Acceptability of different oral formulations in infants and preschool children

Diana A van Riet-Nales,1,2 Barbara J de Neef,3 Alfred F A M Schobben,2 José A Ferreira,4 Toine C G Egberts,2,5 Catharine M A Rademaker5

ABSTRACT

Objective Liquid medicines are easy to swallow. However, they may have disadvantages, such as a bad taste or refrigerated storage conditions. These disadvantages may be avoided by the use of oral solid medicines, such as powders or tablets. The aim of this study was to investigate the acceptability of and preference among four oral formulations in domiciliary infants and preschool children in The Netherlands.

Methods Parents administered four oral placebo dosage forms that were aimed at a neutral taste, at home, to their child (1–4 years of age) twice on one day following a randomised cross-over design: small (4 mm) tablet, powder, suspension and syrup. They were asked to report the child’s acceptability by a score on a 10 cm visual analogue scale (VAS score) and by the result of the intake. At the end of the study, they were asked to report the preference of the child and themselves.

Results 183 children were included and 148 children were evaluated. The data revealed a period/cross-over effect. The estimate of the mean VAS score was significantly higher for the tablet than for the suspension (tablet 9.39/9.01; powder 8.84/8.20, suspension 8.26/7.90, syrup 8.35/8.19; data day 1/all days). The estimate of the mean number of intakes fully swallowed was significantly higher for the tablet than for the other formulations (all p values <0.05). Children and parents preferred the tablet and syrup over the suspension and the suspension over the powder (all p values <0.05).

Conclusions All formulations were well accepted. The tablets were the best accepted formulation; the tablets and syrup the most preferred.

Trial Registration number ISRCTN63138435.

INTRODUCTION

For decades, oral liquid dosage forms, such as syrups and suspensions, have been considered as the favourable type of dosage form in which to administer medicines to young children.1,2 However, oral liquid medicines may have disadvantages, such as a bad taste, portability problems or refrigerated storage conditions.3–6 Therefore, WHO currently favours that young children, particularly in developing countries, be treated with oral solid medicines.7

Oral liquid medicines are more commonly available for use in infants and preschool children than oral solid (flexible) medicines, such as powders or orodispersible tablets.8 Small-sized tablets, also referred to as minitablets, have been identified as a new type of oral solid dosage form in which to administer medicines to young children. However, only few of such tablets have been authorised for children below 4 years of age.1,6–8,11 Nevertheless, small tablets have been widely used in this age group as food supplements, for example, 4 mm sodium fluoride tablets for caries prevention, or 4 mm vitamin AD tablets.12–16

The selection of an oral dosage form and the pharmaceutical aspects of the formulation, such as the palatability of an oral suspension or the size of a tablet, are important factors in the overall acceptability of an oral paediatric medicine.6,10 As adequate child and parent acceptability are prerequisites for good drug adherence, paediatric treatment outcomes may be enhanced by a careful selection of the formulation including the type of the dosage form. Therefore, the aim of this study was to investigate the acceptability of and preference among four oral formulations in domiciliary infants and preschool children in The Netherlands.

METHODS

Study design

A randomised cross-over trial was performed in six Dutch preschool preventive healthcare clinics in Beusichem, Beesd, Culemborg (2 clinics), Maurik and Zaltbommel. Ethical approval was waived by the Central Committee on Research involving human subjects (CCMO) on basis of the Dutch Medical Research Involving Human Subjects Act (WMO). Approval was obtained from the
Drug therapy

Parents were verbally approached by one of four recruiters (a licensed pharmacist and three graduate students) when attending the clinics in 2011. Parents had either received the written information by mail 2 weeks before the appointment, or this information was handed to them at the end of the face-to-face contact. Parents were asked or called by phone for written informed consent and study participation at least 2 weeks after the written information was provided. The results of the selection process were systematically gathered (date when verbally approached, healthcare clinic, recruiter, date of birth, child gender, willingness to participate, reason for exclusion if mentioned voluntarily).

Children were eligible for inclusion in this study if they were between 1 and 4 years of age and if their parents had mastery of the Dutch language. Exclusion criteria were: (1) significant developmental delay; (2) having swallowing difficulties, an eating disorder or a chronic condition requiring oral medication; (3) hypersensitive to lactose, having cow-milk allergy or having an allergy of unknown origin; (4) a member of staff of the preventive healthcare clinic considered that study participation was inappropriate in view of the family situation. During the study, the criterion added was (5) according to the parents' observation, children should not feel ill when the formulations were actually given.

Setting and study participation

The aim of the preschool preventive healthcare clinics is to monitor the mental and physical development of children between 0 and 4 years of age, to advise parents on child-raising issues and to vaccinate children.17 The response rate to the invitation for an appointment is over 99% of children below 2 years of age and over 90% of children between 2 and 4 years of age.18

Sample size

The sample size for acceptability was calculated on basis of a design aimed at detecting a specified difference between theVAS scores of two treatments in a cross-over trial involving four oral formulations on four different days.19 The power was set at 0.8 and the significance level at 0.05. Due to a lack of relevant data from the literature on the acceptability of oral formulations in (young) children, the sample size calculations were based on plausible values for the mean difference and SD of the VAS scores. The sample size for preference was calculated on basis of a statistical design where parents were asked to identify the single most preferred formulation. The same approach was applied as for the calculation of the acceptability. The sample size was set at 150 evaluable children, which would, in most cases, allow a maximum difference of 2 for acceptability and 0.2 for preference to be detected.

Randomisation

The study was randomised for the order of administration of the formulations by an RIVM employee who was not involved in this study. Randomisation was conducted with a random sequence obtained from http://www.random.org. The same sequence was applied to each block of 24 children. Siblings were allocated to the same order to avoid mistakes.

Data analysis

The following analyses were conducted: (1) assessment of systematic differences between the two single VAS scores for a particular formulation (paired Z tests); (2) in case of no significant differences, calculation of the mean VAS scores per child and formulation; (3) evaluation of a potential cross-over or period effect (Z test on the order of the best accepted formulation), in case of such an effect analysis 3 and 4 were done for the administrations of the first formulation only (day 1) and for all data (all 4 days); (4) estimation of the mean VAS score per formulation and computation of the corresponding 95% CIs (Z statistics); (5) testing of differences between the mean VAS scores of two different formulations (Wilcoxon and Mann–Whitney tests); (6) estimation of the mean number of intakes that were fully swallowed by a child per formulation and computation of the corresponding 95% CIs (Z statistics); (7) computation of estimates and associated 95% CIs of the probabilities that the child and parents preferred a particular formulation, and comparison between the four probabilities (Z tests).

All statistics were conducted applying Excel 2007 (Microsoft, Redmond, Washington), R V2.13 (R development core team).

RESULTS

Setting and study participation

Between February and July 2011, 421 children from 373 families were verbally approached; 405 children from 358 families
Table 1  Characteristics of four different oral formulations

<table>
<thead>
<tr>
<th></th>
<th>Tablet</th>
<th>Powder</th>
<th>Suspension</th>
<th>Syrup</th>
</tr>
</thead>
<tbody>
<tr>
<td>Picture</td>
<td><img src="image1" alt="Tablet" /></td>
<td><img src="image2" alt="Powder" /></td>
<td><img src="image3" alt="Suspension" /></td>
<td><img src="image4" alt="Syrup" /></td>
</tr>
<tr>
<td>Appearance</td>
<td>Round biconvex white tablets</td>
<td>Freely flowing powder</td>
<td>Homogeneous and opaque liquid after shaking</td>
<td>Clear solution</td>
</tr>
<tr>
<td>Dosing recommendation</td>
<td>1 tablet of 4 mm (43.0 mg)</td>
<td>250 mg powder (1 sachet)</td>
<td>2.5 mL</td>
<td>2.5 mL</td>
</tr>
<tr>
<td>Taste</td>
<td>Aimed at neutral</td>
<td>Aimed at neutral</td>
<td>Aimed at neutral</td>
<td>Aimed at neutral</td>
</tr>
<tr>
<td>Dosing device</td>
<td>Not applicable</td>
<td>Not applicable (spoon is an example to show the powder and is not dispensed to the participant)</td>
<td>3 mL oral syringe with 0.1 mL graduation. The syringe can be attached to the cap of the container</td>
<td>3 mL oral syringe with 0.1 mL graduation. The syringe can be attached to the cap of the container</td>
</tr>
<tr>
<td>Composition</td>
<td>Lactose monohydrate 34.69 g</td>
<td>Lactose monohydrate 203.7 mg</td>
<td>Methylparahydroxybenzoate 46.0 mg</td>
<td>Methylparahydroxybenzoate 63.1 mg</td>
</tr>
<tr>
<td></td>
<td>Maydis amylum 6.46 g</td>
<td>Maydis amylum 38.0 mg</td>
<td>aluminiummagnesiumsilicate 484.4 mg</td>
<td>Propylyparahydroxybenzoate 10.0 mg</td>
</tr>
<tr>
<td></td>
<td>Maydis amylum pregelificatum 1.42 g</td>
<td>Maydis amylum pregelificatum 8.3 mg</td>
<td>Carboxymethylcellulose 484.5 mg</td>
<td>Citric acid 36.3 mg</td>
</tr>
<tr>
<td></td>
<td>Magnesium stearate 0.43 g</td>
<td>Total 250 mg</td>
<td>Sucrose 12.74 g</td>
<td>Saccharose 8.28 g</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Purified water 37.95 g</td>
<td>Purified water 37.95 g</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Microcrystalline cellulosis 2.50 g</td>
<td>Microcrystalline cellulosis 2.50 g</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Purified water ad 50 mL</td>
<td>Purified water ad 50 mL</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Methylparahydroxybenzoate 46.0 mg
Aluminiummagnesiumsilicate 484.4 mg
Carboxymethylcellulose 484.5 mg
Citric acid 36.3 mg
Sucrose 12.74 g
Purified water 37.95 g
Microcrystalline cellulosis 2.50 g
Purified water ad 50 mL

were eligible for inclusion if their parents would pass the language check. Informed consent was obtained for 183 children from 153 families. Diaries including information on the acceptability and preference of the formulations were returned for 151 children from 124 families (recruitment success rate 45%, loss to follow-up 17%). Three diaries from two families could not be used in the data analysis because it was not clear in which order the formulations were administered (figures 1 and 2). The recruitment success rate in the population eligible for inclusion was similar among all the participating healthcare

**Figure 1** Participant flow through the study.
clinics and recruiters. The age and gender of the children eligible for evaluation were not statistically different from the children eligible for inclusion.

Child and parent acceptability
The data did not indicate systematic differences between the single VAS scores of the two administrations of each formulation to a child. Therefore, the mean VAS scores were used for the further evaluations. The VAS score data indicated a period or cross-over effect by which formulations administered earlier tended to have somewhat higher scores (p value <0.0001). As a consequence, analysis started with the data of the first day only. The estimates of the mean VAS scores of the first day were: tablet 9.39 (32 children); powder 8.84 (45 children); suspension 8.26 (34 children) and 8.35 syrup (37 children); see table 2 for the CIs. The tablet scored better than the suspension even when applying a Bonferroni correction (p=0.001: for correction multiply by 6). The other comparisons were less clear, but there was an indication that the tablet scored better than the syrup as well as the powder.

Using the data from all 4 days, estimates of mean scores per formulation were obtained across the 24 different orders of administrations, each order getting the same weight. Although no clear ranking was visible between the syrup, suspension and powder, the superiority of tablet over the other three forms was more evident than by considering the data from the first day only (table 2).

The estimate of the mean number of administrations that were fully swallowed were 1.96 (tablet), 1.58 (powder), 1.70 (suspension) and 1.67 (syrup) This number was significantly higher for the tablet than for the other formulations (p value <0.05) (table 2). The scatter-plot of the single VAS scores versus the result of the intake (data not shown) clearly illustrated that the VAS score was predictive for the result of the intake. No choking was reported.

Child and parent preference
Children and parents appeared to prefer the tablet and syrup over the suspension and the powder (p values <0.001). There is also some indication (p value=0.082) that parents preferred the tablet to the syrup (table 3).

DISCUSSION
In this randomised cross-over trial, the four formulations investigated can all be considered well accepted by children between 1 and 4 years? The small 4 mm tablet was significantly better

**Table 2 Acceptability of four different oral formulations (n=148 children)**

<table>
<thead>
<tr>
<th>Numerical data</th>
<th>First day (four different groups)</th>
<th>All four days (cross-over design)</th>
<th>Result of the intake</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Mean* (95% CI)</td>
<td>Mean* (95% CI)</td>
<td>Mean† (95% CI)</td>
</tr>
<tr>
<td>Tablet</td>
<td>9.39 (8.85 to 9.93), n=32</td>
<td>9.01 (8.75 to 9.28)</td>
<td>1.96 (1.92 to 2.00)</td>
</tr>
<tr>
<td>Powder</td>
<td>8.84 (8.19 to 9.49), n=45</td>
<td>8.20 (7.84 to 8.56)</td>
<td>1.58 (1.44 to 1.71)</td>
</tr>
<tr>
<td>Suspension</td>
<td>8.26 (7.47 to 9.04), n=34</td>
<td>7.90 (7.42 to 8.38)</td>
<td>1.70 (1.57 to 1.83)</td>
</tr>
<tr>
<td>Syrup</td>
<td>8.35 (7.45 to 9.25), n=37</td>
<td>8.19 (7.73 to 8.64)</td>
<td>1.67 (1.54 to 1.80)</td>
</tr>
<tr>
<td>p Value‡</td>
<td>&lt;0.001</td>
<td>0.054</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Testing for any differences</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Tablet versus powder</td>
<td>&lt;0.001</td>
<td>0.054</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Tablet versus suspension</td>
<td>&lt;0.001</td>
<td>0.001</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Tablet versus syrup</td>
<td>&lt;0.001</td>
<td>0.027</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Powder versus suspension</td>
<td>0.378</td>
<td>0.060</td>
<td>0.081</td>
</tr>
<tr>
<td>Powder versus syrup</td>
<td>0.869</td>
<td>0.611</td>
<td>0.168</td>
</tr>
<tr>
<td>Suspension versus syrup</td>
<td>0.164</td>
<td>0.302</td>
<td>0.513</td>
</tr>
</tbody>
</table>

*Estimate of the mean acceptability as expressed on a 10 cm visual analogue scale (VAS score).
†Estimate of the mean number of administrations of a formulation that were fully swallowed, maximum n=2.00.
‡p Values of the Mann–Whitney/Wilcoxon tests regarding any differences between the mean VAS scores of two different formulations.
§p Values of the Z tests comparing pairs of formulations regarding the mean number of administrations that were fully swallowed.
accepted than the suspension, and there was an indication that the tablet was also better accepted than the powder and syrup. The tablet was significantly more often fully swallowed than the other formulations. Children and parents preferred the tablet and syrup over the suspension, and the suspension over the powder.

Child acceptability of oral medicines has been studied for many years.20–22 Few studies however, have focussed on the acceptability of oral dosage forms as such. Ansah et al23 compared tablet with syrup formulations in 155 children from between birth and five years (of age) for the treatment of malaria, and Bagenda et al24 in 129 children from between 6 months and 12 years of age in case of treatment with highly active anti-retroviral therapy (HAART). Both teams concluded that the tablet formulations resulted in better adherence. Spomer et al3 compared 2 mm uncoated placebo tablets with a sweet syrup in 60 inpatient children aged from between 6 months and 6 years of age, and concluded that the acceptability of the tablet was at least as good as that of the syrup. Despite key differences in the patient population and methodology, the results of our study are consistent with those of the aforementioned authors.

Three studies have been identified on the child acceptability of small tablets.3 9 10 Apart from the study of Spomer et al,3 the study of Van de Vijver et al10 demonstrated that 2 mm medicated tablets were good to excellently swallowed by 16 outpatient Belgium or Dutch cystic fibrosis patients who were between 6 and 30 months of age. Thomson et al10 demonstrated that larger 3 mm tablets could be swallowed by 46 out of 100 children between 6 and 30 months of age. Thomson et al10 also the first study investigating the child acceptability of oral placebo formulations in a domiciliary rather than inpatient setting, with a double rather than single administration of each formulation, a 4 mm, rather than a 2 or 3 mm tablet, and with two different measuring instruments for child acceptability.

In this study, an indication was found that the mean VAS acceptability score of the tablet was higher than that of the syrup, and that the parents preferred the tablet over the syrup. However, when parents were asked to report the child’s preference, no significant difference was found between the syrup and the tablet. Results such as ours provide an argument for the fact that child and parent acceptability and preferences are different outcomes providing complementary information on the suitability of a formulation. Preferably, these outcomes are investigated in the same study.

This study has some limitations. First, the administrations were not supervised by the research team as this would bias normal family routines. Consequently, the evaluation of the child acceptability and preference relied on parental reports. This self-reporting methodology was not validated prior to the start of the study. Therefore, recruiters focused heavily on adequate verbal instructions to the method of administration and reporting.

Second, child acceptability may be influenced by taste aspects. The powder and tablet were manufactured from the same blend, so their taste was identical. However, the taste of the suspension and syrup differed due to the intrinsic nature of these dosage forms. Therefore, it cannot be excluded that any differences in the acceptability and preference among the liquid formulations were also related to taste.

Third, the recruitment was tailored to healthy domiciliary children between 1 and 4 years of age and parent with mastery of the Dutch language. Hence, the applicability of our findings to children outside this population, for example, children who are feeling ill, who are otherwise fractioned or who are from a foreign ethnicity is left for future research. In view of the findings of the teams of Ansah,23 Bagenda,24 and Spomer,3 it is anticipated that our study’s findings will equally hold for older children.

Fourth, chewing was not evaluated as it is common practice in The Netherlands that children may chew on immediate release tablets if they want to. Therefore, the acceptability (swallowability) of tablets that should be taken as a whole, for example, monolithic extended release tablets or tablets with essential taste masking, is left for future research.

Fifth, we did not systematically evaluate the parents’ reasons to decline participation. However, from the voluntary reasons provided, it seemed that parents were mainly ‘too busy’ or having a second name suggesting a non-European ethnicity. It cannot be excluded that parents who did not participate in this study might be more reluctant to administer a particular formulation to their child than those who participated.

This study showed that the acceptability of 4 mm tablets is unlikely to be inferior as those of three currently employed dosage forms in infants and preschool children when aimed at a neutral taste. Thus, there is no reason to further question the acceptability of 4 mm immediate release tablets for children from the age of 1 year. Rather than discussing whether small tablets should be the preferred type of dosage form for the development of future paediatric medicines, pharmaceutical industries are recommended to consider the possibility of developing two essentially different dosage forms alongside each other.

**CONCLUSION**

Oral placebo 4 mm round uncoated tablets, powders, suspensions and syrups may be considered well accepted dosage forms

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**Table 3** Preference of four different oral formulations (n=148 children)

<table>
<thead>
<tr>
<th>Formulation</th>
<th>Child Probability (95% CI)*</th>
<th>Parent Probability (95% CI)*</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tablet</td>
<td>0.40 (0.32 to 0.49)</td>
<td>0.49 (0.40 to 0.58)</td>
</tr>
<tr>
<td>Powder</td>
<td>0.07 (0.03 to 0.11)</td>
<td>0.07 (0.03 to 0.11)</td>
</tr>
<tr>
<td>Suspension</td>
<td>0.27 (0.19 to 0.34)</td>
<td>0.23 (0.17 to 0.30)</td>
</tr>
<tr>
<td>Syrup</td>
<td>0.48 (0.39 to 0.57)</td>
<td>0.36 (0.27 to 0.44)</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Testing for any differences</th>
<th>Difference p Value</th>
<th>Difference p Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tablet versus powder</td>
<td>0.334 &lt; 0.001</td>
<td>0.423 &lt; 0.001</td>
</tr>
<tr>
<td>Tablet versus suspension</td>
<td>0.131 0.046</td>
<td>0.256 &lt; 0.001</td>
</tr>
<tr>
<td>Tablet versus syrup</td>
<td>-0.080 0.306</td>
<td>0.137 0.082</td>
</tr>
<tr>
<td>Powder versus suspension</td>
<td>-0.203 &lt; 0.001</td>
<td>-0.166 &lt; 0.001</td>
</tr>
<tr>
<td>Powder versus syrup</td>
<td>-0.414 &lt; 0.001</td>
<td>-0.285 &lt; 0.001</td>
</tr>
<tr>
<td>Suspension versus syrup</td>
<td>-0.211 &lt; 0.001</td>
<td>-0.256 &lt; 0.001</td>
</tr>
</tbody>
</table>

*Estimate of the probabilities that the parent/child has indicated a preference for the formulation.
†Estimates of differences between the probability that one formulation is preferred and the probability that another formulation is preferred and the corresponding p values of the test that the two probabilities are equal.
Contributors DaRN: conceptualised and designed the study, coordinated the participant recruitment, coordinated the manufacture and dispensing of the placebo formulations, was responsible for the liaison with the statistician (JAF), drafted the initial manuscript and approved the final manuscript as submitted. BjdNt provided support in the design of the study towards the calculation of the sample size; he conducted the data analysis, reviewed and revised the manuscript and approved the final manuscript as submitted. TCGE: supervised the conceptualisation and design of the study, supervised the recruitment process and data collection, reviewed and revised the manuscript and approved the final manuscript as submitted. CMAR: supervised the conceptualisation and design of the study, supervised the recruitment process and data collection, reviewed and revised the manuscript, and approved the final manuscript as submitted. JAF: