The Use of Psychotropic Medications During Breast-Feeding

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Objective: The authors reviewed the risks and benefits regarding the use of psychotropic medications during breast-feeding as they relate to the health and well-being of mothers and their infants. Strategies are discussed to limit infant exposure to a medication while effectively treating the nursing mother.

Method: A MEDLINE search of the literature since 1955 was conducted to determine the use of psychotropic medications in breast-feeding women. Search items included each of the categories of psychopharmacologic agents as well as each of the agents in association with nursing, breast-feeding, postpartum, lactation, and breast milk.

Results: No controlled studies on the safety of psychotropic medications in nursing mothers were found. Case reports and small case series for each of the different psychotropic medications serve as the basis for suggested treatment guidelines for the management of psychiatric illnesses in breast-feeding women. Thus, each case needs to be considered on an individual basis, with a thoughtful analysis of the risks and benefits of nursing and exposure of the infant to medication. The baseline clinical status of the infant should also be reviewed.

Conclusions: Women are vulnerable postpartum to psychiatric disorders and frequently face the need to decide whether to take psychotropic medications while breast-feeding. Should psychiatric medication be indicated, the parents should be provided with the available information regarding the effects of these medications on the neonate. In this way, an informed decision can be made. When psychotropic medication is used during breast-feeding, it is strongly recommended that the infant’s pediatrician be involved in monitoring the infant.

Breast milk offers many advantages to developing infants. The American Academy of Pediatrics endorses breast milk as the best and only source of nutrition necessary for the infant during the first 6 months of life (1). Breast-fed infants have lower rates of gastrointestinal disease, anemia, respiratory ailments, and otitis media (2, 3). In addition, breast-feeding provides a unique opportunity for bonding between infant and mother (4).

The vulnerability for psychiatric illness during the 3 months after delivery raises the possibility that psychotropic medications will be administered (5, 6). Issues to address during analysis of the risks and benefits of psychotropic use during breast-feeding include documented benefits of breast-feeding, the potential adverse impact of untreated maternal mental illness on infant attachment and cognitive and behavioral development, and the effects of untreated mental illness on the mother (7–13). This article reviews the literature on the various classes of psychotropic medications in order to provide the basis for educated treatment planning.

On average, nursing mothers produce 600 to 1,000 ml of milk daily. Factors affecting medication concentration in breast milk are pH, protein content, and lipid content. These vary throughout the postpartum period and at different times during a single feeding, resulting in marked concentration variations in milk aliquots. Milk pH ranges from 6.35 to 7.65 (14). Mature milk is produced approximately 2 weeks after birth. The higher lipid content of hind milk (the milk ejected during the second half of a feeding) makes it likely that the second half will have a higher concentration of maternal medication than the first half (fore milk). Other major factors affecting medication concentration in breast milk include lactose, serum albumin, lysozyme, approximately 30 enzymes, prolactin, and minerals such as calcium and phosphates (15).

The extent to which an infant is exposed to medication is affected by the rate of absorption into maternal circulation, diffusion from maternal circulation to breast milk, and absorption of the agent by the infant. Taking medication immediately after breast-feeding minimizes the amount present in milk and maximizes clearance before the next feeding (16). In vitro studies have demonstrated that full-term neonatal cytochrome P-450 activity is approximately one-half that found in adults (17). Each liver enzyme system matures at a different rate in the developing infant. Thus, different substrates are metabolized at different points in maturation. For example, while glucuronyl transferase activity is mature enough to metabolize bi-
Tricyclic antidepressants. Lipid-soluble agents can be 10–100 times more concentrated in the CSF than in serum (20) and may be higher in infants for a given plasma concentration compared to adults (21). Because body fat storage sites are limited in the neonate, central nervous system concentrations of lipid-soluble substances are greater in newborns than in older infants (22).

Despite the pharmacokinetic complexity and variability of the individual compartmentalized processes just described, most reports do not account for maternal, breast milk, or infant data. Frequently, infant exposure is estimated by measuring breast milk concentrations and assumed average daily milk consumption.

### Method

This review comprises an analysis of 95 studies and reports covering 32 psychotropic agents used by nursing mothers. Of the studies, 66 measured infant serum concentrations. In some cases, infant serum concentrations reflected exposure both in utero and through breast milk. The wide range of variability in sensitivity and reliability of a given assay precluded meaningful statistical grouping of individual case reports. Most reports lacked data for concentrations relative to time of dose administration. The interval between medication dose administration and infant feeding times is rarely reported. Finally, where infant clinical status is documented, behavioral evaluation is rarely based on standardized clinical assessment.

To avoid overinterpretation of data with so many limitations, and since the clinical significance of infant serum concentrations of medications is unclear, detailed discussion is reserved only for data that suggest a possible connection between drug and deleterious effects in offspring. Guidelines for the use of psychotropic agents in breast-feeding mothers will be suggested.

### Results

#### Antidepressants

Data from studies of breast-feeding and antidepressants are summarized in Table 1.

##### Tricyclic antidepressants

Infant serum levels of parent and active metabolites reported in studies of tricyclic antidepressants ranged from nondetectable to less than 28 ng/ml. Despite widely ranging milk-to-plasma ratios and variable infant serum levels of amitriptyline and nortriptyline (either as parent drug or as a metabolite of amitriptyline), there were no reports of adverse effects in the breast-fed infants. Similarly, no adverse effects were reported for the seven infants exposed to imipramine, the five infants exposed to desipramine, or the eight infants exposed to clomipramine.

For doxepin, two single case reports revealed milk-to-plasma ratios of parent and metabolite compounds that were close to or greater than one and measurable serum metabolite concentrations. Respiratory depression occurred in one case (38) but resolved 24 hours after discontinuation of nursing. Since the infant was not exposed to doxepin rechallenge through nursing, it is not possible to state definitively that respiratory compromise was due to

<table>
<thead>
<tr>
<th>Antidepressant</th>
<th>Number of Nursing Infants</th>
<th>Clinical Status of Infants</th>
<th>Adverse Effects</th>
</tr>
</thead>
<tbody>
<tr>
<td>Amitriptyline</td>
<td>7</td>
<td>4</td>
<td>3</td>
</tr>
<tr>
<td>Nortriptyline</td>
<td>18</td>
<td>0</td>
<td>18</td>
</tr>
<tr>
<td>Imipramine</td>
<td>7</td>
<td>0</td>
<td>7</td>
</tr>
<tr>
<td>Desimpramine</td>
<td>5</td>
<td>0</td>
<td>5</td>
</tr>
<tr>
<td>Clomipramine</td>
<td>(23, 33, 36, 37)</td>
<td>8</td>
<td>0</td>
</tr>
<tr>
<td>Doxepin</td>
<td>(38, 39)</td>
<td>2</td>
<td>0</td>
</tr>
</tbody>
</table>

* Respiratory depression that resolved 24 hours after discontinuation of nursing (38).

Data on a nursing infant exposed through nursing to fluoxetine also obtained from V.C. Hendrick (personal communication).

For six of the infants, the adverse effects were unconfirmed and unspecified and resolved spontaneously according to the mothers’ report (45). Colic was reported in three infants (40, 46). One infant experienced an episode of transient seizure-like activity at 3 weeks of age and episodes of unresponsiveness at age 4 months, with one episode of peripheral cyanosis at 5.5 months of age. All events were reported by the mother, and none were witnessed by medical personnel. Results of neurologic monitoring (EEG, brain imaging, developmental milestones) were all within normal limits up to 1 year of age. Maternal regimen of carbamazepine, buspirone, and fluoxetine was discontinued after postpartum day 21 (47).

Data on a nursing infant exposed through nursing to sertraline also obtained from Z. Stowe (personal communication).

“Uneasy sleep” observed at maternal dose of 40 mg/day that normalized when dose was reduced to 20 mg/day (60).

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<tr>
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<th>Adverse Effects</th>
</tr>
</thead>
<tbody>
<tr>
<td>Amitriptyline</td>
<td>(23–28)</td>
<td>7</td>
<td>4</td>
</tr>
<tr>
<td>Nortriptyline</td>
<td>(29–33)</td>
<td>18</td>
<td>0</td>
</tr>
<tr>
<td>Imipramine</td>
<td>(23, 33, 34)</td>
<td>7</td>
<td>0</td>
</tr>
<tr>
<td>Desipramine</td>
<td>(33, 35)</td>
<td>5</td>
<td>0</td>
</tr>
<tr>
<td>Clomipramine</td>
<td>(23, 33, 36, 37)</td>
<td>8</td>
<td>8</td>
</tr>
<tr>
<td>Doxepin</td>
<td>(38, 39)</td>
<td>2</td>
<td>0</td>
</tr>
</tbody>
</table>

Data on a nursing infant exposed through nursing to sertraline also obtained from Z. Stowe (personal communication).

For six of the infants, the adverse effects were unconfirmed and unspecified and resolved spontaneously according to the mothers’ report (45). Colic was reported in three infants (40, 46). One infant experienced an episode of transient seizure-like activity at 3 weeks of age and episodes of unresponsiveness at age 4 months, with one episode of peripheral cyanosis at 5.5 months of age. All events were reported by the mother, and none were witnessed by medical personnel. Results of neurologic monitoring (EEG, brain imaging, developmental milestones) were all within normal limits up to 1 year of age. Maternal regimen of carbamazepine, buspirone, and fluoxetine was discontinued after postpartum day 21 (47).

Data on a nursing infant exposed through nursing to sertraline also obtained from Z. Stowe (personal communication).

“Uneasy sleep” observed at maternal dose of 40 mg/day that normalized when dose was reduced to 20 mg/day (60).
doxepin. In the second case (39), no adverse effects were noted in the infant.

**Selective serotonin reuptake inhibitors (SSRIs).** Fluoxetine use by breast-feeding mothers has been evaluated in 11 published reports of 190 breast-fed infants (Table 1). In 101 cases, infant sera concentrations were not tested. Of the remaining cases, levels of parent compound and metabolite varied from nondetectable to a single report of 340 ng/ml (40). There were no clear associations between levels in infant sera, maternal dose, and infant age. No adverse effects were noted in 180 of the 190 cases. One study of four infants reported normal neurobehavioral development to 1 year of age (44). Of some concern is a single case of a 6-week-old infant with serum concentrations of fluoxetine that were comparable to maternal concentrations (40); adverse effects reported included excessive crying, decreased sleep, vomiting, and diarrhea that dissipated upon discontinuation of nursing. Adverse effects reported in other studies were transient and by maternal report (45, 46) or were confounded by multiple medications (47). Although a retrospective study indicated poorer weight gain for fluoxetine-exposed infants than age-matched control subjects, weights in the exposed infants were not statistically below the national mean (48). These data highlight the importance of professional monitoring of the clinical status of nursing infants rather than relying on laboratory measurements of serum levels and maternal observations.

Fifteen published reports documented the use of the remaining SSRIs (fluvoxamine, paroxetine, sertraline, and citalopram) in breast-feeding mothers. For sertraline, infant serum levels were nondetectable or less than 5 ng/ml for parent compound; the metabolite concentrations that were measured in four reports were less than 10 ng/ml. In the six paroxetine reports, serum levels were measured in 27 of the 37 breast-fed infants: concentrations ranged from nondetectable for 24 of the infants to less than 20 ng/ml in the remaining three. Citalopram and desmethylcitalopram were measured in infant serum in two of the citalopram case reports (60, 61); concentrations ranged from 2.3 ng/ml to 12.7 ng/ml, and metabolite concentrations were nondetectable. Serum levels were not reported for fluvoxamine. No clear adverse effects were noted in the 49 infants exposed to sertraline, the 37 exposed to paroxetine, or the two exposed to fluvoxamine.

**Other antidepressants.** Although five infants exposed through nursing to bupropion, mianserin, or venlafaxine did not experience apparent adverse effects, more data are needed before conclusions can be made regarding their safety in breast-feeding. Since infant serum concentrations and clinical status were not reported for the novel antidepressant trazodone (66) or the reversible monoamine oxidase inhibitor moclobemide (67), the safety of these agents in exposed nurslings is unknown.

**Anxiolytics**

Studies with data on anxiolytic exposure in nursing infants are summarized in Table 2.

For 12 of 13 infants whose mothers were on regimens of clonazepam, exposure was in utero as well as through breast-feeding. In one case of an infant exposed both in utero and through nursing to clonazepam (infant serum level <5 ng/ml), persistent cyanosis was noted at delivery and for the first 10 postpartum days (71). By day 10 breathing normalized, and neurodevelopment appeared to be normal at 5 months (71). For the remaining 11 infants for which clinical status was described, no adverse effects were noted (37).

Only three of 10 infants exposed to diazepam through nursing had data for serum concentrations of parent and metabolite compounds, which varied from nondetectable to 243 ng/ml. Serum levels of both parent and metabolite compounds were lowest in the oldest nursling, an infant of 1 year of age (72), which suggests that older infants have improved metabolic capacity. Clinical status was reported in five infants, with no adverse effects in three (76) and normal development recorded in a fourth despite some apparent sedation (73). In another case report, lethargy and weight loss was reversed following discontinuation of nursing (75). For the 11 infants exposed to oxazepam or temazepam (and its metabolite, oxazepam), no adverse effects were noted.

The remaining reports of medication effects in infants exposed to anxiolytics through breast milk involved treatment with alprazolam, lorazepam, or zolpidem. Infant serum concentrations were not reported, and in the one case where clinical status was reported (69), alprazolam withdrawal appeared upon medication tapering.

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**Table 2. Medication Exposure Effects in Nursing Infants of Mothers Treated With Anxiolytics**

<table>
<thead>
<tr>
<th>Anxiolytic Class and Medication</th>
<th>Number of Nursing Infants</th>
<th>Clinical Status of Infants After Exposure</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Reported</td>
<td>No</td>
</tr>
<tr>
<td>Benztiazepines</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Alprazolam (68, 69)</td>
<td>8</td>
<td>7</td>
</tr>
<tr>
<td>Diazepam (37, 70, 71)</td>
<td>13</td>
<td>1</td>
</tr>
<tr>
<td>Lorazepam (72–76)</td>
<td>10</td>
<td>5</td>
</tr>
<tr>
<td>Oxazepam (78)</td>
<td>4</td>
<td>4</td>
</tr>
<tr>
<td>Temazepam (79)</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>Nonbenztiazepines</td>
<td>10</td>
<td>0</td>
</tr>
<tr>
<td>Zolpidem (80)</td>
<td>5</td>
<td>5</td>
</tr>
</tbody>
</table>

a Irritability, sleep disturbance; symptoms worsened at discontinuation of medication (69).

b Cyanosis, decreased respiration rate, decreased tone, and lethargy in infant with both pre- and postnatal exposure. Symptoms resolved at 10 days, and normal milestones were achieved at age 5 months (71).

c Sedation (73) and lethargy and weight loss that was reversed after nursing was discontinued (75).
TABLE 3. Medication Exposure Effects in Nursing Infants of Mothers Treated With Antipsychotics

<table>
<thead>
<tr>
<th>Antipsychotic Class and Medication</th>
<th>Number of Nursing Infants</th>
<th>Not Reported</th>
<th>No Adverse Effects</th>
<th>Adverse Effects</th>
</tr>
</thead>
<tbody>
<tr>
<td>Phenothiazines</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Chlorpromazine (81–84)</td>
<td>14</td>
<td>0</td>
<td>10</td>
<td>4^ab</td>
</tr>
<tr>
<td>Trifluoperazine (81)</td>
<td>2</td>
<td>0</td>
<td>2</td>
<td>0</td>
</tr>
<tr>
<td>Chlorprothixene (85)</td>
<td>2</td>
<td>0</td>
<td>2</td>
<td>0</td>
</tr>
<tr>
<td>Perphenazine (86)</td>
<td>1</td>
<td>0</td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td>Butyrophenones</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Haloperidol (81, 87, 88)</td>
<td>11</td>
<td>1</td>
<td>7</td>
<td>3^b</td>
</tr>
<tr>
<td>Atypical</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Clozapine (89)</td>
<td>1</td>
<td>1</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Olanzapine (90)</td>
<td>3</td>
<td>0</td>
<td>3</td>
<td>0^c</td>
</tr>
</tbody>
</table>

^a Lethargy in one infant (83).
^b Declining scores on sequential developmental testing in three infants exposed to a combination of chlorpromazine and haloperidol (81).
^c Cardiomegaly at birth in one infant exposed in utero. Nursing was stopped on postpartum day 7; jaundice and sedation persisted.

Antipsychotics

Ten reports have addressed infant exposure to antipsychotic agents through nursing (Table 3). Of 34 clinical status reports of infants exposed to antipsychotics through breast milk, 25 showed no adverse effects. Lethargy was noted in one chlorpromazine-exposed infant (83). Delayed developmental testing at 12–18 months of age was reported for three infants exposed to a combination of haloperidol and chlorpromazine; only one of the three infants had detectable serum levels of neuroleptic (81). In a single case report of clozapine exposure during breastfeeding (89), milk-to-plasma ratios for clozapine revealed that it was concentrated in milk, but since there were no data regarding infant serum levels, the relevance of this is unclear. The early data for olanzapine come from one report of three infants exposed both in utero and through nursing (90); no adverse effects attributable to olanzapine ingestion through breast-feeding were noted.

Mood Stabilizers

A recent study indicated that postpartum bipolar women who discontinued lithium before pregnancy were almost three times as likely to have a recurrence in the first postpartum month as bipolar control subjects (91). Twenty-eight reports of the use of mood stabilizers in nursing mothers include 79 infants exposed to lithium, carbamazepine, valproate, or lamotrigine through breastfeeding (Table 4). Of 25 cases of carbamazepine exposure in nursing infants, there were two cases of transient hepatic dysfunction (93, 94). The etiology of reported instances of seizure-like activity (47) and drowsiness, irritability, and abnormal crying (99) was impossible to ascribe because of the exposure to a combination of agents through breast milk.

Of 36 valproate-exposed infants, no adverse effects were noted for 19, and clinical status was not reported for 16.

TABLE 4. Medication Exposure Effects in Nursing Infants of Mothers Treated With Mood Stabilizers

<table>
<thead>
<tr>
<th>Mood Stabilizer</th>
<th>Number of Nursing Infants</th>
<th>Not Reported</th>
<th>No Adverse Effects</th>
<th>Adverse Effects</th>
</tr>
</thead>
<tbody>
<tr>
<td>Carbamazepine</td>
<td>(47, 92–100)</td>
<td>25</td>
<td>0</td>
<td>16</td>
</tr>
<tr>
<td>Valproic acid (92, 101–109)</td>
<td>36</td>
<td>16</td>
<td>19</td>
<td>1^b</td>
</tr>
<tr>
<td>Lithium (110–115)</td>
<td>13</td>
<td>8</td>
<td>4</td>
<td>1^c</td>
</tr>
<tr>
<td>Lamotrigine (116–118)</td>
<td>5</td>
<td>0</td>
<td>5</td>
<td>0</td>
</tr>
</tbody>
</table>

^a Drowsiness, irritability, and high-pitched crying in a 10-week-old infant of an epileptic mother exposed to multiple agents; symptoms were temporally associated with antihistamine (clemastine) exposure (99). Hyperexcitability in two infants and poor feeding in three infants (100). Transient increase in liver function test results in one infant, with normal development to 6 months of age (93). Hepatic cholestasis in one infant that resolved after nursing was discontinued (94). One infant experienced an episode of transient seizure-like activity at 3 weeks of age and episodes of unresponsiveness at age 4 months, with one episode of peripheral cyanosis at 5.5 months of age. All events were reported by the mother; no events were witnessed by medical personnel. Results of neurologic monitoring (EEG, brain imaging, developmental milestones) were all within normal limits up to 1 year of age. Maternal regimen of carbamazepine, buspirone, and fluoxetine was discontinued after postpartum day 21 (47).

^b Thrombocytopenia and anemia at 3 months of age that reversed upon discontinuation of nursing (106).

^c ECG changes in an infant exposed both in utero and through nursing to age 5 days that resolved after lithium was discontinued (112).

Thrombocytopenia and anemia in an exposed 3-month-old infant reversed upon cessation of nursing (106).

Of the 13 infants exposed to lithium, serum concentrations were found to be very high. One infant with a congenital heart murmur, cyanosis, and hypotonia had been exposed to lithium both in utero and through breast milk (112). There were no adverse effects in four other infants, and clinical status was not reported for eight. No adverse effects were reported in five cases of lamotrigine exposure in nursing infants. We found no published data on gabapentin exposure through nursing.

Treatment Guidelines

General Issues

The decision to breast-feed while taking psychotropic medications is complicated. Considerations include known benefits of breast-feeding to infant and mother, wishes of the mother, risk of infant exposure to the medication, and the possibility that a severely depressed, anxious, or psychotic mother may forego treatment rather than give up breast-feeding.

Parents faced with this decision should be educated about possible side effects in their infants. As the association between infant drug levels and clinical status is unclear, clinical monitoring seems to be the best approach to minimize the risk to infants. Before initiating maternal medication, a pediatric evaluation should assess infant baseline behavior, sleep, feeding, and alertness. Metabo-
Am J Psychiatry 158:7, July 2001 1005

The emerging data for SSRIs is important, since this class comprises the most commonly prescribed antidepressants. The data for fluoxetine is variable and somewhat difficult to interpret. The few adverse effects were generally transient and not verified by medical personnel or objective tests. The lack of adverse effects in 180 fluoxetine-exposed infants justifies its continued use if it has been prescribed antenatally or if there is a history of preferential efficacy with this agent. A previous report suggests that there is little central serotonin reuptake inhibition in infants breast-fed by mothers taking sertraline (120). Although the gradient of antidepressants such as paroxetine tends to increase from fore milk to hind milk (52), the extent to which this is reflected in infant serum depends on the extent of nursing at any single feeding. Furthermore, the lack of adverse effects in 86 breast-fed infants exposed to sertraline or paroxetine and the uniformly low or non-detectable infant serum levels in these infants suggest that these are good choices for nursing mothers with postpartum depression. In the case of sertraline, serum levels peak in infants between 7 and 11 hours after maternal dosing; refraining from nursing during this time period may significantly reduce infant exposure (57). Similar peak levels have not been reported for paroxetine or fluoxetine. As the data for fluvoxamine and citalopram are sparse, these agents are not primary treatment options for breast-feeding mothers.

Postpartum Bipolar Disorder

Treatment of bipolar disorder invariably requires mood stabilizers. Maternal use of lithium and carbamazepine has been associated with serious difficulties in nurseries. There has been a single recorded case of serious blood abnormalities following exposure to sodium valproate through breast milk (106), and this agent has been associated with hepatotoxicity when directly administered to infants (121, 122). Therefore, when administering valproate to breast-feeding mothers, pediatric clinical status, liver enzymes, and platelets should be carefully monitored. Lithium increases the risk of thyroid dysfunction, cyanosis, poor muscle tone, and ECG changes in infants (19, 114, 123). Because renal clearance is decreased in infants up to at least 5 months of age, use of lithium during breast-feeding is not advisable. Nevertheless, for the bipolar patient who invariably decompenses when lithium is discontinued, use by a nursing mother may be reasonable so long as infant clinical status is carefully monitored and serum concentrations of lithium in the infant are followed.

While the American Academy of Pediatrics suggests that carbamazepine exposure appears safe for breast-feeding infants, two cases of hepatic dysfunction (93, 94) and one case of transient seizure-like activity (47) suggest that it is advisable to monitor liver enzymes, bilirubin, and WBC counts frequently and to assay for carbamazepine levels in exposed infants. There are limited (although reassuring) data on lamotrigine exposure through breast milk and no...
data for the use of gabapentin by nursing mothers. Since these agents are not first-line medications for bipolar disorder, their use in breast-feeding should be reserved only when no other alternatives remain for effective treatment.

**Postpartum Anxiety Disorders**

Postpartum anxiety may be ameliorated with nonpharmacologic interventions (e.g., cognitive behavioral therapy, progressive relaxation techniques, environmental stress reduction). When possible, securing a nanny is helpful to reduce sleep deprivation. Occasional low doses of short-acting benzodiazepines such as temazepam or oxazepam are probably safe (124). Reports of adverse side effects for diazepam and withdrawal with alprazolam suggest that these should not be a first option if a benzodiazepine is administered to a breast-feeding mother. Tricyclics (with the exception of doxepin) or SSRIs are also good options for the treatment of panic disorder in breast-feeding mothers.

**Postpartum Psychosis**

Postpartum psychotic symptoms should be treated with antipsychotics. Psychotic women may be so dysfunctional that they are unable to breast-feed their infants. In cases of infant exposure to antipsychotics through breast-feeding, infant clinical status should be regularly monitored for antipsychotic side effects such as somnolence, muscle rigidity, or tremors. The reports of developmental decline in infants nursed by mothers receiving antipsychotic combinations suggest that a monotherapeutic regimen should be maintained (81). Clozapine-induced fatal agranulocytosis in adults and the lack of data on infant clinical status make it imprudent for mothers treated with this agent to breast-feed their infants. Of note, conception is possible while breast-feeding, and the method of birth control should be documented during the risk-benefit discussion.

**Conclusions**

Women with postpartum psychiatric disorders often face the dilemma of whether or not to use psychotropic medication while continuing to breast-feed their infants. In such cases, it is important to safeguard the mental health of the mother while at the same time optimizing the emotional and physical well-being of the infant. All psychotropic agents enter breast milk. While these medications pass into infant circulation to varying degrees, a clear relationship between concentration of these medications (and their active metabolites) on infant physiology, behavior, and development is unknown. Therefore, rather than basing decisions regarding the use of medications during breast-feeding on serum levels, it is prudent to carefully monitor the clinical status of infants who are breast-fed by mothers taking psychiatric medications. In the event that the parents or pediatrician become concerned that the infant’s behavior, activity level, or achievement of developmental milestones may be related to medication exposure, serious consideration should be given to weaning. Studies are needed that relate measured levels of medications and metabolites in the sera of breast-fed infants to clinical status and that carefully note infant age and maternal and infant weights at the time of blood sampling. Accurate assessments may then be made of comparative levels of drug exposure on a per-kilogram basis in infant and mother. Clinical status and behavior of these infants should be carefully observed by using standardized pediatric instruments. In this way, more definitive conclusions may be made about the clinical significance of infant serum concentrations and infant daily exposure.

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PSYCHOTROPICS AND BREAST-FEEDING


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