Successful long-term treatment with azacitidine in patient with chronic myelomonocytic leukemia

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Abstract

The purpose of this article was to present a case of successful long term treatment with azacitidine in patient with Chronic Myelomonocytic Leukemia (CMML) and discussing possible contributing factors for its long term efficacy. Data from our case were compared with similar data available in the literature. Effective treatment with azacitidine resulted in overall survival of 11 years 5 months and we showed that applying multiple cycles of treatment is feasible. Our patient received 71 cycles of treatment with total duration of 7 years and 3 months. Our report about a patient with CMML and a good clinical course revealed, that long term treatment with azacitidine is feasible in some patients. Initially low bone marrow blast count, a relatively small malignant CMML clone, reduction of spleen size and fast platelet response seemed to be factors determining long term response to treatment in our patient. More data on CMML treatment by Hypomethylating Agents and their analysis are needed in order to make firm conclusions.

Introduction

CMML is clinically diverse malignant clonal hematopoietic stem cell disorder characterized by persistent peripheral monocytosis, bone marrow dysplasia and with features of both myelodyplastic and myeloproliferative neoplasm.1 Generally, long term prognosis is poor with a median overall survival of 30 months and a relatively high risk of leukemic transformation, up to 30%.2 Hypomethylating agents belong to a class of drugs which inhibit DNA methyltransferase and consequently reduce DNA methylation. Utilization of HMA in CMML has been reported with seemingly unsatisfying results with overall response rates between 30 and 60% and median overall survival between 12 and 37 months.3,4 Previously only a few predictive factors (increased bone marrow blast count and proliferative features) were found to be associated with shorter survival.5 Surprisingly only age, but none of the other disease characteristics, was a prediction for a response.6

In this case report, we describe an effective long term use of azacitidine in a patient with CMML and tried to find factors, which may contribute to long term azacitidine efficacy.

Case Report

A 58-years-old male patient presented with a moderate anemia (hemoglobin 93 g/L) and thrombocytopenia (platelet count 112x10^9/L). Because he was clinically asymptomatic, we decided for a “watch and wait” approach with a regular outpatient visits. His blood count remained stable without treatment for another 3 years. Later on, a gradual increase in leukocyte count was noticed with predominant monocytosis. Bone marrow work up morphologically revealed CMML with 6% of blast cells. Cytogenetics showed a normal karyotype. For the four following years, he did not need any therapy. In the meantime, he suffered from progressive joint and muscles pain and had platelet count ranging between 50x10^9/L and 140x10^9/L. Fluctuations in platelet count may have suggested accompanied autoimmune mechanism, immune thrombocytopenia.7

At that point the treatment was considered due to increased leucocyte count. His blood count was: WBC 68.5x10^9/L, Hemoglobin 101 g/L and platelets 101x10^9/L. Blood differential showed: 40.8x10^9/L neutrophils (59.6%), 23.1x10^9/L monocytes (33.8%), 4.2x10^9/L lymphocytes (6.1%) without immature granulocytes or blasts. The patient did not have any clinical symptoms, spleen size was 2cm below the left costal margin. After 4.5 years from the initial diagnosis, reexamined bone marrow cytogenetics revealed balanced translocation between homologous chromosomes 11, t(11;11)(p15;q11) in otherwise small clone of metaphases, 15%. The aberration is not recurrent and was not described in CMML before. Based on MD Anderson Prognostic Scoring System (MDAPS) for CMML, the patient was in an intermediate-2 risk group.8 Patient began treatment with azacitidine 75mg/m2 daily 7 days per month. After 2 cycles, treating hematologist tampered the dose to a fixed 75mg for following 4 cycles, due to neutropenia. From cycle 7 onwards he again received a full dose.

Although he never achieved complete remission, we observed clear hematological response and clinical improvement. We evaluated the response according to “International consortium proposal of uniform response criteria for (MDS/MPN)”.9 Clinical improvement was observed after 4 cycles of azacitidine. Besides monocytosis, the patient improved to a 1.4x10^9/L, platelets increased to 250x10^9/L and he became red cells transfusion independent, with hemoglobin level 110 g/L. Dynamic of monocyte and platelet counts were dependent upon azacitidine administrations (Figure 1). Eleven azacitidine cycles resulted in a partial remission.

After 37 cycles (duration of azacitidine treatment 3 years and 9 months), bone marrow histology was typical for CMML, without blasts. Cytogenetics revealed two new clones of comparable size (25% of cells) while the initial clone observed at the diagnosis has disappeared. Both clones had a partial trisomy for long arm of chromosome 1. Besides the loss of chromosome Y, there was an additional derivative chromosome 1 lacking the whole short arm. In the second
Discussion

Latest World Health Organization (WHO) classification of myeloid neoplasms defines CMML as a clonal hematopoietic stem cell disorder characterized by the presence of sustained (i.e., >3 months) Peripheral Blood (PB) monocytosis (≥1x10⁹/L and monocytes being ≥10% of total white blood cell count) along with dysplastic features in the bone marrow. CMML is heterogeneous in nature and patients' clinical course and treatment outcomes are variable. Several prognostic models have been developed.3 For risk stratification, we used MD Anderson Prognostic Scoring System (MDAPS). Otherwise different, similar prognostic models offer comparable results.9 Nevertheless, our patient fitted in the intermediate-2-risk group. The latter means a median overall survival of 9 months, although it should be stressed out, that the scoring system was designed before the era of hypomethylating agents (HMA).10

Cytogenetic abnormalities are present in one-third of the patients with CMML. Firstly, we detected loss of the chromosome Y, which is well known recurrent aberration in MDS.9 On the other hand, other aberrations observed during the course of the disease, i.e. partial trisomy of chromosome 1 is less frequently described in MDS. New cytogenetics abnormalities are a sign of a clonal evolution and this has negative prognostic impact on the overall survival.11 Although the later onset of clonal evolution has a better outcome than an early one.11 According to Mayo Clinic -French Consortium Study there are three cytogenetic risk categories, high (complex and monosomal karyotypes), intermediate and low [normal, sole -Y and sole der(3q)], with adjusted median survival of 3, 20 and 41 months.12 Although our patient started HMA treatment in cytogenetic intermediate risk group, latter disappearance of the initial clone with non-recurrent t(11;11) probably points to relatively benign cytogenetic aberration. However, by clonal evolution the patient acquired a complex high risk karyotype and dyed shortly after that.

Several studies and clinical trials tested efficacy of both azacitidine and decitabine and they revealed overall response rates in the range of 30-60% and a median Overall Survival (OS) between 12 to 37 months.3,4,11,12 Based on latest findings, azacitidine was licensed for non-proliferative CMML-2, whereas there are limited data for proliferative CMML subtypes.4 According to one of the recent studies the overall response rate for azacitidine treatment was 20% and the majority of patients achieved response after a median of 3 cycles of treatment. Duration of the response varied between 3 and 24 months.13 In our case, although the patient had proliferative features of CMML, OS was 11 years and 5 months, which is far longer than OS mentioned in previous publications. Patient’s quality of life on azacitidine was seemingly unimpaired, putting aside some minor respiratory infections and low grades azacitidine related adverse reactions.

When trying to understand long period of azacitidine efficacy in our case, other authors stated, that bone marrow blasts higher than 10% and proliferative features of the disease as an increased WBC and splenomegaly were associated with shorter survival and surprisingly, responders were older (71.4y) than non-responders (66.5y).3 At treatment initiation our patient was 66-years old, had low bone marrow blasts (6%), an increased WBC and an enlarged spleen. Low level of bone marrow infiltration with blasts, might be in relation with an initially relatively small malignant CMML clone. Anyway, karyotyping with its low number of metaphases is not an optimal method. Reduction of spleen size during the treatment
could be another cornerstone on path to continuous improvement. Of interest, one report stated, that doubling of platelets after first round of azacitidine was a positive predictor of overall survival. The latter was observed also in our patient.14

In order to be more precise, we should have determined patient’s molecular genetic profile to find any mutations associated with a better outcome. So far only ASXL1 and SETBP1 mutations were found to have negative prognostic impact on overall survival in CMML. The use of Next-Generation Sequencing (NGS) would probably give us in future more definite answers on prognosis for our patients.15,16 Unfortunately, we had limited access to NGS at that time.

Observed serious adverse effects were consistent with already published ones.17 Myelosuppression and infections were mostly observed and are responsible for treatment delay.13 We reduced azacitidine dose due to neutropenia several times. And postponed some of the applications because of minor respiratory infections which normally occurred in 2.5% of azacitidine treated patients.18 Among more serious side effects, our patient experienced progression of ischemic heart disease which may be attributed to azacitidine. That was not possible to rule out and it manifested clinically after 60 cycles of treatment. Although we can not prove a direct relation between azacitidine and cardiac toxicity, it is highly likely as there were case reports describing possible connection between HMA and acute myocarditis, pericarditis and cardiomyopathy.19,21 Although still rare more common cardiovascular side effects mentioned are chest discomfort, peripheral edema and heart murmurs.20

In conclusion, progression of CMML to acute leukemia occurred as a result of disease natural clinical course. CMML patients have 18% to 63% chance of progress into AML during 5 years.22 Acute leukemia free interval was in our patient almost identical with his OS; 11 years and three months. The latter is much longer than a median acute leukemia free survival of 27.4 months reported for CMML patients within intermediate risk group.23 Nevertheless transformation to acute leukemia has a dismal prognosis with poor outcomes.24

Conclusions

We may conclude, that this was the case report of a CMML patient, with extreme good clinical course. And that is why it should be analyzed in detail in order to understand the malignant disease better.

References