Menstruation is defined as “a woman’s monthly bleeding from the reproductive (vaginal) tract, as a consequence of cyclical changes in hormonal activity”1. Menstruation is a physiological process that, for most women, is a routine part of life. However, menstrual abnormalities are estimated to affect up to one third of women globally and can have a debilitating impact on an individual’s quality of life1–3. Menstrual abnormalities have become increasingly prominent in the latter half of the 20th century, probably due to increased prevalence and recognition4. However, menstrual complaints remain under-reported4.

Currently, women living in high-income countries, for example, the USA or UK, are estimated to have approximately 400–450 ovulations in their lifetime. This estimate is based on the average age at menarche (12.5 years), average age at menopause (50.5 years), age at first birth (29 years), number of children conceived per lifetime (1.8 children per woman) and the average duration of breastfeeding (4 months)5,6. In contrast to these figures, research estimates that hunter–gatherer women of the Palaeolithic era experienced approximately 160 ovulations per lifetime, due to later average age at menarche (16 years) and earlier menopause (47 years). Additionally, women from this era had their first child at an earlier age (19 years), conceived more children over their lifetime (six children on average) and breastfed for much longer (2–3 years per child) than women living now7. A previous study estimated that women in the Palaeolithic era had no more than 40 ovulations per lifetime based on different populations, with different family sizes and length of lactational amenorrhoea7.

In this Review, we define typical menstruation using a standardized system developed by global experts and approved by the International Federation of Gynecology and Obstetrics (FIGO) Menstrual Disorders Committee (MDC). This system facilitates the definition and standardization of what constitutes abnormal uterine bleeding (AUB: FIGO Systems 1 and 2)2. We address the considerable impact of non-malignant AUB across the globe before discussing how delineating the physiology of menstruation could inform and improve the management of non-malignant AUB by optimizing diagnosis and treatment. This Review focuses on the biomedical aspects of menstruation, while acknowledging that cultural, social, environmental and political factors have a notable influence on menstrual health8. Comprehensive discussion of these important non-biomedical aspects can be found elsewhere and are therefore not discussed in detail here9.
Key points

- Menstruation is a phenomenon of repeated tissue injury and repair that is a fine balance between proliferation, decidualization, inflammation, hypoxia, apoptosis, haemostasis, vasoconstriction and, finally, repair and regeneration.
- The endometrium is a dynamic, multicellular tissue highly responsive to sex steroids; subtle variances in the endometrial environment and, therefore, functioning, can lead to abnormal uterine bleeding (AUB).
- AUB is a debilitating symptom that affects up to one third of reproductive-aged women; comprehensive knowledge of menstrual cycle physiology is crucial for understanding and progressing endometrial physiology research.
- There is a high prevalence of iron deficiency and iron deficiency anaemia in those with AUB, on a global scale, and this is often under-recognized and under-reported.
- The terminology and definitions for diagnosing causes of AUB are now standardized in the International Federation of Gynecology and Obstetrics Systems 1 and 2, and should be followed for ease of clinical and research synchrony.
- Treatments for AUB are not specific and a third of patients resort to a hysterectomy for resolution of symptoms, highlighting a clinically unmet need for more targeted and personalized treatments.

Abnormal uterine bleeding
Menstrual bleeding should be assessed using six parameters: frequency, duration, regularity, volume, the presence or absence of intermenstrual bleeding and unscheduled bleeding (in those individuals receiving progestin with or without oestrogen)(2,10). Typical physiological menstruation occurs every 24–38 days, lasts less than 8 days and has a longest to shortest cycle variation of less than 7–9 days. Additionally, the volume of typical menstrual bleeding should not interfere with quality of life. AUB is the widely accepted umbrella term for menstrual abnormalities and is subdivided into either acute or chronic(1,5). Chronic AUB is “bleeding from the uterine corpus that is abnormal in volume, regularity, and/or timing, and has been present for the majority of the past 6 months”(15). Acute AUB is “an episode of heavy bleeding that, in the opinion of the clinician, is of sufficient quantity to require immediate intervention to prevent further blood loss”(16). Acute AUB can occur in the presence of chronic AUB or as an independent episode. AUB can be frequent or infrequent, prolonged, irregular or heavy. Heavy menstrual bleeding (HMB) is defined as “excessive menstrual blood loss which interferes with a woman’s physical, social, emotional and/or material quality of life”(11). Intermenstrual bleeding is the spontaneous bleeding occurring between menstrual cycles that can be either cyclical or random(15). This Review focuses on the biomedical aspects of conditions that lead to HMB.

Classification of AUB: PALM-COEIN
In 2011, the FIGO MDC set out the core classification of conditions that cause AUB, with minor updates in 2018 (REFS2,10). This classification is relevant for primary care physicians as well as specialists and researchers. The nine main categories are arranged in the acronym PALM-COEIN with each letter denoting a cause: Polyp, Adenomyosis, Leiomyoma (uterine fibroids), Malignancy, Coagulopathy, Ovulatory dysfunction, Endometrial, Iatrogenic and Not otherwise classified (FIG. 1; TABLE 1). Patients can therefore be said to have, for instance, AUB-P and/or AUB-I. The group of PALM causes relates to the structural causes of AUB, which are generally discrete entities that can be evaluated or measured.

Fig. 1 | FIGO System 2: the PALM-COEIN classification system. The causes of abnormal uterine bleeding are classified using the acronym PALM-COEIN, with each letter denoting a cause. The structural causes (denoted by yellow letters) are discrete entities and include polyp, adenomyosis, leiomyoma (uterine fibroids) and malignancy. The non-structural causes (denoted by green letters) are depicted for the illustration; however, they cannot be measured or imaged. They include coagulopathy, ovulatory dysfunction, endometrial, iatrogenic and not otherwise classified (for example, a caesarean scar defect)(15,16). Note that the figure is a schematic and is not to scale.

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- Menstruation is a phenomenon of repeated tissue injury and repair that is a fine balance between proliferation, decidualization, inflammation, hypoxia, apoptosis, haemostasis, vasoconstriction and, finally, repair and regeneration.
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- There is a high prevalence of iron deficiency and iron deficiency anaemia in those with AUB, on a global scale, and this is often under-recognized and under-reported.
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Iatrogenic (AUB-I) Medical interventions and devices, pharmacological agents that lead to AUB Dependent on cause Medical interventions and devices such as medicated intrauterine systems (for example, LNG-IUS) or inert intrauterine systems; pharmacological agents such as sex steroids, drugs influencing dopamine metabolism or ovulatory function, or anti-coagulants; it is important to decipher the origin of unscheduled and/or breakthrough bleeding

Not otherwise classified (AUB-N) A spectrum of conditions, only some of which can be evaluated or measured using imaging modalities or histopathology Dependent on cause Includes rarely encountered or ill-defined entities such as uterine arteriovenous malformations, lower segment or upper cervical niche (or an isthmocele) associated with a caesarean section scar defect

AUB, abnormal uterine bleeding; FIGO, International Federation of Gynecology and Obstetrics; HMB, heavy menstrual bleeding; LNG-IUS, levonorgestrel-releasing intrauterine system; PCOS, polycystic ovary syndrome.

The impact of AUB

AUB is a common debilitating symptom that is most often not life-threatening but is life-altering. The reported worldwide prevalence of AUB varies greatly from 3% to 52%, depending on the terminology used, as well as on whether assessments were subjective or objective. Subjective assessments are likely to result in a higher prevalence of AUB than objective assessments as they are based on the woman’s personal perceptions of her bleeding. Objective assessments are conducted either via menstrual diaries or by measuring the quantity of menstrual blood on used menstrual products using laboratory tests, for example the alkaline haematin method.

Subjective assessments can vary depending on cultural and societal viewpoints; therefore it can be difficult to
compare subjective assessments across countries, or even within particular patient demographic groups within a population. However, in view of the definition of HMB, the subjective assessment of a person’s bleeding is what is clinically important. The National Institute for Health and Care Excellence (NICE), Royal College of Obstetricians and Gynaecologists and FIGO estimate that approximately one third of women globally have AUB at some point in their reproductive years\(^2\). A 2019 survey suggested this could be an underestimation due to under-reporting\(^1\). In 2011, in the UK, it was estimated that in England and Wales approximately 1.5 million women were affected by HMB annually\(^17\). HMB is the fourth most common reason for referral to gynaecology services in the UK and approximately 20% of 1.2 million referrals to specialist gynaecology services concern women with HMB in the UK\(^12,18\).

On average, 30% of women globally with AUB are anaemic, and this figure increases to approximately 60% in South Asia, the eastern Mediterranean and Africa, probably due to the combination of dietary deficiencies, AUB and multiple pregnancies\(^19\). Iron deficiency is one of the commonest micronutrient deficiencies worldwide\(^20\). Anaemia is a global health issue that is under-recognized and under-reported in both low-income to middle-income countries and high-income countries\(^2\). In comparison to all other demographic groups, women of reproductive age are most likely to be affected by iron deficiency, which is responsible for approximately 51% of global cases of anaemia\(^2\). The prevalence of iron deficiency worldwide is approximately twice as high as iron deficiency anaemia (IDA)\(^22\). Iron deficiency and IDA lead to impaired oxygen transportation and enzyme reactions, which impacts almost all metabolic pathways in the body. The functional consequences of iron deficiency and IDA include fatigue, reduced exercise capacity, headaches and impaired cognition\(^23,24\). AUB therefore needs to be recognized as an important cause of iron deficiency and IDA.

**Box 1 | The impact of abnormal uterine bleeding**
- One third of women are affected by abnormal uterine bleeding (AUB) at some point in their reproductive years, leading to life-altering effects\(^1\).
- AUB affects all members of society; not only the menstruating individual.
- Physical and mental health effects of AUB include heavy menstrual bleeding, flooding accidents, discomfort, tiredness and depression.
- There is a high global prevalence of iron deficiency and iron deficiency anaemia associated with AUB, which is often under-recognized and under-reported.
- AUB can cause strains on relationships and have a negative impact on an individual’s social network.
- AUB can lead to social isolation due to mental and physical symptoms.
- AUB can have negative effects on work and job prospects, and can have financial implications.
- There is a high cost to the individual, which could be related to the menstrual products themselves, clothes and/or bedding that are damaged by flooding accidents, loss of earnings due to missed work, missed opportunities at work or school, and also treatment costs in countries where health care is privately funded.
- For society, the economic implications relate to the impact of AUB on a substantial proportion of the workforce, as well as the environmental impact of menstrual products, for example, which might have a broader economic impact.

Women with AUB have a health-related quality of life score below the 25th percentile compared with the general female population of the same age\(^25\). Menstrual health complaints have an impact on all aspects of an individual’s life (Box 1), including an increased risk of premature death\(^26\). People are more likely to seek medical advice for HMB for social and physical factors rather than psychological factors\(^1\). However, 50% of women with HMB and mental health concerns describe HMB as the cause of their anxiety and depression\(^27,28\). Structural causes of AUB are more prevalent with advancing reproductive age and might affect reproductive health. Women in high-income countries are, on average, choosing to delay childbirth. For example, in the UK, 50% of babies are born to women ≥30 years of age\(^29\). Therefore, structural causes of AUB, such as uterine fibroids, could have a negative impact on reproductive outcomes. These structural causes of AUB could lead to reduced likelihood of natural conception and less successful assisted reproductive techniques, or even contribute to complications of pregnancy\(^30–32\).

The socioeconomic burden of AUB is high, with total direct and indirect costs in the UK and USA being greater than £1.2 billion and $37 billion, respectively\(^33,34\). Period poverty affects many people due to work absence and cost of additional menstrual products when experiencing AUB. Treatment costs for AUB are likely to increase due to an increase in the number of people seeking medical treatment for the condition. Additionally, the UK NICE guidance for the investigation of AUB\(^1\) changed in 2018 to promote more patients being referred for hysteroscopic investigation, which will potentially contribute to rising costs\(^3\).

During the COVID-19 pandemic, when access to medical care, investigations and treatments were restricted and/or limited, adaptations to care provision for AUB and the deferral of surgical management were essential in ensuring safe and effective care\(^35,36\). In a post-COVID-19 world, it will be important that political commitment and public engagement do not ignore this often neglected area of women’s health\(^31\).

There have been no novel treatments in routine clinical use for the symptoms of AUB and/or HMB for 20 years, and no non-hormonal treatments for AUB and/or HMB for at least 30 years\(^37\). There are also no biomarkers in routine use to manage AUB\(^38\). Individuals presenting with the symptom of AUB often represent a heterogeneous group, resulting in considerable challenges in offering tailored treatments. High-quality research to fill the gaps in our knowledge of the physiology and pathology of the menstrual cycle is therefore critical to improving our understanding of the condition. This knowledge will enable appropriate phenotyping of patients in keeping with the principles of personalized precision medicine and will facilitate more specific treatments. In turn, these changes could improve women’s quality of life and avoid more invasive surgical management.

**Menstrual cycle physiology**

**The endometrium**

The endometrium is a multicellular tissue forming the lining of the uterus. The function of the endometrium is to either prepare for embryonic implantation and
maintain a pregnancy or, in the absence of pregnancy, to shed and repair, that is, menstruate. The uterine endometrium thus undergoes repetitive and physiological cycling of tissue injury and repair every month\(^{36}\). The endometrium is characterized by features of rapid repair without residual scarring or loss of function, similar to a fetus in utero\(^{37,38}\).

The human endometrium is composed of two layers. The upper two thirds, the functional layer, is shed during menstruation. The basal layer, the lower third of the endometrium (adjacent to the myometrium), does not shed during menses. The functional endometrium is composed of columnar surface epithelium, which covers a multilayered stroma. The stroma contains connective tissue with fibroblast-like stromal cells, a vasculature (including the specialized spiral arteries) and a population of tissue-resident endometrial immune cells, as well as a cyclical traffic of innate immune cells. The population of innate immune cells (both cyclical and tissue-resident immune cells) varies across the menstrual cycle, and these endometrial leukocytes play important roles in the breakdown and repair of endometrial tissue during menses\(^{36,40}\) (FIG. 2).

The endometrium is dynamic and highly responsive to the varying circulating levels of the sex steroid hormones ovarian 17β-oestradiol (oestradiol) and pregn-4-ene-3,20-dione (progesterone). The endometrium is exposed to a series of hormonal changes throughout the menstrual cycle, first to systemic oestradiol, then to oestradiol and progesterone and, in the absence of pregnancy, progesterone withdrawal. As a consequence of this sequential exposure, the endometrium adapts in form and function across the menstrual cycle. Understanding this tightly regulated endocrine environment could help identify new therapeutic strategies for AUB.

**The menstrual cycle**

The main phases in the menstrual cycle are the proliferative phase, secretory phase and menstrual phase. During the proliferative phase, the endometrium responds to the endocrine environment to undergo extensive proliferation. This phase is variable in length and oestradiol is the dominant hormone. At ovulation, the oocyte is released from the dominant ovarian follicle. The follicle then transforms into the corpus luteum, which secretes progesterone and leads to a progesterone-dominant secretory phase, typically lasting 14 days. During this phase, in the presence of high levels of oestradiol and progesterone, the cells of the endometrium undergo morphological and functional changes, a process known as decidualization. With the demise of the corpus luteum in the late secretory phase, progesterone and oestradiol levels decline, and the cells of the endometrium are no longer maintained. As a consequence of progesterone withdrawal, there is a coordinated spatial and temporal response in the upper functional layer of the endometrium that culminates in menstruation. Local cellular and molecular events include endometrial apoptosis, inflammatory mediator influx and the induction of matrix metalloproteinase (MMP) expression. The peri-menstrual phase encompasses endometrial transition from the secretory phase, through menstrual breakdown and repair, to regeneration in the proliferative phase. The ovarian equivalent of the peri-menstrual phase is the luteofollicular transition\(^{39,41}\) (FIG. 2).

**Hormonal influences**

The role of oestradiol and progesterone. The endometrium is a target tissue for steroid hormones and its form and function is entirely governed by the prevailing endocrine environment. Therefore, both endogenous and exogenous hormone exposure affects endometrial bleeding patterns. Oestradiol and progesterone induce their physiological effects in the endometrium primarily through their cognate receptors, the oestrogen receptor (ER) and the progesterone receptor (PR). These receptors, along with receptors for androgens, glucocorticoids and mineralocorticoids, belong to a superfamily of nuclear receptors that act as ligand-activated transcription factors\(^{42}\). Current medical treatments for AUB largely focus on hormonal management strategies, using a combination of oestrogen and progestins or progestins alone. Oestrogens and progestins can be delivered through oral preparations, whilst progestins alone can be delivered via medicated intrauterine devices or injections, or as oral forms\(^{43}\). The basic mechanism of action of oral combined hormonal regimens is the suppression of ovulation and oestrogen-mediated endometrial proliferation\(^{44}\). These treatments might not target the underlying cause of AUB; in fact, one in five women using progestin-only treatments might experience unscheduled bleeding (AUB-I)\(^{45}\). Women might also experience hormonal adverse effects due to off-target effects, such as bloating, breast tenderness, acne or mood changes\(^{46}\). Improved knowledge of the underlying mechanism of AUB symptoms, for example in AUB-E, might enable better targeting of the local endometrial hormonal environment and reduce off-target action and adverse effects\(^{47}\).

The PR is encoded in a single gene located on chromosome 11 (11q22-q23). The PR has two main isoforms, PR-A and PR-B. These isoforms are structurally almost identical, except that PR-A is truncated by 164 amino acids at the amino terminal segment compared with PR-B\(^{48}\). PR-A and PR-B act as transcriptional regulators of progesterone responsive genes. PR-A is the dominant isoform and, therefore, the primary mediator of progesterone action and an inhibitor of PR-B. The expression of PR in the uterus is stimulated by oestrogens via the α isoform of ER (ERα), and therefore progesterone responsiveness is dependent on the presence of an adequate oestrogen response\(^{49}\). PR expression exhibits well-described temporal and location expression patterns\(^{50}\). The PR is located in nuclei of the endometrial epithelial and stromal cells during the proliferative phase. Expression of PR persists in the nuclei of the stromal compartment of the functional layer of the endometrium during the secretory phase, particularly in the perivascular region\(^{47}\).

In the genomic signalling pathway, when progesterone (the PR ligand) binds with PR at its ligand-binding domain, a conformational change occurs and the progesterone–PR complex relocates to the nucleus.
**Fig. 2 | Menstrual cycle and ovarian cycle physiology with highlighted aspects of the structure of the endometrium.** The endometrium is a dynamic structure that adapts to the endocrine environment in a cyclic manner, on average every 28 days. It undergoes a process of proliferation during an oestradiol-dominant phase. Following ovulation, the corpus luteum secretes progesterone, leading to a progesterone-dominant stage during which the endometrium decidualizes. In the absence of pregnancy, after the demise of the corpus luteum and progesterone withdrawal, the endometrium is shed during menstruation. This process requires the remaining cells to repair and regenerate without injury or scarring so that the menstrual cycle can repeat. As a multicellular tissue, the endometrium is highly responsive to the endocrine environment. The lower third of the endometrium, adjacent to the myometrium, is known as the basal layer and the upper two thirds, including the luminal surface, as the functional layer. Under the influence of changing levels of oestradiol and progesterone, the cellular structure of the endometrium adapts. The main cellular components within the endometrium are the epithelial cells, stromal cells, vascular cells and a variety of innate immune cells. The numbers of immune cells vary according to the cycle stage (see Figure key). The functional layer is shed during the menstrual phase, leaving behind a denuded basal endometrium. The peri-menstrual phase (also known as the luteo-follicular transition) is the time after which progesterone and oestradiol levels fall, menstruation occurs and the endometrium transitions from a secretory to a proliferative state⁹⁶,⁹⁷. Note that the figure is a schematic and is not to scale.
Two separate ER isoforms have been identified: ERα, encoded by the ESR1 gene, and ERβ, encoded by the ESR2 gene. Expression of variant isoforms of both ER subtypes in the human endometrium, formed by translation of mRNAs generated by alternative splicing of the ESR1 and ESR2 genes, have also been described. ESR1 (ERα) is located on chromosome 6, whilst ESR2 (ERβ) is found on chromosome 10. Oestrogen is a ligand for the ER and downstream effects of oestrogen signalling, such as cellular proliferation and differentiation, occur in a similar manner to progesterone binding with PR. ERα and ERβ are both expressed in the glandular epithelium and the stroma in the proliferative phase, which is when the peak concentrations of ER are observed. During the secretory phase, ERα levels decline in both the endometrial glandular epithelium and stroma. ERβ levels also decline in the glandular epithelium in the secretory phase; however, ERβ expression is maintained in the endometrial stroma.

Sex steroid hormone biosynthesis. The availability of biologically active sex steroids within the endometrium is dependent on the action of local endometrial sex steroid-metabolizing enzymes (Fig. 3). Locally produced sex steroids can exert their action in the cells in which they are synthesized, without release into the extracellular space, a concept known as intracrinology.

Fig. 3 | Major biosynthetic pathways in steroidogenesis. The precursor of all steroid hormones is cholesterol. Steroidogenesis (synthesis of steroid hormones) occurs predominantly in the adrenal glands and gonads (ovaries and testes); however, it also occurs in the placenta. The key enzymes are located in the mitochondria and the endoplasmic reticulum of the cell. The first step in steroidogenesis involves the conversion of cholesterol to pregnenolone (a non-hormonal intermediate) within the cell mitochondria by the cholesterol side chain cleavage (CSCC) enzyme. The major classes of steroid hormones include progestogens, androgens, oestrogens and corticosteroids (glucocorticoids and mineralocorticoids). All steroid hormones have a cyclopentanoperhydrophenanthrene ring, and are further classified based on the number of carbon atoms. Pregnenes have 21 carbon atoms and are called C21 steroids (progestogens and corticosteroids). Oestradiol has 18 carbon atoms (C18 steroids) and include oestrogens, androstanes have 19 carbon atoms (C19 steroids) and include androgens. OH, hydroxylase; HSD, hydroxysteroid dehydrogenase; DHT, dihydrotestosterone; DHEA, dehydroepiandrosterone; DHEAS, dehydroepiandrosterone sulfate.
The local endocrine milieu plays a crucial role in the regulation of endometrial function. For example, pharmacological manipulation of the PR with SPRM treatment could result in molecular and cellular modifications in local sex steroid availability in the endometrium, as well as changes in the availability of local sex steroid metabolizing enzymes. These modifications could result in the creation of a local endometrial oestrogenic microenvironment. Similarly, local delivery of a progestin could consequently also modify the local endometrial steroid microenvironment. For example, the progestin levonorgestrel, which can be delivered via an intrauterine system, modulates endometrial PR expression.

The intracrinology of the human endometrium more generally was comprehensively reviewed in 2018 (ref.62). Endometrial steroid hormone metabolism studies, which extend over nearly 50 years, initially demonstrated steroid sulfatase activity, and these were followed by studies concerning oxidative and reductive 17β-hydroxysteroid dehydrogenase (17β-HSD) activities63. These steroid-metabolizing enzymes are involved in endometrial steroid hormone (ligand) availability and thus are involved with modulation of the local endometrial hormone environment. De novo steroid biosynthesis requires cholesterol, which is shuttled to the inner mitochondrial membrane by steroidalogen acute regulatory protein, where it undergoes side chain cleavage by CYP11A1 (a member of the cytochrome P450 superfamily of enzymes that is involved in lipid synthesis) to yield pregnenolone. Pregnenolone undergoes two enzymatic conversions mediated via CYP17A1, first to 17α-hydroxypregnenolone (17α-hydroxylase activity) and then, following 17,20 lyase action, to dehydroepiandrosterone (DHEA). Expression of steroidalogen acute regulatory protein, CYP11A1 and CYP17A1 has been reported in the endometrium63,64. However, no study has identified definitive de novo steroidogenesis from normal endometrial tissue. DHEA is normally synthesized in the ovary and adrenal glands and released into the circulation.

The endometrium exhibits both oxidative and reductive 17β-HSD activities and the expression of 17β-HSD1, 2, 4, 6, 7, 8, 10, 12 and 14, and 17β-HSD5 (also known as aldo–keto reductase family member C3, AKR1C3) are detected at the mRNA or protein levels. The endometrium also has the capacity to metabolize sulfated compounds and DHEA to form androgens and oestrogens62 (FIG. 3). DHEA is converted to androstenedione by 3β-HSD, which is expressed in endometrial stromal cells. The steroid-metabolizing enzyme, 17β-HSD2, is responsible for conversion of oestradiol and testosterone to oestrone and androstenedione (which has a reduced biological activity), respectively, and conversion of 20-hydroxyprogesterone to biologically active progesterone. The endometrial steroid-metabolizing enzyme 17β-HSD5 has the opposite effects. A balance between the activity of these major steroid-regulating enzymes and others (such as aromatase) play key roles in the local endometrial milieu and endocrine regulation of menstruation. Studies have indicated that serum concentrations of steroid hormones differ from local endometrial concentrations of these hormones and are consistent with local enzyme regulation.

The role of androgens and glucocorticoids. As well as oestradiol and progesterone, androgens65 and glucocorticoids66 play an important role in the regulation of endometrial function. Therefore, hormone equivalents or drugs that modulate the androgen receptor or glucocorticoid receptor could also act as potential therapeutic targets in AUB management. Androgen receptor expression varies throughout the menstrual cycle in the functional layer of the endometrium, both in the stroma and epithelial cells. The androgen receptor is predominantly expressed in the endometrial stroma during the proliferative phase, with reduced expression during the secretory phase. Androgen receptor expression has been observed in the endometrial glands in the late secretory or decidualized phase66. Androgens have been found to impact the proliferation, migration and survival of stromal cells within the endometrium in humans66. Using a mouse model of simulated menses, an injection of the androgen dihydrotestosterone at the time of progesterone withdrawal led to delayed repair of the luminal epithelium, indicating that androgens are involved in impairment of endometrial repair and could be involved in altered menstruation66. Androgens have an antiproliferative effect on both endometrial epithelial and stromal cells66. Androgens have been shown to upregulate decidualization markers on human endometrial stromal cells in vitro and might have a role to play in wound healing66,67. However, the role of androgens in wound healing is not fully understood in humans. In a mouse model of simulated menses, androgens have been shown to alter the expression of MMPs and caspase 3, altering the balance of apoptosis and repair within the endometrium, and therefore endometrial integrity68. The menstruating endometrium is an example of a wounded mucosal surface that rapidly repairs64,68.

Altered androgen signalling has also been identified in endometrial cancer and endometriosis. Insights from 2018 suggest that androgens might have dual roles in the endometrium and could modulate endometrial function by acting directly on the androgen receptor and acting indirectly as precursors to local oestrogens65. The development of selective androgen receptor modulators71 might identify new therapeutic targets for treating AUB.

Glucocorticoids have an anti-inflammatory effect that is mediated by cortisol binding to the glucocorticoid receptor in the stromal, epithelial and uterine natural killer cells72. Cortisol also inhibits angiogenesis by interacting directly with glucocorticoid receptors on vascular endothelium73. The enzyme 11β-HSD1 converts the less potent cortisone into cortisol, a biologically active molecule, and the reverse reaction is catalysed by 11β-HSD2 (ref.74). 11β-HSD1 activates cortisol, and is upregulated at the time of menstruation, which coincides with the maximal endometrial levels of the glucocorticoid receptor, which enables cortisol to have maximal effect75. 11β-HSD2 inactivates cortisol and women with objectively assessed HMB have been reported to have a 2.5-fold increase in levels of 11β-HSD2 compared with women without HMB76. This increase in levels of...
Decidualization

Decidualization is a progesterone-dependent process that ensures the endometrium adapts from a proliferative phenotype to one that will nurture and support a pregnancy. It is a spontaneous occurrence in the human endometrium as it does not require the presence of an embryo. Decidualization occurs due to a rise in progesterone secreted by the corpus luteum following ovulation and the production of cAMP throughout the endometrial stroma. In the absence of embryo implantation, progesterone levels fall and spontaneous decidualization leads to menstrual shedding and cyclic regeneration of the endometrium.

During decidualization, progesterone targets the endometrial stromal cells, resulting in their transformation from an elongated to a rounded morphology. In addition, progesterone induces the expression of progesterone-dependent proteins, such as prolactin, glycogen, tissue factor, insulin-like growth factor binding protein 1 and forkhead box protein O1. This process leads to a unique genetic signature within the individual cell types of the endometrium (stromal fibroblasts, endothelial cells, macrophages, lymphocytes and epithelial cells), and, thus, the unique genetic signature could be a plausible biomarker of endometrial receptivity and progesterone response.

‘Progesterone resistance’ is a term used to describe an abnormal endometrial response to the post-ovulatory increase in progesterone levels, and often manifests as a deficient decidualization response. Progesterone resistance has been described in patients with adenomyosis and it is reported that ERβ expression and PR expression are perturbed in patients with HMB and adenomyosis, which is in contrast to expression levels in those women with HMB and no evidence of adenomyosis. It is interesting that lower levels of HOXA10 expression (a progesterone-regulated gene) have been reported in the secretory endometrium of women with HMB and uterine fibroids than in the secretory endometrium of women without uterine fibroids and with normal menstrual bleeding, suggesting that women with uterine fibroids might have developed progesterone resistance. A blunted or inadequate response to progesterone might therefore interfere with decidualization and expression of progesterone-regulated genes. Progesterone is responsible for endometrial differentiation and decidualization in an oestradiol primed endometrium, and dysregulation of progesterone signalling might therefore contribute to AUB.

Appropriate endometrial cell proliferation and decidualization results in primed decidualized stromal cells that are sensitive to their hormonal environment. Decidualization has been suggested to be a prerequisite for menstruation as spontaneous decidualization does not occur in non-menstruating animals. However, the impact of the decidualization process on menstrual blood loss parameters remains to be determined.

Menstruation

Initiation of menstruation. Menstruation occurs after the demise of the corpus luteum and the withdrawal of progesterone and oestradiol, triggering local endometrial apoptosis, inflammation, hypoxia and tissue breakdown. There are two components to the processes underlying the menstrual phase, the first being progesterone-dependent and the second being progesterone-independent. Once progesterone, an anti-inflammatory hormone, is withdrawn, the endometrium is exposed to a local environment rich in cytokines and prostaglandins. The generation of inflammatory mediators is coordinated by the actions of the NF-κB transcription factor, which regulates genes involved in the inflammatory process. During the progesterone-dominant secretory phase of the menstrual cycle, progesterone inhibits NF-κB, but with progesterone withdrawal in the late secretory phase, this inhibitory effect is no longer present. NF-κB subsequently degrades its inhibitor, IκBα, which leads to a local endometrial increase in cytokines, chemokines and MMP production.

Inhibition of MMP activity in response to circulating oestradiol and progesterone maintains the secretory endometrium. This inhibition is reversed after withdrawal of progesterone, and the reversal of the inhibition of MMP activity upregulates pro-MMPs, at least in part via NF-κB activity. Uregulation of MMPs and associated menstruation is related to progesterone withdrawal rather than oestrogen decline in the late secretory phase of the menstrual cycle. Apoptosis has been demonstrated to occur in the glandular epithelium and then the stromal compartment prior to endometrial shedding in human endometrial samples from the late secretory phase, after progesterone withdrawal. This observation was recapitulated in a mouse model of menstruation, providing support for this preclinical model in menstrual research.

As a result of increased levels of cytokines and chemokines in the endometrium, there is an influx of innate immune cells within the endometrium, activation of MMPs and degradation of the extracellular matrix. Using carefully timed insertion and removal of progesterone implants in non-human primate and
mouse models of menstruation, it has been demonstrated that there is a threshold within the menstrual phase when the endometrium is no longer responsive to the anti-inflammatory effects of progesterone and menstrual bleeding is inevitable\(^{100-102}\). Disruptions in these pathways can lead to increased inflammation, resulting in AUB-E\(^{2}\). Progesterone withdrawal, and the subsequent inflammatory cascade, culminates in menstruation\(^{36}\). Destruction and shedding of the functional layer (bleeding as menstruation) ensue in a piece-meal fashion, leaving a denuded endometrial surface that undergoes concurrent repair\(^{39}\).

Control of endometrial bleeding. In order for tissue regeneration to occur, three factors must coordinate to ensure a timely cessation of menstrual bleeding. First, vasoconstriction of spiral arterioles must occur. The efficient vasoconstriction of endometrial arterioles is necessary to limit menstrual blood flow. According to the Poiseuille equation, a small increase in the vessel radius will dramatically increase flow through the vessel\(^{15}\). For example, a twofold increase in vessel radius leads to a 16-fold decrease in resistance to flow. Second, an effective haemostatic response is required, including repair of damaged vasculature\(^{36}\). Third, appropriately timed re-epithelialization of the denuded basal endometrium is necessary\(^{100-102}\).

Decreased vasoconstriction during menstruation could be due to abnormal production of vasoactive factors following progesterone withdrawal. Examples of these factors include a decreased PGF\(_{2\alpha}\) to PGE\(_2\) ratio, decreased prostaglandin F receptor expression, increased PGE\(_2\) production, decreased levels of endothelin 1 (a potent vasoconstrictor) and increased levels of neural endopeptidase, the enzyme that metabolizes endothelin 1. Spiral arterioles in women with HMB are larger in diameter than those in women without HMB due to altered maturation throughout the menstrual cycle. This altered maturation could be due to increased expression of VEGFA and VEGF receptors 1 and 2 (Ref\(^{38}\)). Women with HMB have reduced vascular smooth muscle (VSM) cell proliferation in spiral arterioles compared with women without HMB\(^{16}\). This reduced proliferation might affect vascular tone and VSM contractility, thereby reducing the hypoxic response and leading to excessive bleeding (AUB-E).

Evidence indicates that the alterations in VSM maturation in women with HMB are linked to reduced myosin heavy chain expression in the spiral arterioles in the early secretory stage. This reduced expression might, in turn, be correlated with expression of specific VSM mitogens (for example, endothelin\(^{100}\)) in the endometrium in women with HMB. Further supporting the importance of endometrial vascular maturation in reducing menstrual blood loss is the identification of roles for the growth factor transforming growth factor β1 (TGFβ1). TGFβ1 is involved with phenotype switching of VSM cells to a contractile phenotype and is also required for repair of the denuded menstrual endometrium\(^{101}\). There is evidence of decreased TGFβ1 bioavailability and signalling in the endometrium of women with HMB\(^{100-102}\), suggesting that TGFβ1 signalling and bioavailability could be potential therapeutic targets for the treatment of HMB in the future.

The role of hypoxia. The presence of hypoxia within the endometrium was proposed 80 years ago after the observation of transplanted endometrium on the anterior chamber of the eye of rhesus macaques\(^{36}\). Transplanted endometrium was observed for a combined total of 432 menstrual cycles, especially during menstruation, and progesterone withdrawal and intense vasoconstriction in the spiral arterioles was considered to induce hypoxia. The presence of hypoxia during endometrial breakdown and repair has also been demonstrated in a mouse model of simulated menstruation\(^{104-106}\). Mice were also placed in hypoxic conditions following progesterone withdrawal, which prevented the development of hypoxia during menstruation. This lack of hypoxia resulted in delayed endometrial repair\(^{101}\). This finding is consistent with hypoxia being necessary for efficient endometrial repair at menses.

Hypoxia-inducible factor (HIF) is composed of an α and a β subunit and is the master regulator of the cellular response to hypoxia\(^{107}\). In normoxia, the α subunit is rapidly degraded by the proteasome and HIF is inactive. Degradation of HIF1α is prevented under hypoxic conditions. HIF1α can then bind to its β subunit and increase transcription of target genes containing a hypoxia response element. This increase in transcription initiates the expression of host of factors involved in metabolism, angiogenesis and tissue repair, including VEGF\(^{108}\). HIF1α is only present in the human endometrium around the time of menstruation\(^{106,109}\). Women with HMB had decreased expression of endometrial HIF1α protein and downstream target genes at menstruation compared with those with normal menstrual blood loss. In addition, women with HMB bled for 2 more days during menstruation than women without HMB\(^{107}\). A mouse model of menstruation with a genetic reduction in HIF1α revealed statistically significantly delayed endometrial repair following menstruation. This finding was consistent with HIF1α having a key role in the hypoxia-mediated menstrual repair process. The delayed repair in a non-hypoxic mouse model of menstruation could be rescued with a pharmacological compound that stabilizes HIF1α, identifying a potential non-hormonal therapeutic target in women with AUB-E\(^{109}\).

Endometrial haemostasis and the coagulation system. Endometrial fibrinolysis plays an important role in normal menstruation and limitation of menstrual blood loss. At menses, disruption of blood vessels after ‘endometrial injury’ results in adherence of platelets to collagen on an injured basement membrane\(^{16}\). Platelet aggregation prompts a coagulation cascade. The coagulation cascade is activated by two pathways: extrinsic and intrinsic. Each cascade results in the conversion of factor X to Xa, which catalyses the conversion of pro-thrombin to thrombin, ultimately leading to the formation of a fibrin clot\(^{110}\). The fibrinolytic system mediates degradation of the fibrin clot. Fibrinolysis involves conversion of plasminogen to active plasmin, promoting the degradation of fibrin deposits.
Plasminogen is converted into plasmin by tissue plasminogen activator during fibrinolysis. Plasmin promotes the degradation of fibrin deposits, which leads to the formation of D-dimers. Menstrual blood contains a high concentration of D-dimers and this component of menstrual blood could be a tool to distinguish menstrual from peripheral blood for forensic purposes.117 Disorders involving altered systemic haemostasis severely affect menstrual blood loss (AUB-C)118. Tissue plasminogen activator and urokinase plasminogen activator orchestrate production of plasmin. Plasminogen activator inhibitor 1 (PAI1) inhibits fibrinolytic activity and has important regulatory functions in haemostasis, turnover of the extracellular matrix and cell adhesion.119

Human endometrium expresses tissue plasminogen activator, urokinase plasminogen activator, PAI1 and the urokinase plasminogen activator receptor115,116. Levels of the PAI1 protein and mRNA are reportedly increased by progesterone and TGFβ1 (REF 115). As discussed in a previous section, decreased TGFβ1 activity might hinder repair of the denuded menstrual endometrium, leading to HMB106,111. Women with HMB are reported to exhibit raised levels of tissue plasminogen activator activity on day 2 of menses, possibly due to over activity of the endometrial fibrinolytic system115. Such observations support a biological basis for the utility of anti-fibrinolytic drugs (such as tranexamic acid) to reduce menstrual bleeding. Such anti-fibrinolytic medications have been used as a therapy for HMB for the past 30 years116.

Tissue repair and regeneration
Tissue formation and remodelling of the endometrium commences alongside menstruation, with the epithelium forming a new luminal surface. These processes occur prior to stromal cell proliferation of the next menstrual cycle. The processes involved in endometrial repair and regeneration have not yet been fully delineated. Three theories exist as to how the endometrium repairs and regenerates.

The first theory is that epithelial cells, from the exposed stumps of endometrial glands (mainly in the basal layer) or from any intact epithelium bordering the denuded areas of stromal tissue, proliferate to create a new functional luminal layer119,127. Another theory is that endometrial stromal cells might be reprogrammed at menstruation to lose their mesenchymal cell characteristics and achieve epithelial cell characteristics through a process called mesenchymal to epithelial transformation (MET). Evidence for MET was found in a mouse model of simulated menses and suggests that both the residual basal layer and the unshed functional layer of the endometrium lead to the re-epithelialization of the remaining denuded epithelium. However, lineage tracing in adult mice showed no definitive evidence for MET in the uterine epithelium118. The third theory of how endometrial repair and regeneration occurs after menstruation is that endometrial stem cells and progenitor cells in the basal layer differentiate into stromal, glandular or luminal epithelial cells during the regeneration process.

Evidence for the existence of endometrial stem cells comes from both human and mouse studies119,122. Bone marrow-derived stem cells have also been found to supplement the endometrial stem cell population and assist in the regeneration of the endometrium121,124. However, their contribution to endometrial regeneration remains controversial, with chimera mouse models revealing no contribution to endometrial cell lineages. These findings suggest that previously identified bone marrow cells in the uterus were misidentified, possibly due to suboptimal staining and imaging of the CD45+ macrophages123–127. Further research to delineate this fundamental endometrial repair process in health and disease is essential to facilitate future pharmacological interventions to limit menstrual blood loss.

The myometrium in menstrual bleeding
The myometrium is composed of smooth muscle that contracts in different directions dependent on the phase of the menstrual cycle. It contracts in an antegrade (fundus to cervix) manner at the time of menstruation to extrude menstrual debris, and then retrograde (cervix to fundus) myometrial contractions might be required to facilitate sperm transport towards the uterine cavity. Myometrial smooth muscle cell contractility decreases with age, suggesting that ageing causes a functional change to the cells128. These myometrial contractions, also known as endometrial waves, originate mainly from the sub-endometrial layer of the myometrium. In the early follicular phase, endometrial waves occur once or twice a minute, last 10–15 s and have a low amplitude. During ovulation, this frequency increases to three to four times per minute. In the luteal phase, the frequency and amplitude of endometrial waves decreases to facilitate embryo implantation. In the absence of implantation, their frequency remains low, but their amplitude increases dramatically129.

Pathological structural features commonly present in the myometrium are uterine fibroids and adenomyosis. The aetiopathogenesis of uterine fibroids is complex and poorly understood. Genetic factors, epigenetic factors, endocrine disrupting chemicals, dietary deficiencies (such as vitamin D deficiency), age, race, ethnicity, family history, ovarian steroids, cytokines and growth factors have all been implicated in the development of uterine fibroids, amongst other factors130,134. The pathogenic mechanisms involved in the formation of adenomyosis are still to be elucidated. The most common theory underpinning the development of adenomyosis is that the basal endometrium invaginates into the myometrium via an altered or interrupted junctional zone132,133. Alternative hypotheses include the mechanism of “Tissue Injury And Repair” (TIAR)134. It is known that the endometrium is ‘injured’ during the process of menstruation but it is unknown exactly how the repair processes occur. During the process of TIAR at the endometrial to myometrial interface adenomyosis might develop as a result of a problem in these TIAR mechanisms and endometrial cells might become incorrectly located in the myometrium134,135. A third hypothesis in the development of adenomyosis is via the involvement of endometrial stem cells132. Adenomyotic foci within the myometrium can be considered as wounds undergoing repeated TIAR with associated epithelial to mesenchymal transition134, fibroblast to myofibroblast transdifferentiation and smooth
Secondary endometrial disorder

A disorder in the phenotype of the endometrium as a result of the presence of a structural cause of abnormal uterine bleeding in the myometrium; for example, fibroids or adenomyosis.

Endometrium fibroids, it is unknown whether endometrial disorders that lead to an impairment in normal menstruation should be considered to be primary endometrial disorders (disorders that are independent of the myometrial condition). Primary endometrial disorders can include: a reduced local hypoxia response with consequent interference with coordinated endometrial repair; an increased inflammatory response after progesterone withdrawal; a reduced haemostasis response due to an overactive endometrial fibrinolytic response; and a decrease in local vasoactive factors limiting vasoconstriction of the spiral arterioles, thereby contributing to excessive bleeding. An alternative, unproven, hypothesis is that the endometrium might behave in a manner consistent with a 'secondary endometrial disorder', in which changes occur in endometrial function as a consequence of the presence of a myometrial condition (for example, uterine fibroids or adenomyosis).

Improving the management of AUB

Nomenclature

Targeting the unmet clinical need related to AUB requires a collective effort from the scientific community. A lack of consistent terminology within reproductive health research has had a negative impact on scientific advances and clinical management. The FIGO MDC updated its terminologies and definitions of normal and AUB in 2018 (REFS 13, 14). These efforts to use consistent nomenclature and definitions have led to widespread acceptance of the term ‘abnormal uterine bleeding’ to describe bleeding that is abnormal in duration, frequency, regularity and volume. Inconsistent terms such as menorrhagia (an out of date term for HMB), oligomenorrhea (an out of date term for infrequent menstruation), dysfunctional uterine bleeding and metrorrhagia (an out of date, confusing term for irregular menstruation) should be abandoned. Consistency in terminology will facilitate data sharing, increase the validity of findings and improve patient care.

In wider society, there is also a need to address the discourse surrounding menstruation to prevent the harmful effects of menstrual stigma. In an online survey of 90,000 people from over 190 countries, over 5,000 euphemisms for the word ‘period’ were identified. Beliefs about menstruation are affected by personal experience, society and culture. If shame and embarrassment prevail, women might not seek the medical care they require.

Diagnosis

Symptoms can be defined according to FIGO System 1. Evaluation for anaemia and iron deficiency should be undertaken, considering the high prevalence of iron deficiency and IDA in patients with HMB on a global scale, which so often is under-recognized and under-reported. Once bleeding has been confirmed to originate from the cervical canal or uterine cavity, a systematic assessment should be made to understand the cause or causes of AUB using FIGO System 2, the PALM-COEIN classification (FIG 5, TABLE 1). This assessment will facilitate specific treatment of the underlying cause, personalizing management to improve effectiveness. If more than one cause of AUB is identified, this has implications for ongoing clinical management. Emphasis is generally placed on detailed history taking and examination as staffing levels, transportation to clinics and imaging modalities provide challenges for diagnosis in low-income to middle-income countries.

To improve the diagnosis of the structural causes of AUB, the Morphological Uterus Sonographic Assessment group have detailed criteria for imaging the myometrium by transvaginal ultrasonography, which is usually the most accessible imaging modality, although it is not 100% sensitive. MRI assessment is therefore indicated, for example, in patients who cannot tolerate or who decline transvaginal ultrasonography. MRI assessment might also be preferred to transvaginal ultrasonography in adolescents, patients with obesity, patients who have never been sexually active, patients with co-existing uterine fibroids and adenomyosis, or in patients with malignancy (such as sarcoma). Requirement for endometrial sampling in patients with AUB is based on risk factors for pre-malignant or malignant changes in the endometrium, including age, BMI, co-morbidities and genetic risk factors.

Personalizing treatments

Effective management of AUB is required to treat and prevent anaemia and IDA and to improve social, emotional and material quality of life. Managing a patient with AUB depends upon the diagnosis (or diagnoses) and, therefore, needs to be tailored for safety and effectiveness alongside patient preferences, presence of co-morbidities, symptoms, impact on the quality of life, fertility desires, age and ongoing expectations.
Fig. 5 | Diagnostic pathway for abnormal uterine bleeding, and iron deficiency and/or iron deficiency anaemia. On presentation with abnormal vaginal bleeding, the patient requires an initial assessment for pregnancy and to determine that the bleeding is coming from within the uterus, to confirm abnormal uterine bleeding (AUB). Such bleeding might either be acute or chronic (or acute on chronic). The time frame in which acute AUB is investigated varies, as the initial clinical priority is to stem the acute haemorrhage. In those with chronic AUB, it is necessary to perform a thorough assessment to determine the cause or causes in order to direct management. These include understanding the precise AUB symptoms using the International Federation of Gynecology and Obstetrics (FIGO) AUB System 1 (green boxes) prior to undertaking the appropriate investigations to identify the cause or causes of AUB using FIGO AUB System 2 classification (red boxes)². Imaging and visualization can occur with one or a combination of transvaginal ultrasonography, sonohysterography, MRI and hysteroscopy. Of note, blood tests for AUB-C would follow a positive screen for coagulopathy. AUB-O is primarily diagnosed on the basis of the structured history and blood tests are generally not necessary, but measurement of serum levels of progesterone might be useful in selected instances. AUB-E is a primary disorder of endometrial haemostasis. The clinician makes this diagnosis if there is no other explanation found on the clinical assessment including history, physical examination, appropriate blood tests and uterine imaging. HMB, heavy menstrual bleeding; FBC, full blood count.

It is debated in the literature whether menstruation is a physiologically necessary process³⁶⁻⁵⁶,⁹⁷,¹³⁸. Menstrual suppression is possible with exogenous sex steroids, as practised by, for example, athletes, military personnel or female astronauts¹⁴⁷. In addition, people without menstrual abnormalities might desire a reduction in menstrual frequency or amenorrhoea on completion of their family¹⁴⁸. However, a woman might also wish to undergo menstruation or retain her fertility and reproductive organs despite having finished her family, and these wishes should always be respected¹⁴⁹. Menstrual suppression might not be a viable option for all women¹⁵⁰.

An audit of HMB management in the UK by the Royal College of Obstetricians and Gynaecologists revealed that most treatments implemented for women with HMB were treating the symptom of excessive bleeding, rather than specific underlying causes¹⁵¹. Patients were often given a choice to try sex steroid therapies, but not all women respond to these agents in the same way, and a third of women felt they needed a hysterectomy to resolve HMB¹⁵¹. Hormone-based therapies increase exposure to sex steroid hormones, which could increase the burden of sex-steroid hormone-dependent conditions, for example, uterine fibroids or adenomyosis. This UK audit highlighted a problem that is most likely global, namely a lack of diagnosis for women presenting with the symptom of HMB. The artificial suppression of ovarian sex steroid hormones does not provide a panacea for women with menstrual abnormalities. Instead, there is a need for appropriate investigation, diagnosis and patient-specific management strategies in the treatment of HMB.

Many of the current medical treatments for AUB are hormonal (such as the levonorgestrel-releasing intrauterine system and cyclical oral progestins) and can work well for those with a non-structural cause of AUB, for example, patients with coagulation disorders (AUB-C), such as von Willebrand disease. However, for individuals with structural causes of AUB, hormonal therapies might be less effective. For example, people with submucosal uterine fibroids (fibroids that deviate the endometrial cavity) might experience increased complications and decreased effectiveness when using an intrauterine hormone delivery system¹⁵². Furthermore, in all people using hormonal treatments there is a risk of adverse effects, such as bloating, mood changes and breast tenderness, which might prove unacceptable. Future development of bedside diagnostics to detect the specific underlying defect in those with non-structural AUB-E (such as excessive inflammation, decreased hypoxia or haemostasis defects) would facilitate personalized treatment to correct specific defects causing AUB.
Surgical treatments can be life changing for people with structural causes of AUB (such as AUB-L or AUB-A) but can temporarily or permanently impair fertility, and therefore necessitate a careful informed discussion of the possible treatment choices. To date, hysterectomy remains the only guaranteed ‘no bleed’ option to prevent AUB for a patient, but this procedure has obvious implications for fertility, highlighting the need to develop more fertility-sparing medical and surgical interventions for AUB.

Gynaecological abnormalities, such as uterine fibroids and adenomyosis, can recur despite surgical treatment or medical adjuvant therapies. As imaging modalities improve, the detection rates of myometrial conditions increase, and strategies to prevent co-morbidities related to these conditions could be developed that would result in earlier diagnosis. Disease-modifying agents are also needed in order to slow disease progression and defer a patient’s need for surgical intervention.

Interestingly, women with uterine fibroids have been found to have more atherosclerotic cardiovascular risk factors than women without fibroids. Women who had undergone a hysterectomy with ovarian conservation were subsequently found to have increased risks of cardiovascular and metabolic disease, including hyperlipidaemia and coronary artery disease, when compared with aged-matched women living in the same geographic location who had not undergone hysterectomy or oophorectomy. These studies highlight that the decision to undergo a hysterectomy is not simple as the uterus might provide additional physiological functions beyond its role in reproduction. A holistic approach to a patient’s health is important to consider when planning management for AUB. Uterine-sparing treatments, and especially new medical treatments, are therefore much needed.

Factors affecting AUB

Age and AUB. It remains unclear whether the endometrium ages across the reproductive lifespan in humans. However, in mice, a blunted steroid hormonal response and reduced DNA methylation leads to defects in the decidualization phase with ageing which, in turn, negatively impacts reproductive outcomes.

The risk of developing structural causes of AUB increases with advancing age. Conversely, during the adolescent years, non-ovulatory cycles are more common, and adolescent girls are more likely to have a coagulopathy, such as von Willebrand disease, rather than a structural cause of AUB when compared with older women. The symptom of HMB in adolescent girls is more likely to result in fatigue, headaches, low cognition affecting verbal learning and memory, poor school attendance and performance, decreased participation in sports and possibly symptoms of depressed mood and/or anxiety.

Environment and AUB. Behaviour and the environment play an important role in a person’s health. How this then affects changes at a genetic level is investigated in epigenetic studies. To date, however, there are no identifiable studies that have specifically investigated epigenetic changes in relation to AUB. Dietary factors, vitamin deficiencies, parity, caesarean section and drug exposure have all been reported to have an impact on the prevalence of the causes of AUB.

It is yet to be determined if SARS-CoV-2 infection has an impact on endometrial physiology and the experience of menstrual bleeding. Data derived during the COVID-19 pandemic have revealed substantial effects of SARS-CoV-2 infection on endothelial cell function and the immune system. In addition, the SARS-CoV-2 receptor, angiotensin-converting enzyme 2, has been found to be present in the human ovary and uterus. Therefore, it is plausible that COVID-19 might directly affect menstrual parameters in those infected, in addition to the expected effects on hypothalamic function with notable illness and/or stress.

Exercise can benefit women in regulating their menstrual cycles, as aerobic and strength training has been found to lead to improved insulin control. Insulin receptors are expressed on the ovary and might influence the production of steroid hormones, including progesterone, thereby potentially impacting menstrual cycle control. However, the efficacy of using exercise as a means to treat the symptom of HMB lacks robust evidence. Traditionally, intense exercise has been linked to hypothalamic amenorrhea; however, elite athletes can also have AUB and HMB.

BMI and AUB. Menstrual disorders are more prevalent in women with overweight (BMI 25–30 kg/m²) or obesity (BMI ≥30 kg/m²) than in those with a BMI in the normal range (BMI 20–25 kg/m²). Obesity is associated with elevated levels of oestrogen through peripheral conversion of androgens to oestrogen by aromatase in adipose tissue (particularly androstenedione). Women with obesity have lower levels of sex hormone binding globulin than women with a BMI in the normal range, which leads to higher levels of circulating or free testosterone. These hormonal changes can result in ovulatory dysfunction (AUB-O), which is infrequent ovulation that extends the proliferative phase of the menstrual cycle. This increased exposure to unopposed oestrogen could support excess proliferation within the endometrium and increase the likelihood of menstrual disturbances.

Structural and non-structural causes of AUB have a higher prevalence in women with a high BMI (≥30 kg/m²) than in those with a normal BMI (20–25 kg/m²). Unfortunately, women with a high BMI have also been found to have less satisfaction with treatments for AUB, such as the levonorgestrel-releasing intrauterine system. Women with a low BMI (≤18.5 kg/m²), either due to diet restrictions or excessive exercise, are more likely to experience menstrual disturbances, particularly amenorrhea, due to functional hypothalamic disorders, than women with a normal BMI. Leptin signalling might also contribute to menstrual disturbances in women with a low BMI, as it has been found that women with a low BMI have reduced synthesis of leptin compared with women with a normal BMI. Elite athletes have an increased incidence of delayed menarche and...
primary amenorrhoea, specifically those participating in sports requiring leanness, for example, running, gymnastics and ballet.  

**Ethnicity and menstrual research.** The majority of research published in the literature related to ethnicity and its impact on the symptom of HMB focuses on conditions that cause AUB in black compared with white populations. There is a paucity of data on how ethnicity impacts the form and function of the female reproductive tract, and in the present context, the uterine endometrium and the menstrual cycle.  

Myometrial conditions, such as uterine fibroids and adenomyosis are more prevalent in black women than in white women. Black women are also more likely to have both of these conditions compared with white women. Black women are at greater risk of developing uterine fibroids at an earlier age, in larger size, greater number and with more severe symptoms compared with white women. This finding stresses the need for more genetically focused studies to further understand the ethnic differences in the progression of these conditions.

**Emerging technologies**

Human health and disease are ideally investigated in vivo; however, this is not always possible. This Review highlights key advances in endometrial physiology research using animal models of the menstrual cycle; for example, the use of non-human primate models or the simulated model of menses in mice. Models to study the endometrium in vitro have been developed in the past few years; for example, the use of endometrial epithelial cell organoid technology. These have the unique advantage of originating from human tissue whilst retaining the phenotype of the individual from whom they originated.

There remain numerous unanswered questions within the field of endometrial physiology research. Emerging technologies will allow us to overcome some of the limitations of human studies and address these outstanding research questions. These include, but are not limited to, the following: How does the presence of a myometrial condition, such as uterine fibroids or adenomyosis, impact the endometrium and lead to the abnormal bleeding phenotype of AUB? Is decidualization prior to menstruation a necessary event for coordinated menstrual loss and repair? Is it possible to characterize the interaction between innate immune cell traffic and the spiral arteriole vasculature within the endometrium and what is the role of the innate immune cell traffic in menstruation? Why does menstruation not result in tissue scarring? If we target key hormone-dependent pathways, would this help regulate menstruation in people with AUB? Is it possible to provide earlier, non-invasive diagnosis of all causes of AUB in a personalized manner, to appropriately treat millions of women around the world with this debilitating symptom? By addressing these key physiological and pathological questions we can implement preventative and therapeutic strategies to improve endometrial physiology.

New and emerging technologies (BOX 2) provide cross-discipline opportunities to offer insights into the biology of the endometrium in health and disease. These technologies have a potential role in diagnostics and could allow the monitoring of responses to treatment in clinical research. Additionally, they could provide avenues to elucidate the pathophysiology of conditions that cause AUB and identify novel targets for disease-modifying drug therapy. These biomedical advances are being made alongside changes in policy and culture (for example, the introduction of free period products in Scotland and New Zealand) and an increasing discourse in society regarding menstruation.

Combined efforts to address menstrual shame, stigma and taboos, create menstrual-friendly policies and advance menstrual research will have considerable benefits for menstrual health as a whole. As an area of huge unmet clinical need, endometrial physiology research provides exciting opportunities for scientific advances whilst also having the potential to lead to a substantial positive impact on a global scale.

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**Box 2 | New and emerging cross-discipline technologies being applied in endometrial physiology research**

**In vitro organoid technology and co-culture models**

Growing endometrial stromal and epithelial cells in vitro in co-culture provides a tool for studying cell to cell communication and endometrial functioning related to different clinical phenotypes.

**Stem cell research**

Endometrial stem cell studies are revealing processes of endometrial regeneration after menstruation.

**Microfluidics**

Via the integration of multiple so-called ‘microphysiological systems’ or 3D models representing part of the endometrium on a microscale, systemic effects that might influence menstruation can be investigated.

**Menstrual fluid research**

This ‘natural tissue biopsy’ provides a potential source of information about the endometrium during normal and abnormal menstruation.

**Bioinformatics**

Through the use of software, large complex datasets related to biological experiments of the endometrium can be analysed, potentially identifying pathways and mechanisms related to normal and abnormal functioning of the endometrium.

**Big data**

Large and varied clinical datasets related to women with abnormal uterine bleeding have the potential to elucidate patterns or subgroups to assist with targeting treatment groups.

**‘-omics’ and precision medicine**

For example, genomics, epigenomics, transcriptomics, proteomics and metabolomics have been applied to gain a deeper understanding of the physiological and pathological functioning of the endometrium.

**Imaging**

It is hoped that the development of imaging capabilities within MRI and ultrasonography (for example, elastography) will provide non-invasive methods for investigation and potentially earlier diagnosis in vivo.

**Artificial intelligence**

Machine learning has the potential to help in the early prediction and identification of women with possible causes of abnormal uterine bleeding, potentially improving future diagnosis and treatment.

**‘FemTech’**

Devices, software and/or other technology related to women’s health have the potential to improve diagnostic capability as well as increase awareness and knowledge surrounding menstrual health.
Conclusions

AUB has a detrimental impact on those it affects and leads to considerable disease burden on a global scale. A large proportion of women with AUB globally also have either iron deficiency or IDA, which often is both under-recognized and under-reported. It is essential that we continue to delineate the physiology of the menstrual cycle using available and emerging technologies. Scientifically combating the knowledge gaps within the field of menstruation will translate to notable benefits across the globe for those who menstruate.

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Author contributions

V.J., H.O.D.C., R.R.C. and J.A.M. researched data for article, contributed substantially to discussion of content, wrote the manuscript and reviewed and edited the manuscript before submission.

Competing interests

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