**Name of Journal:** *World Journal of Gastrointestinal Oncology*

**Manuscript NO:** 83231

**Manuscript Type:** CASE REPORT

**Acute respiratory distress syndrome and severe pneumonitis after atezolizumab plus bevacizumab for hepatocellular carcinoma treatment: A case report**

Cho SH *et al*. Severe pneumonitis after atezolizumab and bevacizumab in advanced HCC

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**Author contributions:** Cho SH, You GR, and Yoon JH contributed to analysis and interpretation of the data and drafting of the article; Park C, Lee JE, and Cho SG contributed to technical and material support; Choi SK and Cho SB contributed to critical revision and final approval of this article; Cho SH and You GR contributed equally to this work as co-first author.

**Supported by** National Research Foundation of Korea, No. NRF-2021R1F1A1061719.

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**Received:** January 13, 2023

**Revised:** February 1, 2023

**Accepted:** April 7, 2023

**Published online:**

**Abstract**

BACKGROUND

Hepatocellular carcinoma (HCC) is one of the most common cancers worldwide and has a high mortality. However, the treatment options for advanced HCC are limited to tyrosine kinase inhibitors, such as sorafenib and lenvatinib. Since previous regimens have an insufficient efficacy, the combination therapy of atezolizumab and bevacizumab (Ate/Bev) has been investigated, which showed an improvement in progression-free and overall survival. However, the adverse events of this combination therapy in advanced HCC have not been established. Herein, we report a novel case of an unresectable HCC and acute respiratory distress syndrome (ARDS) after a combination therapy of Ate/Bev.

CASE SUMMARY

An 82-year-old male visited our outpatient clinic for an incidentally detected liver mass. Liver magnetic resonance imaging and enhanced chest computed tomography (CT) were performed, which showed arterial hyperenhancement with washout in delayed phase suggesting HCC, and a well-defined metastatic solid nodule, respectively. F-18 fluorodeoxyglucose positron emission tomography (PET)-CT exhibited multiple hypermetabolic lesions in the iliac bone, lumbar vertebrae, and femur. Because of the high burden of the intrahepatic tumor, transarterial radioembolization was initially performed; after 37 d, a combination therapy of Ate/Bev was administered. The patient visited the emergency department three days after Ate/Bev treatment complaining of dyspnea. He was diagnosed with severe pneumonitis based on CT. Despite administering oxygen *via* a high-flow nasal cannula, the P/F ratio was only 74; therefore, the patient was diagnosed with ARDS based on the overall examination results. Low tidal volume with high positive end-expiratory pressure, sedative agents combined with a neuromuscular blocker, and a systemic steroid were promptly applied to manage the ARDS. However, the patient did not recover from the hypoxia and expired 31 h after being admitted.

CONCLUSION

Clinicians should be aware of severe pneumonitis due to the immune-related adverse events of this combination therapy, and patients should be closely monitored after therapy.

**Key Words:** Hepatocellular carcinoma; Systemic therapy; Adverse events; Pneumonitis; Atezolizumab; Acute respiratory distress syndrome

Cho SH, You GR, Park C, Cho SG, Lee JE, Choi SK, Cho SB, Yoon JH. Acute respiratory distress syndrome and severe pneumonitis after atezolizumab plus bevacizumab for hepatocellular carcinoma treatment: A case report. *World J Gastrointest Oncol* 2023; In press

**Core Tip:** Nowadays, the combination therapy of atezolizumab and bevacizumab is recommended as the first-line systemic treatment for advanced hepatocellular carcinomas. A global phase III study and recent real-world studies demonstrated rare life-threatening adverse events of atezolizumab and bevacizumab. However, our patient underwent acute respiratory distress syndrome and finally died three days after treatment. To the best of our knowledge, this is the first case of severe respiratory failure resulting in death with a very short interval from atezolizumab and bevacizumab administration. Therefore, we suggest close monitoring of lung toxicity after therapy.

**INTRODUCTION**

Hepatocellular carcinoma (HCC) is the sixth most common cancer and the third leading cause of cancer-related mortality globally[1]. HCC can be treated with various options depending on the tumor stage, remaining liver function, performance status score, and tumor burden[2]. Early-stage HCC has curative treatment options, such as surgical resection, radiofrequency ablation, or liver transplantation. However, only 10%–20% of patients are candidates for curative therapies, and 80% of patients have unresectable HCCs that can only be treated with locoregional therapies, such as transarterial chemoembolization, transarterial radioembolization (TARE), radiotherapy, or palliative management, including systemic therapy[3]. Despite the various locoregional therapies, many patients eventually progress to advanced stages of HCC. Therefore, systemic therapy for managing advanced HCC is clinically significant.

Since 2008, sorafenib, an oral multi-kinase inhibitor, has been used as a first-line systemic chemotherapy based on the SHARP trial, which demonstrated the increased overall survival of patients with advanced HCC compared with placebo-treated advanced HCC patients[4]. In 2018, lenvatinib, another oral multi-kinase inhibitor, was approved as a first-line therapy for advanced HCC according to the REFLECT trial, which demonstrated its non-inferiority over sorafenib in terms of overall survival[5]. However, in large-scale real-world GIDEON study, which comprised of 3,202 HCC patients, the presented median overall survival was 13.6 mo in Child-Pugh A group and 5.2 mo in the Child-Pugh B group. The ELEVATOR study, which investigated the real-world efficacy of lenvatinib, showed that the median overall survival was 10.7 mo in the Child-Pugh A group and 5.3 mo in the Child-Pugh B group. Efforts have been devoted to improve the clinical outcomes of patients with advanced HCC undergoing systemic chemotherapy.

With the introduction of various immune checkpoint inhibitors (ICIs), several clinical trials had attempted to verify their treatment efficacy. The CheckMate 459 trial compared the overall survival between nivolumab and sorafenib, of which nivolumab treatment did not significantly improve overall survival[6]. Also, the KEYNOTE-240 trial compared the overall survival and progression free survival between pembrolizumab and placebo in advanced HCC patients who were previously treated with sorafenib, and failed to reach statistical significance[7]. The IMbrave150 trial investigated the efficacy of the combination therapy of atezolizumab, an anti-PD-L1 antibody, and bevacizumab, a vascular endothelial growth factor (VEGF) -targeting antibody, for treating advanced HCC. The progression-free and overall survival were longer in the combination therapy group than in the sorafenib treatment group, with no differences in grade 3 or 4 adverse events[8,9]. Based on these results, many clinical guidelines have been updated to include atezolizumab and bevacizumab as a first-line systemic treatment for advanced HCC[2]. However, regarding the safety of atezolizumab and bevacizumab, few reports have described the adverse events, especially drug-related pneumonitis.

Herein, we present the novel case of a patient who had an unresectable HCC and experienced acute respiratory distress syndrome (ARDS) due to severe pneumonitis three days after being treated with atezolizumab and bevacizumab.

**CASE PRESENTATION**

***Chief complaints***

An 82-year-old male visited our outpatient clinic for an incidentally detected liver mass discovered on abdominal ultrasonography.

***History of present illness***

He denied fever, night sweats, weight loss, and abdominal pain.

***History of past illness***

The patient was on medication for hypertension and diabetes mellitus for 20 and 10 years, respectively. He also had underlying liver cirrhosis of an unknown cause, which was diagnosed 13 years ago.

***Personal and family history***

The patient had no family history.

***Physical examination***

There were no abnormal findings on physical examination.

***Laboratory examinations***

The laboratory workup showed the following: platelet count, 510 × 109/L (reference range, 130–450 × 109/L); total bilirubin, 0.63 mg/dL (reference range, 0.22–1.3 mg/dL); aspartate aminotransferase level, 37 U/L (reference range, 10–37 U/L); alanine aminotransferase level, 46 U/L (reference range, 10–37 U/L); albumin level, 4.0 g/dL (reference range, 3.5–5.2 g/dL); and prothrombin time, 0.97 international normalized ratio (INR; reference range, 0.8–1.2 INR). In the serum laboratory examinations for viral hepatitis, the hepatitis B surface antigen and hepatitis C antibody were negative, whereas the hepatitis B core antibody and hepatitis B surface antibody were positive, implying a prior hepatitis B viral infection. The alpha-fetoprotein level was 84.97 IU/mL, and the protein level induced by vitamin K absence-II (PIVKA-II) was 60232 mAU/mL.

***Imaging examinations***

Abdominal CT revealed a 16 cm × 11 cm × 10 cm mass with early arterial enhancement and early washout in the delayed phase. Several daughter nodules were found around the main mass. Ascites was not detected (Figure 1). Liver magnetic resonance imaging with Gadoxetate disodium (Primovist®) and enhanced chest CT were performed as the staging workup for the HCC, which showed showed arterial hyperenhancement with washout assessed in the portal venous phase in the enhanced T1-weighted images, and a 7-mm well-defined solid nodule suspected to be a metastatic nodule in the right middle lobe with interstitial lung abnormality (ILA) findings, such as bilateral subpleural reticulation and non-emphysematous cysts with traction bronchiectasis, respectively (Figure 1). Although the patient did not have respiratory symptoms such as dyspnea, pulmonary function tests were performed for ILA evaluation. His forced vital capacity (FVC) and 1-second forced expiratory volume (FEV1) were 1.87 L (52.4% of predicted value) and 1.39 L (63.2% of predicted value), respectively. The FEV1/FVC ratio was 109.7% of predicted value that showed restrictive patterns of lung disease. Because of the suspicious malignant metastatic pulmonary lesions, F-18 fluorodeoxyglucose (FDG) positron emission tomography (PET)-CT was performed to thoroughly evaluate the extrahepatic metastasis, which showed multiple hypermetabolic lesions in both iliac bones, lumbar vertebrae, and the right femur (Figure 2). A solid nodule in the right middle lobe also had mild focal hypermetabolic activity compared to the surrounding lung parenchyma with a maximum standardized uptake value of 1.0.

**FINAL DIAGNOSIS**

The patient was diagnosed with compensated liver cirrhosis, Child-Pugh A (score, 5) and HCC with multiple extrahepatic metastases [modified Union for International Cancer Control Stage (mUICC) IVB, Barcelona Clinic Liver Cancer (BCLC) C]. His performance status score was 0. He was willing to receive anticancer treatment. Although the patient demonstrated an extrahepatic metastasis, TARE of the primary intrahepatic HCC in the right lobe was planned to address the high tumor burden of the mass and to debulk it. Systemic therapy was administered for the remaining lesions.

**TREATMENT**

Prior to the TARE, a Technetium-99m macro aggregated albumin (99mTc-MAA) perfusion lung scan was performed to assess the hepatopulmonary shunt. The calculated lung shunt fraction was only 8.44%, and the TARE was conducted as scheduled. A hypervascular mass (about 15 cm) with several daughter nodules in the right hepatic lobe fed by branches of the right hepatic, left hepatic, and left inferior phrenic arteries was examined *via* angiography. An Yttrium-90 (Y90) infusion was performed by feeding the branches of the right hepatic artery, and the mean dose volumes were 108.419 Gy on the tumor, 85.543 Gy on the liver, and 0.873 Gy on both lungs (0.714 and 0.159 Gy on the right and left lungs, respectively). Additionally, chemoembolization using a doxorubicin-lipiodol mixture was performed through the branches of the right hepatic artery, while bland embolization of the left inferior phrenic artery was simultaneously conducted (Figure 3).

Postoperatively, the patient’s general condition was tolerable without abdominal pain, and no complications, such as liver failure or post-embolization syndrome, were observed. The patient was discharged after two days. A restaging baseline CT was performed 34 d after the initial TARE and before the systemic therapy. Abdominal CT showed more necrotic changes in the known primary HCC lesion, with some sparse lipiodol uptake in several daughter nodules. Chest CT showed no remarkable changes in the 7-mm metastatic nodule. Since the patient’s general condition was tolerable, systemic therapy using the combination of atezolizumab and bevacizumab was administered the day after the CT scan. For the systemic therapy, 1200 mg of atezolizumab and 870 mg (15 mg/kg body weight) of bevacizumab were infused for 60 and 90 min, respectively. No acute side effects occurred during the injections, and the patient returned home in the same day.

**OUTCOME AND FOLLOW-UP**

However, three days after the combination therapy of atezolizumab and bevacizumab, the patient visited the emergency room for severe dyspnea (Modified Medical Research Council Dyspnea Grade 4). The serum laboratory workup showed that the total bilirubin was 0.49 mg/dL (reference range, 0.22–1.3 mg/dL), aspartate aminotransferase level was 59 U/L (reference range, 10–37 U/L), and alanine aminotransferase level was 36 U/L (reference range, 10–37 U/L). The arterial blood gas test showed hypoxemia [partial pressure of oxygen (pO2), 84 mmHg] under 0.9 of the fraction of inspired oxygen (FiO2) level. Chest CT revealed newly detected diffused ground-glass opacities with bilateral septal thickening and consolidations (Figure 4). Infectious pneumonia was ruled out due to lack of related symptoms, such as fever or purulent sputum, and negative microbiological test (*e.g.*, sputum culture and respiratory virus exam) results. Although oxygen was administered *via* a high-flow nasal cannula, the P/F ratio was only 74; therefore, the patient was diagnosed with ARDS based on the overall examination results. The patient was immediately admitted to the intensive care unit after tracheal intubation. Low tidal volume with high positive end-expiratory pressure, sedative agents combined with a neuromuscular blocker, and a systemic steroid (125 mg of methylprednisolone per day) were promptly applied to manage the ARDS. However, the patient’s P/F ratio worsened, and he did not recover from the hypoxia. Unfortunately, he expired 31 h after being admitted.

**DISCUSSION**

The paradigm in the systemic treatment of HCC has shifted from conventional tyrosine kinase inhibitors (TKI) to a combination of an ICI with a TKI or another ICI[10]. According to the IMBrave150 trial, a pivotal study on the use of atezolizumab and bevacizumab for unresectable HCCs, the progression-free survival of this combination treatment was better than that of sorafenib[8]. Follow-up results demonstrated a meaningful survival benefit of the combination therapy of atezolizumab and bevacizumab[9]. Therefore, the HCC treatment guidelines have been updated, including the combination therapy of atezolizumab and bevacizumab as a first-line therapy for advanced HCCs[2,11]. Conventional TKI agents, such as sorafenib, lenvatinib, and cabozantinib, are considered second-line treatments or alternatives to the first-line treatment for ineligible patients for the combination therapy of atezolizumab and bevacizumab.

In addition to treatment efficacy, the safety profile of drug administration is an essential concern. Conventional TKI agents have various treatment-related adverse events, such as diarrhea, hypertension, proteinuria, and skin toxicities, such as rash, desquamation, and hand-foot skin reactions [12]. The IMBrave 150 trial showed a better safety profile for atezolizumab and bevacizumab compared with that of sorafenib. The most common grade 3 or grade 4 adverse events with an incidence of more than 10% in the IMBrave150 trial were hypertension (15.2%), increased aspartate aminotransferase levels (7.0%), and increased alanine aminotransferase levels (3.6%), which were mostly consistent with the follow-up study. Although 1.2% of the patients experienced grade 1–2 pneumonitis, no patients with severe grade toxicity were found. This finding was consistent with the follow-up study (2.0% of patients had grade 1–2 pneumonitis).

Considering the timeline of the combination therapy of atezolizumab and bevacizumab into clinical practice, few real-world studies regarding the treatment efficacy and safety profile have been completed, reporting conflicting results. D'Alessio *et al*[13] conducted a multi-national, retrospective study on 202 patients with HCC treated with the combination of atezolizumab and bevacizumab. They observed that 1.0% of the patients experienced treatment-related pneumonitis of more than grade 3. Tada *et al*[14] compared the results of the combination of atezolizumab and bevacizumab in elderly (≥ 75 years old) and nonelderly (< 75 years old) HCC patients, and revealed that the treatment efficacy and safety did not differ between the groups, and reported no drug-related pneumonitis. However, Ng *et al*[15] performed a study on the immune-related adverse events (irAEs) associated with the use of ICIs, including atezolizumab, in advanced HCCs, and found that pneumonitis occurred in 3.0%–10.0% of the patients (2.4% of which had pneumonitis of grade 3 or higher) and the pneumonitis was lethal in 0.2%–2.0% of the patients. Additionally, Endo *et al*[16] recently reported two fatal cases after applying atezolizumab plus bevacizumab on patients with pre-existing lung diseases. One patient showed honeycomb lungs before treatment and died 5 d after treatment. Another patient previously underwent right lower lobectomy due to lung adenocarcinoma. She died after 11 courses of treatment.

Pneumonitis is an uncommon but possible adverse event closely related to the use of various ICIs in HCC patients. The KEYNOTE-240 trial studied the use of pembrolizumab, an anti-PD-1 monoclonal antibody, in HCC patients, and reported an occurrence of pneumonitis of any grade in 18.3% of the patients, and grade 3 or 4 pneumonitis in 7.2% of the patients[17]. Lung-related irAEs due to ICIs have been investigated in other malignancies, such as lung cancer, renal cell cancer, and melanoma[18]. Lung-related irAEs should be managed based on the grade of pneumonitis. In patients with grade 3 or grade 4 pneumonitis with severe symptoms and life-threatening respiratory compromise, ICI use must be permanently discontinued, and glucocorticoids and empirical antibiotics must be administered[19]. Discontinuing ICI therapy and initiating systemic steroids are highly effective treatments for ICI-related pneumonitis. However, ICI-induced pneumonitis accounts for 35.0% of PD-1/PD-L1 inhibitor-related deaths and may have a similar fatal clinical course to that of our case[20].

Although the associated factors for predicting the development of pneumonitis have been investigated, limited data are available. Decreased pulmonary function and a history of smoking likely increase the risk of pneumonitis[21]. Concurrent radiation therapy, combination immunotherapy, and previous high-dose chemotherapy have also been proposed as potential risk factors[22]. Furthermore, the presence of ILAs is associated with the risk of complications from medical interventions, such as chemotherapy and surgery[23]. Chest CT done on our patient before atezolizumab and bevacizumab therapy showed ILA findings, which may have influenced the rapid onset of ARDS. Although our patient underwent combination therapy of ICI and TKI a meta-analysis by Nishino *et al*[24] reported significantly higher incidences of all-grade pneumonitis (6.6% *vs* 1.6%; *P* < 0.001) and grade 3 or higher pneumonitis (1.5% *vs* 0.2%; *P* = 0.001) in the ICI-combination therapy group than in the monotherapy group. Despite varying reported onset times of pneumonitis after treatment initiation (range, 9 d–19 mo)[21], the median time to onset is reported to be shorter when a combination therapy of ICIs is used[25,26]. The presenting symptoms of ICI-induced pneumonitis are nonspecific, including dyspnea (53.0%), cough (35.0%), and fever (12.0%)[17], and the incidence level for these risk factors in clinical practice in terms of atezolizumab and bevacizumab combination therapy is not yet established. Therefore, clinicians must be aware of the possibilities and characteristics of immune-related pneumonitis associated with these treatments.

This study has some limitations. First, thorough post-treatment pulmonary pathological and laboratory examinations may have clarified the patient’s diagnosis; however, a biopsy was not eligible because of the rapid course of the ARDS, and the family did not want an autopsy. Additionally, several serum biomarkers, such as Krebs von den Lungen-6 (KL-6), were related to the diagnosis of interstitial lung disease. If the results of the biomarkers were available, an acute exacerbation of interstitial lung disease could have been ruled out; unfortunately, these laboratory tests were not performed. Second, radiation pneumonitis due to the initial TARE likely occurred. However, the calculated radiation dose to both lungs was 0.873 Gy (left lung, 0.159 Gy; right lung, 0.714 Gy), which was lower than the recommended dose for preventing pneumonitis[27]. Considering that the CT results were from three days before the combination therapy of atezolizumab and bevacizumab, the possibility of radiation pneumonitis due to the TARE was substantially low. However, although respiratory symptoms such as cough, dyspnea and immune-associated pneumonia are well-known ICI-related adverse events, fatal ARDS with an extremely short duration from therapy, as was in our case, has not been reported; therefore, our case report may be of value.

**CONCLUSION**

Our case implies that the combination therapy of atezolizumab and bevacizumab to treat HCCs might cause fatal pneumonitis leading to ARDS; however, the benefit of this treatment could outweigh the irAEs in terms of survival. Additionally, the severe adverse events of the newly introduced atezolizumab and bevacizumab therapy are uncommon. However, clinicians should be aware of the possible lung toxicity caused by this treatment, especially in newly developed respiratory symptoms of patients. Once the diagnosis is made, management based on the severity of pneumonitis is highly required. Furthermore, future studies should verify the risk factors of developing pneumonitis to identify high-risk patient groups.

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**Footnotes**

**Informed consent statement:** Written informed consent was obtained from the patient’s wife for the publication of this case report and any accompanying images.

**Conflict-of-interest statement:** All the authors report no relevant conflicts of interest for this article.

**CARE Checklist (2016) statement:** The authors have read the CARE Checklist (2016), and the manuscript was prepared and revised according to the checklist.

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**Provenance and peer review:** Unsolicited article; Externally peer reviewed.

**Peer-review model:** Single blind

**Peer-review started:** January 13, 2023

**First decision:** January 21, 2023

**Article in press:**

**Specialty type:** Gastroenterology and hepatology

**Country/Territory of origin:** South Korea

**Peer-review report’s scientific quality classification**

Grade A (Excellent): 0

Grade B (Very good): B

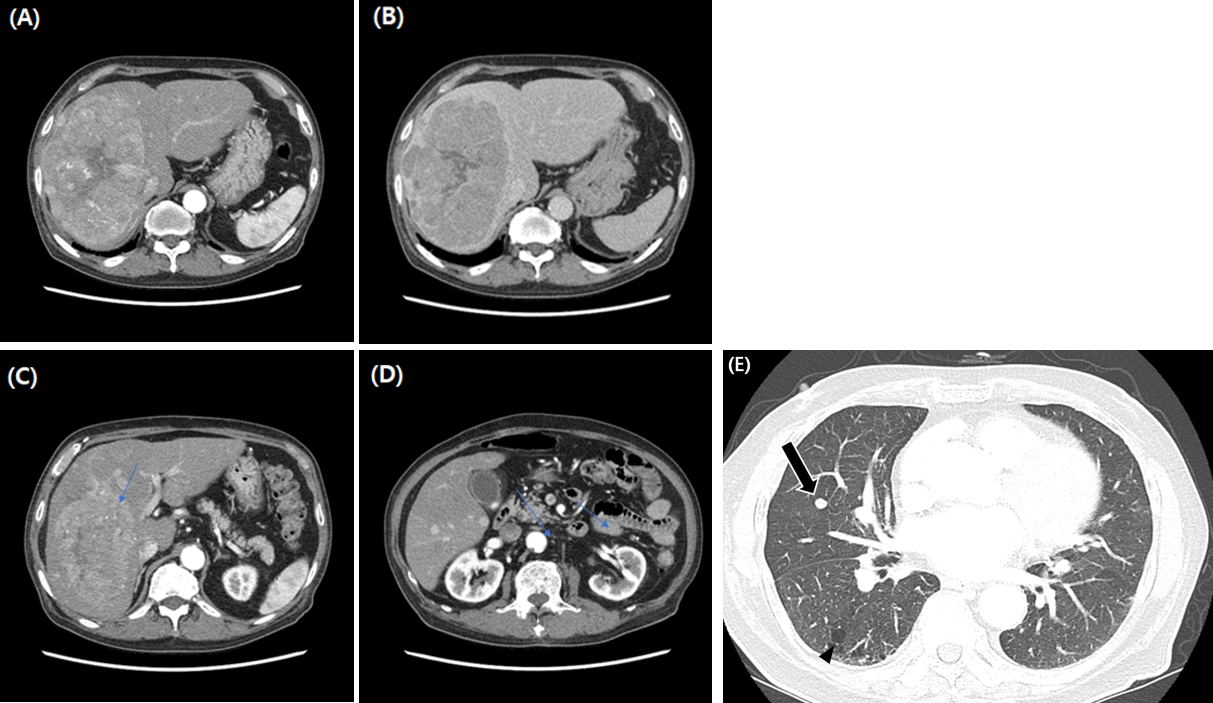
Grade C (Good): C

Grade D (Fair): 0

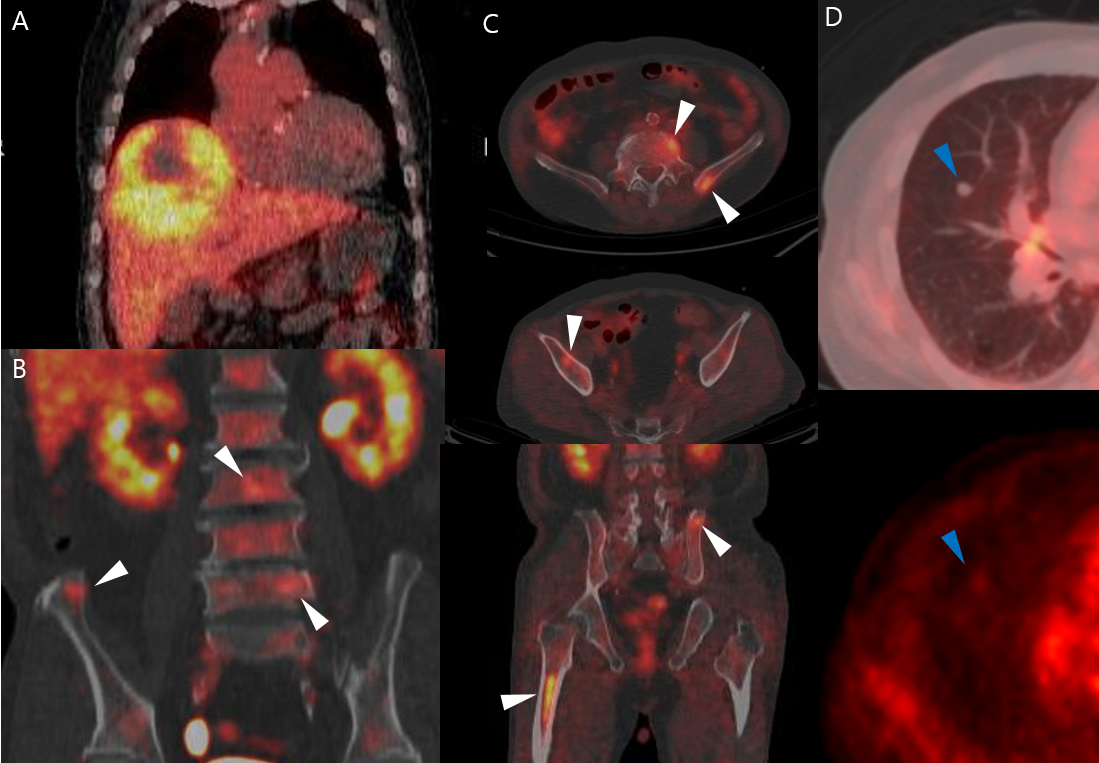
Grade E (Poor): 0

**P-Reviewer:** Lin J, China; Martin M, France **S-Editor:** Liu JH **L-Editor:** A **P-Editor:** Liu JH

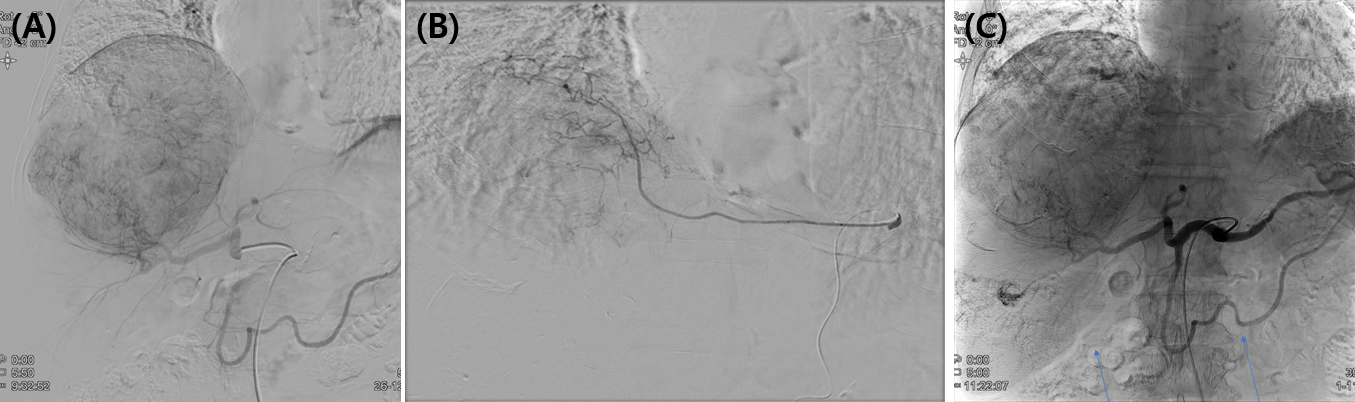
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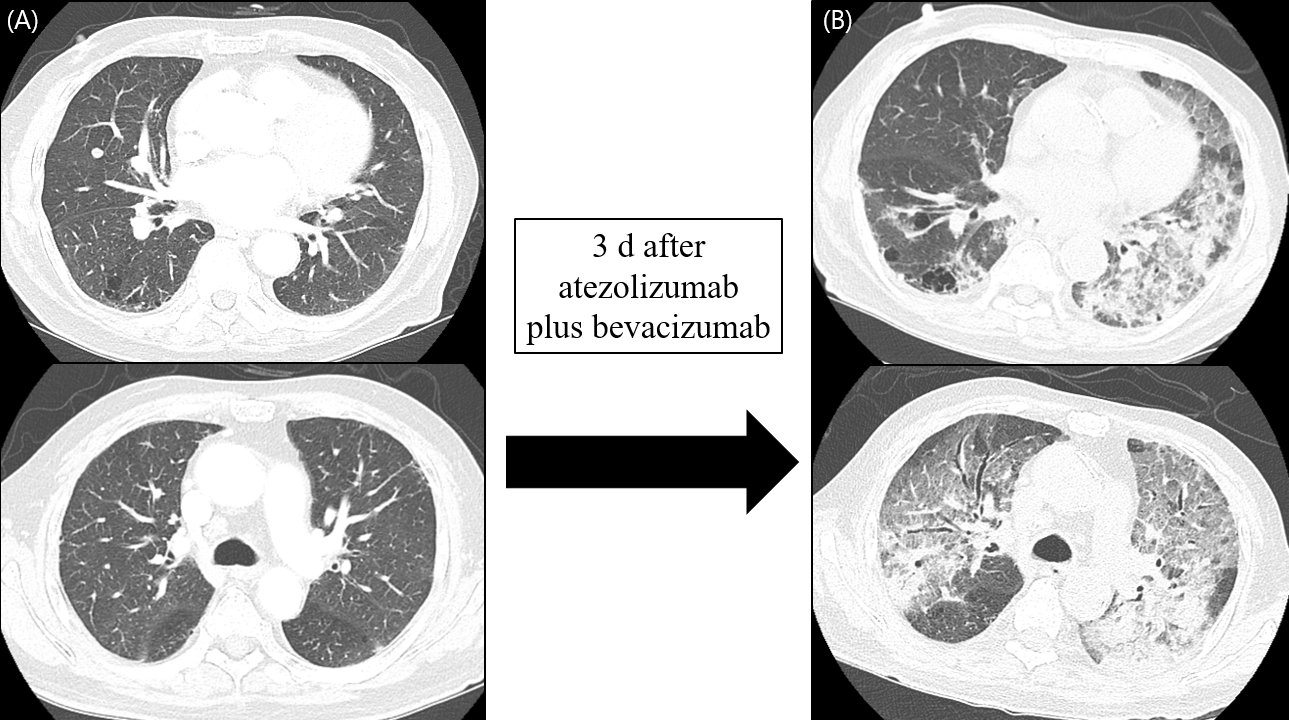
**Figure 1 The patient’s abdominal dynamic computed tomography (CT) at the initial diagnosis.** (A, C, D; arterial phase, B; delayed phase). A and B: An oval mass with a size of 16 cm × 11 cm × 10 cm was located in the right hepatic lobe with early enhancement and delayed washout features; C and D: Several satellite nodules were examined in the liver (blue arrows); E: The lung window of the transverse CT scan obtained at the level of the inferior pulmonary veins shows a well-defined round nodule, suspected to be a metastatic nodule, in the right middle lobe (arrow), as well as subpleural reticulation and non-emphysematous cysts (arrowhead).



**Figure 2 F-18 fluorodeoxyglucose positron emission tomography/computed tomography images.** A: A large hypermetabolic tumor was noted in the right hepatic lobe [maximum standardized uptake value (SUVmax) 5.1]; B and C: Multiple hypermetabolic lesions (SUVmax 4.8) are seen in both iliac bones, lumbar vertebrae, and the right femur (white arrowheads); D: A solid nodule with mild hypermetabolic activity was noted in the right middle lobe (blue arrowheads).



**Figure 3** **Digital subtraction angiography and completion angiography after transarterial radioembolization and chemoembolization.** A: The common hepatic artery angiogram shows a large hypervascular staining of the main mass. The satellite nodules of the right inferior lobe are not identified in this image. No tumor staining is found at the cranial portion of the tumor; B: The left inferior phrenic artery angiogram shows a hypervascular staining of the cranial portion of the tumor. This branch was embolized with tris-acryl gelatin microspheres and gelatin sponge particles; C: The completion angiogram shows the decreased staining of the main mass and lipiodol-laden tumors (blue arrows) in the right hepatic lobe.



**Figure 4 Development of acute respiratory distress syndrome three days after the combination therapy of atezolizumab and bevacizumab.** The lung window of the transverse computed tomography (CT) scan obtained at the level of the left main pulmonary and main pulmonary trunk arteries shows the mixed areas of ground-glass opacity and bilateral consolidation with the anteroposterior gradient (B), compared to baseline CT which was scanned 3 and 4 d prior to atezolizumab and bevacizumab therapy and acute respiratory distress syndrome, respectively, (A) which demonstrated no inflammatory changes and minimal pulmonary fibrosis. A: The common hepatic artery angiogram shows a large hypervascular staining of the main mass. The satellite nodules of the right inferior lobe are not identified in this image. No tumor staining is found at the cranial portion of the tumor; B: The left inferior phrenic artery angiogram shows a hypervascular staining of the cranial portion of the tumor. This branch was embolized with tris-acryl gelatin microspheres and gelatin sponge particles; C: The completion angiogram shows the decreased staining of the main mass and lipiodol-laden tumors (blue arrows) in the right hepatic lobe.