ISSUES

Expanding the evolutionary explanations for sex differences in the human skeleton

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Abstract
While the anatomy and physiology of human reproduction differ between the sexes, the effects of hormones on skeletal growth do not. Human bone growth depends on estrogen. Greater estrogen produced by ovaries causes bones in female bodies to fuse before males' resulting in sex differences in adult height and mass. Female pelvis expand more than males' due to estrogen and relaxin produced and employed by the tissues of the pelvic region and potentially also due to greater internal space occupied by female gonads and genitals. Evolutionary explanations for skeletal sex differences (aka sexual dimorphism) that focus too narrowly on big competitive men and broad birthing women must account for the adaptive biology of skeletal growth and its dependence on the developmental physiology of reproduction. In this case, dichotomizing evolution into proximate-ultimate categories may be impeding the progress of human evolutionary science, as well as enabling the popular misunderstanding and abuse of it.

KEYWORDS
childbirth, estrogen, growth, height, male competition, pelvis, sexual dimorphism

1 | INTRODUCTION

Scholarship on sex differences in the human skeleton, from the focused to the tangential, explains how males are taller due to sexual selection for contest winners, and how females are broader due to natural selection for childbirth (for just two recent examples, see Refs. 1 and 2, respectively; for textbook examples, see Refs. 3–5). While discourse among anthropologists and fellow travelers on skeletal sex differences is often nuanced about causal complexity and unknowns, it does not seem to have affected the discussion beyond these circles, where public perception of evolutionary causality is far simpler. The narrow emphasis on competitive men and birthing women harkens back to the origins of human evolutionary biology, and these explanations for biological sex differences dominate the popular understanding of human evolution today.

Here, “female” refers to humans of all genders with anatomy that is commonly assigned to be female, and the same gender inclusivity applies to “male”—with the understanding that neither sex nor gender divide into uniform, discrete, or binary categories, which is why “sex differences” rather than “sexual dimorphism” is employed throughout this paper.

This paper briefly reviews the complex biology of sex differences in human stature and pelvic dimensions, focusing mainly on the role of estrogen. Investigating how these differences develop expands their evolutionary explanations. Peering from this angle stirs skepticism of the traditional, narrow emphasis on the dominant ideas and creates opportunities for testing them. To be clear, the reigning explanations (male competition for skeletal size differences and childbirth for pelvic differences) are neither extensively reviewed nor rebutted in this paper. Instead, the goal of this paper is to highlight some additional context for the evolution of sex differences in the skeleton. In order to present a fresh approach to a familiar topic, this paper primarily asks why skeletal sex differences exist at all, rather than starting with comparisons of degrees of sex differences between humans and other primates.

Throughout this paper, and perhaps already by now, many readers will be partitioning evidence into the dichotomous realms of proximate and ultimate evolutionary explanations—the ultimate
ones being childbirth and male competition and the proximate ones being mechanisms of growth and development of the skeleton. However, this convention is not a requirement of evolutionary thinking\textsuperscript{12} and it is neither espoused nor endorsed here.

Lastly, in anthropology, the sociocultural consequences of the scientific truth are as equally important as the truth itself. The current pop culture narrative where men are specially built for competition, while women are specially built for reproduction, helps root socioculturally prescribed and proscribed sex roles and stereotypes in “human nature.” Expanding the dominant origins story for sex differences in height and pelvic dimensions will not just improve science but will also help rip human evolution out of the patriarchal playbook.

2 | WHY ARE THERE SEX DIFFERENCES IN HUMAN STATURE?

While human height varies globally, all human populations exhibit the same pattern where mean adult male height is greater than mean adult female height. In the United States (Figure 1),\textsuperscript{13} after nearly the same growth trajectory from 2 years of age, both males and females are roughly 62 in. (157 cm) tall at 13 years. After that, the female growth curve flattens to reach the average final height of about 64 in. (163 cm). Conversely, in males, the growth curve continues on roughly the same trajectory for at least 1.5 more years until it then flattens to reach the average final height of about 70 in. (178 cm). This is an additional 9\% of growth in stature compared to females. Average age of menarche in the United States occurs at about 13 years\textsuperscript{16} (which matches that reported in at least one small-scale subsistence society\textsuperscript{17}). Thus, while males continue to grow in stature, females slow to a stop, and simultaneously begin monthly cycling. Females who reach menarche relatively later continue to grow at the faster prepubertal rate until onset of menses and end up being relatively taller adults.\textsuperscript{18}

The synchrony of menarche with growth deceleration and subsequent arrest is not mere coincidence. Both the menstrual cycle and skeletal growth depend on estrogen.

For males and females, long bone growth and epiphyseal closure are highly dependent on estradiol, the most important of the naturally occurring estrogens, and hereafter also referred to as “estrogen” (the literature cited also employs the terms interchangeably).\textsuperscript{19–21} Estrogen accelerates the loss of progenitor cells in the resting zone of the long bone growth plate, which causes senescence in the growth plate and results in cessation of growth.\textsuperscript{22–24} Estrogen is produced in both the ovaries and testes, where androgens are converted into estrogen by the enzyme aromatase. Aromatase is expressed in the growth plates, too, and so some of the hormone conversion related to bone growth is local.\textsuperscript{25} Estrogen’s effects are biphasic with one-level

\textbf{FIGURE 1} Stature of boys (a) and girls (b) in America.\textsuperscript{13} See also Bogin\textsuperscript{14} and Bogin et al.\textsuperscript{15} [Color figure can be viewed at wileyonlinelibrary.com]
stabilizing bone growth and an even higher-level stimulating epiphyseal closure. Prepubertal females have eight times the estradiol levels of males at the same age, which helps explain both their earlier growth spurt (i.e., peak growth rate, which is not obvious on Figure 1 but see Bogin15) and earlier growth arrest compared to males. An excess of estrogen causes medically diagnosed short stature in both sexes. Estrogen in low doses enhances growth hormone (GH) and insulin-like growth factor 1 (IGF-1) production (the GH-IGF-1 axis) which are key to linear bone growth, but at high doses estrogen inhibits IGF-1. Androgens do stimulate GH, but in their absence, normal growth can occur as long as there is sufficient estrogen. As androgen production increases, males also reach critical levels of estradiol to stimulate the process of growth plate fusion starting around 16 years of age. These levels are likely to be lower than they are in females because at this older age the growth plates are more senescent and require a more brief exposure to estradiol. In addition, levels of estrogen at this time are critical to bone mass maintenance, an important function of estrogen in all humans.

The sex difference in estradiol levels is due to its greater involvement in ovulation and menstruation than in spermatogenesis and related processes, where it is also vital. Estradiol regulates spermatogenesis by testicular Sertoli cells by both inhibiting and stimulating, in ment in ovulation and menstruation than in spermatogenesis and earlier growth arrest compared to males. A excess of estrogen causes medically diagnosed short stature in both sexes. Estrogen in low doses enhances growth hormone (GH) and insulin-like growth factor 1 (IGF-1) production (the GH-IGF-1 axis) which are key to linear bone growth, but at high doses estrogen inhibits IGF-1. Androgens do stimulate GH, but in their absence, normal growth can occur as long as there is sufficient estrogen. As androgen production increases, males also reach critical levels of estradiol to stimulate the process of growth plate fusion starting around 16 years of age. These levels are likely to be lower than they are in females because at this older age the growth plates are more senescent and require a more brief exposure to estradiol. In addition, levels of estrogen at this time are critical to bone mass maintenance, an important function of estrogen in all humans.

The sex difference in estradiol levels is due to its greater involvement in ovulation and menstruation than in spermatogenesis and related processes, where it is also vital. Estradiol regulates spermatogenesis by testicular Sertoli cells by both inhibiting and stimulating, in a dose-dependent and temporally sensitive process. Aromatase activity is higher in motile as opposed to immotile sperm and was found to be significantly decreased in a population of infertile men. In puberty and in adulthood, excess estrogen can inhibit penile erection. In all humans, a delicate balance of estrogen/aromatase is as fundamental to reproduction as it is to skeletal growth. We are safe to assume that Homo sapiens' prolific biology of reproduction is adaptive.

In addition to the effects of estrogen on stature, there may be a pubertal onset of energetic, metabolic, and nutritional costs that force a tradeoff with skeletal growth. When energy intake and physical activity are held constant, basal metabolic rate (BMR) varies significantly across the menstrual cycle, with the lowest BMR occurring approximately 1 week before ovulation, subsequently rising until the beginning of the next menstrual period, then decreasing at menstruation. Several studies have quantified changing energy intake across the menstrual cycle. Peak and minimum intake across the cycle differ by 359 kcal/day (n = 6). Ten days before menstruation, mean energy intake is approximately 500 kcal/day higher than 10 days after (n = 8). Compared to the follicular/ovulatory phase, the luteal phase corresponds to an increased energy intake of 685 KJ/day (or 164 kcal/day). These data point to the metabolic effects of the changing estrogen/progesterone ratios across the cycle, including the costly thickening of the endometrium. These costs may differ not just individually but across the reproductive lifespan. Reiches et al. described a relatively higher cost of menstruation in younger adolescents compared to older adolescents.

As this brief review of the relationship of estrogen to skeletal growth has shown, the reproductive systems of males and females differently affect a skeletal system that is shared by males and females. So, the evolutionary explanation for the existence of sex differences in human height is rooted in the origins of estrogen and its subsequent importance in all vertebrate bodies some 500 million years ago. Also of crucial importance are the origins of internal fertilization and viviparity.

Great apes develop sex differences in body mass like humans do, where both sexes follow similar growth trajectories until the pubertal transition when the females stop growing and the males continue to grow for a longer period of time. Though levels of sex differences in body size differ between species, among the living hominids (great apes and humans) there is likely to be significant shared fundamental biology of reproduction and skeletal growth. Thus, the existence of human sex differences in stature is rooted in ancestry. It remains to be known whether there are important sex- and species-level differences in the biology of skeletal growth among hominids, and whether these could explain the differing degrees of sex differences in body size across primates, even after accounting for allometry and for estrogen production—which is potentially constrained by testes size and may be an important factor in the extended growth of male gorillas and orangutans. (For a discussion of how these “somatic strategies” might occur at primate puberty see Ref. 39.)

However, the traditional and enduring textbook explanation for sex differences in hominid body size is sexual selection—with large ancestral males winning competitions, which boosted their reproductive success compared to smaller males. Because gorillas have both intense male competition and large male bodies, the mere existence of sex differences in human body size serves as evidence of sexual selection being the driver of these differences.

But as Plavcan has cautioned, there is not a straight-forward relationship between sexual selection and primate male body size, largely because the sorts of data that are required to investigate this relationship are difficult to obtain. It is also difficult to tease selection on male body size apart from selection on female body size through the generations, which is sometimes understood within the framework of females as the “ecological sex.” Given the nutritional, energetic, metabolic, and locomotor costs of pregnancy, lactation, and mothering, there are (context-specific) limits to female body size, perhaps leading to biology that favors reproduction over growth.

Yet even within this more complete “ultimate” narrative, with selection optimizing the two sexes’ skeletal growth separately, the sexual selection perspective on male height seems unnecessary. That provocative last sentence is not a claim that the sexual selection explanation is wrong or that it is implausible. But in light of what is known and still unknown about skeletal development and its relationship to the endocrinology of reproduction, suddenly there is room for skepticism about the relevance of male competition and female choice as an explanation for the existence of sex differences in stature, let alone its singular dominance of the narrative. More work is needed if sexual selection is to be held up as the explanation for why male hominids have longer bones than female hominids do.

Given the complex, shared biological systems briefly outlined above, which are intricately tied to successful reproduction and that contribute to terminal height, stature differences within adult males are probably weaker targets of selection than is assumed by sexual...
selection scenarios. In their recent overview of the evolution of human height variation, Stulp and Barrett\textsuperscript{50} made a similar point when they wrote that "height itself is less important as a trait than the underlying components of growth rates and the timing of reproductive maturity that give rise to it. This raises the question of whether height does, in fact, carry any selective advantage independent of its links to life history." (p. 220).

Singly upholding the male competition hypothesis for sex differences in human stature requires, for example, the demonstration that men's estradiol/aromatase production, levels, receptors, and timing are primarily due to the fitness rewards of being taller than females, or primarily due to the fitness rewards of being taller than other males. It also requires, for example, the demonstration that men's estradiol/aromatase production, levels, receptors, and timing are not primarily due to something fundamental to male gonad, genital, and gamete maturation or function, or that they are not primarily due to shared biology with females.

Data from tracking the reproductive success of human males fail to comprehensively answer the question of why there are sex differences in human height.\textsuperscript{51} and further work of this kind, even across primates, will continue to be insufficient for elevating the sexual selection explanation for sex differences in height if it is not integrated with some insightful combination of physiological, endocrinological, developmental, and/or genetic approaches. Perspectives that assume the extended skeletal development of males is a delay in body size maturation and/or is a cost that requires a male-specific selection-based explanation must reckon with the risks that changes to the biology of male skeletal growth would also pose to the biology of male fertility. Furthermore, investigations of these issues need not assume that a lack of sex differences in the skeleton (i.e., "monomorphy") is the biological baseline or default in all primates and, thus, that sex differences in the duration of skeletal growth or in long bone length have been directly driven apart by sex-specific sexual or natural selection on skeletal growth. Free from these assumptions, there is potential for exciting advances including, perhaps, the discovery that sexual selection does indeed play the lead role in this story.

For humans and likely other hominids, male skeletons continue to grow after females' stop because their bodies take longer to produce enough estradiol to surpass the amount that stimulates continued growth and to achieve a level that closes long bone epiphyses. As of now, no advantage to being taller or more massive is required to make sense of this phenomenon which may be largely a by-product of the adaptive reproductive biology that differs between the sexes. Dominance\textsuperscript{52} and competition may be consequences of greater height and mass, but the claim that they cause sex differences in the skeleton requires far more investigation.

3 | WHY ARE THERE SEX DIFFERENCES IN HUMAN PELVIC DIMENSIONS?

On average, human female pelves have longer pubes, more laterally flaring ischial spines and tuberosities, and relatively shorter and wider sacra. Thus, they often have inlets (often measured from sacral promontory to the superior pubic symphysis), midplanes (often measured as the distance between ischial spines), and outlets (often measured from coccyx to inferior pubic symphysis or measured as the distance between ischial tuberosities) that are relatively larger in diameter than those of males.\textsuperscript{53–55} These dimensions together comprise the "true pelvis" or "birth canal" which is relatively larger in females than in males.\textsuperscript{53} So, while there is geographic variation in human pelvic morphology\textsuperscript{56} and while typical female pelvic inlet shape may be "android" like males (contra traditional expectations that they be distinctly "gynecoid"), there are consistent and patterned sex differences in human pelvic morphology pertaining to the size of the space inside the pelvic cavity.

Fetal pelves, between 7 months and birth, display sex differences that already hint at those in adults.\textsuperscript{58} Around the transition to adulthood, female pelves tend to fuse earlier than those of males at all sites. This pattern parallels the sex differences in long bone fusion. The site with the greatest sex difference in closure is the anterior epiphysis of the acetabulum which articulates with the pubis and is actively fusing between 11 and 16 years in females but not until ages 14–17 in males.\textsuperscript{58} Sex differences in pelvic morphology become pronounced during this stage in life. LaVelle\textsuperscript{59} found that between ages 8 and 18, female pelves expand slightly more than males' in the dimensions of the true pelvis. Some of the most conspicuous change occurs in pubis length. Likewise, Greulich and Thoms found greater transverse dimensions in developing and adult female pelves.\textsuperscript{60} Huseynov and colleagues observed, in a cross-sectional sample, that true pelvic dimensions expanded from puberty until the ages of 25–30 years, then after 40 years these dimensions diminished in magnitude.\textsuperscript{61} Whether and how the development of sex differences in the pelvis can be causally linked to intrasex and intersex differences in the timing of the fusion of the pelvic bones remains to be determined.

Because estrogen is produced in greater amounts in female bodies and those amounts change across the life course, estrogen is the established explanation for ontogenetic changes to female pelvic anatomy as compared to that of males.\textsuperscript{60,62} This holds even for the neonatal sex differences because the last few weeks of fetal development occur while estrogen levels are highest in gestation.\textsuperscript{63} But if estrogen is a primary driver of long bone growth and fusion (as discussed above), then how does it act locally just on the bones of the pelvis and only in females?

Rodent experiments from 1929 to 1935 suggest that dosing a male body with estrogen "feminizes" the pelvis,\textsuperscript{66} suggesting the system works on any pelvis. However, a review of skeletal biology in 2005\textsuperscript{26} reported that the stimulatory effects of estrogen on skeletal growth and maturation in humans are poorly reproduced in rodent studies. Regardless of these important issues that may complicate a comparative approach, estrogen's hypothesized localized effects on female pelves warrants deeper consideration. This is especially necessary in light of a recent study of humans that failed to correlate within-individual levels of sex differences in the skull with that in the pelvis, suggesting that a "single systemic influence, such as hormone
levels, is not solely responsible for sex differences in the size and shape of these skeletal elements.64–66

The muscles of the pelvic floor, like the levator ani, the round ligament of the uterus (which is actually a smooth muscle, not a ligament), and other uterine ligaments (the pubocervical, uterosacral, and cardinal [transverse cervical] ligaments) contain estrogen receptors (ER), which suggests they are targets for estrogen.65–69

ER are absent in typical skeletal muscles like the rectus abdominis and erector spine,65,66 supporting the hypothesis that the pelvic muscles are under special hormonal control. Pelvic floor muscles also contain ER in their connective tissue cells, which are the “glue” that fixes the muscles together and to the pelvic bones.66 The round ligament grows during pregnancy and shrinks (not slacks) after parturition, which could influence skeletal remodeling.70 The markedly earlier fusion in females, described above, of the anterior epiphysis of the acetabulum (which forms the ilopubic eminence) could be influenced by its close proximity to the deep inguinal ring, which transmits the round ligament. All of this suggests that the muscles and ligaments of the female pelvis influence the bones to which they are adjacent or anchored in ways that differ from other muscle–bone and ligament–bone interfaces, given the known effects that estrogen has on bone growth and remodeling. In addition, relaxin, which is produced by the ovary and placenta, induces the production of osteoclasts71 which are key to bone resorption and remodeling—a noted phenomenon when it comes to resorption of the human, nonhuman primate, and nonhuman mammal pubis.72 Greater parity increases estrogen and relaxin exposure, which leads to the expectation that greater parity would be correlated to expanding pelvic dimensions, but at least one recent study failed to distinguish nonparous from parous female pelves.61

What is more, the volume occupied by internal female organs—in addition to the bladder and rectum housed within all pelves—may be causing the expansion of the true pelvis. The last few weeks of fetal growth show marked changes in uterus size, position, and angle of flexion.73 Neonatal uteruses are 3.5 cm long and 1.4 cm thick.74 Between the ages of 1–13, uterine volume increases from 0.91 cm³ to 16.15 cm³,75 between ages 16 and 17.5 it is 60 cm³, and between ages 24 and 29 it is 79 cm³.76 The uterus begins a more rapid growth rate around 10 years of age, with the onset of puberty and during concomitant increases in luteinizing hormone (LH), follicle stimulating hormone (FSH), and estradiol.77 At this time, roughly 2–3 years before menarche, the vaginal and vulvar epithelium thicken and, along with the cervix and clitoris, they increase in size.77 Ovarian volume is 1 cm³ in the first year of life (which is larger than the second year),74 from ages 7 to 12.5, it increases from 1.4 to 4.9 cm³, and then from ages 16 to 17.5 it measures 8.9 cm³.76 (Unlike the expanding uterus, ovarian volume from ages 24–29 is reduced, measuring 7.2 cm³.) The size of the uterus and cervix increases over a lifetime with parity.78 During the luteal phase of the menstrual cycle, when the endometrium thickens, the uterus expands to at least 1.6 its volume, likely more.79 Vaginal epithelium reaches its peak thickness mid cycle and vaginal muscle fibers thicken late in pregnancy.77 While these data were collected from small samples, data sets from different sources concur.

Volumetric data on the clitoris proved elusive, but linear dimensions—like those of the bulbs measuring 3–4 cm long when flaccid and 7 cm when erect, and the crura measuring 5–9 cm long—are listed in Ref. 80. In contrast, the only internal organ of comparable size that is specific to male pelves is the prostate which develops from 1.4 cm³ (ages 0–9) to 6.9 cm³ (ages 10–19), and to 15.3 cm³ (ages 20–29), remaining much smaller than the uterus across those age groups.81 In addition, the prostate is nestled under the bladder while the uterus and ovaries are situated higher up, within the pelvic inlet or brim, in direct line between the pubic symphysis and the sacral promontory. That is, the vagina, uterus, and ovaries are not just taking up more volume but are also, arguably, situated within a more skeletally constrained region of the pelvis compared to the prostate. Publically posted pelvic MRIs allow for visual inspection of internal pelvic anatomy and the comparison of one male and one female.82 What, if any, effects that age-related prostate enlargement may have on the male pelvis are apparently unknown, but they are not predicted to mimic what is hypothesized here for developing females because of the difference in context, both in terms of age and estrogen.

The increase in size over the lifetime and the periodic expansion (during intercourse, the menstrual cycle, and pregnancy) of internal gonads and genitals may be spurring changes to the bones that form the cavity they occupy similar to the ways that organs and bones expand together elsewhere in the body. As brains and skulls develop together, signals for growth are recognized by both neural and skeletal tissues in an integrated manner.83 Growth signals may be mediated by tensile strain, caused by mechanical stress on the bones by the growing soft tissue. In a similar fashion, growth of the bony orbit likely responds to the development of the eye.84 Such processes are potentially occurring in the thorax, coupling the developing heart and lungs with an expanding ribcage. Habitual human swimmers provide a natural experiment for investigating this phenomenon. Documented increases in swimmers’ lung volumes, especially in athletes who train intensely from childhood, are correlated with the development of physically wider chests85,86 perhaps due to the increased pressure while actively inhaling and exhaling while immersed in water. The pelvis may be no exception when it comes to the skeleton’s plastic accommodation for developing soft tissue. Furthermore, differences in the shapes of the internal skeletal spaces may vary according to the shapes of the organs within. Variation in uterine shape, existing as early as fetal development, could influence variation in pelvic proportions, and vice versa. Fetal uteruses are cylindrical, pear, heart or hourglass shaped.73 Amount and direction of uterine flexion, and the pace of its development, may also factor into how the pelvis develops, and vice versa. The decrease in uterine volume after peak fertility may help explain why Huseynov and colleagues61 found that older adult female pelves are less expansive.

In sum, sex differences in the dimensions of the true pelvis are influenced by localized effects of estrogen and relaxin within a system of gonads, genitals, ligaments, muscles, and bones in ways that are not fully understood. There is potential for sex differences to arise due to the plasticity of the pelvic bones to accommodate the greater volume of developmentally and functionally dynamic gonads and genitals.
housed within the female pelvis. As with height differences, the explanation is fundamentally rooted in the ancient origins of estrogen, internal fertilization, and pregnancy, the soft tissue differences between the sexes that evolved as a consequence, and how they affect the local skeleton differently in males and females with different hormone levels. Because sex differences in pelvic dimensions are common across primates, this is an ancestral condition in humans.

But, the widespread explanation for sex differences in the human pelvis is merely and simply childbirth. For example, "Females have big pelvises because they give birth to big babies." While intuitive, this explanation for human pelves or for other primes' is no longer a strong one—at least not in isolation as it is frequently provided.

Moffett investigated whether pirmates with greater cephalopelvic proportions (size of neonatal head compared to pelvic inlet) had greater pelvic sex differences and they did, but humans have even more than is explained by cephalopelvic proportions. That is, primes like Hylobates have similar cephalopelvic proportions to humans but exhibit smaller sex differences in the pelvis. Even chimpanzees, which have small enough neonates to fit through the male pelvis (inferred from measures published in Ref. 53 and 88), still have sex differences, with female pelves being more capacious than males.

It is possible that differences in type and magnitude of sex differences in pirmate pelves reflect differences in soft tissue anatomy, reproductive physiology, and effects/amounts of estrogen and relaxin and their receptors. Uterus location, size, flexion, and function could vary in important ways that impact the skeleton. Whether pirmate males have descended testes or not is also likely a factor. Differences across pirmates in clitoral anatomy as well as anatomy involved in estrus swelling could contribute to differences between species in sex differences, too. The round ligament grows during pregnancy and shrinks after parturition in other pirmates as it does in humans, which may contribute to their pelvic remodeling. Interestingly, rodents and lagomorphs have a different system for suspending the uterus and related organs and so if this is affecting their pelvic architecture, it is another reason (added to the estrogen issue mentioned above) that some animal models may be inappropriate for explaining pimate/ human sex differences in the skeleton. When Kurki and also Fischer and Mitteroecker observed that shorter women have relatively large "obstetric" dimensions perhaps it is due to the allometry and/or conservation of size and function of soft tissues, no matter the stature. Finally, human sex differences in the pelvis could be more pronounced than expected compared to other primes' because of the more tubular or constricted construction of the hominin pelvis, and how pelves with internal female organs and greater estrogen exposure develop in the context of bipedalism.

A genital, gonadal, and hormonal view of the evolution of pelvic sex differences (and of the conservation of "obstetric" dimensions even in small bodied females) contrasts the traditional "ultimate" evolutionary approach that downplays developmental dynamics. A developmental perspective has less room for ideations of genetically programmed population- and species-specific tweaks of the space between pelvic bones in females versus males. Investigations of these issues need not assume that a lack of pelvic sex differences is the biological baseline or default and, thus, that skeletal differences have been driven apart by sex-specific selection on adult skeletal morphol-

gy. We need not assume the logic of the obstetrical dilemma hypothesis, where female pelves would be like males' if only selection for childbirth had not forced a compromise.

There is a crucial, constant function of the human female pelvis no matter the sex, age, or parity and that is to house developing, functioning organs. Female bodies and pelves contain tissues during the entire life course that stimulate pelvic bone growth and remodeling. Thus, the internal dimensions of the female pelvis are far more ovarian, uterine, clitoral, and vaginal than they are "obstetric." In the end, it may be that females give birth to big babies because they have big pelves.

4 | CONCLUDING REMARKS

Investigations of sex differences in the human skeleton have faced many of the challenges in evolutionary biology that Smith described: “Some narrative explanations rely on theory-driven assumptions that may not be shared by readers... Some will not use good judgement when taking into account how underdetermination inevitably limits what can be inferred from historical data... Some will make unreasonable assumptions about what the current utility of a feature can tell us about its historical role... Some will make unreasonable assumptions that simplify the contingency of the historical situation... Some will allow coherence and simplicity in narratives to substitute for evidence.”

Sex differences in human height and pelvic dimensions require a bigger, more complicated, and more interesting story than simply "male competition" and "childbirth." Greater estrogen results in the bones in female bodies fusing before males' leading to sex differences in adult height and mass. Female pelves expand more than males due, potentially, to the space taken by vaginas, clitorides, uteruses, and ovaries and because of the estrogen and relaxin produced and employed by the tissues of the pelvic region. Generally speaking, these evolved processes are not unique to Homo sapiens. Understanding the details of the developmental biology of the skeletal and reproductive systems of human males and females, and understanding that development in phylogenetic context, will be crucial to formulating and testing evolutionary hypotheses concerning sex differences in the skeleton. With its focus on just some of the drivers of skeletal development, this paper is only one step towards expanding our evolutionary explanation for sex differences in skeletal growth.

Answers to questions about sex differences in the human skeleton should include what is increasingly known about the evolution of gonads and genitals, their growth at puberty, and their functions during skeletal maturation, sexual intercourse, the menstrual cycle, pregnancy, and menopause, as well as the sensitivities of different tissues to estrogens and androgens. The basis for the existence of sex differences deserves more attention before it will be possible to explain why humans have a certain degree of sex difference compared to other species. This may require some evolutionary research that
breaks free from Mayr’s proximate-ultimate convention. As Laland et al.12 write, “progress within biology demands dismantling of Mayr’s identification of proximate with ontogenetic processes and ultimate with evolutionary processes” (p. 1516). The so-called “proximate” causes of sex differences in the skeleton are not only as much evolutionary ones as the “ultimate,” but they hold great potential to advance investigations into how male competition and childbirth feature in the evolution of skeletal sex differences in humans.

If we do not hold evolutionary hypotheses to higher standards, while also including all biology into the category of “evolution,” then so many “ultimate” answers to important questions in human evolution will loom larger and longer than they deserve.

Finally, a human evolutionary narrative that expands to include the present state of knowledge about skeletal and reproductive biology and their harmonious development is not just better science. It is also less likely than the traditional scientific view to unintentionally evoke or reinforce unscientific beliefs about genetic determinism and genetic essentialism in the zeitgeist. An updated answer to why there are sex differences in the human skeleton is less likely to be interpreted to justify cultural conceptions of masculinity, femininity, and rigid binaries of sex and gender with “human nature.” If we improve the scientific explanations of visible sex differences, then they are less likely to inspire unscientific beliefs about invisible ones. Fewer minds would leap illogically from “men are taller” to “men evolved for competition and dominance.” Likewise, fewer would observe that “women are broader” and conclude that “women evolved for reproduction.” As we advance science and its dissemination, fewer will mistake the human body for a blueprint for the patriarchy.

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