in asthma).

11.1. Estrogens. Estrogen receptor (ERs), important players in allergic lung inflammation, are involved in the increased production of asthma-related cytokines in severe asthma, expressed in majority of immune cells, including macrophages. In asthma, different estrogen receptor α (ERα) and estrogen receptor β (ERβ) isoforms affect the airway remodeling, lung development, and differential expression of ERα, and ERβ results in airway smooth muscle contraction and changes in intracellular calcium [99]. Exaggerated M2 polarization was observed in an ovariectomized mice; however, after estrogen supplementation and macrophage-specific deletion of ERα, it showed impaired M2 polarization [100, 101]. In ASM (airway smooth muscles), ERβ activation mediates estrogen’s proliferation and signaling pathways in asthmatic airway remodeling [102]. Five variants in the
ESTROGEN

C18H24O2

Receptors: ERα, ERβ

• Asthma-related cytokines
• Smooth muscle contraction
• Airway remodeling
• BHR
• M2 macrophage polarization
• Rapid loss of lung function
• NO in BECs
• GATA3, Th2 response
• IL-4, IL-23R, IL-17A
• Th2 CD103+ DCs in BLN

| Ca2+ | i |

PROGESTERONE

C19H28O2

Receptors: PR-A, PR-B

• IL10, IL-1β, IL-5, IL-6, IL-22, IL-4, IL-17A, TNFs, and GATA 3
• Relaxation of bronchial smooth muscle
• PEF (in luteal phase)

ANDROGENS

C19H28O2

Receptors: AR

• Airway relaxation/bronchodilation
• Epithelium-dependent airway relaxation
• FEV1 and FVC
• Foxp3, CD25 (hi) Treg cells

ERα gene (ESR1) in females is associated with BHR and exaggerated loss of lung function [103]. The fall in circulating estrogen and progesterone correlates with the fall in FEV1 from the luteal phase to follicular phase [46] and accompanies with increased BHR [44] and reduced lymphocyte β2 adrenoceptor density [45]. Increased levels of nitric oxide (NO) in the breath generally indicate airway inflammation and asthma. Estrogen increases NO production in bronchial epithelial cells (BECs) of female asthmatics, and its impairment might contribute to altered bronchodilation [104]. In ASM, ERβ activation reduces (Ca2+)i and is involved in the regulation of ASM contraction in asthma [105]. In asthma, more exaggerated airway remodeling is noted in females, suggesting the involvement of sex hormones and gender factors [106].

A study has shown that estrogen mediates Th2 mediated severe asthma by slight upregulation of the GATA-3 expression and IL-4 production [107]. The 17β-estradiol (17β-E2) mediated Th2 cytokine production in allergic asthmatic females is attributed to Th2-oriented CD103+ DCs in the BLN (bronchial lymph node) [108]

ERα upregulates the IL-23R expression, increases the IL-17A production from Th17 cells, and promotes mitochondrial respiration and proliferation [109]. Similarly, 17β-E2 and progesterone together regulate the IL-23R expression partially through let-7f miRNAs and increase the production of IL-17A from Th17 cells in severe asthma [12]. Airway 17β-E2 is increased in postmenopausal asthmatic women with severe asthma, and its measurement may have the possibility of a suitable biomarker in diagnosing neutrophils predominant severe asthma [110].

11.2. Progesterone. Progesterone exerts its effect through the activation of the progesterone receptors (PRs). PRs expressed in airway epithelium can alter the function of epithelial cells by inhibiting ciliary beat frequency and affect mucociliary clearance during menstrual cycle [111]. PR-A and PR-B are the two isoforms of PRs that exhibit distinctive transcriptional patterns on progesterone response promoters [112]. PR-B is the principal gene transcription activator, while PR-A represses PR-B and ERα transcription [113, 114].

The fall in circulating estrogen and progesterone from the luteal phase to the follicular phase accompanies increased BHR [44] and reduced lymphocyte β2 adrenoceptor density [45]. Interestingly, some studies postulated that progesterone increases the relaxation of bronchial smooth muscle and decreases contractility [115] and is considered potent vasodilator in mice pulmonary arteries than estrogen and TES [116]. Luteal phase high progesterone is positively associated with peak expiratory flow rate in women [117]. Improved status of lung function with tissue homeostasis and reduced inflammation is the beneficiary effect of progesterone in the influenza model [118].
Finally, progesterone stimulates the production of proinflammatory cells and upregulates the expression of cytokines and proteins such as IL-10, IL-1β, IL-5, IL-6, IL-22, IL-4, IL-17A, TNFα, and GATA 3 through the regulation of receptors in asthma [12, 107, 118, 119]. AR binds to TES or to active metabolite 5α-DHT and becomes activated, acts as a DNA-binding transcription factor via translocating into the nucleus, and controls gene expression. AR regulate the function of immune cells and also affect the lung gene expression. AR upregulation decreases the expression of Th2 and Th17 cells, reduces IL-4 production in lungs, and decreases neutrophilic inflammation in severe asthma [15]. AR share somewhat similar structural features like PRs, high doses of PRs can antagonize AR [121].

The protective effect of male sex hormones in asthma is established in many studies. Asthmatic female possesses higher number of circulating ILC2 in blood, as TES attenuates the function and proliferation of ILC2 in men [122]. TES decreases the expression of ILC2 stimulating cytokines like IL-33 and thymic stromal lymphopoietin (TSLP) and diminishes the ILC2 that ultimately attenuates the airway inflammation [122]. In ILC2 dominant allergic airway inflammation of Rag1−/− mice, increased ILC2-related type 2 inflammation significantly increased in female mice, and TES downregulated the ILC2 and its cytokines suggesting sex differences in ILC2 dominant inflammation and role of androgen therapy in ILC2 dominant asthma [123]. Androgen supplementation seems to be beneficial to reduce airway inflammation and induce airway relaxation in asthmatic women with low androgen level (DHEA – S < 200 μg/dL) [124].

Interestingly, the adrenal androgen DHEA inhibits phosphoinositide 3-kinase–dependent signal pathway and prevents airway remodeling (bronchial epithelial to mesenchymal transition) [125, 126] and also inhibits human ASM and airway fibroblast proliferation during puberty [127, 128]. A significant improvement in lung function is noted with a trial of weak androgen therapy in boys with poorly controlled asthma having low androgen level and showed higher DHEA-S and greater FEV1% predicted are positively correlated [129]. The serum level of TES and DHEA correlate with higher FEV1 and FVC in middle-aged community-dwelling men. That is why androgen might have potential as a biomarker of lung function [130] and suppressive agent in severe asthma.

The protective role of androgens and their effects on T regulatory cells (Tregs) were also studied in asthma. The CD25(hi) Foxp3(1) and IL-10 producing Tregs prevent the Th2 responses to allergens by an increased level of Foxp3 in CD25(hi) Tregs [131]. But in asthma, both Foxp3 expression and CD25(hi) Treg were decreased [132], and this decrease was believed to be due to low androgen level [133]. However, only menstruating women in the ovulatory phase showed increased Foxp3 expression in human CD25(hi) Tregs by androgen and correlated with androgen response element (ARE) within the Foxp3 locus [134].

### 12. MBD2 and Severe Asthma

MBD2 is an important mediator in Th17 predominant neutrophilic severe asthma pathogenesis and is justified by different studies. Our recent study showed the potential role of MBD2 as a novel biomarker for identifying severe asthma various endotypes [135]. Loss of MBD2 tilts towards Th2 polarization by activating dendritic cells for Th2 immunity from CD4+ T-cell and increases IL-4 and IFN-γ expression from CD4+ T cells [7], and MBD2-/- results in a deficiency in Th17 differentiation [136]. Interestingly, MBD2 was found to suppress Treg function through the promotion of Foxp3 demethylation [137]. Below, we review MBD2 and Th17 severe asthma association from different studies (Table 2 shows MBD2 expression and respiratory outcome in asthma).

MBD2 regulates the differentiation of Th17 cells by promoting the gene expression related to the inflammatory process. A study has shown that MBD2 gene silencing significantly lowers the IL-17 expression as increased expression increases the IL-17 expression from splenic CD4+ T cells in an animal model of asthma [138]. The study also postulated MBD2 regulated Th17 differentiation is mediated via affecting the expression of an interferon regulatory factor 4, needed for developing inflammatory Th17 cells in severe asthma [138, 139]. Similarly, the suppressor of cytokine signaling 3 (SOCS3) is a protein involved in negative regulation of STAT3, a Th17 cell differentiation stimulator, which is

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**Table 2: MBD2 expression and effects on respiratory outcomes.**

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<thead>
<tr>
<th>MBD2</th>
<th>Effects</th>
<th>Respiratory outcome</th>
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<tr>
<td>↓ Treg function by Foxp3 demethylation</td>
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<td>Asthma</td>
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<tr>
<td>↑ IL-17 expression</td>
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<td>Th17 severe asthma</td>
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<td>↓ SOCS3</td>
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<td>Th17 asthma</td>
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<td>↓ Treg functions</td>
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<td>Severe asthma</td>
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<td>↑ HIF-1α</td>
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<td>Th17 asthma</td>
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<td>↑ MBD2</td>
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<td>↓ IL-4, IFN-γ</td>
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<td>TGF-β1 attenuation</td>
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<td>airway remodeling, ↓ M2 macrophage gathering</td>
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MBD2-/-                          | Unsuppressed Tbet/Hlx axis | ↓ Th17 differentiation                      | [136]      |
involved in asthma pathogenesis. In a study, MBD2 down-regulated the SOCS3 expression and promoted the differentiation of Th17 cells in severe asthma, showing its involvement in Th17 mediated neutrophilic predominant severe asthma [140].

A transcription factor regulates hypoxia in severe asthma exacerbations and hypoxia-inducible factor-1 (HIF-1), and deficiency of HIF-1α reduces Th17 cells development with increased Treg differentiation [141], and study has confirmed that less airway inflammation in HIF-1α-/⁻ experimental mice [142] is believed to be due to reduced Th17 cells. MBD2 increases HIF-1α in neutrophil-dominant asthma and is involved in increased differentiation and secretion of Th17 cells and IL17 through regulating the HIF-1α expression [143] and also reduces the Treg function through an increase in HIF-1α suggesting the epigenetic role of MBD2 in severe asthma [141]. Similarly, unsuppressed Tbet/Hlx axis results in deficiency of Th17 cell differentiation in MBD2-/⁻ mice with the protection from experimental autoimmune encephalomyelitis, showing the role of MBD2 in Th17 differentiation [136].

Transforming growth factor-β1 (TGF β1) modulates allergic airway inflammation and remodeling in asthma, correlated with asthma severity, and is probably associated with MBD2 level. MBD2 deficiency attenuates the TGF-β1 production with decreased M2 macrophage gathering in bleomycin-induced lung fibrosis [144], and M2 polarized macrophages are correlated with FEV1 in asthma.

It is worth noting that MBD2 is involved in the differentiation and expression of Th17 and its asthma-related inflammatory cytokines via inducing epigenetic changes through different mechanisms, and MBD2 itself and/or MBD2 mediated Th17 cells can be a potential therapeutic target in severe asthma.

13. Therapeutic Possibility of Sex Hormones in MBD2 Mediated Severe Asthma

Broadly speaking, we observe the two peaks of asthma in human life [145]. The first peak is during childhood; usually, T2 asthma (T2 high) is early-onset allergic and late-onset nonallergic eosinophilic asthma, sensitive to ICS, and blood eosinophils, FeNO, and IgE levels are the common biomarkers [2] and predominantly seen in childhood as the first asthma peak. The second peak is usually adult-onset/old age asthma, non-T2 (T2 low) and is late-onset neutrophilic and paucigranulocytic, as corticosteroids less responsive or resistant, hence also known as severe asthma, and inflammation is driven through T helper 1 (Th1), T helper 17 (Th17) cells, neutrophils, and IL-1β, IL-8, IL-16, IL-17A/F, IFN-γ, and TNF-α. However, these T2 and non-T2 profiles are not always strictly age limited, as children and adults may have both with mixed steroid responses. Adult-onset/old age asthma is generally Th17 predominant, neutrophilic, and severe in nature with MBD2 association [3, 4, 136, 138–144]. The fluctuation curve of sex hormones in the whole life span is shown in the figure. Male sex hormone is beneficial in asthma, as the protective effect from puberty to adulthood reverses in old age due to the reduced androgen level [80]. The expression of MBD2 in humans, as we believe, is shown in figure. It is believed to be at the primary level from birth to adolescence and tends to rise at adulthood, with the plateau probably during old age. At this point, evaluating MBD2, Th17, and the functional role of sex hormones and their correlative association, we believe that male sex hormone regulates the expression of MBD2 and is beneficial in treating Th17 predominant neutrophilic severe asthma of the second peak. ICS: inhaled corticosteroids; FeNO: fractional exhaled nitric oxide; IgE: immunoglobulin E; MBD2: methyl-CpG-binding domain protein 2.

Similarly, the increase of male sex hormone is beneficial in asthma, as the protective effect from puberty to adulthood reverses in old age is believed due to the reduced level of androgen [80], and androgen supplementation induces bronchodilatation [146], decreases Th17 (IL17) expression, and reduces the neutrophilic asthma prevalence and exacerbations. This shows the direct correlation of MBD2 with Th17 and an inverse correlation of sex hormones with Th17 and possibly with MBD2. Does androgen downregulate the MBD2 expression in severe asthma (Figure 4)?

A study has shown that BECs of mice regulate the differentiation of Th2 and Th17 cells [147]. TES relaxes the tracheal smooth muscle [146], but the TES action is annulled with the removal of epithelium, justifying the TES action.
through the epithelium-dependent manner [148]. This shows that androgen possibly affects downregulation of Th17 via MBD2 in BECs, showing BECs as a cellular target in treating severe asthma. However, we need further cellular studies in this aspect.

Through evaluating different studies and reviewing the association and correlation of MBD2, Th17, and sex hormones in non-T2 asthma, we postulate that male sex hormones possibly downregulate the MBD2 expression resulting in decreased MBD2 mediated Th17 differentiation and might have the therapeutic potential.

14. Conclusion and Perspectives

With a better understanding and continuous research on asthma etiology and pathophysiology, asthma heterogeneity and endotypes are exclusively explored. T2 asthma, childhood predominant, is better recognized, and several targeted monoclonal antibodies are also identified for therapy. However, non-T2 asthma, usually adult-onset/old age predominant, is still obscure with neutrophilic, paucigranulocytic, or mixed subtypes, where inflammation is driven through varieties of asthma-related inflammatory cells, and with airway muscular, neural, or comorbidities associated [3, 4]. Non-T2 asthma is less responsiveness with ICS, hence also called severe asthma. However, these profiles also show mixed steroid responses and are not always strictly age-dependent.

Th17 is the kernel of severe asthma, though proinflammatory cytokines, oxidative stress, and neuronal and hormonal responses, and epigenetic regulation are also involved in the pathogenesis.

MBD2 is an essential mediator of asthma epigenetics because it can bind the target gene, induces posttranscriptional histone modification, interprets DNA methylation, and changes the chromatin structure. MBD2, a Th17 regulator, is involved in the pathogenesis of Th17 predominant neutrophilic severe asthma justified by different studies, and asthma severity and MBD2 expression are correlated. That is why MBD2 has the possibility of a therapeutic target in severe asthma.

Sex hormone fluctuates during the normal physiobiological stages of the human body and affects the asthma outcome. Several internal and external etiopathobiological factors confound the effect of sex hormones and affect the asthma severity, prevalence, remission, and outcome. Male sex steroid tends to show a beneficial effect in severe asthma, and fluctuation is related to asthma prevalence. Boys are at higher risk of asthma prevalence before puberty, but with the increase in androgen level and pronounced airway caliber at puberty onset, male are better protected from asthma until adulthood, as females’ prevalence increases at this stage. Later, the tendency of a high prevalence of female adulthood asthma shifts to male at old age due to decreased level of androgen postulating the role of decreased level of male sex hormone in old age Th17 predominant severe asthma.

Is there any downregulatory response on MBD2 by androgens and the role of male sex hormone in MBD2 mediated Th17 predominant severe asthma of old age? This query needs extensive epigenetic and molecular-cellular studies. However, by evaluating the MBD2, Th17, and sex hormone association from various severe asthma studies, the MBD2 expression and Th17 differentiation directly correlate. In contrast, male sex hormone is inversely correlated with MBD2 and Th17 in old age asthma, postulating that the male sex hormone possibly downregulates the MBD2 expression resulting in decreased Th17 differentiation.

Together, given of all studies, we can conclude that evaluation of sex hormone therapeutic potential is a valuable novel approach to solve mysteries surrounding severe asthma therapeutic issues that will guide us towards discovery of new therapeutic agent, reduces the asthma-related complications, and promotes long-term survival that will ultimately reduce the asthma related mortality probably via mechanism distinct from corticosteroid treatment.

Abbreviations

AR: Androgen receptor
ASM: Airway smooth muscle
BECs: Bronchial epithelial cells
BHR: Bronchial hyperresponsiveness
DHEA: Dehydroepiandrosterone
DHT: Dihydrotestosterone
E2: Estradiol
ERs: Estrogen receptor
FEV1: Forced expiratory volume in one second
FVC: Forced vital capacity
HRT: Hormone replacement therapy
ILC2: Type 2 innate lymphoid cells
MBD2: Methyl-CpG-binding domain protein 2
M2: Macrophage
NO: Nitric oxide
OCP: Oral contraceptive pills
PC20: Provocative concentration 20% fall in FEV1
PRs: Progesterone receptor
TES: Testosterone
Th17: T helper 17 cells
17β-E2: 17β-estradiol.

Additional Points

Key Points. T2 asthma (T2 high) and non-T2 (T2 low) are the common asthma endotypes. T2 asthma is better known, while non-T2 is still obscure. Non-T2 asthma is generally neutrophilic, paucigranulocytic, or mixed types with a reduced response (resistance) to ICS (that is why also known as severe asthma) and predominantly affects late adulthood and old age population. Th17 is the kernel of severe asthma because of its important involvement in the pathogenesis of severe asthma. Proinflammatory cytokines, oxidative stress, neuronal and hormonal responses, and epigenetic regulation are also involved in Th17 cell-mediated severe asthma, but the underlying mechanisms are not well understood. MBD2 can mediate asthma epigenetics by interpreting DNA methylation and inducing posttranscriptional histone modification through binding the target gene to change the
chromatin structure and also affects the Th17 expression in severe asthma, showing the possibility of therapeutic target in severe asthma. Sex hormone fluctuates during the normal physiobiological stages of the human body and affects the asthma severity, prevalence, remission, and outcome. Several internal and external etiopathobiological factors also affect the fluctuation status and function of sex hormones. Boys before puberty and old aged male tend to have higher asthma prevalence, which is related to low and decreased androgen level, respectively. Similarly, girls from puberty to adulthood have higher prevalence of asthma that is associated with their hormonal status. Male sex hormone seems to be beneficial in asthma from above statements. The MBD2 and Th17 expression is directly correlated, while male sex hormone is inversely correlated with MBD2 and Th17 in severe asthma. We expect androgen possibly down-regulates the MBD2 expression resulting in decreased Th17 differentiation. Androgen therapy can be valuable novel approach to solve mysteries surrounding severe asthma treatment issues that will guide us towards discovery of new therapeutic agent, reduces the asthma related complications, and promotes long-term survival that will ultimately reduce the asthma-related mortality via novel mechanism distinct from corticosteroid treatment. Queries for future. Role of androgen and inhaled/systemic corticosteroid combined therapy in severe asthma. What will be the potential effect of androgen therapy, as androgen itself changes to hormones with estrogenic properties in some extent? What will be the future of triple therapy with sex hormones, corticosteroids, and peroxisome proliferator-activated receptors γ (PPAR γ) derivatives as they share the near same biochemistry structure?

Consent

This article does not contain any studies with human or animals performed by any of the authors.

Conflicts of Interest

The authors declare no conflicts of interest regarding the publication of this paper.

Authors’ Contributions

Binaya Wasti and Zhifeng Chen contributed equally to this work.

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