Severe dyspnoea in a patient with chronic myelogenous leukaemia on a tyrosine kinase inhibitor

Celalettin Ustun,1 Nicole Randall,1 Eitan Podgaetz,2 Khalid Amin,3 H Erhan Dincer4

NR (Haematology fellow): A 78-year-old male patient with chronic myelogenous leukaemia (CML) diagnosed initially in 2011 presented to the haematology clinic with fatigue and dyspnoea at rest and with exertion. His medical history included benign prostatic hyperplasia and a prior diagnosis of acute myelogenous leukaemia (AML) with translocation 8;21 (good risk AML) diagnosed in 2009, which was in complete remission after standard cytarabine and daunorubicin induction chemotherapy and cytarabine consolidation chemotherapy, requiring no treatment since January 2010. He had also been a smoker, with a 40 pack-year smoking history, who quit 40 years ago. His chest X-ray and CT from 2012 prior to nilotinib therapy demonstrated no focal airspace disease, no evidence of small airways disease and no pleural effusion (figure 1). In July 2013 prior to this current presentation, he had developed a pleural effusion while receiving dasatinib as a second-line agent because of suboptimal response to imatinib. Pleural fluid analysis at that time demonstrated an exudative effusion with 81% of lymphocyte predominance (395 white blood cells, WBCs) and negative staining and cultures for bacterial, fungal and acid fast bacilli. It was consistent with dasatinib-induced pleural effusion. Dasatinib was switched to nilotinib, and pleural effusion resolved until October 2014 when he presented again with fatigue and dyspnoea to the clinic.

On examination, the patient’s heart rate was 82 bpm, blood pressure 115/70 mm Hg, respiratory rate 24 breaths/min, SpO2 94% and temperature 97.3°F. The patient was in mild respiratory distress, actively working to take deep breaths. Respiratory examination was remarkable for poor air entry at the lung bases bilaterally with dullness to percussion to the mid-lung fields posteriorly, right side greater than left. There was no egophony or tactile fremitus. Cardiac examination revealed a regular rhythm with a systolic ejection murmur radiating to the carotid arteries bilaterally.

His WBC count was 7.6×10⁹/L with a normal differential except for an elevated absolute monocyte count of 2.2×10⁹/L. Haemoglobin was 14.0 g/dL and platelet count was 165×10⁹/L. Comprehensive metabolic panel demonstrated normal electrolytes and kidney and liver functions. BCR/ABL1 fusion gene was undetectable by PCR in the peripheral blood (ie, CML was still in molecular remission). Chest CT showed marked diffuse thickening of the pleura in the left hemithorax with a moderate size, slightly loculated pleural effusion (figure 2A).

Echocardiogram revealed normal left ventricular ejection fraction (EF) of 55% with mild diastolic dysfunction and was otherwise unchanged from baseline echocardiogram in October 2013, which also demonstrated mild diastolic dysfunction, normal left ventricular EF of 55%–60%. Baseline ECG in August 2013 demonstrated a normal sinus rhythm, with normal axis and no ST or T wave abnormalities, and current ECG was unchanged.

In addition, rheumatologic work-up was unremarkable, with negative ANA, p-ANCA, c-ANCA, scleroderma antibody, Smith antibody, SSA/SSB, RNP Jo-1 antibody and rheumatoid factor testing.

CU (Staff haematologist): CML is a haematopoietic stem cell malignancy with an incidence rate of 1.5 per 100 000 individuals per year within the USA.1 CML is characterised by the presence of the Philadelphia (Ph) chromosome, which results from a translocation between chromosomes 9 and 22, which results in the pathogenic tyrosine kinase signal transduction protein, BCR/ABL1. The treatment of CML has been dramatically changed by the use of tyrosine kinase inhibitors (TKIs). The first developed was imatinib, which is effective in the majority of patients. Few patients need second-line TKIs (eg, dasatinib or nilotinib) because of resistance or intolerance.2 Second-line TKIs are more effective but also have specific adverse effects, including pleural effusion.

In this patient’s case, CML was in molecular remission, and the current complete blood counts had no indication of disease progression. His haemoglobin was normal, which could not have been responsible for his dyspnoea.

TKIs have been reported to cause pleural effusions, which are predominantly lymphocytic, and can cause fluid retention.3 Lymphocytosis in patients receiving dasatinib has been shown to be associated with pleural effusion and better outcomes.3 Although these are rare side effects, dose reduction, interruption of TKI or switching to another TKI may resolve the recurrence of pleural effusion.4 Pleural effusions are described with TKI therapy, in particular with dasatinib.5 The exact mechanism is not known but postulated to be related to fluid retention from non-specific inhibition of platelet-derived growth factor receptor-β or other kinases,5 or immune related.6 History of cardiac disease, hypertension and use of a twice-daily schedule (vs once daily) were identified as risk factors for the development of pleural effusions.7 Perhaps, we are dealing with recurrence of pleural effusion, although this patient was no longer on dasatinib, but rather on nilotinib. Pleural effusions
are uncommonly associated with nilotinib therapy (1%). However, at this point there were four important questions: (1) Is nilotinib the sole cause of the patient’s pleural effusion? (2) Is the pleural effusion solely responsible for his dyspnoea? (3) What is the reason for the pleural thickening seen on the patient’s CT chest imaging, which to our knowledge has not been reported with TKIs? (4) What is the best diagnostic approach to investigate the aetiology of the pulmonary thickening seen on CT chest imaging?

HED (Staff pulmonologist): Although large pleural effusion and diffuse pleural thickening may cause restrictive pulmonary physiology leading to dyspnoea, we also entertained other possibilities in his case such as pulmonary hypertension related to sleep apnoea. However, his echocardiogram was not suggestive of increased pressures in the pulmonary circulation, and there was no evidence of left ventricular systolic dysfunction but mild diastolic dysfunction. Exudative pleural effusion with diffuse pleural thickening has a broad differential diagnosis of chronic infections, such as tuberculosis or fungal infections, and can be seen in partially treated bacterial infections and malignancies (primarily pleural lymphoma or mesothelioma). Diagnostic yield of pleural fluid cultures or cytology is much less than that of pleural biopsy, tissue cultures or pathological evaluation. Therefore, pleuroscopy, a minimally invasive method, allows for examination of pleural surfaces, both parietal and visceral, and the ability to obtain biopsies from the parietal pleura under direct visualisation.

HED and ER (Staff thoracic surgeon): Pleuroscopy with biopsies and pleural fluid drainage was performed for diagnostic purposes. Intraoperatively, we found diffusely thickened parietal pleura with multiple adhesions and fibrotic changes creating innumerable pockets of loculations. Parietal pleural biopsies showed organising fibrinous pleuritis (OFP) with no evidence of abnormal cellular infiltrate, granuloma or malignancy (figure 3). Tissue and fluid cultures from the pleura and pleural effusion were negative for bacteria, fungi, actinomycetes and acid fast bacilli. The effusion was predominantly composed of lymphocytes. Flow cytometry of pleural fluid was normal and demonstrated polytypic B cells, with no aberrant immunophenotype on T cells and rare to absent blasts.

HED and KA (Staff pathologist): OFP can coexist with various malignant or infectious conditions in the pleural space. Pleuritis is a reactive process which may include inflammatory cell infiltrates. There may be fibrinous exudates on the pleural surface (fibrinous pleuritis), organisation of the fibrin by granulation tissue (organising pleuritis) or maturing connective tissue (fibrous pleuritis). OFP is an acellular fibrous thickening of the pleura with focal plaque-like features. OFP can be associated with various disorders, including cardiovascular disease, tuberculosis, mesothelioma, lymphoma and metastasis.

CU: All the causes of OFP were excluded, and therefore the most likely possible explanation as a cause of pleural thickening was TKI; perhaps, the patient’s OFP started with dasatinib use and worsened with nilotinib use with time. This may represent one of the first cases of this phenomenon; however, it is not possible to definitely identify this finding as this is a diagnosis of exclusion. To our knowledge, this is the first case reported in the English literature. Interestingly, there appears to be no evidence that TKIs, such as nilotinib, contribute to fibrosis in the heart or lung; rather, they may exert antifibrinotic effects, as seen in liver and lung animal models. The possible antifibrinotic...
He was discharged to a rehabilitation centre and was followed up frequently. A month later, his pancytopenia improved; he was off steroids and nasal oxygen support. However, he still had exertional dyspnoea. Chest CT and pulmonary function testing (PFT) were repeated.

HED: His chest CT scan showed minimal pleural effusion, no airspace disease and unchanged pleural thickening (Figure 2B). PFTs showed a restrictive pattern (total lung capacity (TLC) 52% of predicted), which was much worse than a year ago when TLC was 72% of predicted. The pleural thickening seen on chest CT is likely responsible for the restrictive process seen on PFTs and explains his improved, but ongoing dyspnoea.

CU: Although his respiratory symptoms and pleural effusion improved, pleural thickening persisted. It is unknown whether fibrous pleuritis is irreversible at this point of time, but clearly it has not improved in the 2 months since nilotinib was discontinued. Moreover, he developed severe pancytopenia on the alternative drug, omacetaxine. This case demonstrates that pleural complications of TKI can be severe and hinder treatment of CML. Almost half the number of patients with CML on second-generation TKIs experience side effects, including pleural effusion in 26% of patients on dasatinib and 2% on nilotinib. Side effects are often a reason for the difficulty patients have with adhering to these lifelong drugs. Unfortunately, although there are multiple TKIs, there can be cross reactions between them regarding adverse effects. Other non-TKI options are limited and have different but severe toxicities, as demonstrated in our patient. Allogeneic haematopoietic cell transplantation can be considered in younger patients who cannot tolerate TKIs; however, this is not a good option for this 78-year-old patient.

However, long-term outcomes in patients who are able to tolerate TKI therapy are good with an overall 5-year survival of 93%, but this decreases in older adults (defined as age 30–85 years) to 82%.

Contributors All contributing authors were involved in the care of the patient. CU planned and prepared the first draft of the manuscript. All authors contributed to the development of the final version of the manuscript. CU is responsible for the overall content of the manuscript as guarantor.

Competing interests None declared.

Patient consent Obtained.

Provenance and peer review Not commissioned; externally peer reviewed.

REFERENCES


