Pharmacotherapy during pregnancy, childbirth and lactation: points and principles to consider (a 2015 update)

G Schellack,* N Schellack,b M Krielc

*Clinical Research, South Africa
bDepartment of Pharmacy, Sefako Makgatho Health Sciences University, Pretoria, South Africa
cFemina Hospital, Netcare, Pretoria, South Africa
*Corresponding author, email: natalie.schellack@smu.ac.za

Abstract

Pregnancy, childbirth and lactation pose unique challenges in terms of drug therapy. The pregnant mother and her unborn child are exceptionally vulnerable from a physiological, clinical and ethical standpoint. This warrants careful consideration with respect to a number of important aspects, which could firstly influence the decision to opt for drug therapy, and secondly, could influence the specific agent selected for each indication. The US Food and Drug Administration has introduced changes to the content and format of information presented in prescription drug labelling to assist healthcare providers when assessing benefit versus risk, and in the subsequent counselling of pregnant women and nursing mothers who need to take medication. This change came into effect at the end of June 2015. This article provides an overview of these important aspects.

Keywords: embryo, foetus, lactation, neonate, pregnancy

Introduction

Obstetrics and neonatology drug therapy, also referred to as pharmacotherapy, represents a major medical challenge to practising healthcare workers. This is because drug therapy, i.e. the use of medication to attain a certain clinical outcome, poses a significant risk during vulnerable periods of the human reproductive cycle.

Vulnerable periods of the human reproductive cycle

Fertilisation of the ripened ovum and the attainment of complete implantation

The two greatest risks during fertilisation of the ripened ovum and the attainment of complete implantation are spontaneous abortion or the reabsorption of the products of conception,1,2 both of which would probably go unnoticed. This period lasts for approximately two weeks (Figure 1).

Unborn child

Drugs may cross the placental barrier and reach the systemic blood circulation of the unborn child.

Embryonic development and organogenesis

The embryonic period lasts until the end of the eighth week after fertilisation, during which the foetus is exceptionally vulnerable to structural abnormalities.1,4

Foetal development and maturation

During the second and third trimester, drugs usually only affect the growth and maturation of the foetus, since organogenesis is completed by the end of the embryonic period, although the development of the external genitalia continues into the second trimester, and the development of the central nervous system is an ongoing process for the duration of the pregnancy and beyond1,2,5,6 (Figure 1).

Mother and infant

The birthing process

Drugs used to manage the intrapartum period may have a direct effect on the infant during and directly after the birth, e.g. the result could be a suppressed level of consciousness or even apnoea in the newborn infant when a general anaesthetic agent is used to facilitate a Caesarean section.7

Breastfeeding

Drugs may be excreted in the mother’s breast milk.5

The development of the newborn child

Newborn infants are vulnerable, whether or not they are of normal gestational age and birthweight. Various factors contribute to the challenges created by pharmacotherapy in infants. These include their smaller body size, higher percentage of body water, the fact that liver biotransformation, for example, is slower in the neonate, and that the neonate also displays a slower rate of renal elimination of certain drugs. Premature infants have an even higher percentage of body water than neonates.3,6

Therefore, drugs can exert a potentially harmful effect on the unborn infant during any stage of the pregnancy, albeit in varying degrees.2,3
Pharmacology is a vast and complex subject. Therefore, this article only focuses on the principles of pharmacotherapy with respect to how they apply to pregnancy, childbirth and lactation. The basic pharmacokinetic and pharmacodynamics terminology referred to in this chapter is summarised in Table 1.

**Table 1: Basic pharmacokinetic and pharmacodynamic terminology**

<table>
<thead>
<tr>
<th>Term</th>
<th>Definition</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pharmacokinetics</td>
<td>The study of the kinetics of drug absorption, distribution, metabolism and elimination or excretion in humans and animals. In other words, it describes what happens to drugs in the body</td>
</tr>
<tr>
<td>Pharmacodynamics</td>
<td>The study of the biological effects resulting from an interaction between drugs and biological (body) systems. In other words, it describes the effect that drugs have on the body</td>
</tr>
<tr>
<td>Biopharmaceutical properties</td>
<td>The relationship between the physical and chemical properties of a drug, in a specific dosage form, and the pharmacological, toxicological or clinical effects which are observed following its administration. This information can be used to optimise drug availability at the site of action of the drug</td>
</tr>
<tr>
<td>Therapeutic drug monitoring</td>
<td>The mathematical relationship between a drug-dosing regimen and its resulting serum concentrations (pharmacokinetics)</td>
</tr>
<tr>
<td></td>
<td>The relationship between the drug concentration at the site of action of the drug and the resultant pharmacological response (pharmacodynamics)</td>
</tr>
<tr>
<td></td>
<td>The use of serum drug concentrations to optimise drug therapy in individual patients, in conjunction with their clinical status and response to therapy</td>
</tr>
<tr>
<td>Apparent volume of distribution</td>
<td>The apparent volume into which a drug distributes in the body's fluid compartments at equilibrium. This is the volume into which the specific drug dosage will need to be dissolved in order for it to reach the same concentration as it does in the plasma</td>
</tr>
<tr>
<td>Clearance</td>
<td>The volume of body fluid which is totally cleared of the drug per unit of time</td>
</tr>
<tr>
<td>Systemic bioavailability</td>
<td>Defined as the fraction of an orally administered dosage which reaches the systemic blood circulation of the patient. It reflects the extent of absorption and presystemic elimination</td>
</tr>
<tr>
<td>Elimination half-life</td>
<td>This is the time it takes for the drug's plasma concentration to be reduced by 50%</td>
</tr>
<tr>
<td>Plasma steady-state concentration</td>
<td>Implies a stable plasma drug level, or plateau concentration, during which the drug's rate of absorption is equal to its rate of elimination, i.e. the drug's &quot;input&quot; equals its &quot;output&quot;. It takes approximately 4–5 half-lives before the steady state is reached</td>
</tr>
<tr>
<td>Absorption</td>
<td>This may be defined as the process by which a drug proceeds from its site of administration to the central blood circulation (the site of measurement within the body). This process is not restricted to oral administration only, but is equally applicable to events which follow other routes of administration, i.e. intramuscular and subcutaneous injection and rectal administration. However, intravenously injected drugs enter the bloodstream directly, and therefore absorption is not required to take place</td>
</tr>
</tbody>
</table>

**Pharmacotherapy during pregnancy, childbirth and lactation**

Pharmacotherapy during pregnancy, childbirth and lactation may be required for a number of reasons, including acute illness or trauma during the course of a pregnancy, and chronic illness or disability, including a few conditions which are of particular importance in this setting. These are human immunodeficiency
It is of vital importance in these settings to carefully weigh up the possible benefits and risks of pharmacotherapy of not treating the condition at all against the possible outcomes for both mother and child. The decision to opt for drug therapy should always be a sound and rational one. The outcomes which the clinician aims to achieve should be realistic. It should be considered what the available drugs, under the specified conditions, may be reasonably expected to achieve when given to the patient in question. Furthermore, treatment cannot necessarily be interrupted, postponed or avoided altogether merely because a female is pregnant or breastfeeding. This poses a complex clinical challenge for which expert opinion merely because a female is pregnant or breastfeeding. This to the patient in question. Furthermore, treatment cannot be considered what the available drugs, under the specified conditions, may be reasonably expected to achieve when given to the mother, as part of a foetal therapy regimen, in a few instances.

The following aspects need to be considered.

**Physiological changes during pregnancy that may affect drug action and kinetics**

Certain physiological changes during pregnancy have implications for drug therapy, and may affect any of the four basic kinetic processes of absorption, distribution, metabolism and elimination or excretion. Therefore, the following aspects could alter the way in which drug molecules are handled by the body, i.e. alter their pharmacokinetic profile.

1. Illness, emergency or other condition requiring drug treatment
2. Decision to treat YES or NO
3. Consider what the available drugs, under the specified conditions, may be reasonably expected to do when given to the patient in question
4. Drugs given to a pregnant or lactating female
5. Effects on the mother, as well as the unborn and breastfed child
   - Positive or negative

**Drug distribution may be affected due to the increased plasma volume which accompanies pregnancy, and which may result in an increased volume of distribution of certain drugs. Furthermore, pregnancy results in a decreased blood albumin level, which can result in an increased fraction of free drug molecules. This is especially significant in the case of drugs which are highly protein bound, also increasing their volume of distribution, and altering other kinetic properties. Only the free, unbound fraction is able to cross the placental barrier. The higher fraction of free drug molecules in these instances implies that there are higher levels of the active drug in circulation for both mother and unborn child, and this increases the likelihood of drug toxicity.**

**Altered liver functioning may affect the plasma concentrations of drugs which follow hepatic metabolism.**

The increased plasma volume, in turn increases cardiac output, renal blood flow and glomerular filtration rate. This can increase the renal excretion of drugs which are significantly eliminated via this route. Drugs and their metabolites are also excreted in breast milk.

**Figure 2: Pharmacotherapeutic decision-making during pregnancy, childbirth and lactation**
Drugs may be toxic to the developing embryo and foetus. The first trimester of pregnancy, i.e. the stage of embryonic development and organogenesis, is of particular importance since the teratogenic effects of certain drugs influence the normal development of the unborn child on a structural or functional level (Figure 1). A teratogen is a drug, or other chemical substance, which may affect normal embryonic development and cause recognisable congenital (birth) defects. The expectant mother may not even be aware of the fact that she is carrying a developing embryo during the very early stages of pregnancy. She may unknowingly harm her unborn child through the careless or indifferent use of drugs. Therefore, this is an important topic to include in preconception care. However, many pregnancies are unplanned or unexpected, implying that preconception care would not have been rendered at all. Sufficient precaution needs to be taken in the form of patient education, and highly effective contraceptive methods instituted, when treating women of childbearing age or potential in the event of drugs which have been shown to pose a significant risk to an unborn child, or drugs with insufficient evidence of relative safety during pregnancy. Some drugs even need to be avoided in men who may father children while being treated with them. 

Cross-placental transfer of drug molecules (and their metabolites)

Drug treatment during pregnancy inadvertently implies that the unborn child will be exposed to, either the effects of the drug on the mother, the direct effects of the drug on the embryo or foetus, or a combination of both. The placenta acts as a barrier between the circulatory systems of the mother and child throughout the duration of the pregnancy. However, this barrier is not very efficient in terms of drug molecules. This implies that when such molecules enter the maternal blood circulation, they have the potential of crossing this barrier, and entering the foetal circulation as well. Lipid-soluble drugs are capable of crossing the placenta via simple diffusion. Most water-soluble drugs can also cross the placenta because of the relative inefficiency of the barrier. Heparin is an exception. The four characteristics of drug molecules which are most likely to cross the placenta are highlighted in Table 2.

Excretion in breast milk

During lactation, drugs may pass from the bloodstream to the breast milk, especially if they are lipid soluble or basic drugs. (Basic drugs tend to ionise in the breast milk since it is more acidic than blood). Drugs can also pass from the bloodstream to the breast milk if they contain water-soluble molecules with a relative molecular mass of less than 100 daltons (Da). The Da is the unit of measurement used to express molecular mass, and is indicative of the size of a molecule.

Drug safety during pregnancy and lactation

There are limited data at disposal on the actual safety profiles of many drugs during pregnancy and lactation. There are many reasons for this, including the difficulties in conducting suitable clinical trials, both on ethical and technical grounds. However, many women still take 3–5 medications during their pregnancy. The prescriber must make pharmacotherapeutical decisions that pertain to each individual patient. Due consideration must be given to their unique maternal, foetal and infant risk-benefit profile. This highlights the importance of healthcare providers having access to a system which enables them to make better and safer drug choices for pregnant and lactating patients.

Towards the end of 2014, the US Food and Drug Administration (FDA) announced an amendment to the content and format of information presented in prescription drug labelling. This refers to the new Pregnancy and Lactation Labeling Final Rule. The FDA’s goal with this amendment is to provide pregnant or breastfeeding mothers, as well as their healthcare providers, with the best possible information on the use of available agents during the various stages of pregnancy and lactation. These amendments came into effect on 30 June 2015.

The old FDA requirements consisted of the five-letter categories, A, B, C, D and X. These categories attempted to broadly identify risk, while using a specific medication. This simplified system did not provide sufficient information to allow healthcare providers to make the most informed decisions on drug choices for pregnant or lactating mothers. This system was misinterpreted as a grading system, and the lack of information presented a potential risk to both mothers and their foetuses. The new rule (Figure 3) consists of up-to-date and well-organised information to provide healthcare providers with better information during the different stages of pregnancy.

This labelling also assists healthcare providers during the counselling of patients on the correct use of their medication during pregnancy and lactation.

The FDA also amended its requirements in terms of the relevant labelling subsections and their content. Therefore, the subheadings of “Pregnancy”, “Labor and delivery” and “Nursing mothers” have been replaced or updated with “Pregnancy (to include labour and delivery)”, “Lactation (to include nursing mothers)” and “Females and males of reproductive potential”. Both the pregnancy and lactation sections are further divided into a risk summary, clinical considerations and data section. The guideline also assists with the labelling of new products, and reversing or amending existing labelling.

Information on the use of a specific drug during all three trimesters of pregnancy is also provided in the “Pregnancy” section.
subsection. This subsection also includes registers in which data are collected and maintained on how the said drug can affect pregnancy during the certain trimester, or after birth and into childhood. The new label also includes information on physiological changes during pregnancy, and how these affect dosage and risk during the stages of pregnancy, as well as the postpartum period. Information is provided on maternal adverse reactions, foetal or neonatal adverse reactions, and the effects of drugs during delivery or labour. Disease-associated risks to mothers, embryos and foetuses are also included.

The “Lactation” subsection provides more information on whether or not the active ingredient will have an effect on the infant during breastfeeding. The new “Females and males of reproductive potential” subsection contains information on how certain medications can affect fertility, pregnancy testing and contraception.

It is of great importance to consult suitable drug references when prescribing medicines to pregnant or lactating mothers. Known teratogens should obviously be avoided during pregnancy. However, situations may arise in which the benefits of treating the mother with a certain drug may outweigh the possible harm which the drug may or may not do. The package inserts and patient information leaflets of registered medicines should also be consulted for specific information on the use of such agents in the treatment of pregnant and lactating women, or even in women who are of childbearing potential, but who are not yet pregnant.

Well known examples of teratogenic drugs include:

- Isotretinoin, used in the treatment of severe forms of acne.
- Methotrexate, an antineoplastic and immunosuppressant agent.
- Antiepileptic agents, phenytoin and valproate (the latter is also an antiretroviral agent).
- Warfarin, an oral anticoagulant.

Using ethanol (drinking alcohol) during pregnancy may cause foetal alcohol syndrome. There are many more examples (Table 3).

### Table 3: Examples of known teratogenic drugs to avoid during pregnancy

<table>
<thead>
<tr>
<th>Teratogen</th>
<th>Effect</th>
</tr>
</thead>
<tbody>
<tr>
<td>Anticonvulsants</td>
<td>Valproate and carbamazepine are associated with neural tube defects. Phenytoin may cause malformations in the central nervous system and adversely affect foetal growth</td>
</tr>
<tr>
<td>Anticoagulants</td>
<td>Warfarin is associated with haemorrhage in the foetus, as well as malformations in the central nervous and and skeletal systems</td>
</tr>
<tr>
<td>Antihypertensive agents</td>
<td>Angiotensin-converting enzyme inhibitors cause renal damage, and may restrict normal growth patterns in the unborn child</td>
</tr>
<tr>
<td>Antineoplastic agents</td>
<td>Antineoplastic agents are linked to a high risk of multiple congenital malformations</td>
</tr>
<tr>
<td>Ethanol (drinking alcohol)</td>
<td>The effects of ethanol may be cumulative, and include foetal alcohol syndrome, abnormal functioning of the central nervous system and disturbances in behaviour</td>
</tr>
<tr>
<td>Isotretinoin</td>
<td>Isotretinoin is linked to a very high risk of multiple congenital malformations</td>
</tr>
<tr>
<td>Misoprostol</td>
<td>Misoprostol is associated with malformations of the central nervous system and limbs</td>
</tr>
<tr>
<td>Neuroleptic drugs</td>
<td>Lithium is associated with congenital defects of the cardiovascular system</td>
</tr>
<tr>
<td>NSAIDs</td>
<td>NSAIDs are linked to premature closing of the ductus arteriosus</td>
</tr>
<tr>
<td>Tetracycline</td>
<td>Tetracycline is associated with malformations of the teeth (including permanent discoloration) and bone</td>
</tr>
<tr>
<td>Thalidomide</td>
<td>Thalidomide is associated with malformations of the internal organs and limbs</td>
</tr>
</tbody>
</table>

In addition, the following drugs are associated with withdrawal symptoms in the newborn infant:

- Barbiturates
- Benzodiazepines
- Opioid analogs
- Tricyclic antidepressants

NSAIDs: nonsteroidal anti-inflammatory drugs

* Drugs that are not listed here are not necessarily safe to use

Figure 3: A comparison between the previous prescription drug labelling requirements and the new Pregnancy and Lactation Labeling Final Rule of the US Food and Drug Administration (subsections 8.1–8.3)
Many different factors determine the choice and possible outcomes of drug therapy during pregnancy, childbirth and lactation.

These factors include:
- Age, obstetric history and gestation.
- Physical characteristics, e.g. size and body mass index.
- Diet and nutritional status.
- Gene pool and genetic factors.
- Previous responses and reactions to drug treatment, including allergic reactions and anaphylaxis.
- Other drugs already in use, which may give rise to drug interactions, including over-the-counter medicines and herbal remedies.
- The influence of current and pre-existing illness.
- Health status and general standard of living.
- Unwanted and toxic effects of the drug in question.
- Patient compliance.
- The pharmacological profile of the drug in question, since the altered physiology during pregnancy may affect the way in which the body responds to the drug, as well as how the body deals with the drug molecules and their metabolites.2,5

The following treatment principles may be applied when considering or continuing drug therapy during pregnancy and lactation. Preferably, drugs should be selected that are well known to be safe and effective during pregnancy and lactation. The drug with the shortest plasma half-life should be used at the lowest possible dosage, for the shortest possible duration of treatment. New drugs should be avoided owing to the lack of associated data and clinical experience. Known teratogens should be avoided, even when not yet pregnant, in women of childbearing potential, and sometimes even in fertile males. Self-medicating practices must be discouraged, together with the indiscriminate use of over-the-counter drugs, herbal remedies and nutritional supplements. Possible drug interactions must be carefully considered. Commonly used concomitant medications in pregnancy include antacids, for example, which may interfere with drug absorption from the gastrointestinal tract. The most up-to-date drug information should always be consulted.1,2,4,7

Conclusion

The decision to choose pharmacotherapy during pregnancy, lactation or childbirth is not always optional. Drug treatment may be unavoidable, but will inevitably expose the unborn child to the effects, whether pharmacological or toxic, of the drug itself. Effects on the foetus may be warranted and desired in a few instances. Irrespective of the reason for exposing the pregnant mother and her unborn child to drug therapy, certain aspects and principles need to be considered before commencing the pharmacotherapy to ensure that the intervention is safe, rational and scientifically sound.

References