Hormone supplementation for pubertal induction in girls
Matthews, Debbie; Bath, Louise; Högler, Wolfgang; Mason, Avril; Smyth, Arlene; Skae, Mars

DOI:
10.1136/archdischild-2016-311372

License:
None: All rights reserved

Document Version
Peer reviewed version

Citation for published version (Harvard):
https://doi.org/10.1136/archdischild-2016-311372

Link to publication on Research at Birmingham portal

Publisher Rights Statement:
Eligibility for repository: Checked on 10/4/2017

This article has been accepted for publication in Archives of Disease in Childhood, 2017 following peer review, and the Version of Record can be accessed online at http://dx.doi.org/10.1136/archdischild-2016-311372

General rights
Unless a licence is specified above, all rights (including copyright and moral rights) in this document are retained by the authors and/or the copyright holders. The express permission of the copyright holder must be obtained for any use of this material other than for purposes permitted by law.

• Users may freely distribute the URL that is used to identify this publication.
• Users may download and/or print one copy of the publication from the University of Birmingham research portal for the purpose of private study or non-commercial research.
• Users may use extracts from the document in line with the concept of 'fair dealing' under the Copyright, Designs and Patents Act 1988 (?)
• Users may not further distribute the material nor use it for the purposes of commercial gain.

Where a licence is displayed above, please note the terms and conditions of the licence govern your use of this document.

When citing, please reference the published version.

Take down policy
While the University of Birmingham exercises care and attention in making items available there are rare occasions when an item has been uploaded in error or has been deemed to be commercially or otherwise sensitive.

If you believe that this is the case for this document, please contact UBIRA@lists.bham.ac.uk providing details and we will remove access to the work immediately and investigate.
Hormone Supplementation For Pubertal Induction In Girls

<table>
<thead>
<tr>
<th>Journal:</th>
<th>Archives of Disease in Childhood</th>
</tr>
</thead>
<tbody>
<tr>
<td>Manuscript ID</td>
<td>archdischild-2016-311372.R1</td>
</tr>
<tr>
<td>Article Type:</td>
<td>Review</td>
</tr>
<tr>
<td>Edition:</td>
<td>not in use</td>
</tr>
<tr>
<td>Date Submitted by the Author:</td>
<td>28-Mar-2017</td>
</tr>
<tr>
<td>Complete List of Authors:</td>
<td>Matthews, Debbie; Royal Victoria Infirmary, Childrens BATH, LOUISE; Royal Hospital for Sick Children, 9 Scienies Road Hogler, Wolfgang; Birmingham Childrens Hospital, Endocrinology Mason, Avril; Royal Hospital for Children, Paediatric Endocrinology Smyth, Arlene; Turner Syndrome Support Society UK Skae, Mars; Royal Manchester Children's Hospital, Paediatric Endocrinology</td>
</tr>
<tr>
<td>Keywords:</td>
<td>Puberty, Oestrogen, Progestogen</td>
</tr>
</tbody>
</table>
Title:

Hormone Supplementation For Pubertal Induction In Girls

Authors: Debbie Matthews\textsuperscript{1}, Louise Bath\textsuperscript{2}, Wolfgang Högler\textsuperscript{3,4}, Avril Mason\textsuperscript{5}, Arlene Smyth\textsuperscript{6}, Mars Skae\textsuperscript{7,8}

Affiliations:

1 Department of Child Health, Newcastle upon Tyne Hospitals NHS Foundation Trust, Newcastle upon Tyne, UK

2 Department of Endocrinology and Diabetes, Royal Hospital for Sick Children, Edinburgh

3 Department of Endocrinology and Diabetes, Birmingham Children’s Hospital, Birmingham, UK

4 Institute of Metabolism and Systems Research, University of Birmingham, Birmingham, UK

5 Developmental Endocrinology Research Group, Royal Hospital for Children, Glasgow, UK

6 Executive Officer Turner Syndrome Support Society, Clydebank Business Park, Glasgow, UK

7 University of Manchester, Manchester, UK
8 Department of Paediatric Endocrinology, Royal Manchester Children's Hospital, Manchester, UK

Corresponding Author:-

Debbie S. F. Matthews

Consultant Paediatric Endocrinologist

Department of Child Health

Newcastle upon Tyne Hospitals NHS Foundation Trust

Queen Victoria Road

Newcastle upon Tyne NE1 4LP, UK.

Debbie.matthews@nuth.nhs.uk

01912825321

Word count 3,356
SUMMARY

Pubertal induction in girls with ovarian insufficiency aims to mimic normal puberty, a highly complex process. Here we amalgamate the sparse global evidence and propose three options for pubertal induction regimens including oral ethinyloestradiol, and oral and transdermal 17β-oestradiol. The introduction of progestogens is discussed and the transition to hormone supplementation for adult women. The merits and disadvantages of the different options are detailed. The available evidence indicates that transdermal 17β-oestradiol has the most favourable efficacy, safety, and cost profile but randomised controlled trials are urgently required to determine which regimen provides the best clinical outcomes.

INTRODUCTION

Girls with primary or secondary ovarian insufficiency require supplementation with oestrogen to induce puberty. However, there are no licensed hormone preparations for children, resulting in off-label prescribing of formulations licensed for adults. Traditionally, most clinicians in the UK have used low dose synthetic ethinyloestradiol with variable clinical outcomes.[1] More recently, the escalating cost, poor availability and unfavourable outcomes associated with low dose ethinyloestradiol has necessitated the use of alternatives. In continental Europe and USA, there is greater experience of using natural transdermal or oral 17β-oestradiol.[2] Following critical review of the available literature and extensive consultation by a working group of the British Paediatric Endocrine Society (BSPED), this guidance document for pubertal induction has been produced. The document provides the full range of therapeutic options including oral and transdermal 17β-oestradiol and oral ethinyloestradiol.
Pubertal induction aims to achieve normal tempo and magnitude of breast and uterine development and adolescent growth spurt by mimicking the natural pubertal process. These outcomes are achieved with low starting doses of oestrogen which are then gradually increased, whilst monitoring the clinical response in linear growth and breast staging. High dose oestrogen early in puberty or rapid dose escalation may result in reduced final height and poor breast development, such as prominent nipple development with poor supporting breast tissue. There are also concerns that unphysiological supplementation may adversely affect uterine development and bone mass accrual.[3]

Girls are known to prefer progress in tandem with their peers. Therefore, the recommended age for commencing pubertal induction is around 11-12 years, although some girls may actually present much later.

The aims of pubertal induction are to:-

- Allow the natural development of secondary sexual characteristics (particularly breast shape and size).
- Allow normal uterine growth to an adult size and shape.
- Achieve a good adolescent growth spurt.
- Achieve normal peak bone mass.
- Support psychological maturation and adjustment.

LITERATURE REVIEW

There is a real paucity of carefully constructed, randomised controlled clinical trials in girls undergoing induction of puberty. The evidence base is derived mainly from
expert experience, a small number of observational studies and very few controlled trials on small study populations. In addition, studies of treatment acceptability and patient adherence are lacking.

**Transdermal/Oral 17β-oestradiol for Induction of Puberty**

17β-oestradiol is the most physiological form of oestrogen, being identical to ovarian secreted oestrogen and measurable in serum. Whilst oral 17β-oestradiol is metabolized in the liver to the weaker oestrogen oestrone, transdermal 17β-oestradiol does not undergo this first-pass effect. The available evidence on the use of 17β-oestradiol is sparse on all clinical outcomes mentioned above.

**Transdermal 17β-oestradiol**

Three regimens using transdermal 17β-oestradiol were reviewed. In a carefully monitored observational study, Ankarberg-Lindgren et al., used a low dose transdermal 17β-oestradiol regimen in 15 girls with primary ovarian insufficiency (POI) which resulted in pubertal development to breast stage 2-3 (B2-3) over a period of 3.5-29 months (median 10 months).[4] However, this dosing regimen was based on body weight with complicated cutting of patches into small fractions and did not extend beyond early to mid-puberty. Davenport provided a regimen for pubertal induction in girls with Turner syndrome basing transdermal 17β-oestradiol dose on body weight in early puberty, and adjusting patch size to target serum oestradiol levels.[5] Nabhan et al. used much higher doses of transdermal 17β-oestradiol starting with 25µg/24hrs.[6]
There is very limited but encouraging evidence on breast development [7], uterine growth [8, 9] and bone accrual [10] in females treated with transdermal 17β-oestradiol. Because of its mode of administration, transdermal 17β-oestradiol does not lower IGF-1 levels since there is no effect on hepatic metabolism.[11] Therefore, it might enhance linear growth although this has not been studied to date.[8]

Oral 17β-oestradiol

Four regimens using oral 17β-oestradiol for pubertal induction were reviewed. Zacharin used a starting dose of 0.5mg 17β-oestradiol every second day, increasing doses over 2 years to an adult dose of 2mgs daily.[12] Delemarre et al.[13] and Bannink et al.[14] based 17β-oestradiol dose on body weight, starting with 5µg/kg/day and increasing to an adult dose over 2 and 3 years respectively. Labarta et al. used 0.2mg 17β-oestradiol daily for one year, followed by 0.5mgs daily for the second year.[15]

In terms of clinical outcomes, Bannink et al. treated 56 girls with Turner syndrome with incremental oral 17β-oestradiol and described normal breast development up to B4-5 in 49 girls (87%).[14] Labarta et al. treated 48 girls with Turner syndrome over 2 years using individualised or fixed dose 17β-oestradiol and described breast development to B4 in 42% and 65% of girls respectively.[15] The results from studies looking at uterine growth are variable.[14, 16, 17] Torres-Santiago et al. found significant but similar improvements in whole-body and lumbar BMD Z-scores over 12 months in 40 girls with Turner syndrome randomised to either oral or transdermal 17β-oestradiol.[18]
Dose titration against serum oestradiol levels

An ultrasensitive assay may be used to monitor serum 17β-oestradiol concentrations in early puberty for both transdermal and oral 17β-oestradiol induction regimens. These results may assist in adjusting doses of oestradiol aiming for serum concentrations in the early pubertal range (10-40pmols/L).\[4\] Ankarberg-Lindgren et al. showed that standard doses of transdermal oestradiol based on body weight resulted in considerable inter-individual variation in serum 17β-oestradiol concentrations highlighting the clinical value of using serum levels to guide dosing regimens.\[19\] Conversely, the conclusion from Bannink’s low-dose oral 17β-oestradiol study was that serum oestradiol concentrations do not provide additional information on the progression through puberty.\[14\] For those girls completing puberty, a pharmacokinetic and pharmacodynamic study of oral and transdermal 17β-oestradiol in girls with Turner syndrome suggested an adult target 17β-oestradiol concentration of 350pmol/l, as derived from healthy menstruating adult women using integrated mean levels over the natural cycle.\[20\]

Oral Ethinyloestradiol for Induction of Puberty

Three published regimens for pubertal induction using ethinyloestradiol were reviewed.\[13, 21, 22\] These regimens share a gradual increase in ethinyloestradiol dose, either from a starting dose of 2µg daily \[22\] or 0.1 µg/kg/d \[13, 21\] followed by the addition of a progestogen after 2-2.5 years of unopposed oestrogen.

Ethinyloestradiol is synthetic, cannot be measured in serum, and undergoes first-pass metabolism in the liver. Similar to 17β-oestradiol, published data regarding the clinical efficacy of oral ethinyloestradiol in pubertal induction are very limited.
Suboptimal breast development is reported in girls treated with ethinyloestradiol and oestradiol valerate.[23, 24] However, it is unclear whether this is secondary to problems with the formulation, dose effect, rate of dose escalation or timing of start of therapy.[23] One study of 38 girls treated with incremental oral ethinyloestradiol showed that only 50% developed mature, heart-shaped uterine configurations.[24] Similarly, another study suggested that replacement therapy with ethinyloestradiol gave rise to poor uterine growth and development.[3]

Few studies assessed the effect of ethinyloestradiol on bone mass accrual and there are significant pitfalls in the interpretation of dual energy X-ray absorptiometry results of children and adults with short stature such as in Turner syndrome.[25] A randomised crossover trial in 18 young women with POI showed no significant change in lumbar spine BMD z-score following ethinyloestradiol treatment, raising concerns that this agent may not be effective in increasing bone mass.[10] Another randomised crossover study in 17 young women with Turner Syndrome demonstrated that ethinyloestradiol treatment was associated with high urinary deoxypyridinoline cross-link concentrations, suggesting an unfavourable effect on bone turnover.[26]

Oral ethinyloestradiol is associated with lower IGF-1 concentrations due to its first pass hepatic effect although no studies have compared growth rates and final height using different female hormone replacement strategies. A small synergistic effect on final height was found between low dose childhood ethinyloestradiol and growth hormone treatment in girls with Turner syndrome.[21]

Introduction of Progestogens during Late Puberty
Much of the literature considers the role of oestrogen in pubertal induction with little discussion about the regimen, timing and choice of progestogens.

Progestogens are usually introduced after a suitable duration of unopposed oestrogen (2-3 years) or if more than one episode of significant breakthrough bleeding occurs. Progestogens are usually given in 12-14 day blocks, each inducing withdrawal bleeding. The frequency of blocks may be adjusted according to the patient’s wishes, but at least every 2-3 months, which avoids endometrial hypertrophy. Introducing a progestogen too soon, especially using one of the more androgenic agents such as norethisterone, may potentially compromise uterine growth and development.[3]

Options for treatment include oral Utrogestan (200mg once daily) or oral Medroxyprogesterone acetate (5mg once daily). Utrogestan is a natural micronized progesterone which can be given orally and gives good cycle control without significant side effects. Medroxyprogesterone acetate is a synthetic derivative of 17α-hydroxyprogesterone and is less androgenic than derivatives of 19-nortestosterone such as norethisterone.[27]

**Oestrogen and Progestogen Replacement Therapy in Adult Women**

Adolescent girls undergoing pubertal induction require adult regimens at the end of puberty. Similar to children and adolescents, no product is designed for long term use in women with POI. Options include oral 17β-oestradiol, transdermal 17β-oestradiol, the combined oral contraceptive pill (COCP) and equine conjugated oestrogens (popular in the USA). Typical daily adult regimens for these preparations are oral 17β-oestradiol 2mg daily, transdermal 17β-oestradiol 50-100µg patches
applied twice weekly and left in place until replaced, and COCP daily containing 20-30µg of ethinyloestradiol.

In adult women, progestogen may be given cyclically or continuously depending on whether women wish to experience withdrawal bleeds. Oral progestogens are available either as single agents (e.g. Provera®) or in user-friendly combined packs (e.g. Elleste-Duet®, Elleste-Duet Conti®) or as the COCP (e.g. Microgynon®, Marvelon®). Combined transdermal patches with 17β-oestradiol and progestogen are available, e.g. Evorel Sequi® & Evorel Conti®. However, breakthrough bleeding may be more common in young women using transdermal progestogens.

Unfavourable cardiovascular risk of ethinyloestradiol/COCP

The COCP is cheap (free via the NHS in the UK) and readily available. However, if taken as prescribed on a monthly cycle, women with POI lack oestrogen supplementation 1 in 4 weeks. In addition, the COCP has an adverse cardiovascular and metabolic profile. Specifically, the addition of the ethinyl side chain in ethinyloestradiol induces renin substrate at a much greater rate than natural products and increases the risk of hypertension, particularly in susceptible groups such as women with Turner syndrome.[28] Use of the COCP is also linked with an increased risk of venous thromboembolism.[29] Metabolic studies in adults suggest that ethinyloestradiol treatment gives rise to increased SHBG, decreased IGF-1 and increased insulin resistance.[11] In addition, C-reactive protein and other acute phase reactants may increase which independently predict cardiovascular disease.[30]
Studies comparing transdermal 17β-oestradiol with the COCP demonstrate reduced blood pressure, better renal function and less activation of the renin-angiotensin system in women using 17β-oestradiol.[31]

Women with POI are a heterogeneous group with different risk profiles. Women with Turner syndrome seem to have a reduced risk of breast cancer (relative risk 0.3) whereas women who have had whole body irradiation as conditioning for bone marrow transplantation have an increased risk (relative risk 6.5).[32, 33] Both groups of women are at greater risk of hypertension and Type 2 diabetes than their normal peers and their choice of oestrogen replacement therapy needs to minimise these risks. Women who have had cranial irradiation for brain tumours have an increased risk of stroke and treatment with transdermal 17β-oestradiol is preferred.[34]

Summary of Literature Review

In summary, pubertal induction using oral or transdermal 17β-oestradiol is well described but there is some concern regarding appropriate starting doses and individual variation in response. The transdermal route of administration leads to lower peak serum 17β-oestradiol concentrations, lower hepatic metabolism and more stable steady state profiles compared with the oral route.[20] However, transdermal products may be less acceptable to patients. Possibly, the choice of oestrogen matters little at the start of pubertal induction where low dosage is of great importance. While there is good evidence that puberty can be induced too quickly leading to reduced final height, the question whether it can be induced too slowly cannot be answered with confidence.[35] Review of the current literature suggests that for long-term hormone replacement, oral/transdermal 17β-oestradiol needs to be
favoured over preparations containing ethinyloestradiol, given their higher cardiovascular risk.

Ultimately, the most important requirement is that girls with ovarian insufficiency are treated with oestrogen in a timely manner and that treatment is continued through to natural menopausal age. Oestradiol deficiency causes cancellous bone loss, endothelial dysfunction, reduced insulin production, abnormal lipid patterns, increased central adiposity and early atheroma. Hence, it is concerning that at a large UK adult Turner clinic, 24% of patients were not receiving oestrogen treatment at all at their first clinic attendance, highlighting the importance of treatment acceptability, adherence and patient education.

Finally, there is very little information on specific dose response (breast and uterine growth and shape, bone accrual, growth) to oral or transdermal 17β-oestradiol and oral ethinyloestradiol, and the bioequivalency of preparations. Estimated daily dose equivalence from the current literature (depending on assays and clinical endpoints) are: 50/100µg transdermal (applied twice weekly until replaced) = 2mg oral 17β-oestradiol (per day) = 20µg ethinyloestradiol (per day).

**PROPOSED REGIMENS FOR PUBERTAL INDUCTION**

Pubertal induction should be individualised taking the girl’s and family’s views into consideration as well as parameters such as height, age, pubertal stage and co-morbidities. The optimal oestrogen treatment comprising route, drug, and dose increments should be determined for each girl. Amongst the paediatric endocrine community, there is general agreement that oestrogen starting doses for pubertal

https://mc.manuscriptcentral.com/adc
induction should be about 10% of adult replacement doses. Following extensive literature review, consultation with external experts and the UK Turner Syndrome support group, we present below optimised regimens for pubertal induction in girls with ovarian insufficiency. All induction regimens will take girls into later pubertal stages (Tanner 3-5) over 2.5 years, following which adult hormone replacement options should be used.

**Transdermal 17β-oestradiol**

The published regimens for pubertal induction using transdermal 17β-oestradiol were considered impractical to administer and implement.[4-6] Therefore, a pragmatic approach to the transdermal regimen was taken, ensuring the use of low doses of 17β-oestradiol, particularly in early puberty.[12] (personal communication with M Zacharin and T Randell, 2016).

Regimen using 25µg 17β-oestradiol matrix patch ([Table 1](#))

Matrix patches are self-adhesive and release approximately 25µg 17β-oestradiol /24 hours. Since the oestradiol is evenly distributed throughout the patch, the patches can be cut to provide the required dose. Practically, patches are cut into ½ or ¼ as more complex divisions would be prone to inaccuracies and impracticable. Unused patch fractions may be stored in their packaging in the fridge for up to one week. The patch (or patch fraction) should be applied to clean dry skin over the buttocks or hips using Opsite® (a transparent adhesive film) if necessary to ensure good adhesion.
Table 1 Regimen for pubertal induction using 25µg/24h 17β-oestradiol matrix patch applied once or twice weekly and left in situ for 3-4 days

<table>
<thead>
<tr>
<th>Monday to Thursday</th>
<th>Friday to Sunday</th>
<th>Duration (months)</th>
</tr>
</thead>
<tbody>
<tr>
<td>¼ patch</td>
<td>No patch</td>
<td>6</td>
</tr>
<tr>
<td>¼ patch</td>
<td>¼ patch</td>
<td>6</td>
</tr>
<tr>
<td>½ patch</td>
<td>¼ patch</td>
<td>6</td>
</tr>
<tr>
<td>½ patch</td>
<td>½ patch</td>
<td>6</td>
</tr>
<tr>
<td>1 patch</td>
<td>1 patch</td>
<td>6</td>
</tr>
</tbody>
</table>

Progress to adult oestrogen / progestogen replacement therapy

Oral 17β-oestradiol

Whilst oral 17β-oestradiol is indeed used globally for pubertal induction in various preparations, there is a lack of published evidence. Here, we adopt a modification of the regimens published by Labarta and Zacharin.[12, 15] 17β-oestradiol is only commercially available in 1mg tablets and this regimen involves breaking the 1mg tablets (Table 2).

Table 2 Regimen for pubertal induction using 1mg 17β-oestradiol tablets

<table>
<thead>
<tr>
<th>Dose</th>
<th>Tablets</th>
<th>Frequency</th>
<th>Equivalent daily dose</th>
<th>Duration (months)</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.5mg</td>
<td>½</td>
<td>Alternate days</td>
<td>0.25mg</td>
<td>12</td>
</tr>
<tr>
<td>0.5mg</td>
<td>½</td>
<td>Daily</td>
<td>0.5mg</td>
<td>6</td>
</tr>
<tr>
<td>0.5mg/1mg</td>
<td>½, 1</td>
<td>Alternate days</td>
<td>0.75mg</td>
<td>6</td>
</tr>
<tr>
<td>1mg</td>
<td>1</td>
<td>Daily</td>
<td>1mg</td>
<td>6</td>
</tr>
</tbody>
</table>

Progress to adult oestrogen / progestogen replacement therapy

Girls and young women taking natural 17β-oestradiol for pubertal induction may have serum oestradiol levels measured to monitor therapy. Ideally, serum oestradiol
levels should be maintained <50pmols/L during the first 18-24 months of pubertal induction to accelerate linear growth without rapidly advancing bone maturation.[5] However, serum oestradiol levels <60pmols/L are not measurable by most clinical laboratory methods. If a girl seems to be making either too slow or too rapid progress through puberty, ultra-sensitive oestradiol assays should be employed. Such assays are based on liquid chromatography tandem mass spectrometry with limits of detection of 4-8pmols/L.[38]

**Oral Ethinyloestradiol**

This regimen is derived from Hindmarsh.[22] (Table 3) Ethinyloestradiol is not easily measured in serum.

### Table 3 Regimen for pubertal induction using 2µg ethinyloestradiol tablets

<table>
<thead>
<tr>
<th>Dose (µg)</th>
<th>Tablets</th>
<th>Frequency</th>
<th>Duration (months)</th>
</tr>
</thead>
<tbody>
<tr>
<td>2</td>
<td>1</td>
<td>Daily</td>
<td>6</td>
</tr>
<tr>
<td>4</td>
<td>2</td>
<td>Daily</td>
<td>6</td>
</tr>
<tr>
<td>6</td>
<td>3</td>
<td>Daily</td>
<td>6</td>
</tr>
<tr>
<td>8</td>
<td>4</td>
<td>Daily</td>
<td>6</td>
</tr>
<tr>
<td>10</td>
<td>5 x 2µg or 1 x 10µg</td>
<td>Daily</td>
<td>6</td>
</tr>
</tbody>
</table>

Progress to **adult oestrogen / progestogen replacement therapy**

**Table 4** provides a summary of the suggested oestrogen replacement regimens for pubertal induction
Table 4 Summary of the suggested oestrogen regimens

<table>
<thead>
<tr>
<th>Months from start of induction</th>
<th>25µg 17β-oestradiol matrix patch (e.g. Evorel® 25)</th>
<th>17β-oestradiol (Oestradiol valerate) 1mg tablets</th>
<th>Ethinyloestradiol 2 µg tablets</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>¼ patch for 3-4 days, no patch 3-4 days</td>
<td>0.5mg (½ tablet) alternate days</td>
<td>2µg (1 tablet) daily</td>
</tr>
<tr>
<td>6</td>
<td>¼ patch all week (change every 3-4 days)</td>
<td>0.5mg (½ tablet) alternate days</td>
<td>4µg (2 tablets) daily</td>
</tr>
<tr>
<td>12</td>
<td>¼ patch for 3-4 days, ½ patch for 3-4 days</td>
<td>0.5mg (½ tablet) daily</td>
<td>6µg (3 tablets) daily</td>
</tr>
<tr>
<td>18</td>
<td>½ patch all week (change every 3-4 days)</td>
<td>0.5mg and 1mg alternate days</td>
<td>8µg (4 tablets) daily</td>
</tr>
<tr>
<td>24*</td>
<td>1 patch all week (change every 3-4 days)</td>
<td>1mg (1 tablet) daily</td>
<td>10µg (5 tablets) daily</td>
</tr>
<tr>
<td>30*</td>
<td>Adult COCP or HRT</td>
<td>Adult COCP or HRT</td>
<td>Adult COCP or HRT</td>
</tr>
</tbody>
</table>

* Progestogens should be introduced only after a suitable duration of unopposed oestrogen (usually 2-3 years) or if more than one episode of significant breakthrough bleeding occurs.

Clinical monitoring of progress for all pubertal induction regimens

To ensure safety and efficacy of the suggested approaches, the following clinical data should be collected (Table 5):
Table 5 Clinical monitoring of progress for all pubertal induction regimens

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Pre-Puberty</th>
<th>During puberty (every 6/12)</th>
<th>Post puberty</th>
</tr>
</thead>
<tbody>
<tr>
<td>Blood Pressure</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes (every 6/12)</td>
</tr>
<tr>
<td>Height velocity (HV)</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes (until HV&lt;2 cms/year)</td>
</tr>
<tr>
<td>Pubertal staging</td>
<td>Yes</td>
<td>Yes</td>
<td>No</td>
</tr>
<tr>
<td>Pelvic USS</td>
<td>Yes</td>
<td>No</td>
<td>Yes. Document uterine size and shape</td>
</tr>
<tr>
<td>Bone age</td>
<td>Yes</td>
<td>Yes (annually)</td>
<td>No</td>
</tr>
<tr>
<td>Bone density scan</td>
<td>No</td>
<td>No</td>
<td>One year post menarche; in case of low size-corrected BMD or non-compliance measure again in 3-5 years</td>
</tr>
</tbody>
</table>

PROGRESSION TO ADULT HORMONE REPLACEMENT THERAPY

A progestogen will be introduced for all patients for 12-14 days every 1-3 months at breakthrough bleeding or after 2.5 years of treatment with oestrogen. The preferred progestogen is utrogestan 200mg once daily. Alternatively, medroxyprogesterone acetate 5mg daily may be used. Norethisterone 5mg daily is available but is more androgenic than the other preparations and is linked to a higher incidence of dysmenorrhoea.

Once a dose of transdermal 17β-oestradiol 25µg/24h, oral 17β-oestradiol 1mg or oral ethinyloestradiol 10µg is reached and the girls are receiving a cyclical progestogen, there are further options for their longer term management as young women.
Transdermal / oral 17β-oestradiol

Transdermal 17β-oestradiol may be continued as a matrix patch or an oestrogen gel (e.g. Sandrena®). Adult doses of transdermal oestradiol via patch vary between 50-100µg/24 hours and adult doses of gel vary between 0.5-1mg oestradiol daily. Some young women may prefer oral medication and there are a number of different proprietary preparations which provide 1-2mg of oral 17β-oestradiol daily according to requirement (e.g. Elleste Solo®).

Many preparations are produced in user-friendly packs with patches/tablets containing oestrogen alone, followed by patches/tablets containing both oestrogen & progestogen combined (e.g. Evorel Sequi® [patch], Elleste Duet® [oral]). Many oral preparations contain norethisterone as the progestogen but in low doses of 0.5-1.0mg. Similarly, the dose of medroxyprogesterone acetate is low in these preparations (1-2mgs).

Women wishing to avoid withdrawal bleeds may be given continuous combined preparations, either transdermal patches (e.g. Evorel Conti®) or oral tablets (e.g. Elleste Duet Conti®). However, women with any residual ovarian function may experience troublesome breakthrough bleeding on these preparations.

In young adult women standard laboratory oestradiol assays are used to monitor serum oestradiol levels, aiming for a target 17β-oestradiol level of 350pmol/l.[20]

Progestogen may also be provided using a levonorgestrel-releasing intrauterine device (e.g. Mirena® coil). It is important to ensure that the uterus is of adult dimensions (about 7.5 x 5 x 2.5cms) by ultrasound before use and girls who are not sexually active may require a brief general anaesthetic for insertion.[39]
Women with any potential residual ovarian function in whom pregnancy is not desired should be counselled about the need for additional contraception if using these preparations.

**Combined Oral Contraceptive Pill (COCP)**

Advice on the use of the COCP for adult replacement therapy should be guided by a risk assessment as set out by the Faculty of Sexual & Reproductive Healthcare.[40] The contained oestrogen is usually ethinyloestradiol (e.g. Microgynon30®, Marvelon®) although 17β-oestradiol is used occasionally (e.g. Qlaira®). In order to maximise oestrogen exposure, girls are advised to take at least 3 packs of pills “back-to-back” to reduce “oestrogen-free” weeks. This has the additional benefit of reducing the frequency of withdrawal bleeds. Preparations can also be taken continuously to avoid withdrawal bleeding although initially there may be some breakthrough bleeding until the endometrial lining is atrophied.

The full scope of oestrogen/progestogen replacement for adult women is beyond the remit of this guideline. It is anticipated that young women with ovarian insufficiency will be reviewed in a Transition clinic alongside an adult Gynaecologist or an adult Endocrinologist and kept under review throughout adult life.

**CONCLUSION AND RESEARCH NEEDS**

The induction regimens proposed here are based on synthesis of the literature, expert views, consultation, pragmatism and practicability. The evidence suggests that transdermal 17β-oestradiol has the most favourable efficacy, safety and cost
profile. Therefore, this BSPED working group recommends it as first choice for
pubertal induction in girls.

Nevertheless, how the essential clinical outcomes (breast and uterine size and
shape, final height, bone mass, safety and acceptance) compare between regimens
has not been well studied. The lack of randomised controlled studies on pubertal
induction in girls needs to be addressed. It is anticipated that prospective collection
of data from these proposed regimens will provide valuable information about their
efficacy and acceptability.
Pros and Cons of oral/transdermal 17β-oestradiol versus oral ethinyloestradiol for pubertal induction in girls

Pros for oral / transdermal 17β-oestradiol

- 17β-oestradiol is more physiological than synthetic ethinyloestradiol especially when administered transdermally since the first pass hepatic effect is abolished.
  - Observational studies suggest that oral or transdermal 17β-oestradiol is effective at inducing puberty. Treatment using transdermal 17β-oestradiol can be individualised and can mimic normal puberty closely.
  - Oral 17β-oestradiol tablets and transdermal matrix patches are readily available, cheap and have got a favourable cardiovascular risk profile compared to ethinyloestradiol.

Cons for oral / transdermal 17β-oestradiol

- Transdermal patches may be more difficult to use particularly when cutting patches to small sizes as they may fall off and require tape support.
- Transdermal patches may be less acceptable to girls undergoing pubertal induction, particularly if the patch becomes visible or they have a reaction to the adhesive.
- There is some suggestion of inter-individual variation in response to oral 17β-oestradiol tablets and transdermal patches.

Pros for oral ethinyloestradiol

- Oral ethinyloestradiol has been used extensively for pubertal induction, particularly in the UK & USA.
- The tablet preparations are acceptable and easy to take.

- Millions of women worldwide use ethinyloestradiol in the form of the COCP which has a good safety profile.

Cons for oral ethinyloestradiol

- In recent times, low dose ethinyloestradiol tablets (2 and 10 µg) have escalated in cost significantly. They are no longer always readily available.
- Although effective at inducing puberty, the outcomes may be suboptimal and more physiological agents such as 17β-oestradiol may be preferable.
- Oral ethinyloestradiol is associated with an increased risk of hypertension and venous thromboembolism in adults and this risk may be present also in children.
ACKNOWLEDGEMENTS

The authors would like to thank Professor Margaret Zacharin for her helpful advice and Dr Tabitha Randell for sharing her personal experience. We acknowledge the BSPED Clinical Practice Committee for their valuable role as a consultation body.

REFERENCES


https://mc.manuscriptcentral.com/adc


40. Faculty of Sexual & Reproductive Healthcare of the Royal College of Obstetricians & Gynaecologists. UK Medical Eligibility Criteria for Contraceptive Use. 2016.