Hyperphagia in neonates withdrawing from methadone

Alma Martinez, Beth Kastner, H William Taeusch

Abstract

Aims—To examine whether hyperphagia is a clinically significant problem in infants born to women receiving methadone maintenance.

Methods—The volume of feeds, changes in infant body weight, as well as occurrence of adverse clinical effects in infants withdrawing from methadone were studied during the first month of life. A retrospective chart review was conducted for all infants at San Francisco General between 1992 and 1995, born to women receiving methadone maintenance during their pregnancy. Forty-four infants were identified and the data obtained from hospital medical records. The daily oral intake of these infants was recorded during the first month of life. The incidence of hyperphagia (oral intake > 190 cc/kg/day) was measured. Associations between infant oral intake and maternal methadone dose were studied using correlation analysis as well as ANOVA for repeated measures. Adverse clinical symptoms were also recorded. A subset of premature infants was studied separately.

Results—The incidence of hyperphagia was 26% by day 8 and 56% by day 16 of life in the infants. Hyperphagia was not associated with maternal methadone dose or with infant withdrawal scores. Infants who were hyperphagic lost significantly more weight during the first week of life than those who were not. Despite significantly greater intake, the hyperphagic infants did not gain weight more rapidly during the first month of life compared with those infants with lower oral intake. Infants who were hyperphagic (maximum intake of 290 cc/kg/day) did not experience increased vomiting, aspiration, diarrhoea, or abdominal distention.

Conclusions—Hyperphagia is commonly found in infants withdrawing from methadone and can be persistent in a significant number. Hyperphagia was not associated with either increased neonatal weight gain or with adverse gastrointestinal consequences. Hyperphagia may occur in infants withdrawing from methadone who have high metabolic demands due to clinical signs not controlled by opiate treatment.


Keywords: methadone withdrawal; hyperphagia; metabolism

Excessive oral intake, or hyperphagia, is listed as one of many clinical findings in infants withdrawing from opiates.1–3 Conversely, other reports list frantic, disorganised sucking and poor feeding as symptoms of opiate withdrawal.1–3 It is unclear from these previous reports how often excessive oral intake is found in infants withdrawing from opiates and whether hyperphagia is associated with adverse clinical consequences for the neonate, such as vomiting, aspiration, diarrhoea, or abdominal distention.

This study aimed to evaluate infants exposed in utero to methadone, to quantify their volume of feeds during inpatient stay, to determine the incidence of hyperphagia in these infants (compared with published norms of feeding patterns in normal neonates), and to quantify weight change as well as the occurrence of adverse clinical effects in infants withdrawing from methadone.

Methods

Data were collected by retrospective chart review after obtaining approval from the University of California at San Francisco Institutional Review Board for Human Research. The patients were identified from a database of all births at San Francisco General Hospital between 1992 and 1995.

The study population included all infants born at San Francisco General Hospital to women enrolled in a methadone programme during their pregnancy. Infants exposed to other opiates or other illicit drugs in addition to methadone were included in the study. Maternal use of any additional drugs was noted. Infants were treated for signs of withdrawal primarily with tincture of opium, as recommended by the American Academy of Pediatrics.1 Diazepam was also used occasionally, but never without the concomitant use of tincture of opium. Infants were excluded from this study if the mother used opiates but was not enrolled in a methadone programme.

All infants in this study remained in hospital after birth for treatment of withdrawal. A significant number of women were prescribed large daily methadone doses, and their infants often required lengthy hospital stays. In the group of infants studied only seven were discharged home before they were 12 days old, and a few remained in hospital for longer than 30 days for management of opiate withdrawal. Data were collected at prospectively chosen time points during the infants’ hospital stay (days 8, 12, 16, 20, 24, 28 and 30 of life). Data collected included maternal methadone drug dose, birthweight, gestational age, and daily
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Finnegan withdrawal scores,* assigned by the infant’s nurse at each data point. The volume of oral intake, caloric intake, and weight gain were also obtained at the same time points during this period. All infants received infant formula, none was breastfed, and no infant required tube feedings during the study. Drug treatment required for withdrawal was recorded, as were episodes of emesis, drooling, ineffective feeding, diarrhoea, and abdominal distention.

Investigators have carefully documented daily formula intakes for healthy newborn infants living at home. These investigators have shown that during the first month of life, normal, term infants eat between 140 (SEM 22) cc/kg/day and 170 (25) cc/kg/day. Using these previous data, we defined hyperphagia for the purposes of this study, as a daily oral intake greater than 190 cc/kg/day.

Most values are presented as mean (SEM). Withdrawal scores are shown as median scores and range. Anova for repeated measures was used to compare differences in oral intake between infants exposed to differing methadone doses, and unpaired two-tailed t test was used to compare hyperphagics with non-hyperphagics for weight gain. The Wilcoxon Rank Sum Test was used to compare withdrawal scores of hyperphagics with those of non-hyperphagics. Linear regression analysis was performed to determine the predictive value of maternal methadone dose, birthweight, and gestational age to oral intake or to withdrawal scores. The data were analysed using the True Epistat Data Analysis Program (Richardson, Texas). The level of statistical significance was chosen as p<0.05.

Results

Forty four infants were identified during the study period who were born to women enrolled in a methadone maintenance programme. Thirty nine per cent of these women were polydrug users, with 60% also using heroin. Fifty five per cent of the women reported using cocaine or tested positive during the current pregnancy, and 10% had a history of alcohol use during the current pregnancy.

The mean birthweight for all infants studied was 2804 (112) g (range 1340 to 4100). The mean gestational age was 37 (0.5) weeks (range 30 to 42 ). The mean maternal methadone dose was 50 (3.2) mg (range 15 to 95). The formula oral intake for all infants as well as the number of infants studied at each time point studied is shown in table 1. By day 20 of life, the mean oral intake of infants withdrawing from methadone, had reached 202 cc/kg/day. Using previous data for normal formula intake of healthy, term infants, we defined hyperphagia as a daily oral intake greater than 190 cc/kg/day. Using these criteria, 26% of the infants were hyperphagic by day 8 of life, and 56% by day 16 of life. (fig 1).

Using a multiple linear regression model, we found no significant correlation between infant oral intake at any time period studied and maternal methadone dose at delivery, birthweight, or gestational age (on day 8 of life, r = 0.27, p = 0.4). We stratified the infants into three groups of methadone exposure, based on daily maternal dose. The “low” dose of exposure was defined as 0–30 mg daily, “medium” as 35–50 mg daily, and “high” as 55–95 mg daily. Using these stratified data, we found no difference in the daily intake between the infants born to mothers with low, medium, or high methadone doses at each time point measured (p>0.05, Anova) (fig 2).

As shown in fig 3, despite significantly greater oral intake in hyperphagics than in infants with lower oral intake, the hyperphagics lost significantly more weight during the first week of life. During the rest of their hospital stay, the hyperphagic infants did not show differences in weight gain when compared to the lower oral intake group (fig 3).

Withdrawal scores of infants who were hyperphagic did not differ from infants who had lower levels of oral intake except for day 28 of life (Wilcoxon Rank Sum Test). At this late date, the median withdrawal score for hyperphagics was 12 compared with five for non-hyperphagics; p= 0.03.

There were no increased gastrointestinal symptoms, such as vomiting, diarrhoea, drooling, abdominal distention or aspiration found in infants who were hyperphagic compared with infants with lower oral intake (p>0.05;
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Figure 2  Formula intake in infants withdrawing from methadone during the first month of life. Oral intake was stratified by maternal methadone dose. Comparisons between low dose (< 30 mg/day), medium dose (35–50 mg/day), and high dose (35–95 mg/day) are shown. Values are mean (SEM); p>0.05, Anova.

Figure 3  Weight change (g/day), as measured from the previous time point, for infants withdrawing from methadone during the first month of life. Comparisons made between infants with normal intake vs high (hyperphagia) intake; *p<0.05, Anova.

Anova). Contrary to our expectations, no hyperphagic infant had a clinical complication attributable to increased oral intake.

Fifteen preterm infants were included in this study. The patient characteristics of the preterm infants are shown in table 2. The mean gestational age for the preterm infants was 33 weeks (range 30–36), the mean birthweight was 2136 g (range 1340–3365). There was no statistical difference between the maternal methadone dose in preterm or term infants. The oral intake for these preterm infants was 56 cc/kg/day during the first week of life. The smallest infant studied and in this instance, the infant’s oral intake was deliberately limited by the physicians caring for the child. Beginning in the second week of life, this infant had increased oral intake that continued for the rest of his inpatient stay. The lack of ineffectual feeding in this cohort of infants withdrawing from methadone is an interesting finding. Perhaps the use of neonatal scoring systems to quantify the severity of withdrawal, as well as the prompt use of medications to control withdrawal, enable these infants to eat effectively.

The use of a historical control group for this study is a serious consideration in evaluating these findings. The elegant data published by previous workers detailing the normal infant oral intake or feeding problems in the preterm infants when compared with term infants at any time point (p > 0.05; Anova).

Discussion

While hyperphagia or excessive oral intake has been cited before as a clinical finding in neonates undergoing opiate withdrawal, the actual incidence of this finding has not been reported.1–3 Hyperphagia is common in infants withdrawing from methadone, occurring in 56% of infants by the second week of life. Infants with hyperphagia lost significantly more weight during the first week of life, and had similar weight gains thereafter, when compared with non-hyperphagic infants withdrawing from methadone. No significant correlation between maternal methadone dose and infant oral intake or withdrawal scores was found during the first month of life. The infants who were hyperphagic did not have a greater incidence of ineffective feeding or sucking, vomiting, or diarrhoea. A smaller subset of preterm infants was studied separately and they were no different from full term infants with regards to all outcomes measured for this study.

Excessive sucking or disorganised and ineffectual sucking associated with poor oral intake has been reported before in infants withdrawing from opiates.4–8 We found the opposite—that is, by the second week of life, a large percentage of infants are hyperphagic. In our study there was only one infant who had low oral intake (56 cc/kg/day) during the first week of life. This was the smallest infant studied and in this instance, the infant’s oral intake was deliberately limited by the physicians caring for the child. Despite prematurity in this subset of infants, there were no statistical differences in

Table 2  Characteristics of preterm infants exposed to methadone

<table>
<thead>
<tr>
<th>Preterm infants</th>
<th>Range</th>
</tr>
</thead>
<tbody>
<tr>
<td>Birthweight (g)</td>
<td>2136 (138) – 1340–3365</td>
</tr>
<tr>
<td>Gestational age (weeks)</td>
<td>33.4 (0.4) – 30–36</td>
</tr>
<tr>
<td>Methadone dose (mg)</td>
<td>47.4 (3) – 30–95</td>
</tr>
</tbody>
</table>

All newborn data shown are mean (SEM). For each time point, there was no significant difference between oral intake for premature infants vs term infants (p>0.05; Anova).
oral intake would be costly and difficult to replicate.10 11 These investigators followed a cohort of healthy, term infants for 120 days, and measured formula intake, caloric intake, and growth parameters. It would also not be possible to use a “healthy” hospital control group of infants, as healthy infants are not admitted for lengthy periods as were these infants withdrawing from methadone. This remains an acknowledged limitation of the current study. A natural extension for future studies would be to look at the feeding patterns of infants similarly treated for withdrawal who are exposed to other opiates.

The question of whether large volumes of oral intake are harmful for these infants was a primary reason for the initiation of this study. We found no evidence of adverse gastrointestinal clinical occurrences in the term or preterm hyperphagic infants. The finding of greater weight loss in the first four days of life followed by similar weight gains in the hyperphagic infants compared with the infants on lower oral intake, suggests higher metabolic demands (or greater caloric losses) in the hyperphagic infants. Perhaps administration of higher doses of medications in hyperphagic infants might lead to lower oral intake as well as more effective treatment of other signs of withdrawal.

Our finding of a lack of correlation between maternal methadone dose and infant’s oral intake, as well as a lack of correlation between withdrawal scores and hyperphagia during the first month of life, is consistent with results from some previous studies but disagrees the work of others. Mack et al found no correlation between neonatal serum concentrations of methadone and the severity of withdrawal in the neonates.13 Previous workers similarly found no correlation with neonatal withdrawal and maternal methadone dose.12 Conversely, others have found a significant correlation between the severity of neonatal withdrawal and maternal methadone dose.14 15 These same authors, however, found no correlation found between neonatal serum concentrations and the severity of withdrawal in the infant.16 Doberczak and colleagues reported that clinical signs of withdrawal correlated with the rate of decline of the neonatal plasma drug concentrations during the first few days of life.14 Lastly, others have reported that the need for medical treatment to infants after birth is highest in those born to mothers receiving higher doses of methadone at delivery.15 17 These divergent findings have been used to argue for either weaning pregnant women to a low methadone maintenance dose14 15 17 to attempt complete maternal detoxification during pregnancy,18 or for maintaining the maternal methadone dose throughout the pregnancy to avoid fetal distress and death associated with maternal detoxification and withdrawal.1 11 12 13 14 16 17 The medical management of pregnant women addicted to opiates remains controversial.

The incidence of low birthweight and prematurity found in this study population seems similar to results reported before. The mean birthweight of the infants in this current study was 2804 g, with 33% of infants weighing less than 2500 g at birth. This supports previous observations of a 25 to 50% incidence of low birthweight associated with opiate exposure during gestation.18 20 23

Most of the infants in this study were exposed to multiple illicit drugs during gestation, and we acknowledge that the results in this paper can not be solely attributed to the effects of maternal methadone use. The problem of multiple drug exposure in utero continues to be a significant limitation in interpreting the results of clinical research in this area. As many investigators have shown, women who use illicit drugs during pregnancy frequently expose themselves to multiple drugs. In a recent study from the Netherlands, over 56% of women in a methadone programme had used other opiates, and 57% had used cocaine.24 Findings such as these preclude clinical investigations of infants exposed to only one illicit drug during pregnancy. Exposure to additional drugs may have affected the infants in this study by adding to their clinical signs. Infants exposed to methadone alone, or in combination with other drugs during gestation, might show qualitative and/or quantitative differences in the clinical signs of withdrawal as well as in other neonatal characteristics, as has been described before by others.21 25

In summary, hyperphagia was common among a group of infants withdrawing from methadone. By the second week of life, over half of the infants studied were eating more than 190 cc/kg/day. The hyperphagia seemed to be well tolerated by infants, and there were no increased adverse clinical findings associated with high intake in this relatively small number of infants. The infants who exhibited hyperphagia lost more weight in the first few days of life and also did not show greater weight gain than other infants when examined in the first month of life. We did not find that any infants had poor or ineffectual feeding. Further studies should be undertaken to determine whether it is beneficial for infants withdrawing from opiates to have medical management of hyperphagia, and whether this can be achieved without adversely affecting hospital stay.

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