The Treatment of Colorectal Cancer During Pregnancy: Cytotoxic Chemotherapy and Targeted Therapy Challenges

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Disclosures of potential conflicts of interest may be found at the end of this article.

Key Words. Pregnancy • Antineoplastics • Colorectal neoplasms • Monoclonal antibodies

ABSTRACT

Cancer diagnosed during pregnancy has increased because of delayed child-bearing and the known occurrence of age-dependent malignancies. Cases of colorectal cancer (CRC) in pregnancy have recently been reported. With the expected rise in CRC diagnosed in young adults coupled with the current trend of delayed child-bearing, CRC during pregnancy is likely to increase. Treating pregnant women with CRC by using antineoplastics presents a dilemma because there are many unknowns to guide treatment decisions. We review the issues regarding the use of 10 CRC-approved agents in pregnancy. The Oncologist 2016; 21:563–570

Implications for Practice: Colorectal cancer (CRC) in pregnancy is likely to become more common because of the current population trend in delayed child-bearing and the increase in CRC incidence expected among young adults. Practitioners should become familiar with the challenges associated with systemic treatment of a pregnant patient with CRC. This review addresses concerns surrounding the 10 systemic agents approved for CRC to help provide treatment guidance when such a case arises.

INTRODUCTION

Cancer diagnosed during pregnancy has increased during the past several decades and currently has a reported incidence of 1 in 1,000 pregnancies [1–12]. This increase is linked to women delaying child-bearing to the third and fourth decades of life, coupled with the known occurrence of age-dependent malignancies [1, 2, 6, 10–13]. Breast, ovarian, and cervical cancer; melanoma; and lymphoma are among the more common cancers diagnosed during pregnancy [1–3, 8, 9, 11, 12]. Colorectal cancer (CRC) during pregnancy is rare, with an incidence of 1 in 13,000 pregnancies [2, 10, 14]. Although infrequent, CRC cases during pregnancy have been reported in the past decade [15–31]. With the trend in delayed child-bearing, the rare situation of CRC during pregnancy will likely rise in incidence.

Ten systemic agents have been approved for CRC. Using these antineoplastics in a pregnant patient continues to be an undefined area, with many ethical, safety, and efficacy concerns. Physicians must weigh the risks to the fetus against the benefits of the mother’s treatment. Confounding treatment decisions are cancer stage, the mother’s outcomes, harm that can accompany treatment delays, treatment timing with regard to gestation, potential teratogenic effects, and, in the worst-case scenario, the consideration of termination of the pregnancy. Physicians making treatment decisions face many unknowns, thus further complicating an already sensitive discussion with the patient, the patient’s significant other, and the patient’s family. Unknown factors exist because of limited chemotherapy exposure in pregnancy, multiple agents used simultaneously, varying teratogenic potential, other teratogenic exposures (chemical, environmental, medication), data limited to retrospective cases, and pharmacokinetic changes that occur during pregnancy. Our review focuses on CRC antineoplastics and the challenges that are present when their use is considered during pregnancy.

COLORECTAL CANCER IN THE YOUNG ADULT POPULATION

As mentioned previously, CRC during pregnancy may become more of a reality because of delayed child-bearing. Adding to this potential is the increase of young adults diagnosed with CRC. CRC is the third most common cancer diagnosed among women in the U.S. [32]. In women, it represents 8% of all new cancer cases, with 63,610 estimated new CRC cases expected in 2015. CRC is a malignancy that occurs in people at an older age; the median age of diagnosis is 68 years [33]. Although most CRC cases are diagnosed beyond age 45 years, 5.4% are diagnosed in patients younger than 45 years. Recent Surveillance, Epidemiology, and End Results (SEER) database reviews have revealed an increase in younger CRC patients [34–36]. Recently, a SEER...
Unknown factors exist because of limited chemotherapy exposure in pregnancy, multiple agents used simultaneously, varying teratogenic potential, other teratogenic exposures (chemical, environmental, medication), data limited to retrospective cases, and pharmacokinetic changes that occur during pregnancy.

### OVERVIEW OF PREGNANCY AND CHEMOTHERAPY/TARGETED THERAPY

#### Drug Exposure During Pregnancy

Drug exposure to the fetus during pregnancy and teratogenic effects depend on medication characteristics that allow for placental transfer, the timing of gestation, dosing, administration route, and maternal drug metabolism. Most drugs cross the placenta by passive diffusion [6, 37–40]. Unionized, low-molecular-weight, low-protein-bound, and highly lipophilic agents will cross the placenta more readily [1, 38–40]. Drugs with a molecular weight of less than 500–600 cross the placenta, whereas those with molecular weights greater than 1,000 are reported to cross poorly [6, 11, 38, 39]. Most traditional antineoplastics have a weight below 400, resulting in potential chemotherapy exposure to the fetus [11]. Larger molecules, such as IgG, require specific receptor-mediated active transfer across the placenta [7, 8, 41, 42].

The U.S. Food and Drug Administration (FDA) has historically defined and classified medications into risk categories on the basis of known or potential effects on the fetus (Table 1) [43]. CRC antineoplastics are listed as category C and D because of a lack of human evidence, potential, or known harm [37, 44–54]. The FDA has recently put forth an initiative to transition from using these categories because of their simplicity [55]. Medication prescribing information will soon report an alternative and more descriptive method for pregnancy, lactation, and fertility statements. To date, however, these categories are still in place as this classification transitions into current practice with approved medications; therefore, practitioners must use caution and consider the limitations with this classification. Individual characteristics for CRC agents are summarized in Table 2 [13, 15–23, 37, 44–54].

Pharmacokinetic changes during pregnancy may affect chemotherapy metabolism and exposure [1, 39, 56–61]. During pregnancy, the volume of distribution increases, protein binding decreases, hepatic clearance changes, and renal elimination increases [1, 6, 56–58]. Medications that are excreted unchanged by the kidneys can be eliminated more rapidly because of an approximate 50% increase in creatinine clearance, with increases seen as early as 9–14 weeks’ gestation and peaking particularly in the second trimester [1, 39, 56, 60, 61]. A decrease in albumin allows for more unbound free drug; this is particularly concerning for those highly protein-bound antineoplastics, such as oxaliplatin, which is greater than 90% protein bound [39, 46, 56, 57, 61]. Hepatic alterations include increase in activity of certain hepatic enzymes, such as UGT, CYP3A4, CYP2C9, and CYP2A6, which can lead to changes in drug metabolism [56, 57, 61–63]. Irinotecan has a complex metabolism involving both conversions to its active metabolite (SN38) by carboxylesterase and to inactive metabolites via CYP3A4 [47]. SN38 is conjugated by UGT1A1 to SN38G, a much less active form. Irinotecan may follow an increased drug clearance pattern during pregnancy, similar to that of labetalol, that has increased clearance correlated to UGT1A1 upregulation [57, 62, 63]. Limited chemotherapy pharmacokinetic evaluations during pregnancy and the physiologic changes that occur throughout pregnancy make it difficult to extrapolate these alterations in determining dose, metabolism, and clearance.

#### Timing of Drug Administration

Major congenital malformations are reported at 3%–4% of the general population; approximately 2%–3% of these are correlated to drug administration [9, 13, 38–40, 64]. Congenital malformations rates related to chemotherapy administration inversely relate to gestational age. These malformations have been reported at 10%–20% in the first trimester, 8% in the second trimester, and 6% in the third

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**Table 1.** Food and Drug Administration pregnancy risk categories

<table>
<thead>
<tr>
<th>Category</th>
<th>Definition</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>Adequate and well-controlled studies have failed to demonstrate a risk to the fetus in the first trimester (and there is no evidence of risk in later trimesters)</td>
</tr>
<tr>
<td>B</td>
<td>Animal-reproduction studies have not demonstrated a fetal risk but there are no well-controlled studies in pregnant women (and there is no evidence of a risk in later trimesters)</td>
</tr>
<tr>
<td>C</td>
<td>Animal reproduction studies have shown an adverse effect on the fetus and there are no adequate and well-controlled studies in humans, but potential benefits may warrant use of the drug in pregnant women despite potential risk</td>
</tr>
<tr>
<td>D</td>
<td>There is positive evidence of human fetal risk based on adverse reaction data from investigational or marketing experience or studies in humans, but potential benefit may warrant use of the drug in pregnant women despite potential risks</td>
</tr>
<tr>
<td>X</td>
<td>Studies in animals or humans have demonstrated fetal abnormalities or there is positive evidence of fetal risk based on adverse reaction reports from investigational or marketing experience, or both, and the risks of use of the drug in pregnant women clearly outweigh the potential benefits</td>
</tr>
</tbody>
</table>

From [43].
trimester [2, 4, 11, 42]. Selig et al. found adverse pregnancy outcomes with chemotherapy to be 33%, 27%, and 25% with the first, second, and third trimester, respectively [4]. In this review, adverse pregnancy outcomes included congenital malformations, stillbirths, spontaneous abortions, functional defects, and blood or electrolyte abnormalities.

As evidenced by these citations, the most concerning time period for physical malformation from chemotherapy exposure is the first trimester [1, 2, 4, 9, 11, 38, 65]. Malformations correlate with the organ differentiation during the gestational time period with the neural tube, heart, limbs, and lips developing earlier followed by the ears and palate [1, 9, 65]. After organogenesis, the fetus continues to be vulnerable because of continued maturation of the eyes, teeth, ears, palate, genitalia, hematopoietic system, and central nervous system with growth and functional maturation continuing until term. Intrauterine growth restrictions, low birthweight, premature delivery, functional defects, and adverse effects of antineoplastics on the mother, such as myelosuppression, are risks to the fetus during the second and third trimesters [1, 8–11, 65]. Because of potential adverse effects, chemotherapy administration is not recommended 3 weeks before the expected delivery or beyond 35 weeks of gestational age; possible complications include myelosuppression, bleeding, and death during delivery [1, 2, 5, 8, 10, 11, 13]. Effects related to function, intellect, and behavior are difficult to correlate to a specific cause because they generally are not present at the time of birth and may present several years after the exposure [6, 11, 37, 40].

Recent reviews on monoclonal antibodies in pregnancy suggest an opposing time frame risk for administration [7, 8, 37, 41, 42]. Monoclonal antibodies are large hydrophilic compounds with molecular weights that well exceed placental diffusion transfer. To reach the fetus, active transport is required. During the first trimester, transfer of IgG is minimal; however, fetal IgG concentration begins to rise in the second trimester, indicating active transport across the placenta [7, 37, 41, 42]. Trastuzumab is an IgG subclass monoclonal antibody targeting human epidermal growth factor receptor 2. Trastuzumab in pregnancy develops in oligohydramnios, increasing the risk for preterm delivery and fetal mortality, particularly when given after the first trimester [7, 8, 13, 41, 42, 65]. Cetuximab, panitumumab, bevacizumab, ramucirumab, and ziv-aflibercept are IgG subclass monoclonal antibodies or fusion proteins and may follow this same fetal exposure pattern [7, 37, 49–51, 53, 54].

Table 2. Characteristics of colorectal cancer therapies

<table>
<thead>
<tr>
<th>Agent [reference]</th>
<th>Molecular weight</th>
<th>Pregnancy risk category</th>
<th>Current pregnancy data</th>
</tr>
</thead>
<tbody>
<tr>
<td>5-Fluorouracil (5-FU) [13, 44]</td>
<td>130.1</td>
<td>D</td>
<td>Human data: cases showing low percentage for congenital malformations in the 2nd and 3rd trimester Animal data: embryolethal and teratogenic</td>
</tr>
<tr>
<td>Capecitabine [23, 45]</td>
<td>359.3</td>
<td>D</td>
<td>Human data: 1 case in combination with oxaliplatin; infant reported normal; however, limited information Animal data: embryolethal and teratogenic</td>
</tr>
<tr>
<td>Irinotecan [20, 21, 47, 48]</td>
<td>587–677.19 (SN-38 = 392)</td>
<td>D</td>
<td>Human data: 2 cases given 5-FU in 2nd and 3rd trimester report no congenital malformations Animal data: embryotoxic, teratogenic, decreased learning, and low birth weights</td>
</tr>
<tr>
<td>Oxaliplatin [15–19, 46]</td>
<td>397.3</td>
<td>D</td>
<td>Human data: 5 cases given 5-FU in 2nd and 3rd trimester report limited congenital malformations (1 case of hypothyroidism) Animal data: embryo-fetal toxicity and teratogenic</td>
</tr>
<tr>
<td>Bevacizumab [37, 49]</td>
<td>149,000</td>
<td>C</td>
<td>Human data: no i.v. administration data Animal data: teratogenic, fetal resorptions, reduced maternal and fetal weights</td>
</tr>
<tr>
<td>Ziv-Aflibercept [50]</td>
<td>97,000</td>
<td>C</td>
<td>Human data: none Animal data: embryo-fetal toxicity and teratogenic effects</td>
</tr>
<tr>
<td>Cetuximab [37, 53]</td>
<td>152,000</td>
<td>C</td>
<td>Human data: none Animal data: embryolethal and abortifacient effects</td>
</tr>
<tr>
<td>Panitumumab [37, 54]</td>
<td>147,000</td>
<td>C</td>
<td>Human data: none Animal data: embryolethal and abortifacient effects</td>
</tr>
<tr>
<td>Regorafenib [52]</td>
<td>500.83</td>
<td>D</td>
<td>Human data: none Animal data: embryolethal and teratogenic effects</td>
</tr>
<tr>
<td>Ramucirumab [51]</td>
<td>147,000</td>
<td>C</td>
<td>Human data: none Animal data: none</td>
</tr>
</tbody>
</table>

*Table 1 describes the pregnancy categories.*
CANCER THERAPIES AND PREGNANCY

Chemotherapy
The fluoropyrimidines, 5-fluorouracil (5-FU) and capecitabine, are the backbone for CRC treatment when used as monotherapy or in combination with additional chemotherapy, with or without targeted therapy. Both fluoropyrimidines have low molecular weights, and 5-FU has negligible protein binding; capecitabine, a prodrug to 5-FU, is moderately protein-bound (35% bound to albumin) [44, 45]. Drug exposure would likely be high given these drug properties, and animal studies in mice have demonstrated placental transfer [13, 37, 44, 45].

Many cases of human 5-FU exposure in pregnancy exist. According to the 2013 National Toxicology Program (NTP) monograph regarding chemotherapy and pregnancy, 178 cases with 5-FU have been reported [13]. A majority of these cases occurred in breast cancer, and in most (172 cases) the drug was given with additional therapy. Seventeen pregnant women received 5-FU in the first trimester; 161 did so in the second and third trimesters. The NTP monograph reports 4 cases out of 13 (31%) with major malformations when 5-FU was given in the first trimester; only 2 of 161 (1.2%) cases occurred in the second or third trimester. Malformations reported in the first trimester included microcephaly, low-set ears, hypertelorism, a right palmar simian crease, ventriculomegaly, colpocephaly, and skeletal deformities. These cases were coexposed to additional chemotherapy (methotrexate, cyclophosphamide, doxorubicin) or other teratogenic exposures (radiation or radiographic imaging). Clubfoot and hemihypertrophy of the lower extremity were malformations occurring in the second- and third-trimester group.

A follow-up of the children of 81 patients with breast cancer who had been given bolus 5-FU in combination with doxorubicin and cyclophosphamide (FAC) in the second and third trimester has been recently published; most patients (86%) had received >4 cycles of FAC [64]. Seventy-eight percent were available to answer the follow-up survey, and 3 cases of congenital malformations were reported. These abnormalities were Down syndrome, clubfoot, and ureteral reflux. The children were a median of 7 years (range, <1–21 years). Fifty patients responding to the survey answered questions about postneonatal outcomes; 98% considered the child healthy. Reported health concerns included developmental milestone delays, difficulties in school, asthma, vision, heart murmur, lazy eye, absence seizures, ear/nose/throat conditions, and gastroesophageal reflux disease; however, the authors correlated these percentages to the population prevalence. The rates of allergies and eczema were higher than those in the population, but the authors attributed this to over-reporting.

A concern with extrapolation from these cases is the route of administration; bolus dosing is used with FAC, whereas 5-FU in CRC is given as both a bolus and continuous infusion. No congenital abnormalities, except for hypothyroidism, were found in 8 pregnant women with CRC who received 5-FU alone or in combination (with oxaliplatin or irinotecan) after the first trimester (Table 3) [15–22]. Capecitabine was given in one case of CRC during pregnancy, along with oxaliplatin, in the first trimester [13, 23]. The authors report that the patient was exposed during the first trimester, but once the pregnancy was discovered treatment was stopped [23]. Infant follow-up at 2 years was reported as normal; however, there is limited information regarding the amount of exposure, the delivery, and infant follow-up.

5-FU given in the second or third trimester appears to present a low concern for congenital malformations and has long-term follow-up in the bolus-dosing setting. Consideration for dihydropyrimidine dehydrogenase testing before administration should be considered for this population because significant adverse effects with 5-FU can occur when deficient in this enzyme. Literature regarding capecitabine is sparse; therefore, if a fluoropyrimidine is needed, 5-FU should be used.

Oxaliplatin, a third-generation platinum agent, is used in locally advanced and metastatic colorectal cancer (mCRC) in combination with a fluoropyrimidine, with or without targeted therapy. Oxaliplatin’s molecular weight is less than 400, indicating exposure to the fetus by placental transfer [46]. It is highly protein-bound (>90%); therefore, with lower albumin levels seen in pregnancy, this may indicate more free active drug. It is approximately 50% eliminated by the kidney and may have a higher clearance in pregnancy with standard dosing. Studies in pregnant rats given oxaliplatin from day 6 to 16 showed early resorptions, decreased fetal weight, and delayed ossification.

Seven humans receiving oxaliplatin in pregnancy have been described [15–19, 23, 66]. In the five CRC patients administered 5-FU, oxaliplatin, and leucovorin (FOLFOX), treatment was given after the first trimester (initiated at 13–23 weeks’ gestation) (Table 3) [15–19]. In one infant, hypothyroidism was reported, but none of the remaining infants had congenital malformations reported at birth. Two infants were described as small for gestational age (born at 31 and 36 weeks’ gestation, respectively) [16, 19]. No developmental deficits have been reported in the follow-up of these infants exposed to FOLFOX; however, height and weight for some of the children remain in a lower percentile [16, 18, 19]. One infant with congenital malformations, including a cleft lip, cleft palate, and esophageal atresia with a tracheoesophageal fistula, was exposed in the first trimester to a combination of vinorelbine, irinotecan, and oxaliplatin [13]. It is difficult to determine the exact agent that caused these malformations or whether the cause was the combination, the first-trimester administration, or genetic predisposition. Infants born without congenital malformations (with the exception of hypothyroidism) have been reported after administration of FOLFOX after the first trimester [15–19]. Although limited to six exposed infants, healthy infant follow-up has been reported, with the exception of lower growth percentiles in some children [16, 18, 19].

Irinotecan, a topoisomerase I inhibitor, is used in mCRC alone or in combination with targeted therapy with or without additional chemotherapy. Irinotecan, as mentioned previously, has a complex metabolism. Irinotecan is converted to SN38 (irinotecan’s active metabolite, which is responsible for most antitumor activity), and irinotecan is converted inactively via CYP3A4 [47]. SN38 elimination is through conjugation via the UGT1A1 enzyme. Pregnancy enzyme increases in CYP3A4, and UGT may lead to an increased conversion to inactive metabolites; this makes dosing
and exposure variable [56, 57, 62, 63]. Irinotecan is moderately protein-bound (30%–68%), whereas SN38 is highly protein-bound (95%); therefore, more unbound SN38 is a potential concern [37, 47]. Both molecular weights are within the range that allows placental transfer [47, 48].

Human pregnancy exposure to irinotecan is limited to two cases in addition to the preceding case described in combination with oxaliplatin and vinorelbine [20, 21]. In both cases, irinotecan was combined with 5-FU and leucovorin and therapy was initiated after the first trimester (at 18 and 23 weeks’ gestation). No congenital malformations were reported; however, in one case intrauterine growth restriction at birth was reported. Although infant follow-up was not long-term, the authors reported healthy infants in both cases.

Animal cases are not as favorable. In rats and rabbits during organogenesis, a range of i.v. doses showed embryolethal and structural defects [37, 47]. Perhaps one reason for these differences in animal and human experience is the timing of administration; in the animals, the doses were given

### Table 3. Cases of CRC in pregnancy treated with chemotherapy

<table>
<thead>
<tr>
<th>Case [reference]</th>
<th>Chemotherapy regimen</th>
<th>Infant at birth</th>
<th>Infant follow-up</th>
<th>Mother follow-up</th>
</tr>
</thead>
<tbody>
<tr>
<td>mCRC, age 33 yr [15] Chemotherapy initiated: 22 weeks’ gestation</td>
<td>FOLFOX; total cycles: 6</td>
<td>Vaginal delivery at 38 weeks’ gestation • Reported healthy • No developmental complications</td>
<td>Infant follow-up (2 yr): • Met developmental milestones • No recognizable complications</td>
<td>Receiving palliative chemotherapy</td>
</tr>
<tr>
<td>mCRC, age 40 yr [16] Chemotherapy initiated: 23–24 weeks’ gestation</td>
<td>FOLFOX; total cycles: 4</td>
<td>Cesarean delivery at 31.5 weeks’ gestation • Admitted to neonatal unit • Small for gestational age • Hypothyroidism</td>
<td>Infant follow-up (approximately 11 mo) • Weight (50th percentile corrected for prematurity) • Length (90th percentile) • Head circumference (90th percentile) • Result of Denver developmental screening test was normal for adjusted age</td>
<td>Deceased 5 mo after mCRC diagnosis (declined more CRC treatment)</td>
</tr>
<tr>
<td>mCRC, age 26 yr [17] Chemotherapy initiated: 13 weeks’ gestation</td>
<td>FOLFOX; total cycles: 10</td>
<td>Cesarean delivery at 33 weeks’ gestation • No malformations reported</td>
<td>Twins’ follow-up (2 yr) • Developing normally</td>
<td>Deceased 1 yr after mCRC diagnosis</td>
</tr>
<tr>
<td>mCRC, age 25 yr [18] Chemotherapy initiated: 20 weeks’ gestation</td>
<td>FOLFOX; total cycles: 6</td>
<td>Vaginal delivery at 33.6 weeks’ gestation • No malformations reported</td>
<td>Infant follow-up (3.5 yr) • Height: 60th percentile • Weight: 45th percentile • No deficits reported</td>
<td>1 yr, no evidence of disease after metastatic liver resection</td>
</tr>
<tr>
<td>mCRC, age 38 yr [19] Chemotherapy initiated: 19 weeks’ gestation</td>
<td>FOLFOX; total cycles: 3</td>
<td>Cesarean delivery at 36 weeks’ gestation • Small for gestational age • Normal neurological examination • No malformations reported</td>
<td>Infant follow-up (10 mo) • Weight: 10th–25th percentile • Head circumference: 10th–25th percentile • No deficits reported</td>
<td>Receiving treatment at 13 mo since diagnosis</td>
</tr>
<tr>
<td>mCRC, age 33 yr [20] Chemotherapy initiated: 23 weeks’ gestation</td>
<td>FOLFIRI; total cycles: 3 cycles</td>
<td>Cesarean delivery at 30 weeks’ gestation • Admitted to neonatal unit because of prematurity and intrauterine growth restriction • No malformations reported</td>
<td>Infant follow-up (13 mo) • Healthy • Achieved appropriate growth and development milestones</td>
<td>FOLFOX with progression after 4 cycles followed by best supportive care</td>
</tr>
<tr>
<td>Krukenberg tumor primary, determined to be CRC, age 34 yr [21] Chemotherapy initiated: 18 weeks’ gestation</td>
<td>FOLFIRI; total cycles: 10</td>
<td>Vaginal delivery at approximately 37 weeks’ gestation • No malformations reported</td>
<td>Infant follow-up (4 mo) • Normal development</td>
<td>Adjuvant chemotherapy</td>
</tr>
<tr>
<td>CRC, age 31 yr [22] Chemotherapy initiated: 29 weeks’ gestation</td>
<td>5-FU; total cycles: NR</td>
<td>Delivery: 39 weeks’ gestation with uneventful delivery • Reported healthy • No malformations reported</td>
<td>NR</td>
<td>NR</td>
</tr>
</tbody>
</table>

Abbreviations: CRC, colorectal cancer; FOLFIRI, 5-fluorouracil, irinotecan, folinic acid; FOLFOX, 5-fluorouracil, oxaliplatin, and leucovorin; mCRC, metastatic colorectal cancer; NR, not reported.
during organogenesis. However, in rats a dose given after organogenesis showed decreased learning ability and decreased female body weights. Effects of irinotecan and SN38 exposure to the human fetus are not known; however, given their pharmacokinetic properties, exposure is likely. Because there are only two human pregnancy cases, it would be premature to use this as a basis for recommending irinotecan use in pregnancy. If irinotecan must be used, practitioners should consider testing for UGT1A1 homozygous allele before administration because of increased neutropenia related to more exposure seen at standard dosing. If the patient is determined to be deficient in this enzyme or to have Gilbert’s disease, doses should be reduced.

If 5-FU, oxaliplatin, and irinotecan are used during pregnancy (after a thorough discussion with the family and after the patient provides consent), these agents should be given only in the second or third trimester; they should be held 3 weeks before delivery or at 35 weeks’ gestation, as recommended for other antineoplastics [1, 2, 5, 8, 10, 11, 13].

Angiogenesis Inhibitors
The complex placental vasculature is critical for fetal development and survival because it is responsible for the maternal-fetal blood supply, gas exchange, and nutrient supply [67, 68]. The vascular network requires vasculogenesis and angiogenesis, which begins 18–21 days after conception and continues through the gestational period. Vascular endothelial growth factor (VEGF) is involved in both processes. VEGF-A, VEGF-B, VEGF-C, VEGF-D, and placental growth factor are among the regulators of angiogenesis [42, 67, 68]. Inhibition and disruption of key regulators involved in this vascular network can produce pregnancy complications, including preeclampsia; intrauterine growth restriction; stillbirth; preterm delivery; miscarriage; and abnormal development of the heart, blood vessels, forelimbs, cranial region, and forebrain [42, 51, 52, 67, 68]. Thalidomide, an antineoplastic used primarily in multiple myeloma, is predominantly known for its teratogenic effects and initial removal from the market in the early 1960s [7, 8]. Thalidomide, an antineoplastic used primarily in multiple myeloma, is predominantly known for its teratogenic effects and initial removal from the market in the early 1960s [7, 8].

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The complex placental vasculature is critical for fetal development and survival because it is responsible for the maternal-fetal blood supply, gas exchange, and nutrient supply [67, 68]. The vascular network requires vasculogenesis and angiogenesis, which begins 18–21 days after conception and continues through the gestational period. Vascular endothelial growth factor (VEGF) is involved in both processes. VEGF-A, VEGF-B, VEGF-C, VEGF-D, and placental growth factor are among the regulators of angiogenesis [42, 67, 68]. Inhibition and disruption of key regulators involved in this vascular network can produce pregnancy complications, including preeclampsia; intrauterine growth restriction; stillbirth; preterm delivery; miscarriage; and abnormal development of the heart, blood vessels, forelimbs, cranial region, and forebrain [42, 51, 52, 67, 68]. Thalidomide, an antineoplastic used primarily in multiple myeloma, is predominantly known for its teratogenic effects and initial removal from the market in the early 1960s [7, 8].

use by pregnant women for morning sickness resulted in an increased female body weights. Effects of irinotecan and SN38 exposure to the human fetus are not known; however, given their pharmacokinetic properties, exposure is likely. Because there are only two human pregnancy cases, it would be premature to use this as a basis for recommending irinotecan use in pregnancy. If irinotecan must be used, practitioners should consider testing for UGT1A1 homozygous allele before administration because of increased neutropenia related to more exposure seen at standard dosing. If the patient is determined to be deficient in this enzyme or to have Gilbert’s disease, doses should be reduced.

If 5-FU, oxaliplatin, and irinotecan are used during pregnancy (after a thorough discussion with the family and after the patient provides consent), these agents should be given only in the second or third trimester; they should be held 3 weeks before delivery or at 35 weeks’ gestation, as recommended for other antineoplastics [1, 2, 5, 8, 10, 11, 13].

Angiogenesis Inhibitors
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Cetuximab and panitumumab should be avoided in pregnancy given the mechanism of inhibition on EGFR, a player in fetal growth and maintenance of pregnancy, teratogenic animal effects, and no human exposure. Furthermore, these agents are known to cause magnesium wasting, which may directly affect the fetus.

**CONCLUSION**

CRC is accepted as one of the most common malignancies globally, but the rising incidence in young patients suggests the challenges that may lie ahead for a patient who is pregnant. CRC in young adults is aggressive [73]; therefore, postponing treatment until full pregnancy term in a pregnant CRC patient is unlikely to be feasible. The purpose of this review was to provide guidance to clinicians faced with difficult decisions when pregnancy arises in a patient with locally advanced or metastatic CRC. Because of organogenesis, chemotherapy should be avoided during the first trimester, as this is the most concerning time period for congenital malformations; chemotherapy administration should be reserved for the second and third trimesters. Patients and families should receive thorough counseling before any chemotherapy administration, including the consideration of fertility counseling. CRC monoclonal antibodies and multikinase inhibitors should be avoided in pregnancy because of a lack of human evidence and concerns regarding mechanism of action. Follow-up of infants who were exposed to chemotherapy in utero, particularly long-term follow-up regarding cognitive dysfunction, is an area that needs to be strengthened through reporting. Therefore, we stress the importance of reporting outcomes in all pregnant patients exposed to chemotherapy in order to obtain more information to make sound decisions and provide future guidance to other physicians.

**REFERENCES**

Colorectal Cancer Treatment in Pregnancy