Quality of compounded hydrocortisone capsules used in the treatment of children

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Abstract

Objectives: Due to the lack of paediatric-licensed formulations, children are often treated with individualized pharmacy-compounded adult medication. An international web-based survey about the types of medication in children with adrenal insufficiency (AI) revealed that the majority of paediatric physicians are using pharmacy-compounded medication to treat children with AI. Observations of loss of therapy control in children with congenital adrenal hyperplasia with compounded hydrocortisone capsules and regained control after prescribing a new hydrocortisone batch led to this ‘real world’ evaluation of pharmacy-compounded paediatric hydrocortisone capsules.

Methods: Capsule samples were collected randomly from volunteering parents of treated children suffering from congenital adrenal hyperplasia from all over Germany. Analysis of net mass and hydrocortisone content by high-performance liquid chromatography with ultraviolet (HPLC-UV) detection method was performed based on the European Pharmacopeia.

Results: In a total of 61 batches that were sent, 5 batches could not be analysed because of missing dose information, insufficient number of capsules or were not possible to be evaluated. Fifty-six batches containing 1125 capsules were evaluated. 21.4% of the batches revealed insufficiency in uniformity of net mass or drug content and additional 3.6% failed because they did not contain the labelled drug.

Conclusions: Compounded medication is a possible cause of variation of steroid doses in children with adrenal insufficiency or congenital adrenal hyperplasia, putting these vulnerable patients at risk of poor disease control and adrenal crisis. These data may apply to other individualized compounded oral medication as well, emphasizing the need for development of licensed paediatric formulations approved by regulatory authorities.

Introduction

A five-year-old girl with congenital adrenal hyperplasia (CAH) and no history of non-compliance developed accelerated growth and elevated androgens. Visual inspection of her compounded hydrocortisone capsules showed variable capsule filling (Fig. 1); assuming similar fluctuations in hydrocortisone content as in filling, a new batch was prescribed which immediately improved disease control. Based on this observation, we undertook a ‘real world’ evaluation of pharmacy-compounded paediatric hydrocortisone capsules.

Compounding adult medication is commonly used to provide paediatric doses. An international web-based survey was distributed through the newsletter of the European Society of Paediatric Endocrinology (ESPE) to specify their current treatment of adrenal insufficiency (AI) in neonates and children (1). Sixty-seven participants
Quality of compounded hydrocortisone from 16 countries, mostly from Europe, stated the use of divided licensed adult medication (60%) as off-label and prescription of individualized capsules (55%) as unlicensed use (1). The prescribed dosage varied from 0.5 mg to 5.0 mg with the vast majority of doses being 1.0 mg and 2.0 mg. Compounding is regulated by less stringent criteria than those used for licensed drugs as compounded drugs are neither approved by the EMA (European Medicines Agency) or FDA (Food and Drug Administration) nor required to demonstrate efficacy (2). There is also exemption from Good Manufacturing Practice (GMP) requirements in manufacture of compounded medicines which increases the risks of inaccurate dosing (2). Compounding is by definition an unlicensed use which accounts for up to 60% of medicines administered to children in Europe and the United States (3, 4). The European Pharmacopeia (EP) is a reference work giving common standards for the quality of medicines and their components to provide a basis for the safe use of medicines. The EP contains acceptance criteria for mass and content uniformity to evaluate batches or single capsules (5).

Here, we report a problem with dosing when using compounded hydrocortisone in children suffering from AI and CAH. These patients require hydrocortisone replacement therapy to avoid life-threatening adrenal crises and careful titration of therapy to control androgen excess and optimize growth and development. The lowest dose available of hydrocortisone in the USA is a 5.0 mg tablet and in Europe a 10.0 mg tablet. However, the average dose used in paediatrics is 2.0 mg with doses as low as 0.5 mg required for neonates and infants (1). There is therefore a necessity to compound hydrocortisone for paediatric use. Use of compounded hydrocortisone medication in children is thereby unlicensed as well as off-label treatment. Hydrocortisone has a narrow therapeutic index, and patients with CAH have poor health outcomes in adulthood on current glucocorticoid treatment in part related to treatment in childhood (6, 7). Poor compliance is often blamed for poor disease control in children with CAH. However, as we report here, this might also be due to the lack of content uniformity in compounded hydrocortisone preparations.

**Methods**

A newsletter from the German CAH-patient organisation as well as information provided by treating physicians during regular appointments was used to ask parents and caregivers to participate in this study. Parents or caregivers were invited to send 20 capsules of their child’s compounded hydrocortisone medication to our laboratory. The capsules received for this study were prescribed for CAH-patients aged 0–14 years. No information about sex or other clinical variables was collected. Sixty-one batches (n_capsules: 1210, mean 20/batch) from 11 of 16 federal states in Germany from 35 families and produced in 37 different pharmacies were examined. Net mass of capsule content was calculated by weighing each capsule and subtracting the nominal casing mass from the total mass. To calculate uniformity, mean net mass of each batch and deviations from this mean net mass per capsule were calculated. To evaluate the hydrocortisone content, a HPLC-UV method was developed (8) and validated according to EMA Guideline (http://www.ema.europa.eu/docs/en_GB/document_library/Scientific_guideline/2011/08/WC500109686.pdf Published 2012. Accessed September 2, 2013) (9). To calculate uniformity of hydrocortisone content, mean batch content and deviations from these means per capsule were calculated. The results were evaluated based on the EP acceptance criteria for mass and content uniformity (5). A batch failed if >10% exceeds tolerable limits and ≥1 capsule exceeds non-tolerable limits (5). The limits used were 10 and 20% for mass and 15 and 25% for content. The study was approved by the institutional ethics committee (EA2/134/12). Study participants received a results information letter relating to their supplied batch.
Results

Of 61 capsule batches that were received for analysis, 5 batches were excluded from all further analyses – one of them because surprisingly olive oil had been used as excipient in the capsule which leaked out of the capsules and neither net mass nor drug content could be measured; one batch had no dosing information provided, and in three batches, the number of capsules was too small for evaluation. Fifty-six batches containing 1125 capsules were further evaluated in detail, and the results for individual capsules evaluated are shown in Table 1. Two batches contained no hydrocortisone at all but as the primary criteria are based on uniformity analysis of mass and content, these were included in the analysis and passed the content criteria with a homogenous result of 0% hydrocortisone in all capsules. Batch analysis revealed the following results: for drug content, 49 batches (87.5%) fulfilled the acceptance criteria based on the EP guidelines, whilst 7 batches (12.5%) failed. Extreme hydrocortisone-containing batches showed deviations from mean drug content of ±30–200%. For net mass, 47 batches (84%) fulfilled the acceptance criteria, whilst 9 batches (16%) failed. Thus, of the compounded capsules that contained hydrocortisone, 21.4% of batches revealed a failure in the acceptance criteria in net mass or drug content: 3 batches with content deviations only (5.4%), 5 batches with net mass deviation only (9.0%) and 4 batches with both content and net mass deviation (7.1%). In summary, 21.4% failed the uniformity analysis, and 3.6% did not contain the labelled steroid at all, totalling up to 25% of batches that cannot ensure safe therapy for the patients.

Discussion

In our ‘real world’ analysis, every 4th batch of compounded hydrocortisone revealed a failure of the acceptance criteria based on the EP. In extreme batches that contained hydrocortisone, the drug content deviated from mean drug content by ±30–200% and some batches did not contain any detectable hydrocortisone, but another glucocorticoid, despite being labelled as such.

Here, we report a problem with dosing when using compounded hydrocortisone in children suffering from AI and CAH. These patients require hydrocortisone replacement therapy to avoid life-threatening adrenal crises and careful titration of therapy to control androgen excess and optimize growth and development. The lowest dose available for hydrocortisone in the USA is a 5.0mg tablet and a 10.0mg tablet in Europe. However, the average dose used in paediatrics is 2.0mg with doses as low as 0.5mg required for neonates and infants (1). There is therefore a necessity to compound hydrocortisone for paediatric use. Hydrocortisone has a narrow therapeutic index, and patients with CAH have poor health outcomes in adulthood on current glucocorticoid treatment in part related to treatment in childhood (6, 7). Poor compliance is often blamed for poor disease control in children with CAH, and this causes considerable distress to children and parents. However, as we show here, this might also be due to the lack of content uniformity in compounded hydrocortisone.

The advantage of our study is that it was done as a ‘real world’ analysis of what medication patients receive from their local pharmacy. A limitation of our study is that it was undertaken in only one country, however, the regulations used apply across Europe and are similar to those in USA. Based on our international web-based survey of medication, it is common paediatric practice in most countries to use divided licensed adult medication (60%) or prescribe individualized capsules (55%) (1). Sample collection was voluntary and not designed to be representative, thereby we cannot exclude a possible sampling bias because parents in doubt (e.g. receiving oily capsules) of the child’s medication might be more likely to participate.

There are no obligatory quality checks of pharmacy-compounded medication regarding drug content. Pharmacies may take part in a voluntary external control programme, but these tests are expensive and participation is not monitored or reported. As the regulations for pharmacy-compounded hydrocortisone are identical to those for other pharmacy-produced medication, our findings may also be applicable to other compounded medication. Further studies on a broader range of compounded paediatric medication are needed to evaluate the inherent risk of significant dosing errors caused by e.g. adherence or the pharmacuetic compounding process. This risk may frequently escape
paediatrician’s notice when dealing with therapy failure in paediatric patients. Our results demonstrate that variations in pharmacy-compounded medication might cause poor disease control in children. In newborns and infants, precise dosing of hydrocortisone at low dose is essential; otherwise, these vulnerable patients are at risk of poor disease control and potential adrenal crisis. There is a need for the development of licensed paediatric formulations approved by regulatory authorities such as the EMA and FDA.

Declaration of interest
R J R and M J W are Directors of Diurnal Ltd, UK. The other authors have indicated that they have no potential conflicts of interest to disclose.

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Author contribution statement
U Neumann and S Spielmann conceptualised and designed the study, collected the capsules, coordinated the responses to the parents, carried out initial data analysis, drafted the initial manuscript and approved the final manuscript as submitted. D Burau and C Kloft carried out the capsule analyses as well as the data analysis, drafted the initial manuscript and approved the final manuscript as submitted. M J Whitaker and R J Ross participated in conceptualising the study, drafting and reviewing the manuscript, and approved the final manuscript as submitted. O Blankenstein conceptualised and designed the study, participated in capsule collection, carried out the initial analysis, drafted the initial manuscript and approved the final manuscript as submitted.

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References