Avatrombopag Optimizes Response to Niraparib by Managing Thrombocytopenia Associated with Poly-ADP Ribose Polymerase (PARP) Inhibition in Ovarian Cancer and Breast Cancer: A Case Series

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Case series
Patients: —
Final Diagnosis: Breast cancer • ovarian cancer
Symptoms: Thrombocytopenia
Medication: —
Clinical Procedure: —
Specialty: Oncology

Objective: Unusual or unexpected effect of treatment
Background: Thrombocytopenia is a potentially treatment-limiting adverse event of particular interest with the PARP inhibitor niraparib. This adverse event may necessitate niraparib dose reduction or treatment discontinuation, resulting in suboptimal treatment outcomes. Here, we report on niraparib dose optimization in 2 patients with breast cancer and 4 patients with ovarian cancer through concurrent administration of the thrombopoietin receptor stimulating agent avatrombopag to mitigate thrombocytopenia, enabling niraparib reescalation and improved clinical response.

Case Reports: Three of 6 patients received niraparib 300 mg daily, the highest recommended dose, for a sustained period. Avatrombopag therapy enabled niraparib dose escalation that led to reductions in biomarkers associated with disease progression. Before initiation of avatrombopag, increases in CA-125 levels, a marker for ovarian cancer, were observed in association with niraparib dose interruption, and in 2 patients with ovarian cancer CA-125 levels fell in response to niraparib dose escalation enabled by concurrent avatrombopag therapy. Further, in 2 patients with metastatic breast cancer, intracranial response was observed in association with avatrombopag-enabled niraparib therapy. In 1 patient with metastatic breast cancer, niraparib induced an intracranial response, while previous use of talazoparib had not, confirming preclinical findings of superior blood-brain-barrier penetration with niraparib.

Conclusions: Avatrombopag is currently approved for use in chronic immune thrombocytopenia and thrombocytopenia associated with chronic liver disease in patients undergoing a surgical procedure. A clinical trial of avatrombopag for chemotherapy-induced thrombocytopenia is ongoing. Preliminary results in these 6 patient cases demonstrate the need for a confirmatory trial of avatrombopag for optimizing the dose of niraparib.

MeSH Keywords: Breast Neoplasms • Ovarian Neoplasms • Poly(ADP-ribose) Polymerases • Receptors, Thrombopoietin • Thrombocytopenia

Full-text PDF: https://www.amjcaserep.com/abstract/index/idArt/927008
Background

By interfering with DNA repair mechanisms, such as the homologous recombination repair pathway [1,2], poly-ADP ribose polymerase (PARP) inhibitors have been shown to be effective in the maintenance treatment of advanced ovarian cancer [3–8] and in advanced breast cancer with germline BRCA1/2 mutations [9]. Clinical trials of niraparib indicate responses in patients with both germline and somatic BRCA mutations, with a smaller degree of benefit in BRCA-negative homologous recombination deficiency (HRD)-positive patients, and the least benefit in BRCA-negative HRD-negative patients [3,6,10]. Most recently, on April 29, 2020, niraparib was approved for first-line maintenance treatment of ovarian cancer following complete or partial response to platinum-based chemotherapy [10,11]. In an analysis of the NOVA trial, niraparib improved progression-free survival outcomes in a dose-related manner [12]. Moreover, the unique pharmacology of niraparib, including improved blood-brain-barrier penetration over other PARP inhibitors, has been shown to induce intracranial response in animal models [13].

Although niraparib has unique characteristics, its therapeutic efficacy may be limited by adverse events such as thrombocytopenia [14]. The FDA-approved labeling for niraparib advises interruption of niraparib therapy or dose reduction for management of thrombocytopenia, which may compromise efficacy [12,14]. Unfortunately, there are no medications currently approved for management of this potential treatment-limiting adverse event [14]. Although one agent, oprelvekin, was approved for the management of chemotherapy-induced thrombocytopenia, this agent has been discontinued [15]. Based on the mechanism of action of thrombopoietin (TPO) receptor agonists, these agents have the potential to treat thrombocytopenia associated with therapeutic agents for cancer, such as niraparib [16–18].

TPO receptor agonists increase platelet levels by interacting with the TPO receptor on megakaryocytes, the endogenous target of thrombopoietin. Given that TPO receptor agonists increase platelet production, use of these agents may counteract or mitigate PARP-induced thrombocytopenia [16]. Approved TPO receptor agonists are avatrombopag, eltrombopag, and lusutrombopag, all of which are administered orally, as well as romiplostim, which is administered as a subcutaneous injection [17–20]. Unlike eltrombopag, avatrombopag does not require regular monitoring for liver test abnormalities during therapy, which can reduce the complexity of monitoring in highly comorbid patients with cancer receiving maintenance treatment with niraparib. In addition, use of avatrombopag offers flexibility in oral dosing. Unlike eltrombopag, avatrombopag can be taken with polyvalent cations such as calcium, magnesium, and iron [17,18]. As a result, based on its mechanism of action, safety, and flexible dosing, avatrombopag is a rational choice for the prevention and management of thrombocytopenia associated with niraparib therapy.

This case series reviewed the treatment of 2 patients with breast cancer and 4 patients with ovarian cancer. All patients received niraparib for the maintenance treatment of advanced breast cancer or advanced ovarian cancer in combination with avatrombopag to reduce the risk of thrombocytopenia and improve the dose-intensity of niraparib therapy. By optimizing the dose-intensity of therapy, avatrombopag has the potential to optimize treatment outcomes, consistent with the dose-response relationship observed in multiple clinical trials of niraparib [3,6,10]. In this case series, outcomes were evaluated prospectively over the course of treatment for breast or ovarian cancer in 6 patients.

Case Reports

Case 1: Breast cancer with brain metastases

A patient with breast cancer, born in 1977, was diagnosed with ER-positive HER2-negative stage III breast cancer in August 2016. From September 2016 to January 2017, she received 4 cycles of doxorubicin and cyclophosphamide followed by maintenance letrozole from a different provider. By January 2019, she had progressive disease with brain metastases. She received radiosurgery (Gamma Knife®) and subsequently started abemaciclib, which she did not tolerate well. Before presenting in July 2019 for her fourth opinion, she had previously been seen at multiple other centers.

From July 2019 to October 2019, she received talazoparib and experienced progressive disease in the brain with no response to therapy. Radiosurgery (Gamma Knife®) was not feasible and she declined whole-brain radiotherapy. The patient was then switched to a different PARP inhibitor, niraparib. Niraparib was chosen based on its ability to cross the blood-brain barrier and favorable results in regression of intracranial tumors in animal models [13]. The patient started treatment with niraparib 300 mg daily in November 2019, with CT-verified response in early January and late February 2020. This response was accompanied by clinical improvement, including successful discontinuation of steroids and absence of reported seizures between December 2019 and March 2020. As of her latest CT scan performed April 16, 2020, this patient has experienced complete intracranial response (Figure 1).

Use of avatrombopag in this patient

Before starting therapy with avatrombopag, this patient experienced a dramatic drop in platelet count to 31 000/μL on
January 8, 2020, shortly after initiating niraparib 300 mg daily on December 29, 2019. After starting treatment with avatrombopag 20 mg daily, the patient experienced a dramatic increase in platelet counts, which peaked at a supranormal level of 757,000/µL on January 23, 2020. As this patient was extremely responsive to avatrombopag, avatrombopag was held and reinitiated at a lower total weekly dose of 60 mg (20 mg administered on Monday, Wednesday, and Friday each week) starting on February 5, 2020. The patient was able to take niraparib at the maximal dose of 300 mg daily while receiving avatrombopag. Before initiating avatrombopag, niraparib 300 mg daily had previously caused profound thrombocytopenia. After

Figure 1. Case 1, change in brain CT scan in a patient with breast cancer and brain metastases.

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initiating avatrombopag, platelet counts exceeded 250 000/µL on 3 of 5 occasions from February 5, 2020 to March 11, 2020. During this time, the patient received the maximal dose of niraparib (300 mg daily) in combination with avatrombopag 60 mg weekly (Figure 2).

**Figure 2.** Case 1, platelet count, niraparib dose, and avatrombopag dose over time.

**Case 2: Breast cancer with brain metastases**

A female patient born in 1976 was diagnosed with locally advanced estrogen receptor (ER)-positive HER2-negative breast cancer in February 2016. She initially received alternative medicine. In November 2017, she sought conventional medical treatment. She had stage IV disease with liver metastases and pleural effusion and was enrolled in a clinical trial of
carboplatin in combination with paclitaxel and a checkpoint inhibitor. On the clinical trial, the patient achieved partial response, until she developed progressive disease in July 2018. She received treatment with fulvestrant, palbociclib, leupro-lide, and exemestane from July 2018 until the onset of pro-gressive disease with brain metastases in April 2019. She refused cranial radiotherapy and experienced no response with single-agent doxorubicin. On May 30, 2019, she was started on niraparib and achieved partial response in the brain, liver, and lungs.

**Use of avatrombopag in this patient**

From May 30, 2019 to March 16, 2020, the patient received weekly dose of niraparib ranging from 200 mg to 2100 mg, co-administered with avatrombopag weekly doses ranging from 140 mg weekly to 280 mg weekly. Of note, following an interruption in niraparib dosing the week of June 15, 2019, there were a total of 3 weeks in which avatrombopag was not administered (the weeks of July 29, 2019, August 26, 2019, and November 25, 2019). The gap in use of avatrombopag during July and August led to subsequent low platelet levels on of 140 000/µL on September 26, 2019 and 148 000/µL on October 21, 2019. These low platelet levels due to a gap in avatrombopag therapy led to 2 additional interruptions in niraparib therapy the weeks of November 25, 2019, and December 26, 2019. Despite the interruption of avatrombopag therapy, use of avatrombopag in this patient aided in escalating the dose of niraparib from 1800 mg weekly to 2000 mg weekly from August 29, 2019 to October 27, 2019. Most recently, the dose of avatrombopag was increased from 140 mg weekly to 280 mg weekly, resulting in a rebound in platelet counts from 48 000/µL on February 25, 2020 to 94 000/µL on March 2, 2020. The increase in avatrombopag dose also enabled escalation of the weekly dose of niraparib from 200 mg weekly to 1400 mg weekly as of March 16, 2020. On August 26, 2019 and on January 8, 2020, CT assessments indicated intracranial partial response (Figure 3). In addition to 2 standard-dose platelet transfusions, continuous administration of avatrombopag for all but 3 weeks during the dose titration helped enable administration of the optimal dose of niraparib in this patient.

**Case 3: Ovarian cancer**

A patient with stage IV ovarian cancer, born in 1959, was diagnosed in October 2018. She received carboplatin, paclitaxel, and bevacizumab from November 2018 to February 2019. After experiencing partial response to primary treatment, she received maintenance niraparib therapy. On her last CT scan, January 26, 2020, she was found to have stable disease. Her CA-125 levels indicate response to treatment.
Case 5: Ovarian cancer

A female patient born in 1966 was diagnosed with stage III ovarian cancer in October 2018. She received 6 cycles of carboplatin, paclitaxel, and bevacizumab from November 2018 to April 2019 followed by niraparib maintenance therapy. She declined dual maintenance therapy with bevacizumab and niraparib. Her CA-125 levels showed a rise when niraparib maintenance was interrupted, and a subsequent decline when niraparib therapy was reintroduced and the dose was escalated.

Use of avatrombopag in this patient

Before starting avatrombopag, the patient experienced 3 episodes of platelet count declines in association with niraparib use from April 2019 to August 2019 (platelet counts were 1000/µL on May 20 and May 28, 2019, 10 000/µL on July 31, 2019, and 65 000/µL on August 28, 2019). After starting avatrombopag at a weekly dose of 140 mg, the patient was able to tolerate an increase in her weekly niraparib dose from 500 mg to 900 mg on December 2, 2019 and a further increase to 1000 mg weekly on March 18, 2020. On February 5, 2019, the dose of avatrombopag increased from 140 mg weekly to 280 mg weekly. Despite continued up-titratin in the dose of niraparib, platelet counts continued to rise. CA-125 levels fell from 43 U/mL on February 5, 2020 to 37 U/mL by March 4, 2020 concurrent with continuous use of niraparib and avatrombopag in this patient.

Case 6: Ovarian cancer

A patient born in 1944 was diagnosed with stage III ovarian cancer in March 2019. From March 2019 to May 2019, she received 4 cycles of carboplatin, paclitaxel, and bevacizumab, followed by debulking surgery in June 2019. Following disease progression, she received 4 cycles of carboplatin and paclitaxel until September 2019, followed by rucaparib maintenance. CA-125 levels in this patient had normalized since surgical intervention and have remained stable throughout treatment. As of March 2020, she was disease-free.

Use of avatrombopag in this patient

In this patient, niraparib therapy was initiated at a dose of 700 mg weekly on December 4, 2019, which was gradually increased to 2100 mg weekly by January 7, 2020. With upward dose titration, platelet counts fell from a peak of 199 000/µL on December 11, 2019 to 106 000/µL by December 26, 2019. As a result of this drop in platelet counts, avatrombopag was also initiated on December 26, 2019 at a daily dose of 20 mg (140 mg weekly). As a result of avatrombopag coadministration, the patient experienced an increase in platelet counts to 163 000/µL on January 7, 2020. To support further dose...
Figure 4. Case 3, platelet count, niraparib dose, and avatrombopag dose over time.
Therapeutic doses used*  
Sustained maximal dose of niraparib** (Y/N)  
Biomarkers confirming clinical response  
Key notes on case

** Table 1. Outcomes in patients receiving avatrombopag to optimize therapeutic outcomes with niraparib therapy in breast cancer and ovarian cancer.

| Case 1 (breast cancer with brain metastases) | Avatrombopag: 0–140 mg/week; Niraparib: 0–2100 mg/week | Y | CT-confirmed intracranial response and successful discontinuation of steroids and absence of reported seizures after intracranial response enabled by niraparib and avatrombopag | Use of niraparib in this patient initially led to profound thrombocytopenia (31,000/µL), which was rapidly transformed to a supranormal platelet count (757,000/µL) when the patient received avatrombopag 20 mg daily. Dosing of avatrombopag 20 mg two to three times weekly (total weekly dose: 40 mg to 60 mg) enabled continued therapy with niraparib and clinical response. |
| Case 2 (breast cancer with brain metastases) | Avatrombopag: 140–280 mg/week; Niraparib: 0–2100 mg/week | N | CT-confirmed intracranial partial response confirmed on 2 occasions | Continuous administration of avatrombopag for all but 3 weeks enabled titration to the maximal dose of niraparib, enabling intracranial partial response. Of note, a period when avatrombopag was interrupted led to a reduction in platelet counts in subsequent months. |
| Case 3 (ovarian cancer) | Avatrombopag: 140 mg/week; Niraparib: 0–2100 mg/week | Y | Reduction in CA-125 levels in association with treatment with niraparib enabled by avatrombopag therapy | The patient has been able to receive the maximal dose of niraparib while receiving avatrombopag. Notably, the maximal dose of niraparib caused profound thrombocytopenia (36,000/µL) when it was administered without avatrombopag. CA-125 levels increased upon niraparib dose interruption and increased upon niraparib dose escalation enabled by avatrombopag therapy. |
| Case 4 (ovarian cancer) | Avatrombopag: 140–280 mg/week; Niraparib: 0–2100 mg/week | Y | CA-125 levels were stable during continued dosing of niraparib and avatrombopag | Use of avatrombopag enabled the patient to tolerate a maximal dose of niraparib. CA-125 levels were stable during continued dosing of niraparib and avatrombopag. |
| Case 5 (ovarian cancer) | Avatrombopag: 140–280 mg/week; Niraparib: 0–2100 mg/week | N | Drop in CA-125 levels in association with continued dosing of niraparib and avatrombopag | Use of avatrombopag enabled the patient to tolerate a weekly dose of 500 mg of niraparib, which was later escalated to 1000 mg of niraparib. A reduction in CA-125 levels was observed with niraparib dose optimization. |
| Case 6 (ovarian cancer) | Avatrombopag: 0–280 mg/week; Niraparib: 0–2100 mg/week | N | CA-125 levels were stable during continued dosing of niraparib and avatrombopag. The patient experienced complete response and is disease-free as of March 2020 | While receiving concurrent avatrombopag, the patient was able to maintain platelet counts above 100,000/µL while on niraparib with some niraparib dose reductions. Avatrombopag therapy enabled this patient to tolerate niraparib, which may not have been possible in the absence of avatrombopag. |

* Total weekly dose of treatments administered daily; ** defined as 3 or more consecutive weeks receiving niraparib 300 mg daily.
Results

Three of 6 patients were able to sustain treatment with the maximum dose of niraparib (300 mg daily) for at least 3 continuous weeks. Continuous administration of avatrombopag mitigated thrombocytopenia and enabled dose escalation of niraparib to induce CT-confirmed response in 2 patients with breast cancer (cases 1 and 2). Reductions in CA-125 levels were observed in association with niraparib therapy enabled by avatrombopag therapy in 2 of 4 patients with ovarian cancer (cases 3 and 5). The remaining 2 patients had stable CA-125 levels (cases 4 and 6). One of these patients (case 6) has experienced complete response and was disease-free as of March 2020. One patient with breast cancer who experienced profound thrombocytopenia (31 000/µL) on niraparib attained supranormal platelet counts (757 000/µL) within 3 weeks of initiating avatrombopag 20 mg daily and was transitioned to a reduced dose (20 mg 3 times weekly), which enabled continued therapy with niraparib and continued response (Table 1).

Discussion

Use of niraparib is limited by potential adverse events, including thrombocytopenia [14], which can lead to dose interruption or treatment discontinuation, which may reduce efficacy [12,14]. With a lack of approved medications for management of thrombocytopenia, there is a need for new therapeutic options [14,15]. As demonstrated in this case series, the TPO receptor agonist avatrombopag can counteract thrombocytopenia by interacting with the TPO receptor on megakaryocytes, the endogenous target of thrombopoietin [16–18]. In this case series of 2 patients with breast cancer and 4 patients with ovarian cancer, concurrent administration of the TPO receptor agonist avatrombopag mitigated thrombocytopenia and enabled both niraparib dose reescalation and improved clinical response.

In 2 patients with metastatic breast cancer, intracranial response was observed in association with avatrombopag-enabled niraparib therapy. Notably, in 1 patient with metastatic breast cancer, niraparib induced intracranial response, while previous use of talazoparib had not. This observation is confirmatory of preclinical findings with niraparib indicating superior blood-brain-barrier penetrance over other PARP inhibitors [13].

To reduce the risk of adverse events with niraparib, including thrombocytopenia, the starting dose of niraparib may be adjusted based on baseline bodyweight and platelet count. Specifically, patients with a baseline bodyweight <77 kg or with a baseline platelet count <150 000/µL may receive a reduced starting dose of 200 mg of niraparib daily, rather than a starting dose of 300 mg daily. In the PRIMA trial evaluating niraparib in the first-line maintenance setting of ovarian cancer, this strategy was introduced as a protocol amendment, and was shown to reduce the risk of thrombocytopenia from 52.4% with fixed starting dose niraparib to 33.7% with individualized starting dose niraparib. However, the initial dose adjustment of niraparib does not eliminate the issue of thrombocytopenia, as 14.8% of patients receiving individualized starting dose niraparib in PRIMA experienced thrombocytopenia of grade 3 severity or higher. Based on this case series, the risk of developing treatment-limiting thrombocytopenia may be further reduced through the use of avatrombopag in appropriate patients [19,20].

In patients with ovarian cancer, prior to avatrombopag initiation, increases in CA-125 levels in association with niraparib dose interruption occurred as a result of suboptimal niraparib dosing in association with thrombocytopenia. Additionally, in 2 patients with ovarian cancer, CA-125 levels fell in response to niraparib dose escalation enabled by concurrent avatrombopag therapy. This result indicates a potential for improved clinical outcomes in association with niraparib dose optimization enabled by avatrombopag therapy. Preliminary results in these 6 patient cases demonstrate the need for a confirmatory trial of avatrombopag for optimizing the dose of niraparib.

Conclusions

Responses in this small series of patients indicate a role for avatrombopag in optimizing the dose of niraparib in patients receiving maintenance treatment for ovarian cancer and breast cancer. In this case series, the adverse event profile of niraparib did not differ from those noted for avatrombopag in trials supporting its indication for use in immune thrombocytopenia. One potential limitation of this case series is the fact that thrombocytopenia induced by PARP inhibition might not share the same bone marrow suppression effect observed with chemotherapy-induced thrombocytopenia. Avatrombopag should be further investigated in thrombocytopenia induced by niraparib in a trial assessing bone marrow for megakaryocyte count over the course of niraparib therapy. These preliminary results, presented in the form of a case series, demonstrate the need for a large prospective trial to confirm our findings.

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