

<p>Certainty of evidence: very low</p>	<p>One uncontrolled, prospective observational longitudinal study (de Vries et al. 2011) provided evidence relating to the impact on anxiety in children and adolescents with gender dysphoria. Anxiety was measured using the Trait Anxiety Scale of the State-Trait Personality Inventory (STAI). This is a validated and commonly used measure of trait and state anxiety. It has 20 items and can be used in clinical settings to diagnose anxiety and to distinguish it from depressive illness. Higher scores indicate greater anxiety.</p> <p>The study provided evidence for anxiety at 2 time points:</p> <ul style="list-style-type: none"> • before starting a GnRH analogue (mean [\pmSD] age: 14.75 [\pm1.92] years), and • shortly before starting gender-affirming hormones (mean [\pmSD] age: 16.64 [\pm1.90] years). <p>The mean (\pmSD) anxiety (STAI) score was not statistically significantly different at baseline compared with follow-up (n=41, 39.43 [\pm10.07] versus 37.95 [\pm9.38], p=0.276) (VERY LOW).</p> <p>This study provides very low certainty evidence that treatment with GnRH analogues, before starting gender-affirming hormones, does not affect levels of anxiety.</p>
<p>Quality of life</p>	<p>This is a critical outcome because gender dysphoria in children and adolescents may be associated with a significant reduction in health-related quality of life.</p> <p>No evidence was identified.</p>
<p>Important outcomes</p>	
<p>Impact on body image</p> <p>Certainty of evidence: very low</p>	<p>This is an important outcome because some children and adolescents with gender dysphoria may want to take steps to suppress features of their physical appearance associated with their sex assigned at birth or accentuate physical features of their desired gender.</p> <p>One uncontrolled, prospective observational longitudinal study provided evidence relating to the impact on body image (de Vries et al. 2011). Body image was measured using the Body Image Scale (BIS) which is a validated 30-item scale covering 3 aspects: primary, secondary and neutral body characteristics. Higher scores represent a higher degree of body dissatisfaction.</p> <p>The study (de Vries et al. 2011) provided evidence for body image measured at 2 time points:</p> <ul style="list-style-type: none"> • before starting a GnRH analogue (mean [\pmSD] age: 14.75 [\pm1.92] years), and • shortly before starting gender-affirming hormones (mean [\pmSD] age: 16.64 [\pm1.90] years). <p>The mean (\pmSD) body image (BIS) scores for were not statistically significantly different from baseline compared with follow-up for:</p> <ul style="list-style-type: none"> • primary sexual characteristics (n=57, 4.10 [\pm0.56] versus 3.98 [\pm0.71], p=0.145) • secondary sexual characteristics (n=57, 2.74 [\pm0.65] versus 2.82 [\pm0.68], p=0.569)

	<ul style="list-style-type: none"> neutral body characteristics (n=57, 2.41 [±0.63] versus 2.47 [±0.56], p=0.620) (VERY LOW). <p>This study provides very low certainty evidence that treatment with GnRH analogues, before starting gender affirming hormones, does not affect body image.</p>
<p>Psychosocial impact: global functioning</p> <p>Certainty of evidence: very low</p>	<p>This is an important outcome because gender dysphoria in children and adolescents is associated with internalising and externalising behaviours, and emotional and behavioural problems which may impact on social and occupational functioning.</p> <p>One uncontrolled, observational, prospective cohort study (de Vries et al 2011) and one prospective cross-sectional cohort study (Costa et al. 2015) provided evidence relating to psychosocial impact in terms of global functioning. Global functioning was measured using the Children's Global Assessment Scale (CGAS). The CGAS tool is a validated measure of global functioning on a single rating scale from 1 to 100. Lower scores indicate poorer functioning.</p> <p>One study (de Vries et al. 2011) provided evidence for global functioning (CGAS) at 2 time points:</p> <ul style="list-style-type: none"> before starting a GnRH analogue (mean [±SD] age: 14.75 [±1.92] years), and shortly before starting gender-affirming hormones (mean [±SD] age: 16.64 [±1.90] years). <p>The mean (±SD) CGAS score was statistically significantly higher (improved) from baseline compared with follow-up (n=41, 70.24 [±10.12] versus 73.90 [±9.63], p=0.005) (VERY LOW).</p> <p>One study (Costa et al. 2015) in adolescents with gender dysphoria who had 6 months of psychological support followed by either GnRH analogues and continued psychological support (the immediately eligible group) or continued psychological support only (the delayed eligible group who did not receive GnRH analogues) provided evidence for global functioning (CGAS) measured at 4 time points:</p> <ul style="list-style-type: none"> at baseline (T0) in both groups, after 6 months of psychological support in both groups (T1), after 6 months of GnRH analogues and 12 months of psychological support in the immediately eligible group and 12 months of psychological support only in the delayed eligible group (T2), and after 18 months of psychological support and 12 months of GnRH analogues in the immediately eligible group and after 18 months of psychological support only in the delayed eligible group (T3). <p>The mean [±SD] CGAS score was statistically significantly higher (improved) for all adolescents (including those not receiving GnRH analogues) at T1, T2 or T3 compared with baseline (T0).</p> <p>For the immediately eligible group (who received GnRH analogues) versus the delayed eligible group (who did not receive GnRH analogues) there were no statistically significant differences in CGAS</p>

	<p>scores between the 2 groups at baseline T0 (n=201, p=0.23), T1 (n=201, p=0.73), T2 (n=121, p=0.49) or T3 (n=71, p=0.14) time points.</p> <p>For the immediately eligible group (who received GnRH analogues), the mean (\pmSD) CGAS score was not statistically significantly different at:</p> <ul style="list-style-type: none"> • T1 compared with T0 • T2 compared with T1 • T3 compared with T2. <p>The mean (\pmSD) CGAS score was statistically significantly higher (improved) at:</p> <ul style="list-style-type: none"> • T2 compared with T0 (n=60, 64.70 [\pm13.34] versus n=101, 58.72 [\pm11.38], p=0.003) • T3 compared with T0 (n=35, 67.40 [\pm13.39] versus n=101, 58.72 [\pm11.38], p<0.001) • T3 compared with T1 (n=35, 67.40 [\pm13.93] versus n=101, 60.89 [\pm12.17], p<0.001) (VERY LOW). <p>These studies provide very low certainty evidence that during treatment with GnRH analogues, global functioning may improve over time. However, there was no statistically significant difference in global functioning between GnRH analogues plus psychological support compared with psychological support only at any time point.</p>
<p>Psychosocial impact: psychosocial functioning</p> <p>Certainty of evidence: very low</p>	<p>This is an important outcome because gender dysphoria in children and adolescents is associated with internalising and externalising behaviours, and emotional and behavioural problems which may impact on social and occupational functioning.</p> <p>Two studies provided evidence for this outcome. One uncontrolled, observational, prospective cohort study (de Vries et al, 2011) and 1 cross-sectional observational study (Staphorsius et al. 2015) assessed psychosocial functioning using the Child Behaviour Checklist (CBCL) and the self-administered Youth Self-Report (YSR). The CBCL is a checklist parents complete to detect emotional and behavioural problems in children and adolescents. YSR is similar but is self-completed by the child or adolescent. The scales consist of a Total problems score, which is the sum of the scores of all the problem items. An internalising problem scale sums the anxious/depressed, withdrawn-depressed, and somatic complaints scores while the externalising problem scale combines rule-breaking and aggressive behaviour. The standard scores are scaled so that 50 is average for the child or adolescent's age and gender, with a SD of 10 points. Higher scores indicate greater problems, with a T-score above 63 considered to be in the clinical range.</p> <p>One study (de Vries et al. 2011) provided evidence for psychosocial functioning (CBCL and YSR scores) at 2 time points:</p> <ul style="list-style-type: none"> • before starting a GnRH analogue (mean [\pmSD] age: 14.75 [\pm1.92] years), and • shortly before starting gender-affirming hormones (mean [\pmSD] age: 16.64 [\pm1.90] years).

	<p>At follow up, the mean (\pmSD) CBCL scores were statistically significantly lower (improved) compared with baseline for:</p> <ul style="list-style-type: none"> • Total T score (n=54, 60.70 [\pm12.76] versus 54.46 [\pm11.23], $p < 0.001$) • Internalising T score (n=54, 61.00 [\pm12.21] versus 52.17 [\pm9.81], $p < 0.001$) • Externalising T score (n=54, 58.04 [\pm12.99] versus 53.81 [\pm11.86], $p = 0.001$). <p>At follow up, the mean (\pmSD) YSR scores were statistically significantly lower (improved) compared with baseline for:</p> <ul style="list-style-type: none"> • Total T score (n=54, 55.46 [\pm11.56] versus 50.00 [\pm10.56], $p < 0.001$) • Internalising T score (n=54, 56.04 [\pm12.49] versus 49.78 [\pm11.63], $p < 0.001$) • Externalising T score (n=54, 53.30 [\pm11.87] versus 49.98 [\pm9.35], $p = 0.009$). <p>The proportion of adolescents scoring in the clinical range decreased from baseline to follow up on the CBCL total problem scale (44.4% versus 22.2%, $p = 0.001$) and the internalising scale of the YSR (29.6% versus 11.1%, $p = 0.017$) (VERY LOW).</p> <p>One study (Staphorsius et al. 2015) assessed CBCL in a cohort of adolescents with gender dysphoria (transfemale: n=18, mean [\pmSD] age 15.1 [\pm2.4] years and transmale: n=22, mean [\pmSD] age 15.8 [\pm1.9] years) either receiving GnRH analogues (transfemale, n=8 and transmale, n=12), or not receiving GnRH analogues (transfemale, n=10 and transmale, n=10).</p> <p>The mean (\pmSD) CBCL scores for each group were (statistical analysis unclear):</p> <ul style="list-style-type: none"> • transfemales (total) 57.8 [\pm9.2] • transfemales receiving GnRH analogues 57.4 [\pm9.8] • transfemales not receiving GnRH analogues 58.2 [\pm9.3] • transmales (total) 60.4 [\pm10.2] • transmales receiving GnRH analogues 57.5 [\pm9.4] • transmales not receiving GnRH analogues 63.9 [\pm10.5] (VERY LOW). <p>These studies provide very low certainty evidence that during treatment with GnRH analogues psychosocial functioning may improve, with the proportion of adolescents in the clinical range for some CBCL and YSR scores decreasing over time.</p>
<p>Engagement with health care services</p> <p>Certainty of evidence: very low</p>	<p>This is an important outcome because patient engagement with health care services will impact on their clinical outcomes.</p> <p>Two uncontrolled observational cohort studies provided evidence relating to loss to follow up, which could be a marker of engagement with health care services (Brik et al. 2018 and Costa et al. 2015).</p> <p>In one retrospective study (Brik et al. 2018), 9 adolescents (9/214, 4.2%) who had stopped attending appointments were excluded from the study between November 2010 and July 2019 (VERY LOW).</p>

	<p>One prospective study (Costa et al. 2015) had evidence for a large loss to follow-up over time. The sample size at baseline (T0) and 6 months (T1) was 201, which dropped by 39.8% to 121 after 12 months (T2) and by 64.7% to 71 at 18 months follow-up (T3). No explanation of the reasons for loss to follow-up are reported (VERY LOW).</p> <p>Due to their design there was no reported loss to follow-up in the other 3 effectiveness studies (de Vries et al 2011; Khatchadourian et al. 2014; Staphorsius et al. 2015).</p> <p>These studies provide very low certainty evidence about loss to follow up, which could be a marker of engagement with health care services, during treatment with GnRH analogues. Due to the large variation in rates between studies no conclusions could be drawn.</p>
<p>Impact on extent of and satisfaction with surgery</p>	<p>This is an important outcome because some children and adolescents with gender dysphoria may proceed to transitioning surgery.</p> <p>No evidence was identified.</p>
<p>Stopping treatment</p> <p>Certainty of evidence: very low</p>	<p>This is an important outcome because there is uncertainty about the short- and long-term safety and adverse effects of GnRH analogues in children and adolescents with gender dysphoria.</p> <p>Two uncontrolled, retrospective, observational cohort studies provided evidence relating to stopping GnRH analogues. One study had complete reporting of the cohort (Brik et al. 2018), the other (Khatchadourian et al. 2014) had incomplete reporting of its cohort, particularly for transfemales where outcomes for only 4/11 were reported.</p> <p>Brik et al. 2018 narratively reported the reasons for stopping GnRH analogues in a cohort of 143 adolescents (38 transfemales and 105 transmales). Median age at the start of GnRH analogues was 15.0 years (range, 11.1–18.6 years) in transfemales and 16.1 years (range, 10.1–17.9 years) in transmales. Of these adolescents, 125 (87%, 36 transfemales, 89 transmales) subsequently started gender-affirming hormones after 1.0 (0.5–3.8) and 0.8 (0.3–3.7) years of GnRH analogues. At the time of data collection, the median duration of GnRH analogue use was 2.1 years (1.6–2.8).</p> <p>During the follow-up period 6.3% (9/143) of adolescents had discontinued GnRH analogues after a median duration of 0.8 years (range 0.1 to 3.0). The percentages and reasons for stopping were:</p> <ul style="list-style-type: none"> • 2.8% (4/143) stopped GnRH analogues although they wanted to continue endocrine treatments for gender dysphoria: <ul style="list-style-type: none"> ○ 1 transmale stopped due to increase in mood problems, suicidal thoughts and confusion attributed to GnRH analogues ○ 1 transmale had hot flushes, increased migraines, fear of injections, stress at school and unrelated medical issues, and temporarily stopped treatment (after 4 months) and restarted 5 months later. ○ 1 transmale had mood swings 4 months after starting GnRH analogues. After 2.2 years had unexplained severe nausea and rapid weight loss and discontinued GnRH analogues after 2.4 years

	<ul style="list-style-type: none"> ○ 1 transmale stopped GnRH analogues because of inability to regularly collect medication and attend appointments for injections. ● 3.5% (5/143) stopped treatment because they no longer wished to receive gender-affirming treatment for various reasons (VERY LOW). <p>Khatchadourian et al. 2014 narratively reported the reasons for stopping GnRH analogues in a cohort of 26 adolescents (15 transmales and 11 transfemales), 42% (11/26) discontinued GnRH analogues during follow-up between 1998 and 2011.</p> <p>Of 15 transmales receiving GnRH analogues, 14 received testosterone during the observation period, of which:</p> <ul style="list-style-type: none"> ● 7 continued GnRH analogues after starting testosterone ● 7 stopped GnRH analogues after a median of 3.0 years (range 0.2 to 9.2 years), of which: <ul style="list-style-type: none"> ○ 5 stopped after hysterectomy and salpingo-oophorectomy ○ 1 stopped after 2.2 years (transitioned to gender-affirming hormones) ○ 1 stopped after <2 months due to mood and emotional lability (VERY LOW). <p>Of 11 transfemales receiving GnRH analogues, 5 received oestrogen during the observation period, of which:</p> <ul style="list-style-type: none"> ● 4 continued GnRH analogues after starting oestrogen ● 1 stopped GnRH analogues when taking oestrogen (no reason reported) (VERY LOW). <p>Of the remaining 6 transfemales taking GnRH analogues:</p> <ul style="list-style-type: none"> ● 1 stopped GnRH analogues after a few months due to emotional lability ● 1 stopped GnRH analogues before taking oestrogen (the following year delayed due to heavy smoking) ● 1 stopped GnRH analogues after 13 months due not to pursuing transition (VERY LOW). <p>These studies provide very low certainty evidence for the number of adolescents who stop GnRH analogues and the reasons for this.</p>
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Abbreviations: GnRH, gonadotrophin releasing hormone; SD, standard deviation.

In children and adolescents with gender dysphoria, what is the short-term and long-term safety of GnRH analogues compared with one or a combination of psychological support, social transitioning to the desired gender or no intervention?

Outcome	Evidence statement
Safety	
Change in bone density: lumbar	This is an important outcome because puberty is an important time for bone development and puberty suppression may affect bone development, as shown by changes in lumbar bone density.

<p>Certainty of evidence: very low</p>	<p>Three uncontrolled, observational, retrospective studies provided evidence relating to the effect of GnRH analogues on bone density (based on lumbar BMAD) between starting with a GnRH analogue and at 1 and 2 year intervals (Joseph et al. 2019), and between starting GnRH analogues and starting gender-affirming hormones (Klink et al. 2015 and Vlot et al. 2017). All outcomes were reported separately for transfemales and transmales; also see subgroups table below.</p> <p>BMAD is a size adjusted value of BMD incorporating body size measurements using UK norms in growing adolescents. It was reported as g/cm³ and as z-scores. Z-scores report how many standard deviations from the mean a measurement sits. A z-score of 0 is equal to the mean, a z-score of -1 is equal to 1 standard deviation below the mean, and a z-score of +1 is equal to 1 standard deviation above the mean.</p> <p>One retrospective observational study (Joseph et al. 2019, n=70) provided non-comparative evidence on change in lumbar BMAD increase using z-scores.</p> <ul style="list-style-type: none"> • The z-score for lumbar BMAD was statistically significantly lower at 2 years compared with baseline in transfemales (z-score [±SD]: baseline 0.486 [0.809], 2 years -0.279 [0.930], p=0.000) and transmales (baseline -0.361 [1.439], 2 years -0.913 [1.318], p=0.001) (VERY LOW). • The z-score for lumbar BMAD was statistically significantly lower at 1 year compared with baseline in transfemales (baseline 0.859 [0.154], 1 year -0.228 [1.027], p=0.000) and transmales (baseline -0.186 [1.230], 1 year -0.541 [1.396], p=0.006) (VERY LOW). • Actual lumbar BMAD values in g/cm³ were not statistically significantly different between baseline and 1 or 2 years in transfemales or transmales (VERY LOW). <p>Two retrospective observational studies (Klink et al. 2015 and Vlot et al. 2017, n=104 in total) provided non-comparative evidence on change in lumbar BMAD between starting GnRH analogues and starting gender-affirming hormones. All outcomes were reported separately for transfemales and transmales; also see subgroups table below.</p> <p>In Klink et al. 2015 the z-score for lumbar BMAD was not statistically significantly different between starting GnRH analogues and starting gender-affirming hormones in transfemales but was statistically significantly lower when starting gender-affirming hormones in transmales (z-score mean [±SD]: GnRH analogue 0.28 [±0.90], gender-affirming hormone -0.50 [±0.81], p=0.004). Actual lumbar BMAD values in g/cm³ were not statistically significantly different between starting GnRH analogues and starting gender-affirming hormones in transfemales or transmales (VERY LOW).</p> <p>Vlot et al. 2017 reported change from starting GnRH analogues to starting gender-affirming hormones in lumbar BMAD by bone age.</p> <ul style="list-style-type: none"> • The z-score for lumbar BMAD in transfemales with a bone age of <15 years was statistically significantly lower at starting gender-affirming hormone treatment than at starting GnRH
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	<p>analogues (z-score median [range]: GnRH analogue -0.20 [-1.82 to 1.18], gender-affirming hormone -1.52 [-2.36 to 0.42], $p=0.001$) but was not statistically significantly different in transfemales with a bone age ≥ 15 years (VERY LOW).</p> <ul style="list-style-type: none"> • The z-score for lumbar BMAD in transmales with a bone age of < 14 years was statistically significantly lower at starting gender-affirming hormone treatment than at starting GnRH analogues (z-score median [range]: GnRH analogue -0.05 [-0.78 to 2.94], gender-affirming hormone -0.84 [-2.20 to 0.87], $p=0.003$) and in transmales with a bone age ≥ 14 years (GnRH analogue 0.27 [-1.60 to 1.80], gender-affirming hormone -0.29 [-2.28 to 0.90], $p \leq 0.0001$) (VERY LOW). • Actual lumbar BMAD values in g/cm^3 were not statistically significantly different between starting GnRH analogues and starting gender-affirming hormones in transfemales or transmales with young or old bone age (VERY LOW). <p>Two uncontrolled, observational, retrospective studies provided evidence for the effect of GnRH analogues on bone density (based on lumbar BMD) between starting GnRH analogues and either at 1 or 2 year intervals (Joseph et al. 2019), or starting gender-affirming hormones (Klink et al. 2015). All outcomes were reported separately for transfemales and transmales; also see subgroups table below.</p> <p>One retrospective observational study (Joseph et al. 2019, $n=70$) provided non-comparative evidence on change in lumbar BMD increase using z-scores.</p> <ul style="list-style-type: none"> • The z-score for lumbar BMD was statistically significantly lower at 2 years compared with baseline in transfemales (z-score mean [\pmSD]: baseline 0.130 [0.972], 2 years -0.890 [± 1.075], $p=0.000$) and transmales (baseline -0.715 [± 1.406], 2 years -2.000 [1.384], $p=0.000$) (VERY LOW). • The z-score for lumbar BMD was statistically significantly lower at 1 year compared with baseline in transfemales (z-score mean [\pmSD]: baseline -0.016 [± 1.106], 1 year -0.461 [± 1.121], $p=0.003$) and transmales (baseline -0.395 [± 1.428], 1 year -1.276 [± 1.410], $p=0.000$) (VERY LOW). • With the exception of transmales, where lumbar BMD in kg/m^2 increased between baseline and 1 year (mean [\pmSD]: baseline 0.694 [± 0.149], 1 year 0.718 [± 0.124], $p=0.006$), actual lumbar BMD values were not statistically significantly different between baseline and 1 or 2 years in transfemales or between 0 and 2 years in transmales (VERY LOW). <p>One retrospective observational study (Klink et al. 2015, $n=34$) provided non-comparative evidence on change in lumbar BMD between starting GnRH analogues and starting gender-affirming hormones.</p> <ul style="list-style-type: none"> • The z-score for lumbar BMD was not statistically significantly different between starting GnRH analogue and starting gender-affirming hormone treatment in transfemales, but was statistically significantly lower when starting gender-affirming hormones in transmales (z-score mean [\pmSD]: GnRH analogue 0.17 [± 1.18], gender-affirming hormone -0.72 [± 0.99], $p < 0.001$) (VERY LOW).
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	<ul style="list-style-type: none"> Actual lumbar BMD in g/cm² was not statistically significantly different between starting GnRH analogues and starting gender-affirming hormones in transfemales but was statistically significantly lower when starting gender-affirming hormones in transmales (mean [±SD]: GnRH analogues 0.95 [±0.12], gender-affirming hormones 0.91 [±0.10], p=0.006) (VERY LOW). <p>These studies provide very low certainty evidence that GnRH analogues reduce the expected increase in lumbar bone density (BMAD or BMD) compared with baseline (although some findings were not statistically significant). These studies also show that GnRH analogues do not statistically significantly decrease actual lumbar bone density (BMAD or BMD).</p>
<p>Change in bone density: femoral</p> <p>Certainty of evidence: very low</p>	<p>This is an important outcome because puberty is an important time for bone development and puberty suppression may affect bone development, as shown by changes in femoral bone density.</p> <p>Two uncontrolled, observational, retrospective studies provided evidence relating to the effect of GnRH analogues on bone density (based on femoral BMAD) between starting treatment with a GnRH analogue and starting gender-affirming hormones (Klink et al. 2015 and Vlot et al. 2017). All outcomes were reported separately for transfemales and transmales; also see subgroups table below.</p> <p>One retrospective observational study (Klink et al. 2015, n=34) provided non-comparative evidence on change in femoral area BMAD between starting GnRH analogues and starting gender-affirming hormones. All outcomes were reported separately for transfemales and transmales.</p> <ul style="list-style-type: none"> The z-score for femoral area BMAD was not statistically significantly different between starting GnRH analogues and starting gender-affirming hormones in transfemales or transmales (VERY LOW). Actual femoral area BMAD values were not statistically significantly different between starting GnRH analogues and starting gender-affirming hormones in transmales or transfemales (VERY LOW). <p>One retrospective observational study (Vlot et al. 2017, n=70) provided non-comparative evidence on change in femoral neck (hip) BMAD between starting GnRH analogues and starting gender-affirming hormones. All outcomes were reported separately for transfemales and transmales; also see subgroups table below.</p> <ul style="list-style-type: none"> The z-score for femoral neck BMAD in transfemales with a bone age of <15 years was not statistically significantly lower at starting gender-affirming hormones than at starting GnRH analogues (z-score median [range]: GnRH analogue -0.71 [-3.35 to 0.37], gender-affirming hormone -1.32 [-3.39 to 0.21], p≤0.1) or in transfemales with a bone age ≥15 years (GnRH analogue -0.44 [-1.37 to 0.93], gender-affirming hormone -0.36 [-1.50 to 0.46]) (VERY LOW). The z-score for femoral neck BMAD in transmales with a bone age of <14 years was not statistically significantly lower at starting gender-affirming hormones than at starting GnRH analogues (z-score median [range]: GnRH analogue -0.01

	<p>[-1.30 to 0.91], gender-affirming hormone -0.37 [-2.28 to 0.47]) but was statistically significantly lower in transmales with a bone age ≥ 14 years (GnRH analogue 0.27 [-1.39 to 1.32], gender-affirming hormone -0.27 [-1.91 to 1.29], $p=0.002$) (VERY LOW).</p> <ul style="list-style-type: none"> Actual femoral neck BMAD values were not statistically significantly different between starting GnRH analogues and starting gender-affirming hormones in transfemales or in transmales with a young bone age, but were statistically significantly lower in transmales with a bone age ≥ 14 years (GnRH analogue 0.33 [0.25 to 0.39], gender-affirming hormone 0.30 [0.23 to 0.41], $p \leq 0.01$) (VERY LOW). <p>Two uncontrolled, observational, retrospective studies provided evidence for the effect of GnRH analogues on bone density (based on femoral BMD) between starting GnRH analogues and either at 1 or 2 year intervals (Joseph et al. 2019), or starting gender-affirming hormones (Klink et al. 2015). All outcomes were reported separately for transfemales and transmales; also see subgroups table below.</p> <p>One retrospective observational study (Joseph et al. 2019, $n=70$) provided non-comparative evidence on change in femoral neck BMD increase using z-scores. All outcomes were reported separately for transfemales and transmales.</p> <ul style="list-style-type: none"> The z-score for femoral neck BMD was statistically significantly lower at 2 years compared with baseline in transfemales (z-score mean [\pmSD]: baseline 0.0450 [± 0.781], 2 years -0.600 [± 1.059], $p=0.002$) and transmales (baseline -1.075 [± 1.145], 2 years -1.779 [± 0.816], $p=0.001$) (VERY LOW). The z-score for femoral neck BMD was statistically significantly lower at 1 year compared with baseline in transfemales (z-score mean [\pmSD]: baseline 0.157 [± 0.905], 1 year -0.340 [± 0.816], $p=0.002$) and transmales (baseline -0.863 [± 1.215], 1 year -1.440 [± 1.075], $p=0.000$) (VERY LOW). Actual femoral neck BMD values in kg/m^2 were not statistically significantly different between baseline and 1 or 2 years in transmales or transfemales (VERY LOW). <p>One retrospective observational study (Klink et al. 2015, $n=34$) provided non-comparative evidence on change in femoral area BMD between starting GnRH analogues and starting gender-affirming hormones. All outcomes were reported separately for transfemales and transmales.</p> <ul style="list-style-type: none"> The z-score for femoral area BMD was not statistically significantly different between starting GnRH analogues and starting gender-affirming hormones in transfemales, but was statistically significantly lower in transmales (z-score mean [\pmSD]: GnRH analogue 0.36 [± 0.88], gender-affirming hormone -0.35 [± 0.79], $p=0.001$) (VERY LOW). Actual femoral area BMD values were not statistically significantly different between starting GnRH analogues and starting gender-affirming hormones in transfemales, but were statistically significantly lower in transmales (mean [\pmSD] GnRH analogue 0.92 [± 0.10], gender-affirming hormone 0.88 [± 0.09], $p=0.005$) (VERY LOW).
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	<p>These studies provide very low certainty evidence that GnRH analogues may reduce the expected increase in femoral bone density (femoral neck or area BMAD or BMD) compared with baseline (although some findings were not statistically significant). These studies also show that GnRH analogues do not statistically significantly decrease actual femoral bone density (femoral area BMAD or femoral neck BMD), apart from actual femoral area BMD in transmales.</p>
<p>Cognitive development or functioning</p> <p>Certainty of evidence: very low</p>	<p>This is an important outcome because puberty is an important time for cognitive development and puberty suppression may affect cognitive development or functioning.</p> <p>One cross-sectional observational study (Staphorsius et al. 2015, n=70) provided comparative evidence on cognitive development or functioning in adolescents with gender dysphoria on GnRH analogues compared with adolescents with gender dysphoria not on GnRH analogues. Cognitive functioning was measured using an IQ test. Reaction time (in seconds) and accuracy (percentage of correct trials) were measured using the Tower of London (ToL) task. All outcomes were reported separately for transfemales and transmales; also see subgroups table below. No statistical analyses or interpretation of the results in these groups were reported:</p> <ul style="list-style-type: none"> • IQ in transfemales (mean [±SD] GnRH analogue 94.0 [±10.3], control 109.4 [±21.2]). IQ transmales (GnRH analogue 95.8 [±15.6], control 98.5 [±15.9]). • Reaction time in transfemales (mean [±SD] GnRH analogue 10.9 [±4.1], control: 9.9 [±3.1]). Reaction time transmales (GnRH analogue 9.9 [±3.1], control 10.0 [±2.0]). • Accuracy score in transfemales (GnRH analogue 73.9 [±9.1], control 83.4 [±9.5]). Accuracy score in transmales (GnRH analogue 85.7 [±10.5], control 88.8 [±9.7]). <p>This study provides very low certainty evidence (with no statistical analysis) on the effects of GnRH analogues on cognitive development or functioning. No conclusions could be drawn.</p>
<p>Other safety outcomes: kidney function</p> <p>Certainty of evidence: very low</p>	<p>This is an important outcome because if renal damage (raised serum creatinine is a marker of this) is suspected, GnRH analogues may need to be stopped.</p> <p>One prospective observational study (Schagen et al. 2016, n=116) provided non-comparative evidence on change in serum creatinine between starting GnRH analogues and at 1 year. All outcomes were reported separately for transfemales and transmales; also see subgroups table below.</p> <ul style="list-style-type: none"> • There was no statistically significant difference between baseline and 1 year for serum creatinine in transfemales (mean [±SD] baseline 70 [±12], 1 year 66 [±13], p=0.20). • There was a statistically significant decrease between baseline and 1 year for serum creatinine in transmales (baseline 73 [±8], 1 year 68 [±13], p=0.01). <p>This study provides very low certainty evidence that GnRH analogues do not affect renal function.</p>

<p>Other safety outcomes: liver function</p> <p>Certainty of evidence: very low</p>	<p>This is an important outcome because if treatment-induced liver injury (raised liver enzymes are a marker of this) is suspected, GnRH analogues may need to be stopped.</p> <p>One prospective observational study (Schagen et al. 2016, n=116) provided non-comparative evidence on elevated liver enzymes between starting GnRH analogues and during use. No comparative values or statistical analyses were reported.</p> <ul style="list-style-type: none"> • Glutamyl transferase was not elevated at baseline or during use in any person. • Mild elevations of AST and ALT above the reference range were present at baseline but were not more prevalent during use than at baseline. • Glutamyl transferase, AST, and ALT levels did not significantly change from baseline to 12 months of use. <p>This study provides very low certainty evidence (with no statistical analysis) that GnRH analogues do not affect liver function.</p>
<p>Other safety outcomes: adverse effects</p> <p>Certainty of evidence: very low</p>	<p>This is an important outcome because if there are adverse effects, GnRH analogues may need to be stopped.</p> <p>One uncontrolled, retrospective, observational cohort study (Khatchadourian et al. 2014) provided evidence relating to adverse effects from GnRH analogues. It had incomplete reporting of its cohort, particularly for transfemales where outcomes for only 4/11 were reported.</p> <p>Khatchadourian et al. 2014 reported adverse effects in a cohort of 26 adolescents (15 transmales and 11 transfemales) receiving GnRH analogues. Of these:</p> <ul style="list-style-type: none"> • 1 transmale developed sterile abscesses; they were switched from leuprolide acetate to triptorelin, and this was well tolerated. • 1 transmale developed leg pains and headaches, which eventually resolved • 1 participant gained 19 kg within 9 months of starting GnRH analogues. <p>This study provides very low certainty evidence about potential adverse effects of GnRH analogues. No conclusions could be drawn.</p>

Abbreviations: ALT, alanine aminotransferase; AST, aspartate aminotransferase; BMAD, bone mineral apparent density; BMD, bone mineral density; GnRH, gonadotrophin releasing hormone; IQ, intelligence quotient; NS, not significant; SD, standard deviation.

In children and adolescents with gender dysphoria, what is the cost-effectiveness of GnRH analogues compared to one or a combination of psychological support, social transitioning to the desired gender or no intervention?

Outcome	Evidence statement
Cost-effectiveness	No studies were identified to assess the cost-effectiveness of GnRH analogues for children and adolescents with gender dysphoria.

From the evidence selected, are there any subgroups of children and adolescents with gender dysphoria that may benefit from GnRH analogues more than the wider population of interest?

Subgroup	Evidence statement
<p>Sex assigned at birth males (transfemales)</p> <p>Certainty of evidence: Very low</p>	<p>Some studies reported data separately for sex assigned at birth males (transfemales). This included some direct comparisons with sex assigned at birth females (transmales).</p> <p>Impact on gender dysphoria</p> <p>One uncontrolled prospective observational longitudinal study (de Vries et al. 2011) provided evidence for gender dysphoria in sex assigned at birth males. See the clinical effectiveness results table above for a full description of the study.</p> <p>The mean (\pmSD) UGDS score was statistically significantly lower (improved) in sex assigned at birth males compared with sex assigned at birth females at both baseline (T0) (n=not reported, mean UGDS score [\pmSD]: 47.95 [\pm9.70] versus 56.57 [\pm3.89]) and T1 (n=not reported, 49.67 [\pm9.47] versus 56.62 [\pm4.00]); between sex difference $p < 0.001$ (VERY LOW).</p> <p>One further prospective observational longitudinal study (Costa et al. 2015) provided evidence for the impact on gender dysphoria in sex assigned at birth males. See the clinical effectiveness results table above for a full description of the study. Sex assigned at birth males had a statistically significantly lower (improved) mean (\pmSD) UGDS score of 51.6 [\pm9.7] compared with sex assigned at birth females (56.1 [\pm4.3], $p < 0.001$). However, it was not reported if this was baseline or follow-up (VERY LOW).</p> <p>These studies provide very low certainty evidence that in sex assigned at birth males (transfemales), gender dysphoria is lower than in sex assigned at birth females (transmales).</p> <p>Impact on mental health</p> <p>One uncontrolled prospective observational longitudinal study (de Vries et al. 2011) provided evidence for the impact on mental health (depression, anger and anxiety) in sex assigned at birth males. See the clinical effectiveness results table above for a full description of the study.</p> <ul style="list-style-type: none"> • The mean (\pmSD) depression (BDI-II) score was not statistically significantly different in sex assigned at birth males compared with sex assigned at birth females at both baseline (T0) (n=not reported, mean BDI score [\pmSD]: 5.71 [\pm4.31] versus 10.34 [\pm8.24]) and T1 (n=not reported, 3.50 [\pm4.58] versus 6.09 [\pm7.93]), between sex difference $p = 0.057$ • The mean (\pmSD) anger (TPI) score was statistically significantly lower (improved) in sex assigned at birth males compared with sex assigned at birth females at both baseline (T0) (n=not reported, mean TPI score [\pmSD]: 5.22 [\pm2.76] versus 6.43 [\pm2.78]) and T1 (n=not reported, 5.00 [\pm3.07] versus 6.39 [\pm2.59]), between sex difference $p = 0.022$ • The mean (\pmSD) anxiety (STAI) score was statistically significantly lower (improved) in sex assigned at birth males

compared with sex assigned at birth females at both baseline (T0) (n=not reported, mean STAI score [\pm SD]: 4.33 [\pm 2.68] versus 7.00 [\pm 2.36]) and T1 (n=not reported, 4.39 [\pm 2.64] versus 6.17 [\pm 2.69]), between sex difference $p < 0.001$ (**VERY LOW**).

This study provides very low certainty evidence that the impact on mental health (depression, anger and anxiety) may be different in sex assigned at birth males (transfemales) compared with sex assigned at birth females (transmales). Over time there was no statistically significant difference between sex assigned at birth males and sex assigned at birth females for depression. However, sex assigned at birth males had statistically significantly lower levels of anger and anxiety than sex assigned at birth females at both baseline and follow up.

Impact on body image

One uncontrolled prospective observational longitudinal study ([de Vries et al. 2011](#)) provided evidence relating to the impact on body image in sex assigned at birth males.

- The mean (\pm SD) BIS score for primary sex characteristics was statistically significantly lower (improved) in sex assigned at birth males compared with sex assigned at birth females at both baseline (T0) (n=not reported, mean BIS score [\pm SD]: 4.02 [\pm 0.61] versus 4.16 [\pm 0.52]) and T1 (n=not reported, 3.74 [\pm 0.78] versus 4.17 [\pm 0.58]), between sex difference $p = 0.047$
- The mean (\pm SD) BIS score for secondary sex was statistically significantly lower (improved) in sex assigned at birth males compared with sex assigned at birth females at both baseline (T0) (n=not reported, mean BIS score [\pm SD]: 2.66 [\pm 0.50] versus 2.81 [\pm 0.76]) and T1 (n=not reported, 2.39 [\pm 0.69] versus 3.18 [\pm 0.42]), between sex difference $p = 0.001$
- The mean (\pm SD) BIS score for neutral body characteristics was not statistically significantly different in sex assigned at birth males compared with sex assigned at birth females at both baseline (T0) (n=not reported, mean BIS score [\pm SD]: 2.60 [\pm 0.58] versus 2.24 [\pm 0.62]) and T1 (n=not reported, 2.32 [\pm 0.59] versus 2.61 [\pm 0.50]), between sex difference $p = 0.777$ (**VERY LOW**).

This study provides very low certainty evidence that the impact on body image may be different in sex assigned at birth males (transfemales) compared with sex assigned at birth females (transmales). Sex assigned at birth males are less dissatisfied with their primary and secondary sex characteristics than sex assigned at birth females at both baseline and follow up, but the satisfaction with neutral body characteristics is not different.

Psychosocial impact

One uncontrolled prospective observational longitudinal study ([de Vries et al. 2011](#)) provided evidence for psychosocial impact in terms of global functioning (CGAS) and psychosocial functioning (CBCL and YSR) in sex assigned at birth males.

- Sex assigned at birth males had statistically higher mean (\pm SD) CGAS scores compared with sex assigned at birth

	<p>females at both baseline (T0) (n=54, 73.10 [\pm8.44] versus 67.25 [\pm11.06]) and T1 (n=54, 77.33 [\pm8.69] versus 70.30 [\pm9.44]), between sex difference p=0.021</p> <ul style="list-style-type: none"> • There was no statistically significant difference between sex assigned at birth males and sex assigned at birth females for the CBCL Total T score at T0 or T1 (n=54, p=0.110) • There was no statistically significant difference between sex assigned at birth males and sex assigned at birth females for the CBCL internalising T score at T0 or T1 (n=54, p=0.286) • Sex assigned at birth males had statistically lower mean (\pmSD) CBCL externalising T scores compared with sex assigned at birth females at both T0 (n=54, 54.71 [\pm12.91] versus 60.70 [\pm12.64]) and T1 (n=54, 48.75 [\pm10.22] versus 57.87 [\pm11.66]), between sex difference p=0.015 • There was no statistically significant difference between sex assigned at birth males and sex assigned at birth females for the YSR Total T score at T0 or T1 (n=54, p=0.164) • There was no statistically significant difference between sex assigned at birth males and sex assigned at birth females for the YSR internalising T score at T0 or T1 (n=54, p=0.825) • Sex assigned at birth males had statistically lower mean (\pmSD) YSR externalising T scores compared with sex assigned at birth females at both T0 (n=54, 48.72 [\pm11.38] versus 57.24 [\pm10.59]) and T1 (n=54, 46.52 [\pm9.23] versus 52.97 [\pm8.51]), between sex difference p=0.004 (VERY LOW). <p>One uncontrolled, observational, prospective cohort study (Costa et al. 2015) provided evidence for psychosocial impact in terms of global functioning (CGAS) in sex assigned at birth males.</p> <ul style="list-style-type: none"> • Sex assigned at birth males had statistically significant lower mean (\pmSD CGAS scores at baseline) compared with sex assigned at birth females (n=201, 55.4 [\pm12.7] versus 59.2 [\pm11.8], p=0.03) (VERY LOW). <p>These studies provide very low certainty evidence that psychosocial impact may be different in sex assigned at birth males (transfemales) compared with sex assigned at birth females (transmales). However, no conclusions could be drawn.</p> <p>Change in bone density: lumbar</p> <p>Three uncontrolled, observational, retrospective studies provided evidence relating to the effect of GnRH analogues on lumbar bone density in sex assigned at birth males (Joseph et al. 2019, Klink et al. 2015 and Vlot et al. 2017). See the safety results table above for a full description of the results.</p> <p>These studies provide very low certainty evidence that GnRH analogues reduce the expected increase in lumbar bone density (BMAD or BMD) in sex assigned at birth males (transfemales; although some findings were not statistically significant). These studies also show that GnRH analogues do not statistically significantly decrease actual lumbar bone density (BMAD or BMD) in sex assigned at birth males (transfemales).</p> <p>Change in bone density: femoral</p>
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	<p>Three uncontrolled, observational, retrospective studies provided evidence for the effect of GnRH analogues on femoral bone density in sex assigned at birth males (Joseph et al. 2019, Klink et al. 2015 and Vlot et al. 2017). See the safety results table above for a full description of the results.</p> <p>These studies provide very low certainty evidence that GnRH analogues may reduce the expected increase in femoral bone density (femoral neck or area BMAD or BMD) in sex assigned at birth males (transfemales; although some findings were not statistically significant). These studies also show that GnRH analogues do not statistically significantly decrease actual femoral bone density (femoral area BMAD or femoral neck BMD) in sex assigned at birth males (transfemales).</p> <p>Cognitive development or functioning One cross-sectional observational study (Staphorsius et al. 2015) provided comparative evidence on cognitive development or functioning in sex assigned at birth males. See the safety results table above for a full description of the results.</p> <p>This study provides very low certainty evidence (with no statistical analysis) on the effects of GnRH analogues on cognitive development or functioning in sex assigned at birth males (transfemales). No conclusions could be drawn.</p> <p>Other safety outcomes: kidney function One prospective observational study (Schagen et al. 2016) provided non-comparative evidence on change in serum creatinine in sex assigned at birth males. See the safety results table above for a full description of the results.</p> <p>This study provides very low certainty evidence that GnRH analogues do not affect renal function in sex assigned at birth males (transfemales).</p>
<p>Sex assigned at birth females (transmales)</p> <p>Certainty of evidence: Very low</p>	<p>Some studies reported data separately for sex assigned at birth females (transmales). This included some direct comparisons with sex assigned at birth males (transfemales).</p> <p>Impact on gender dysphoria One uncontrolled prospective observational longitudinal study (de Vries et al. 2011) and one prospective observational longitudinal study (Costa et al. 2015) provided evidence for gender dysphoria in sex assigned at birth females. See the sex assigned at birth males (transfemales) row above for a full description of the results.</p> <p>These studies provide very low certainty evidence that in sex assigned at birth females (transmales), gender dysphoria is higher than in sex assigned at birth males (transfemales) at both baseline and follow up.</p> <p>Impact on mental health One uncontrolled prospective observational longitudinal study (de Vries et al. 2011) provided evidence relating to the impact on mental health (depression, anger and anxiety) in sex assigned at birth</p>

females. See the sex assigned at birth males (transfemales) row above for a full description of the results.

This study provides very low certainty evidence that the impact on mental health (depression, anger and anxiety) may be different in sex assigned at birth females (transmales) compared with sex assigned at birth males (transfemales). Over time there was no statistically significant difference between sex assigned at birth females and sex assigned at birth males for depression. However, sex assigned at birth females had statistically significantly greater levels of anger and anxiety than sex assigned at birth males at baseline and follow up.

Impact on body image

One uncontrolled prospective observational longitudinal study ([de Vries et al. 2011](#)) provided evidence relating to the impact on body image in sex assigned at birth females. See the sex assigned at birth males (transfemales) row above for a full description of the results.

This study provides very low certainty evidence that the impact on body image may be different in sex assigned at birth females (transmales) compared with sex assigned at birth males (transfemales). Sex assigned at birth females are more dissatisfied with their primary and secondary sex characteristics than sex assigned at birth males at both baseline and follow up, but the satisfaction with neutral body characteristics is not different.

Psychosocial impact

One uncontrolled prospective observational longitudinal study ([de Vries et al. 2011](#)) provided evidence for psychosocial impact in terms of global functioning (CGAS) and psychosocial functioning (CBCL and YSR) in sex assigned at birth females. One uncontrolled, observational, prospective cohort study ([Costa et al. 2015](#)) provided evidence for psychosocial impact in terms of global functioning (CGAS) in sex assigned at birth females. See the sex assigned at birth males (transfemales) row above for a full description of the results.

These studies provide very low certainty evidence that psychosocial impact may be different in sex assigned at birth females (transmales) compared with sex assigned at birth males (transfemales). However, no conclusions could be drawn.

Change in bone density: lumbar

Three uncontrolled, observational, retrospective studies provided evidence relating to the effect of GnRH analogues on lumbar bone density in sex assigned at birth females ([Joseph et al. 2019](#), [Klink et al. 2015](#) and [Vlot et al. 2017](#)). See the safety results table above for a full description of the results.

These studies provide very low certainty evidence that GnRH analogues reduce the expected increase in lumbar bone density (BMAD or BMD) in sex assigned at birth females (transmales; although some findings were not statistically significant). These studies also show that GnRH analogues do not statistically

	<p>significantly decrease actual lumbar bone density (BMAD or BMD) in sex assigned at birth females (transmales).</p> <p>Change in bone density: femoral Three uncontrolled, observational, retrospective studies provided evidence relating to the effect of GnRH analogues on femoral bone density in sex assigned at birth females (Joseph et al. 2019, Klink et al. 2015 and Vlot et al. 2017). See the safety results table above for a full description of the results.</p> <p>These studies provide very low certainty evidence that GnRH analogues may reduce the expected increase in femoral bone density (femoral neck or area BMAD or BMD) in sex assigned at birth females (transmales; although some findings were not statistically significant). These studies also show that GnRH analogues do not statistically significantly decrease actual femoral bone density (femoral area BMAD or femoral neck BMD) in sex assigned at birth females (transmales), apart from actual femoral area.</p> <p>Cognitive development or functioning One cross-sectional observational study (Staphorsius et al. 2015) provided comparative evidence on cognitive development or functioning in sex assigned at birth females. See the safety results table above for a full description of the results.</p> <p>This study provides very low certainty evidence (with no statistical analysis) on the effects of GnRH analogues on cognitive development or functioning in sex assigned at birth females (transmales). No conclusions could be drawn.</p> <p>Other safety outcomes: kidney function One prospective observational study (Schagen et al. 2016) provided non-comparative evidence on change in serum creatinine in sex assigned at birth females (transmales). See the safety results table above for a full description of the results.</p> <p>This study provides very low certainty evidence that GnRH analogues do not affect renal function in sex assigned at birth females (transmales).</p>
Duration of gender dysphoria	No evidence was identified.
Age at onset of gender dysphoria	No evidence was identified.
Age at which GnRH analogue started	No evidence was identified.
Age at onset of puberty	No evidence was identified.
Tanner stage at which GnRH analogue started	No evidence was identified.
Diagnosis of autistic spectrum disorder	No evidence was identified.

Diagnosis of mental health condition	No evidence was identified.
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Abbreviations: BDI-II, Beck Depression Inventory-II; BIS, Body Image Scale; CBCL, Child Behaviour Checklist; CGAS, Children's Global Assessment Scale; SD, standard deviation; STAI, Trait Anxiety Scale of the State-Trait Personality Inventory; TPI, Trait Anger Scale of the State-Trait Personality Inventory; UGDS, Utrecht Gender Dysphoria Scale; YSR, Youth Self-Report

From the evidence selected,

- (a) what are the criteria used by the research studies to define gender dysphoria, gender identity disorder and gender incongruence of childhood?
- (b) what were the ages at which participants commenced treatment with GnRH analogues?
- (c) what was the duration of treatment with GnRH analogues?

Outcome	Evidence statement																
Diagnostic criteria	<p>In 5 studies (Costa et al. 2015, Klink et al. 2015, Schagen et al. 2016, Staphorsius et al. 2015 and Vlot et al. 2017) the DSM-IV-TR criteria of gender identity disorder was used.</p> <p>The study by Brik et al. 2020 used DSM-V criteria. The DSM-V has one overarching definition of gender dysphoria with separate specific criteria for children and for adolescents and adults. The general definition describes a conflict associated with significant distress and/or problems functioning associated with this conflict between the way they feel and the way they think of themselves which must have lasted at least 6 months.</p> <p>It was not reported how gender dysphoria was defined in the remaining 3 studies (VERY LOW).</p> <p>From the evidence selected, all studies that reported diagnostic criteria for gender dysphoria (6/9 studies) used the DSM criteria in use at the time the study was conducted.</p>																
Age when GnRH analogues started	<p>8/9 studies reported the age at which participants started GnRH analogues, either as the mean age (with SD) or median age (with the range):</p> <table border="1"> <thead> <tr> <th>Study</th> <th>Mean age (±SD)</th> </tr> </thead> <tbody> <tr> <td>Costa et al. 2015</td> <td>16.5 years (±1.3)</td> </tr> <tr> <td>de Vries et al. 2011</td> <td>13.6 years (±1.8)</td> </tr> <tr> <td>Joseph et al. 2019</td> <td>13.2 years (±1.4) in transfemales 12.6 years (±1.0) in transmales</td> </tr> <tr> <td>Khatchadourian et al. 2014</td> <td>14.7 years (±1.9)</td> </tr> <tr> <td>Klink et al. 2015</td> <td>14.9 years (±1.9) in transfemales 15.0 years (±2.0) in transmales</td> </tr> </tbody> </table> <table border="1"> <thead> <tr> <th>Study</th> <th>Median age (range)</th> </tr> </thead> <tbody> <tr> <td>Brik et al. 2020</td> <td>15.5 years (11.1–18.6) in transfemales 16.1 years (10.1–17.9) in transmales</td> </tr> </tbody> </table>	Study	Mean age (±SD)	Costa et al. 2015	16.5 years (±1.3)	de Vries et al. 2011	13.6 years (±1.8)	Joseph et al. 2019	13.2 years (±1.4) in transfemales 12.6 years (±1.0) in transmales	Khatchadourian et al. 2014	14.7 years (±1.9)	Klink et al. 2015	14.9 years (±1.9) in transfemales 15.0 years (±2.0) in transmales	Study	Median age (range)	Brik et al. 2020	15.5 years (11.1–18.6) in transfemales 16.1 years (10.1–17.9) in transmales
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	Vlot et al. 2017	13.5 years (11.5–18.3) in transfemales 15.1 years (11.7–18.6) in transmales
	Age at the start of GnRH analogues was not reported in Staphorsius et al. 2015, but participants were required to be at least 12 years (VERY LOW).	
	The evidence included showed wide variation in the age (11 to 18 years old) at which children and adolescents with gender dysphoria started GnRH analogues.	
Duration of treatment	The duration of treatment with GnRH analogues was reported in 3/9 studies. The median duration was: <ul style="list-style-type: none"> • 2.1 years (range 1.6–2.8) in Brik et al. 2020. • 1.3 years (range 0.5–3.8) in transfemales and 1.5 years (range 0.25–5.2) in transmales in Klink et al. 2015. <p>In Staphorsius et al. 2015, the mean duration was 1.6 years (SD ±1.0).</p> <p>In de Vries et al. 2011, the mean duration of time between starting GnRH analogues and gender-affirming hormones was 1.88 years (SD ±1.05).</p> <p>The evidence included showed wide variation in the duration of treatment with GnRH analogues, but most studies did not report this information. Treatment duration ranged from a few months up to about 5 years.</p>	

Abbreviations: DSM, Diagnostic and Statistical Manual of Mental Disorders criteria; SD, standard deviation.

6. Discussion

A key limitation to identifying the effectiveness and safety of GnRH analogues for children and adolescents with gender dysphoria is the lack of reliable comparative studies. The lack of clear, expected outcomes from treatment with a GnRH analogue (the purpose of which is to suppress secondary sexual characteristics which may cause distress from unwanted pubertal changes) also makes interpreting the evidence difficult. The size of the population with gender dysphoria means conducting a prospective trial may be unrealistic, at least on a single centre basis. There may also be ethical issues with a 'no treatment arm' in comparative trials of GnRH analogues, where there may be poor mental health outcomes if treatment is withheld. However, the use of an active comparator such as close psychological support may reduce ethical concerns in future trials.

The studies included in this evidence review are all small, uncontrolled observational studies, which are subject to bias and confounding, and are of very low certainty as assessed using modified GRADE. All the included studies reported physical and mental health comorbidities and concomitant treatments very poorly. For example, very little data are reported on how many children and adolescents needed additional mental health support, and for what reasons, or whether additional interventions, and what form and duration (for example drug treatment or counselling) that took. This is a possible confounder for the treatment outcomes in the studies because changes in critical and important

outcomes may be attributable to external care rather than the psychological support or GnRH analogues used in the studies.

The studies that reported diagnostic criteria for gender dysphoria (6/9 studies) used the Diagnostic and Statistical Manual of Mental Disorders (DSM) criteria in use at the time the study was conducted (either DSM-IV-TR or DSM-V). The definition was unclear in the remaining studies. There was wide variation in the ages at which participants started a GnRH analogue, typically ranging from about 11 to 18 years. Similarly, there was a wide variation in the duration of use, but few studies reported this.

Changes in outcome scores for clinical effectiveness were assessed for statistical significance in the 3 studies reporting these outcomes ([Costa et al. 2015](#); [de Vries et al. 2011](#); [Staphorsius et al. 2015](#)). However, there is relatively little interpretation of whether the changes in outcome scores seen in these studies are clinically meaningful.

For some outcomes there was no statistically significant difference from before starting GnRH analogues until just before starting gender-affirming hormones. These were the Utrecht Gender Dysphoria Scale (UGDS) (which was assessed in 1 study [de Vries et al. 2011](#)), the Trait Anger (TPI) and Trait Anxiety (STAI) Scales (which were assessed in 1 study [de Vries et al. 2011](#)), and Body Image Scale (BIS) which was assessed in 1 study ([de Vries et al. 2011](#)).

The Beck Depression Inventory (BDI-II) was used in 1 study ([de Vries et al. 2011](#)) to assess change in depression from before starting GnRH analogues to just before starting gender-affirming hormones. The result is statistically significant, with the mean (\pm SD) BDI-II score decreasing from 8.31 (\pm 7.12) at baseline to 4.95 (\pm 6.27) at follow up ($p=0.004$). However, both scores fall into the minimal range using the general guidelines for interpretation of BDI-II (0 to 13 minimal, 14 to 19 mild depression, 20 to 28 moderate depression and 29 to 63 severe depression), suggesting that while statistically significant, it is unclear if this is a clinically meaningful change.

Psychosocial outcomes were assessed in 3 studies ([Costa et al. 2015](#); [de Vries et al. 2011](#); [Staphorsius et al. 2015](#)) using the Children's Global Assessment Scale (CGAS) and Child Behavior Checklist/Youth Self-Report (CBCL/YSR). The CGAS score was assessed in 2 studies ([Costa et al. 2015](#); [de Vries et al. 2011](#)). In [de Vries et al. 2011](#) the mean (\pm SD) CGAS score statistically significantly increased over time from 70.24 [\pm 10.12] at baseline to 73.90 [\pm 9.63] at follow up. CGAS scores are clinically categorised into 10 categories (10 to 1, 20 to 11 and so on until 100 to 91) and both scores reported were in a single category (71 to 80, no more than slight impairment) suggesting that while statistically significant, it is unclear if this is a clinically meaningful change. The [Costa et al. 2015](#) study does highlight a larger change in CGAS scores from baseline to follow-up (mean [\pm SD] 58.72 [\pm 11.38] compared with 67.40 [\pm 13.39]), but whether this is clinically meaningful is unclear. The average score moved from the clinical category of 60 to 51 (variable functioning with sporadic difficulties) at baseline to 70 to 61 (some difficulty in a single area, but generally functioning pretty well) at follow up, but the large standard deviations suggest clinically significant overlaps between the scores from baseline to follow-up.

Psychosocial functioning using the CBCL/YSR was assessed in 2 studies ([de Vries et al. 2011](#); [Staphorsius et al. 2015](#)). In [de Vries et al. 2011](#) there was a statistically significant reduction in both CBCL and YSR scores from before starting GnRH analogues to just before

starting gender-affirming hormones. The study interpreted the CBCL/YSR with a proportion of adolescents who scored in the clinical range (a T-score above 63), which allows changes in clinically meaningful scores to be assessed, and proportions of adolescents in the clinical range for some CBCL and YSR scores decreased over time. One cross-sectional study ([Staphorsius et al. 2015](#)) assessed CBCL scores only, but it was unclear if this was the Total T score, or whether subscales of internalising or externalising scores were also assessed, and whether the results were statistically significant.

The 2 prospective observational studies ([Costa et al. 2015](#); [de Vries et al. 2011](#)) are confounded by a number of common factors. Firstly, the single assessment of scores at baseline means it is unclear if scores were stable, already improving or declining before starting treatment. Secondly, in an uncontrolled study any changes in scores from baseline to follow-up could be attributed to a regression-to-mean, for example getting older has been positively associated with maturity and wellbeing. The studies use mean and standard deviations in the descriptive statistics and analyses; however, they do not report testing the normality of data which would support the use of parametric measures. The study by de Vries et al. 2011 used general linear models (regression) to examine between and within group variances (changes in outcomes). In using such models, the data is assumed to be balanced (measured at regular intervals and without missing data), but the large ranges in ages at which participants were assessed and started on various interventions suggests that ascertainment of outcome was unlikely to be regular and missing data was likely. Missing data was handled through listwise deletion (omits those cases with the missing data and analyses the remaining data) which is acceptable if data loss is completely random but for some outcomes where there was incomplete data for individual items this was not random (items were introduced by the authors after the first eligible adolescents had started GnRH analogues). The study provided no detail on whether these assumptions for the modeling were met, they also provided no adequate assessment of whether any regression diagnostics (analysis that seek to assess the validity of a model) or model fit (how much of the variance in outcome is explained by the between and within group variance) were undertaken.

The 2 retrospective observational studies ([Brik et al. 2020](#); [Khatchadourian et al. 2014](#)) both only report absolute numbers for each trajectory along with reasons for stopping GnRH analogues. It is difficult to assess outcomes from such single centre studies because there is little comparative data for outcomes from other such services. A lack of any critical or other important outcomes also means the success of the treatment across all the participants is difficult to judge.

Three uncontrolled, observational, retrospective studies provided evidence relating to the effect of GnRH analogues on bone density ([Joseph et al. 2019](#); [Klink et al. 2015](#); [Vlot et al. 2017](#)). In all 3 studies, the participants acted as their own controls and change in bone density was determined between starting GnRH analogues and either after 1 and 2 year follow-up timepoints (Joseph et al. 2019) or when gender-affirming hormones were started (Klink et al. 2015 and Vlot et al. 2017). Observational studies such as these can only show an association with GnRH analogues and bone density; they cannot show that GnRH analogues caused any differences in bone density seen. Because there was no comparator group and participants acted as their own controls, it is unclear whether the findings are associated with GnRH analogues or due to changes over time. The authors reported z-scores which allows for comparison with the expected increase in bone density in the

general population. However, because no concomitant treatments or comorbidities were reported it is possible that the findings may not be because of GnRH analogues and there is another way in which the study population differs from the general population.

All the studies are from a limited number of, mainly European, care facilities. They are described as either tertiary referral or expert services but the low number of services providing such care and publishing evidence may bias the results towards the outcomes in these services only and limit extrapolation.

The first study ([Brik et al. 2020](#)) was an uncontrolled, retrospective, observational study that assessed the outcome trajectories of adolescents receiving GnRH analogues for gender dysphoria. This study followed-up 143 individuals who had received GnRH analogues (38 transfemales and 105 transmales) using clinical records to show outcomes for up to 9 years (continuing use of GnRH analogues, reasons for stopping GnRH analogues and onward care such as gender-affirming hormone use). The methods and results are well reported, but no analysis of data was undertaken. The views of adolescents and their parents are particularly difficult to interpret because no data on how many responded to each question and in what ways are reported.

The second study ([Costa et al. 2015](#)) was an uncontrolled, prospective observational study which assessed global functioning in adolescents with gender dysphoria using CGAS every 6 months, including during the first 6 months where statistically significant improvements were seen without GnRH analogues. The study is confounded by significant unexplained loss to follow-up (64.7%: from n=201 adolescents to n=71 after 18 months). Missing data for those lost to follow-up maybe more than sufficient to change the direction of effects seen in the study if the reasons for loss to follow-up are systematic (such as deriving little or no benefit from treatment). The study uses clustered data in its analysis, a single outcome (CGAS) measured in clusters (at different visits), and the analysis does not take account of the correlation of scores (data at different time points are not independent) as a significant change in scores early in the study means the successive changes measured against baseline were also significant. The study relies on multiple (>20) pairwise independent *t*-tests to examine change in CGAS between the 4 time points, increasing the possibility of type-I error (a false positive which occurs when a researcher incorrectly rejects a true null hypothesis) because the more tests performed the more likely a statistically significant result will be observed by chance alone.

The [Costa et al. 2015](#) study compares immediately eligible and delayed eligible cohorts, however, it is highly likely that they are non-comparable groups because the immediately eligible group were those able to start GnRH analogues straight away whilst those in the delayed eligible group were either not ready to make a decision about starting treatment (no age comparison was made between the 2 groups so it is unclear if they were a younger cohort than the immediately eligible group) or had comorbid mental health or psychological difficulties. The authors report that those with concomitant problems (such as mental health problems, substantial problems with peers, or conflicts with parents or siblings) were referred to local mental health services but no details are provided.

The third study ([de Vries et al. 2011](#)) was an uncontrolled, prospective observational study which assessed gender dysphoria and psychological functioning before and after puberty suppression in adolescents with gender dysphoria. Although the study mentions the DSM-IV-TR there is no explicit discussion of this, or any other criteria, being used as the