LETTERS TO THE EDITOR

Chronic Myeloproliferative Neoplasm with Concurrent BCR-ABL Translocation and JAK2 Mutation

Sir,

Chronic myeloproliferative neoplasms (CMNs) are clonal disorders of haematopoietic stem cells and are classified according to their clinical and phenotypical features and genetic aberations.1,2 CMNs are divided according to the presence of a Philadelphia chromosome/BCR-ABL fusion as chronic myeloid leukaemia (CML Ph+). Philadelphia chromosome negative (Ph-CMNs) conditions are polycythemia vera (PV), essential thrombocythemia (ET) and primary myelofibrosis (PMF). More than 95% cases of PV and 60% of ET and PMF are positive for JAK2 mutations.3

Among more than 100 published cases of Philadelphia chromosome positive chronic myeloid leukaemia (Ph+ CML) investigated for Jans kinase mutation, no mutated case has been found.4,5 Selected patients with Ph+ CML with marked thrombocytosis prove to be negative for the JAK2 mutation.5 We report here a case of untreated patient of chronic myeloproliferative neoplasm with concurrent BCR-ABL translocation and JAK2 mutation. This 50-year-old man presented with splenomegaly, leuкоcito sis (white blood cell count 27300/ul), haemoglobin (Hb) level of 12.7 g/dl and thrombocytes (platelet count 248000/ul), neutrophils at 57%, myelocytes at 30%, metamyelocytes at 6% and blasts at 2%.

Bone marrow aspiration revealed diluted cellularity, normoblastic erythropoiesis, myelopoiesis hyperplastic with left shift and megakaryocytosis. Sections of bone marrow trephine showed mostly bony trabeculae, however, haematopoietic tissue was fairly adequate. Architecture was preserved in the larger portions of the marrow with normal haematopoiesis. Other areas however, showed increased fibrosis and fibroblastic activity with loss of normal marrow architecture. In those areas, megakaryocytes were increased, whereas other haematopoietic elements were uniformly depressed. Plasma cells were prominent. Reticulin was increased Grade-II/III. BCR-ABL translocation by FISH was detected in 25% of the 500 nuclei counted. PCR was positive for JAK2 mutation.

This case demonstrates that in untreated patient of chronic myeloproliferative disorder BCR-ABL and JAK2, mutation may be concomitantly detectable in haematopoietic cells of a single patient. Four well documented cases with concurrent JAK2 and BCR-ABL translocation were reported by Hussain et al.6 It has been hypothesized by Kralovics et al.7 that, a varying combination of different molecular defects in a pathologic stem cell might be responsible for the phenotypic heterogeneity as seen in this case. The frequency of BCR-ABL translocation and concurrent JAK2 mutation might be low. Further studies on large cohorts are needed to clarify the co-existence of JAK2 mutation and BCR-ABL translocation in CMNs, as the sensitivity of newer technique of mutated alleles improve. At present, patients with concurrent BCR-ABL translocation are being treated with TKI’s imatinib, nilotinib and desatinib. No drug is available to treat JAK2 mutation. The natural history of these patients and their prognosis is unknown and needs further follow-up.

REFERENCES


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Interferences or Real ECG Changes?

Sir,

A 76-year-old female with previous history of hypertension and hypercholesterolemia was admitted. She had 6 months history of intermittent episodes of complete loss of consciousness, however, she denied any symptoms suggestive of epilepsy, shortness of breath, palpitations, light-headedness, dizziness or chest pains either preceding or after the episodes of blackouts. She did however sustain bruises with each episode and was, therefore, increasingly scared to live on her own. During her admission, she was extensively investigated with normal baseline blood tests, normal CXR, a normal CT brain, a normal 12 lead ECG, several normal ambulatory ECG recordings and a normal trans-thoracic echocardiogram. She underwent a treadmill exercise stress test (standard Bruce protocol) and managed 96% of her age predicted heart rate, with no ECG changes or symptoms. In view of an unclear diagnosis, she was recommended for implantable loop recorder (ILR). Whilst awaiting ILR implantation, a review of her notes by a senior doctor in the team revealed a rhythm strip recording (obtained at 04:27 hours on the day of admission (Figure 1).

A junior doctor had labelled Figure 1 as “interference” only, following which it was filed in the notes! The strip actually showed a 7-second pause with an escape ventricular beat, followed by restoration of normal sinus rhythm. In view of this rhythm strip, she was recommended for implantation of a dual chamber permanent pacemaker. Patient remained asymptomatic on 9 months follow-up with no further episodes of blackouts.

ECG interference has been reported with the use of electrocautery during the surgical procedure, use of mobile phones, those with hearing aid, permanent pacemaker and the use of portable digital media player in patients with implantable loop recorders. In this case the patient had no external devices and the interference was probably motion artifacts.

The ECG rhythm strip had been wrongly labelled as interference. An appropriate early interpretation could have saved valuable time and cost to the patient.

REFERENCES


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Figure 1: Rhythm strip recorded on ECG.