Medications as a Source of Human Exposure to Phthalates.

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(Article begins on next page)
Phthalates are a group of multifunctional chemicals used in consumer and personal care products, plastics, and medical devices. Laboratory studies show that some phthalates are reproductive and developmental toxicants. Recently, human studies have shown measurable levels of several phthalates in most of the U.S. general population. Despite their widespread use and the consistent toxicologic data on phthalates, information is limited on sources and pathways of human exposure to phthalates. One potential source of exposure is medications. The need for site-specific dosage medications has led to the use of enteric coatings that allow the release of the active ingredients into the small intestine or in the colon. The enteric coatings generally consist of various polymers that contain plasticizers, including triethyl citrate, dibutyl sebacate, and phthalates such as diethyl phthalate (DEP) and dibutyl phthalate (DBP). In this article we report on medications as a potential source of exposure to DBP in a man who took Asacol (active ingredient mesalamine (mesalazine)) for the treatment of ulcerative colitis. In a spot urine sample from this man collected 3 months after he started taking Asacol, the concentration of monobutyl phthalate, a DBP metabolite, was 16,868 ng/mL (6.180 µg/g creatinine). This concentration was more than two orders of magnitude higher than the 95th percentile for males reported in the 1999–2000 National Health and Nutrition Examination Survey (NHANES). The patient’s urinary concentrations of monoethyl phthalate (443.7 ng/mL, 162.6 µg/g creatinine), mono-2-ethylhexyl phthalate (3.0 ng/mL, 1.1 µg/g creatinine), and monobenzyl phthalate (9.3 ng/mL, 3.4 µg/g creatinine) were unremarkable compared with the NHANES 1999–2000 values. Before this report, the highest estimated human exposure to DBP was more than two orders of magnitude lower than the no observable adverse effect level from animal studies. Further research is necessary to determine the proportional contribution of medications, as well as personal care and consumer products, to a person’s total phthalate burden. Key words: biomarkers, environmental health, medications, phthalates, reproductive health. Environ Health Perspect 112:751–753 (2004). doi:10.1289/ehp.6804 available via http://dx.doi.org/ [Online 29 January 2004]

Case Presentation

As part of an ongoing study on environmental agents and male reproductive health, male partners of subfertile couples who presented to the Vincent Burnand Andrology Laboratory at Massachusetts General Hospital (MGH; Boston, MA) provided a semen and a spot urine sample at their clinic visits. Men also completed a questionnaire including information on lifestyle factors, medical history, and medication use. The results of the relationship between phthalates and testicular function have been reported elsewhere (Duty et al. 2003a, 2003b). In this article we report on medication use as a likely source of dibutyl phthalate (DBP) exposure in one of the subjects from this ongoing study who had unusually high urinary concentrations of monobutyl phthalate (MBP), the primary metabolite of DBP.

The study was approved by the Harvard School of Public Health and MGH Human Subjects Committees. Information on medication use was collected with a nurse-administered questionnaire at the time of the visit to the clinic to provide the semen and urine samples. The subject was asked, “Have you recently (past few weeks) taken any medications? If yes, please provide types and date(s) last taken.”

Several phthalate monoesters were measured in a single spot urine sample collected in a sterile specimen cup. The analytical approach has been described in detail elsewhere (Blount et al. 2000a; Silva et al. 2003). Briefly, the determination of phthalate metabolites in urine involved enzymatic deconjugation of the metabolites from the glucuronidated form, solid-phase extraction, separation with HPLC, and detection by tandem mass spectrometry. Detection limits were in the low nanogram per milliliter range. 13C4-labeled internal standards were used to increase the precision of measurements. One method blank, two quality control samples (human urine spiked with phthalates), and two sets of standards were analyzed along with every 21 unknown urine samples. Analysts at the Centers for Disease Control and Prevention (CDC; Atlanta, GA) were blind to all information concerning the subjects. Creatinine adjustment was used to correct for urine dilution.

The case patient was in his early thirties and had an unremarkable occupational history, without any known workplace exposures. He had a medical history of ulcerative colitis and reported taking Asacol (Proctor & Gamble, Cincinnati, OH), which contains the active ingredient mesalamine (5-aminosalicylic acid), also known as 5-ASA or mesalazine in Europe. He took twelve 400 mg Asacol tablets daily during the 3 months before the collection of his urine sample. Although not fully understood, the anti-inflammatory action of mesalazine is thought to be through blocking cyclooxygenase and inhibiting prostaglandin production in the colon (Proctor & Gamble Pharmaceuticals 2000; Schroeder 2002). The Asacol delayed-release tablets are coated with methacrylic acid copolymer B (Eudragit-S; Rhône Gmbh & Co. KG, Darmstadt, Germany), which dissolves at pH 7, releasing mesalazine in the terminal ileum and beyond for topical anti-inflammatory action in the colon. Other inactive ingredients in Asacol tablets are colloidal silicon dioxide, edible black ink, iron oxide red, iron oxide yellow, lactose, magnesium stearate, polyethylene glycol, povidone, sodium starch glycolate, talc, and DBP (Proctor & Gamble Pharmaceuticals 2000).

The patient’s urinary MBP concentration was 16,868 ng/mL (6.180 µg/g creatinine), whereas monoethyl phthalate (MEP), mono-2-ethylhexyl phthalate (MEHP), and monobenzyl phthalate (MBzP) concentrations were 443.7 ng/mL (162.6 µg/g creatinine), 3.0 ng/mL (1.1 µg/g creatinine), and 9.3 ng/mL (3.4 µg/g creatinine), respectively.

Discussion

There is scientific and public concern about potential human health risks from exposure to phthalates, diesters of phthalic acid. These concerns stem from studies showing that a large proportion of the U.S. general population are exposed to phthalates (Blount et al. 2000b; CDC 2003), as well as from animal studies consistently showing that some phthalates are developmental and reproductive toxicants (Agarwal et al. 1985; Cater et al.

Address correspondence to R. Hauser, Occupational Health Program, Harvard School of Public Health, Building 1, Room 1405, 665 Huntington Ave., Boston, MA 02115 USA. Telephone: (617) 432-3326. Fax: (617) 432-0219. E-mail: rhauser@hsp.harvard.edu

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Human exposure to phthalates can occur via ingestion, inhalation, and dermal routes, as well as through parenteral exposure from medical devices containing phthalates. Currently, no human data on the proportional contribution of the various sources of phthalates to human body burden are available. Until recently, the primary exposure of the general population to phthalates was believed to result from ingestion of foods, especially fatty foods such as milk, butter, and meats. Recent data show that the low-molecular-weight phthalates (DEP, DBP, BBzP) may also be dermally absorbed and that the more volatile phthalates can be inhaled (ATSDR 1995, 2001).

The need for site-specific dosage medica-
tions has led to the use of enteric coatings (Ashford and Fell 1994; Marvola et al. 1999). Enteric-coated medications remain intact in the stomach and release the active ingredients of the underlying medication core into the lower part of the small intestine or in the colon. Enteric coatings generally consist of various polymers, such as cellulose acetate phthalate, cellulose acetate butyrate, ethylcellulose, polyvinyl acetate phthalate, and methacrylate copolymers. These polymer coatings are plasti-
cized with compounds such as triethyl citrate, dibutyl sebacate, and phthalates such as DEP and DBP (Frohoff-Hulsmann et al. 1999; Harris and Ghebre-Sellasse 1989). DEP and DBP metabolize to MEP and MBP, respec-
tively, which can be measured in urine. In addi-
tion to tablets and capsules, other medications and preparations may contain phthalates as plasticizers. Specifically, cellulose acetate–free films for transdermal use may contain DBP as a plasticizer (Rao and Diwan 1997).

The patient’s urinary phthalate concentra-
tions were compared with the concentra-
tions measured for the Second National Report on Human Exposure to Environmental Chemicals (CDC 2003; Table 1). CDC (2003) reported data on phthalate monooester levels from partici-
pants in NHANES 1999–2000. NHANES is an ongoing survey designed to measure the health and nutrition status of the civilian non-institutionalized U.S. population [National Center for Health Statistics (NCHS) 2003]. The samples in the present study and the NHANES 1999–2000 samples were analyzed by the same CDC laboratory.

Compared with the NHANES 1999–2000 data set, the patient’s urinary MEP, MEHP, MBzP levels were unremarkable. However, the patient’s concentration of MBP in urine was two orders of magnitude higher than the U.S. population 95th percentile for males reported in the NHANES 1999–2000 data set. To date, the highest estimated DBP exposure in a subset of the U.S. population, specifically women of childbearing age (Kohn et al. 2000), was considered more than two orders of magnitude lower than the lowest no observable adverse effect level (NOAEL) of DBP from animal studies (Mylchreest et al. 2002). Our results suggest that medications taken chronically can contribute to DBP expo-
sure that approaches the NOAEL measured in animal studies. Further research is necessary to determine whether any health risks are associated with human exposure to high levels of MBP.

Conclusion

In the present study, we identified an individual with a urinary MBP level two orders of magnitude higher than the U.S. population 95th percentile and linked this unusually high urinary MBP concentration with the use of a specific medication that contained DBP. However, because this is a case report on a single patient, replication of this finding in other populations is needed to definitively conclude that the medication was the main contributor to the very high urinary concentra-
tion of MBP. Additionally, further research is needed to determine the proportional con-
tribution of medications, as well as other con-
sumer products, to a person’s total phthalate burden. Adding to the complexity of apportioning sources of phthalates is that individu-
als may be exposed orally as well as by inhalation or dermal routes. When databases on consumer products and foods containing phthalates become available, these databases should be linked with levels of urinary phtha-
lates measured in ongoing epidemiologic

<table>
<thead>
<tr>
<th>Phthalate metabolite</th>
<th>Patient</th>
<th><strong>Males</strong></th>
<th><strong>Adults</strong></th>
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<tbody>
<tr>
<td>MEP</td>
<td>443.7 (162.6)</td>
<td>54 (120)</td>
<td>3,480 (1,940)</td>
</tr>
<tr>
<td>MBP</td>
<td>16,888 (6,180)</td>
<td>23.1 (17.0)</td>
<td>115 (63.6)</td>
</tr>
<tr>
<td>MBzP</td>
<td>3.3 (3.4)</td>
<td>17.7 (12.3)</td>
<td>108 (73.5)</td>
</tr>
<tr>
<td>MEHP</td>
<td>3.0 (1.1)</td>
<td>3.40 (2.76)</td>
<td>26.3 (21.6)</td>
</tr>
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Table 1. Concentrations [ng/mL (µg/g creatinine)] of MEP, MBP, MBzP, and MEHP in the patient compared with selected demographic groups in NHANES 1999–2000.
investigations. Until this time, individual case reports such as this one will provide clues on relevant sources of human exposure and their potential contribution to total body burden.

REFERENCES


