A Case of Ischemic Colitis Associated with Paclitaxel Loaded Polymeric Micelle (Genexol-PM®) Chemotherapy

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Introduction

Paclitaxel has been widely used for treating many solid tumors. Although colonic toxicity is an unusual complication of paclitaxel-based chemotherapy, the reported toxicities include pseudomembranous colitis, neutropenic enterocolitis, and on rare occasions ischemic colitis. Genexol-PM®, which is a recently developed cremophor-free, polymeric micelle-formulated paclitaxel, has shown a more potent antitumor effect because it can increase the usual dose of paclitaxel due to that Genexol-PM® does not include the toxic cremophor compound. We report here on a case of a 57-year-old man with advanced non-small cell lung cancer and who developed ischemic colitis after chemotherapy with Genexol-PM® and cisplatin. He complained of hematochezia with abdominal pain on the left lower quadrant. Colonoscopy revealed diffuse mucosal hemorrhage and edema from the sigmoid colon to the splenic flexure. After bowel rest, he recovered from his symptoms and the follow-up colonoscopic findings showed that the mucosa was healing. Since then, he was treated with pemetrexed monotherapy instead of a paclitaxel compound and platinum.

Key Words: Paclitaxel; Genexol-PM; Colitis, Ischemic

Case Report

A 57-year-old man was admitted to our hospital with intermittent loose stool for 3 days and subsequent bloody diarrhea with abdominal pain on left lower quadrant. He had been diagnosed as advanced NSCLC in April 2009, which revealed adenocarcinoma in histology, confirmed by endobronchial ultrasonography-guided transbronchial needle aspiration (EBUS-TBNA),
and multiple metastases to brain in imaging studies. After whole brain radiotherapy, he was treated with 230 mg/m² of polymeric micelle-formulated paclitaxel (Genexol-PM®) and 60 mg/m² of cisplatin. He was on medication due to diabetes mellitus and hypertension for 1 year, and he did not have any antplatelet agents or anticoagulants. He had never been treated for peptic ulcer, liver cirrhosis, or other hereditary disease that could have a tendency of hemorrhage.

On day 5 after the second cycle of chemotherapy, he complained of bloody diarrhea (up to 4 times, each amount of 100 mL) with abdominal pain. Vital sign was stable, and there was not any hemorrhagic appearance except hematochezia. Physical examination showed abdominal tenderness on left lower quadrant with mild distension, but did not reveal rebound tenderness or muscle guarding. Laboratory findings revealed: white blood cell count 9,900/mm³, absolute neutrophil count 7,870/mm³, hemoglobin 13.3 g/dL, platelet count 194,000/mm³, C-reactive protein (CRP) 0.50 mg/dL (0 ~ 0.3), serum protein 7.0 g/dL, serum albumin 4.2 g/dL, serum lactate dehydrogenase 573 U/L (218 ~ 472), serum aspartate transaminase 17 U/L, serum alanine transaminase 34 U/L, prothrombin time INR 1.29, and activated partial thromboplastin time 29.4 second. Afterwards, the white blood cell was not significantly changed, but CRP level was increased (21.88 mg/dL).

Plain radiography of abdomen did not show the obstructive pattern of small intestine or colon, and there was no evidence of free-air in abdominal cavity. Computed tomography (CT) of abdomen revealed long segmental and concentric wall thickening, called “double halo sign,” from sigmoid to distal transverse colon, without significant stenosis or obstruction in mesenteric vessels (Figure 1A). Gastroduodenoscopy showed shallow ulcer on antrum, but there was no evidence of hemorrhage. Colonoscopy revealed continuous, circular, and diffuse mucosal hemorrhage and edema from sigmoid colon to splenic flexure (Figure 1B). There were not the cobble stone-like appearance suggestive of “pseudomembrane” on examination. Biopsies were obtained separately each from involved mucosa in sigmoid to transverse colon. Pathologic examination of specimens revealed a mild edema, focal atrophic change of mucosal glands, and infiltration of erythrocytes into lamina propria, suggestive of early phase of ischemic colitis (Figure 1C). Stool culture and Clostridium difficile toxin assay were negative.

Then, from these colonoscopic and CT findings, a diagnosis of ischemic colitis was made. We maintained the fasting state with total parenteral nutrition for 12 days. During the bowel rest, his symptom of abdominal pain was gradually recovered and bloody diarrhea was completely resolved. CRP level was improved nearly to

Figure 1. (A) Computed tomography finding shows long segmental and concentric wall thickening, called “double halo sign” (arrow), from sigmoid to distal transverse colon, without significant stenosis or obstruction in mesenteric vessels; (B) Colonoscopy finding reveals continuous circular diffuse mucosal hemorrhage and edema from sigmoid colon to splenic flexure; (C) Pathologic examination of specimens revealed a mild edema, focal atrophic change of mucosal glands, and infiltration of erythrocytes into lamina propria, suggestive of early phase of ischemic colitis (H&E stain, ×400).
normal range (1.1 mg/dL). After 1 month from the onset, follow-up colonoscopic finding showed healing state of mucosa. Since then, he was treated with pemetrexed monotherapy instead of paclitaxel compound and platinum.

Discussion

This case showed a ischemic colitis as the colonic toxicity after chemotherapy with polymeric micelle-formulated paclitaxel and cisplatin in a patient with advanced NSCLC, and reported case of this adverse events in chemotherapy with newly developed agent, Genexol-PM®, is absent in Korea, so far.

Paclitaxel (Taxol®; Bristol-Myers Squibb, Wallingford, CT, USA) is a chemotherapeutic agent effective for many solid tumors. Dose-limiting toxicities by paclitaxel, such as neuropathy and neutropenia, may be related with the taxane compound itself or with the vehicle necessary to formulate the drug. Owing to water insolubility of the drug, paclitaxel is formulated with the micelle-forming vehicle, Cremophor-EL (CrEL), to enhance drug solubility. But, as mentioned above, the addition of CrEL has been reported to have relevance to hypersensitivity reaction, neuropathy, cytotoxic effect, inconvenience in preparation or alteration in pharmacokinetics of the drug. To overcome the unfavorable effects due to CrEL, the development of new CrEL-free formulations of paclitaxel have been attempted to use some of other drug delivery systems, such as emulsion, micelles, water-soluble prodrugs and conjugates. Genexol-PM®, as a outcome of these efforts, is a novel formulation of paclitaxel, a sterile lyophilized polymeric micellar formulation without CrEL. Genexol-PM® was found to have higher maximum tolerated doses and higher levels of biodistribution in a variety of tissues including liver, spleen, kidney, and lung and more importantly tumors. Also, the in vivo antitumor efficacy has been shown to be greater than that of Taxol®. In a phase II clinical trial, several adverse events associated with Genexol-PM® and cisplatin combination regimen were reported, which included neutropenia, peripheral sensory neuropathy, hypersensitivity reaction, and so on. But, in spite of higher paclitaxel doses, the side effects were comparable with conventional chemotherapy of paclitaxel and cisplatin. Moreover, the response rate and survival duration were more favorable than in most phase II or phase III clinical trials using Taxol® combined with cisplatin. Therefore, a cremophor-free, polymeric micelle-formulated paclitaxel (Genexol-PM®) combined with cisplatin has been reported to show significant antitumor effect with low incidence and severity of toxicity in spite of administration of higher doses of paclitaxel compared with CrEL-containing formulation in advanced NSCLC.

Colon toxicities associated with chemotherapy have the low incidence, but among them, pseudomembranous colitis and neutropenic enterocolitis are relatively common after paclitaxel combined chemotherapy. A few cases of ischemic colitis after taxane compound-containing chemotherapy, such as paclitaxel or docetaxel, Ischemic colitis caused by certain medications or drugs is unusual, and necessary to differentiate with other disease with bloody diarrhea in colon, such as gastrointestinal infections including Escherichia coli O157:H7, Clostridium difficile, and cytomegalovirus, diverticular diseases, Crohn’s disease, or malignancy with stricture. In this case, colonoscopic findings did not show the typical appearances of pseudomembrane, circular ulceration or crypt abscess suggestive of inflammatory bowel diseases, and stool culture and toxin assay revealed negative. Biopsy did not reveal the malignant lesion or viral infection, and CT findings did not show the evidence of diverticuli. Besides, white blood cell was within normal range at the time of admission. Therefore, we could rule out several diseases and make the diagnosis of ischemic colitis.

The exact mechanism by which paclitaxel-associated gastrointestinal toxicity occurs is poorly understood, but there are several postulations as to the onset of the complication. Paclitaxel may induce the necrosis of the gastrointestinal mucosa by taxane-based effect, arresting the mitosis of rapidly dividing colonic mucosal cells and also synergistic interaction with compromised bowel...
due to previous chemotherapy may play a part. Furthermore, in a study, paclitaxel had antiangiogenic properties in mice, interfering with vascular smooth muscle cell proliferation, migration, and neointimal accumulation. In fact, the patient had a history of diabetes mellitus and hypertension, and he was treated with the chemotherapeutic agent previously. So, the vascular insufficiency with underlying disease and the fragility of intestine due to previous chemotherapy would be the risk factors for the colonic toxicity.

Reported adverse effects on the gastrointestinal tract in chemotherapy with Genexol-PM are mild and transient. Because both Taxol® and Genexol-PM® have common component of paclitaxel, the nature of colonic toxicity may not reveal the dissimilarity. So, in escalating the usual dosage of the polymeric micelle-formulated paclitaxel, the monitoring for toxicities of gastrointestinal tract as well as hematologic abnormalities should be considered. Furthermore, as in this case, at the time of disease onset, suspicious drugs including polymeric micelle-formulated paclitaxel should be stopped, and the shift to other line of chemotherapeutic agent is considerable after the complete recovery of adverse events.

References