Gatekeeping hormone replacement therapy for transgender patients is dehumanising

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ABSTRACT
Although informed consent models for prescribing hormone replacement therapy are becoming increasingly prevalent, many physicians continue to require an assessment and referral letter from a mental health professional prior to prescription. Drawing on personal and communal experience, the author argues that assessment and referral requirements are dehumanising and unethical, foregrounding the ways in which these requirements evidence a mistrust of trans people, suppress the diversity of their experiences and sustain an unjustified double standard in contrast to other forms of clinical care. Physicians should abandon this unethical requirement in favour of an informed consent approach to transgender care.

INTRODUCTION
When I decided that I wanted to take hormones to feminise my body, the last thing I wanted to do was to go in front of a psychologist to justify my decision. Deciding to take hormones was not a decision I made in haste. Moreover, like many others, I had socially transitioned months before pursing hormone replacement therapy (HRT). Thankfully, my educational privileges provided protection: I did not have to see a psychologist for a letter of referral to get a HRT prescription because the university health clinic practiced the informed consent model, which is becoming increasingly common across the USA.1 I was able to see a physician who would prescribe me oestrogen and antiandrogens without such a letter, although it is typically required under the Word Professional Association for Transgender Health (WPATH) Standards of Care (Coleman et al, p180).2 Many others are not so lucky—and my own luck ran out when I had to seek two referral letters for genital surgery.

The assessment of gender dysphoria left me feeling exposed, naked and dehumanised. Although the assessment process is alone a difficult experience, it is only made worse by the apparent conflation of gender dysphoria under the WPATH Standards of Care and Gender Dysphoria as a psychiatric diagnosis defined in the Diagnostic and Statistical Manual of Mental Disorders, Fifth Edition (DSM-5).3 Discussions of the ethics of referral requirements for HRT have tended to emphasise traditional approaches to bioethics, with arguments against those requirements primarily relying on autonomy and beneficence.4 5 Though ethically sound, those approaches fail to foreground the lived experiences of the trans people who undergo these assessments. I hope that my contribution, which will draw on my personal and communal experiences to illuminate the dehumanising nature of HRT gatekeeping, will motivate physicians to provide HRT on an informed consent model for older teenagers and adults.6

The informed consent model shifts focus away from the assessment of gender dysphoria and the provision of mental healthcare and instead sees the obtention of appropriate informed consent as the primary role of hormone providers (Coleman et al, p188).2 Providers working under the informed consent model typically record consent on a document listing the potential benefits, risks and limitations of HRT. Working under the informed consent model includes discussions of the patient’s expectations, decision-making process, understanding of risks and benefits, support structures and general health, but does not involve an evaluation of their gender or whether they are truly trans (Chang, Singh and dickey, p143).7 The goal is not to assess but to facilitate thoughtful decision-making. The model ensures that the decision was not made in haste without appropriate knowledge and forethought, while preserving patients’ own authority over their experiences and avoiding the dehumanising process criticised in the present article.

MISTRUSTING TRANS VOICES
We generally trust what other people say about their own mental states. If someone says, ‘my arm hurts’, we typically grant credence to their claim. We have this trust in people’s self-reports of their mental states because we hold mental states to be within the purview of people’s epistemic authority—authority over knowledge.8 This authority is defeasible: I do not need to demonstrate the authoritative nature of my assertion and probing questions would typically be unjustified, but if a serious reason to doubt it is present, it would be legitimate to doubt the claim and ask further questions (Bettcher and Shrage, p100).9 If someone says that their arm hurts but are laughing at the same time, we may have a good reason to doubt, yet in the absence of such conflicting indication, it would be illegitimate to doubt them. Defeasibility addresses the epistemic tension between our privileged access to our own mental states and the fallibility of that self-knowledge.10 If I were to doubt that person’s claim without serious reasons to believe otherwise, I would be committing an injustice because I would unjustly fail to recognise their authoritative knowledge of their own experience of the world.11

Not all knowledge relating to mental states can be authoritatively reported by individuals. We do not typically grant credence to mental health self-diagnoses, especially not by non-professionals—though perhaps we should more often.
However, it is important to note that self-reports of gender dysphoria do not fall within this type of specialised knowledge about mental health which is reserved for professionals. As WPATH defines it, gender dysphoria refers to ‘discomfort or distress that is caused by a discrepancy between a person’s gender identity and that person’s sex assigned at birth (and the associated gender role and/or primary and secondary sex characteristics)’ (Coleman et al, p166). The assessment of gender dysphoria is not an assessment of a mental health condition—the WPATH Standards of Care predates the DSM-5 Gender Dysphoria diagnosis by a few years.

Experiences of gender dysphoria are part of the mental experiences over which we have epistemic authority. The simple report of having gender dysphoria to a physician, combined with an informed consent process to ensure that expectations match the actual effects of HRT, should suffice to obtain a prescription since gender dysphoria is distress or discomfort towards those very features that people seeking HRT want to change. By requiring that trans people submit to an assessment of gender dysphoria instead of satisfying themselves with the patient’s affirmations, physicians deny the authority trans people have over their own mental experiences, an authority that should be granted to everyone by virtue of being persons. To unjustifiably deny that authority is dehumanising. Referral requirements for HRT treat self-reports of gender dysphoria not as one would deny that authority is dehumanising by those who do not fall under a pathologising model. Although the shift to the terminology of gender incongruence in the ICD-11 partly addresses this concern, as it is more respectful of the varied experiences of trans embodiment, requiring a gender incongruence diagnosis, which is solely available on the basis of being trans, would, unfortunately, perpetuate a pathologising model.

DOUBLE STANDARDS
In their work on the ethics of informed consent in transgender care, Cavanaugh et al have pointed out that ‘[t]here is no scientific evidence of the benefit of (referral letter) requirements’ (p1151). The requirement is based on expert consensus instead of studies linking letter requirements to positive outcomes. On the contrary, studies have shown no adverse outcomes associated with the informed consent model.

I think their point warrants more probing. If trans bodies and lives are equally morally valuable to cis bodies—bodies belonging to cis people which have not been altered by transition-related interventions such as HRT—and lives, why should a psychological assessment be required in the absence of clear evidence of an important and overriding risk? Within a framework that sees trans people as mentally ill, the risk of misdiagnosis may seem to warrant such a requirement. But if we agree that being trans is not pathological and that trans people’s desire for HRT is simply part of normal human variance, can clear evidence of such a risk be established? I do not think so. For those who, like me, do not believe that such a risk is clearly evidenced, the maintenance of assessment requirements for HRT expresses a dehumanising devaluation of trans lives and bodies.

What counts as clear evidence of an important and overriding risk is debatable, but I would argue that HRT poses no more risk than various other medical interventions for which no psychological assessment is required. We do not typically think that it is ethical to require psychological assessments prior to abortions, for instance, an intervention which bears some parallels to transition-related care. Both are frequently justified by reference to personal autonomy and are frequently but not always motivated by distress, and yet neither pregnancy nor being trans is illness. I invite physicians to answer this question for themselves and
inquire into how double standards in clinical practices may reflect an unconscious hostility towards trans lives and experiences.

CONCLUSION
Physicians are slowly but surely moving towards providing HRT on an informed consent basis, without requiring psychological assessments. I hope that this article, informed by my personal and communal experiences as a trans scholar, will shed light on the ethics of requiring an assessment of gender dysphoria by a mental health professional prior to prescribing HRT. More than just unjustified, these requirements are dehumanising and pathologising for trans people and should be abandoned.

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REFERENCES