Mitotane (op’DDD) restores growth and puberty in nine children with Cushing’s disease

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Abstract

To investigate whether low-dose mitotane (up to 2 g/day) could be a temporary therapeutic alternative to transsphenoidal surgery (TSS) in pediatric Cushing’s disease (CD). Twenty-eight patients with CD aged 12.2 years (± 2.2) were referred to our center. We compared nine patients treated with mitotane alone for at least 6 months to 13 patients cured after surgery. Primary outcomes were changes in growth velocity, BMI and pubertal development. The following results were obtained: (1) Mitotane improved growth velocity z-scores (−3.8 (±0.3) vs −0.2 (±0.6)), BMI z-scores (2.1 (±0.5) vs 1.2 (±0.5) s.d.) and pubertal development. After 1 year on mitotane, the mean BMI z-score was not significantly different in both groups of patients. (2) Control of cortisol secretion was delayed and inconsistent with mitotane used as monotherapy. (3) Side effects were similar to those previously reported, reversible and dose dependent: unspecific digestive symptoms, concentration or memory problems, physical exhaustion, adrenal insufficiency and hepatitis. (4) In one patient, progressive growth of a pituitary adenoma was observed over 40 months of mitotane treatment, allowing selective adenomectomy by TSS. In conclusions, low-dose mitotane can restore growth velocity and pubertal development and decrease BMI in children with CD, even without optimal control of cortisol secretion. It may promote pituitary tumor growth thus facilitating second-line TSS. However, given its possibly life-threatening side effects (transient adrenal insufficiency and hepatitis), and in the absence of any reliable follow-up procedures, this therapy may be difficult to manage and should always be initiated and monitored by specialized teams.

Key Words
- Cushing’s disease
- children
- growth
- puberty
- op’ddd
- mitotane

Introduction

Cushing’s disease (CD) is characterized by adrenocorticotropic (ACTH)-dependent cortisol excess originating from a pituitary adenoma and accounts for approximately 85% of cases of pediatric Cushing’s syndrome (1, 2, 3, 4). The resulting hypercortisolism causes considerable morbidity in childhood and adolescence, predominantly affecting growth and pubertal development (1, 4, 5, 6, 7, 8). The diagnosis is often delayed by at least 2 years after the first symptoms, likely because of the lack of obvious signs such as buffalo hump, facial erythrosis, myopathy or metabolic syndrome. Metabolic complications are rarely reported. Further, although the diagnosis of CD is usually easy to make once it has been suspected (9), the treatment remains a challenge. The gold

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standard treatment for both pediatric and adult patients with CD is transphenoidal surgery (TSS) with selective microadenomectomy, but the failure rate is about 25–50% (10, 11, 12, 13, 14, 15, 16), prompting physicians to use alternative therapies such as pituitary radiotherapy or adrenolytic agents.

O,p′-dichlorodiphenyldichloroethane (mitotane, Lysodren 500mg; HRA Pharma, Paris, France) is an adrenolytic drug with a direct cytotoxic effect on the zona reticularis of adrenal glands (17, 18, 19). Its therapeutic use has been validated in adults with metastatic adrenocortical carcinoma, with a target concentration of 14–20mg/L. At lower concentrations, mitotane reduces cortisol levels both through inhibition of steroidogenesis without cell destruction (inhibiting cholesterol side-chain cleavage enzyme (20, 21) and 11-beta-hydroxylase (22)) and through enhanced cortisol clearance by induction of CYP3A4 activity (23, 24). About 40% of mitotane is absorbed through the intestine, and then binds to lipophilic proteins and is metabolized in the liver. It accumulates in adipose tissue from where it can be released for weeks to months (24, 25), making it difficult to adjust the dosage correctly. Mitotane is associated with various side effects that are always dose-dependent and are related to its pharmacodynamic properties in some cases (modification of lipid profile, hepatic cytolysis, cholestasis, adrenal insufficiency) but not in others (neurological disorders, leucopenia, nausea and vomiting).

Currently, mitotane is relatively rarely used for the treatment of CD. When used in adults with CD, it has tended to be administered as first-line therapy alone (>6g/day) (26, 27), in combination with other adrenolytic agents (28) or as second-line therapy after failed surgery (29, 30, 31), achieving suppressed cortisol secretion (Table 1). Although reviews and textbooks mention mitotane as an adjuvant therapy for CD in children, its efficacy and safety in this context has not been reported so far. We describe here the efficacy and safety of mitotane as treatment of CD in nine children in comparison to 13 patients cured by TSS.

### Patients and methods

#### Patients

Between 1978 and 2014, 28 patients with CD (15F/13M), aged 12.2 (±2.2) (7–16.2) years, were referred to our center. CD was diagnosed when hypercortisolism (defined as midnight plasma cortisol >4.4µg/dL and/or urinary-free cortisol (UFC) >70µg/m²/24h, salivary cortisol was not routinely used in our center during the study period to define hypercortisolism (9)) was present in combination with elevated or non-suppressed ACTH levels (>20pg/mL), and either an adenoma on pituitary imaging or an increase in ACTH and cortisol levels after synthetic corticotropin-releasing hormone or vasopressin infusion. Bilateral inferior sinus sampling was not performed in most patients.

We retrospectively collected data on clinical and biological parameters describing the efficacy and safety of treatments (surgery, radiotherapy, adrenolytic drugs). According to the Jardé law in France, the French National Data Processing and Liberties Commission (CNIL) approved the study. Consent has been obtained from each patient after full explanation of the purpose and nature of all procedures used. All patients but one (patient #28, whose case is described in detail below) were considered to be cured or in sustained remission. A total of nine out of the 28 patients were treated with mitotane alone for at least 6 months before the end of their growth period. Mitotane was initiated with a 1g/day dose and titrated to achieve normalization of UFC levels and the fewest possible side effects (serum mitotane levels, UFC levels and liver function tests were followed up after 1 month of therapy, then every 3–6 months). We excluded from the analysis the patients who received mitotane either after the age of 16 years or for less than 6 months (Fig. 1). A total of 13 patients, who never received mitotane, were considered to be cured (absence of hypercortisolism, defined as disappearance of clinical signs of cortisol excess and midnight plasma cortisol ≤4.4µg/dL and/or UFC<70µg/m²/24h, 5 years after surgery) by TSS and

### Table 1  Synopsis of publications mentioning op’DDD in CD.

<table>
<thead>
<tr>
<th>Author</th>
<th>Year</th>
<th>N</th>
<th>Op’DDD dosage (g/day)</th>
<th>Administration</th>
<th>Duration</th>
<th>% of patients with UFC &lt; 100 µg/24 h</th>
</tr>
</thead>
<tbody>
<tr>
<td>Luton (27)</td>
<td>1978</td>
<td>46</td>
<td>6.00</td>
<td>Continuous</td>
<td>8 months</td>
<td>83</td>
</tr>
<tr>
<td>Schteingart (29)</td>
<td>1980</td>
<td>36</td>
<td>Not reported</td>
<td>Continuous: 19</td>
<td>Not reported</td>
<td>81</td>
</tr>
<tr>
<td>Benecke (30)</td>
<td>1991</td>
<td>2</td>
<td>0.5–2</td>
<td>Continuous</td>
<td>5 and 8 years</td>
<td>100</td>
</tr>
<tr>
<td>Kawai (26)</td>
<td>1999</td>
<td>1</td>
<td>1–4</td>
<td>Discontinuous</td>
<td>18 years</td>
<td>100</td>
</tr>
<tr>
<td>Baudry (31)</td>
<td>2012</td>
<td>76</td>
<td>4–6</td>
<td>Continuous</td>
<td>6 months–15 years</td>
<td>100</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>100 when mitotane &gt; 8 mg/L</td>
<td></td>
</tr>
</tbody>
</table>

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were used as the comparison group. We assessed the efficacy of the mitotane therapy in terms of changes in growth velocity (GV) z-scores, BMI z-scores, pubertal development and biological control of cortisol secretion (24 h UFC).

**Statistical analysis**

GraphPad Prism 4 software was used for the statistical analysis. Data are expressed as mean (±s.d.). Means were compared between groups with non-parametric t-tests.

**Results**

**Patients**

The two groups (mitotane treated, n=9; cured by TSS only, n=13) were comparable in age, BMI, height z-scores, GV z-scores, midnight plasma cortisol, UFC and ACTH at diagnosis. The characteristics of the patients are summarized in Table 2 and Supplementary Table 1 (see section on supplementary data given at the end of this article).

**Efficacy of mitotane**

**Comparison between baseline and after 1 year of mitotane therapy**

In patients who received mitotane as a treatment for their CD, we observed significant improvements in the GV z-score (−3.8 (±0.3) vs −0.2 (±0.6); P=0.0001) and BMI z-score (2.1 (±0.5) vs 1.2 (±0.5) s.d.; P=0.02) after 1 year of treatment (Fig. 2). The increase in GV allowed maintenance in height z-scores rather than catch-up growth (−0.9 (±0.4) after 1 year of mitotane vs −0.8 (±0.4); P=0.85). Final height data were missing for some patients.

**Table 2** Summary of characteristics of the 28 patients at diagnosis of CD.

<table>
<thead>
<tr>
<th></th>
<th>All 15F/13M</th>
<th>op’DDD group</th>
<th>TSS group</th>
<th>op’DDD vs TSS</th>
</tr>
</thead>
<tbody>
<tr>
<td>N</td>
<td>28</td>
<td>9</td>
<td>13</td>
<td></td>
</tr>
<tr>
<td>Age (year (s.d.))</td>
<td>12.1 (2.2)</td>
<td>12.0 (1.5)</td>
<td>12.0 (2.6)</td>
<td>NS</td>
</tr>
<tr>
<td>Height (z-score (s.d.))</td>
<td>−1.7 (1.6)</td>
<td>−0.8 (1.2)</td>
<td>−2.2 (1.4)</td>
<td>NS</td>
</tr>
<tr>
<td>Growth velocity (z-score (s.d.))</td>
<td>−4.0 (0.9)</td>
<td>−3.7 (0.9)</td>
<td>−4.1 (0.9)</td>
<td>NS</td>
</tr>
<tr>
<td>BMI (z-score (s.d.))</td>
<td>2.6 (2.2)</td>
<td>2.3 (1.4)</td>
<td>21 (1.4)</td>
<td>NS</td>
</tr>
<tr>
<td>MPC (µg/dL (s.d.)) N &lt; 3.5</td>
<td>18.9 (6.2)</td>
<td>16.8 (5.7)</td>
<td>17.6 (5.8)</td>
<td>NS</td>
</tr>
<tr>
<td>UFC (µg/24 h (s.d.)) N &lt; 60</td>
<td>285 (148)</td>
<td>263 (159)</td>
<td>268 (140)</td>
<td>NS</td>
</tr>
<tr>
<td>ACTH (pg/mL (s.d.)) N 10–60</td>
<td>48.5 (35.7)</td>
<td>68.8 (53.6)</td>
<td>64.2 (52)</td>
<td>NS</td>
</tr>
</tbody>
</table>

Mitotane group: nine patients who received more than 6 months of mitotane at initial presentation. TSS group: 13 patients cured by transsphenoidal surgery. BMI, body mass index; GV, growth velocity; MPC, midnight plasma cortisol; NS, means are not significantly different; UFC, urinary free cortisol.
In all cases, several weeks of mitotane treatment were necessary for the normalization or near-normalization of cortisol secretion. We found an inverse correlation between serum mitotane and UFC levels ($P=0.049$). Moreover, two thresholds may be highlighted: first, when mitotane levels were above 6 mg/L, UFC levels were <150 µg/24 h in all cases (17/17, 100%) and <80 µg/24 h in 9/17 cases (53%); and second, when mitotane levels were >10 mg/L, UFC levels were <80 µg/24 h in all cases (6/6, 100%). Figure 3 and Supplementary Table 2 illustrate the inter- and intra-patient variability in the control of cortisol secretion.

Comparison between mitotane treatment and successful TSS after 1 year
At 1 year, the mean BMI $z$-score was not statistically different in mitotane-treated patients to that observed after successful TSS (1.2 (±0.5) vs 0.8 (±0.4), respectively). The GV $z$-score, however, was higher 1 year after surgery than after 1 year on mitotane (1.6 (±0.4) vs −0.2 (±0.6), respectively; $P=0.006$).

Among the 17 patients who had reached the average age of onset of puberty (girls over 11 and boys over 12 years of age) at diagnosis, 15 presented with an abnormal pubertal development according to Tanner stages: either absent (6/15) or impaired (9/15). Of these, six received mitotane and resumed their pubertal development. In addition, two patients younger than the average age of puberty onset (11 and 12 years for girls and boys respectively) were treated with mitotane. They went through puberty during the treatment. One girl from this group had an uneventful pregnancy and gave birth to a healthy infant 3 years after the end of mitotane treatment. For one patient, we did not have any information about puberty.

Safety of mitotane
Various side effects were observed (see Table 3, detailing side effects documented in the seven most recent entries in patient medical records). These are similar to those previously described. All patients complained of digestive symptoms such as nausea and abdominal pain during the first weeks of treatment. Symptoms improved with the fractionation of the daily dose and the concomitant absorption of lipid-rich meals. The neurocognitive impact was not quantitatively assessed with standardized performance tests but parents and patients commonly reported asthenia (4/6) and difficulties in school (2/6) when mitotane levels reached 10 mg/L.

Acute adrenal insufficiency was suspected in patient #15, with asthma, abdominal pain and undetectable serum and free urinary cortisol (mitotane levels of 27 mg/L in the acute phase), after 16 months of treatment. Mitotane was suspended; patient received appropriate hydrocortisone therapy and made a prompt clinical recovery. He exhibited adrenal insufficiency requiring hydrocortisone supplementation for 2 months, and then relapsed with an increase in UFC levels up to more than 100 µg/dL concomitantly with mitotane levels above 3 mg/L. Mitotane was started again at lower dose.

The most serious side effect was severe acute hepatitis in patient #28. This patient was a girl aged 11.8 years at diagnosis of CD. She presented with a low GV $z$-score ($−4$ s.d.), delayed puberty (Tanner B2 for more than 1 year), a UFC level of 152 µg/24 h, a midnight plasma cortisol level of 10.3 µg/dL and a rise in ACTH levels after a corticotropin-releasing hormone stimulation test (30–118 pg/mL). The pituitary MRI failed to visualize an adenoma. Mitotane was instituted at 1.5 g/24 h. After 6 months of treatment, her GV $z$-score had greatly improved (+2 s.d.) and pubertal development resumed (Tanner B3). The acute hepatitis occurred after 7 months on mitotane, revealed by abdominal pain and icterus (aspartate aminotransferase 452 U/L, alanine aminotransferase 528 U/L, prothrombin 22%). Serum levels of mitotane were found to be elevated (36 mg/L) during the period of liver failure likely because of lipid storage and progressive release and decreased...
to undetectable, cortisol secretion resumed, CD became clinically evident again and the patient underwent TSS, 38 months after diagnosis.

Changes in pituitary adenoma

In patient #26, we observed the progressive growth of an adenoma in the left pituitary lobe over the 40 months of mitotane treatment (Fig. 4). A complete selective adenomectomy by TSS was performed 3 years and 7 months after the diagnosis; and the diagnosis of CD induced by a corticotroph adenoma was confirmed through ACTH immunohistostaining. The surgery was uneventful, and followed by an immediate corticotroph deficiency. One year after surgery, the patient still required glucocorticoid replacement. He was treated with growth hormone between the ages of 14 and 17 years because of severe growth retardation. His final height z-score is −1.5. During the mitotane treatment, he began puberty and achieved appropriate pubertal development (testicular volume 25 mL and serum testosterone 3.9 ng/mL at the age of 17 years).

Discussion

Treatment of CD in children is a challenge that requires an experienced multidisciplinary team, defined by the Pituitary Society as Pituitary Tumor Centers of Excellence (PTCOE) (32). Like in adults, the ideal treatment is based on excision of the ACTH-secreting corticotroph adenoma; however, several factors may prevent the removal of the pituitary adenoma and lead to the use of alternative therapies. When pituitary surgery is not possible, the therapeutic options depend on the expertise and habits of the treating center/country, drug availability in children and the patients' tolerance of the drugs. So far, the choice appears much more limited in children than in adults, being restricted to pituitary radiotherapy and drugs that inhibit cortisol production by the adrenal glands, such as mitotane, ketoconazole and metyrapone.

Mitotane is approved for the treatment of adrenocortical carcinoma. It has been used as a cortisol-lowering drug for the treatment of CD in adults (27, 28, 29, 30, 31). A case report describes the use of mitotane for 18 years for intractable CD, leading to the steady control of hypercortisolism (26). In one study in adults with CD, low-dose mitotane (0.5 to 2 g daily) resulted in physiological cortisol levels with minimal side effects, and the authors found no correlation between plasma mitotane levels and
Table 3  Various side effects observed in seven patients who received prolonged low-dose op’DDD monotherapy.

<table>
<thead>
<tr>
<th>Side effects</th>
<th>Prevalence</th>
<th>Description</th>
<th>Outcome</th>
</tr>
</thead>
<tbody>
<tr>
<td>Adrenal deficiency</td>
<td>2/6</td>
<td>Undetectable serum and urinary cortisol</td>
<td>Normalization after dose decrease</td>
</tr>
<tr>
<td>Digestive toxicity</td>
<td>6/6</td>
<td>Nausea, vomiting</td>
<td>Improved after a couple weeks of treatment</td>
</tr>
<tr>
<td>Hepatic toxicity</td>
<td>2/6</td>
<td>Isolated cytolysis</td>
<td>Very slow recovery</td>
</tr>
<tr>
<td>Neurologic toxicity</td>
<td>4/6</td>
<td>Asthenia</td>
<td>Improvement after dose decrease</td>
</tr>
<tr>
<td></td>
<td>2/6</td>
<td>Difficulties at school</td>
<td></td>
</tr>
</tbody>
</table>

Regarding growth, previous studies suggest that effective therapy allows height catch-up of around one z-score, yet children rarely reach their target height (8, 11, 33, 34, 35, 36, 37). In children given mitotane, we observed an improvement in GV, despite not normalizing UFC (UFC levels did not always drop below 80µg/24h). Likewise, pubertal development and BMI improved. These results may catch with data recently published by Ceccato et al., illustrating an improvement of body composition after remission of CD in adults, fat mass reduction being higher after surgical remission (38). In addition, mitotane may facilitate an increase in adenoma volume through the stimulation of pituitary ACTH secretion driven by the adrenal insufficiency. Indeed, some decades ago, it was suggested that Nelson’s syndrome could develop as a consequence of adrenalectomy in children (39).

Side effects were similar to those previously reported, i.e., non-specific digestive symptoms, concentration or memory problems, exhaustion and adrenal insufficiency. One individual, who had a high plasma concentration of mitotane, developed acute severe hepatitis with transient liver failure. As our observations support the view that side effects are likely to be reversible and dose dependent, we propose that the use of mitotane should be considered in
Mitotane to treat pediatric Cushing’s disease

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Pediatric CD, always in the context of close clinical and biochemical follow-up as well as thorough training of patients and their families. Notably, growth and puberty were restored despite non-optimal control of cortisol secretion (UFC between 80 and 150 µg/24h). In most cases, UFC levels were normalized when mitotane concentrations reached 10mg/mL. This suggests that low doses of mitotane are likely sufficient for the control of the disease and would also avoid side effects. Nevertheless, mitotane should be considered only as an adjuvant therapy, to buy time and prepare better conditions for TSS. Though it remains to be demonstrated whether the treatment favors growth of the pituitary adenoma, follow-up of patients given mitotane should include pituitary MRI.

In conclusion, CD in children is a rare and severe condition that affects major biological phenomena, in particular, growth and puberty. Mitotane may be used as an alternative therapy when transsphenoidal pituitary surgery is not feasible. Nevertheless, close surveillance of potential side effects, plasma concentrations of the drug and pituitary MRI is required, in order to use the lowest clinically effective dose while waiting for a definitive cure for the disease.

Supplementary data
This is linked to the online version of the paper at https://doi.org/10.1530/EC-18-0215.

Declaration of interest
A Linglart is a member of the editorial board of Endocrine Connections, but was not involved in the review or editorial process for this paper. None of the other authors has any conflict of interest that could be perceived as prejudicing the impartiality of the research reported.

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Author contribution statement
E M and A L conceived the study. N L performed the biochemical assays. A R, C T, R C and A L recruited patients and provided clinical data. E M and A L analyzed the results and wrote the first version of the manuscript. All authors contributed to the writing and edition of the manuscript.

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