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Managing acute promyelocytic leukemia in patients belonging to the Jehovah’s Witness congregation

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Abstract

Acute promyelocytic leukemia (APL) is a hyper-acute leukemia and presents with cytopenias and disseminated intravascular coagulation. Jehovah’s Witnesses with APL offer a unique challenge during induction by refusing transfusion and pose a difficult challenge in this curable disease. Our focus over the last 8 years has been decreasing early deaths in APL in both academic and community centers. As a result we have extensive experience in APL induction with a proven improvement in early deaths. Three patients with APL belonging to the Jehovah’s Witness congregation were treated in our practice and published literature in treating Witnesses with APL was reviewed. It is highly imperative to prevent induction mortality in this patient population. The goal of treatment among the Witnesses is to prevent death during induction and subsequently cure them. We discuss the management and proactive measures to prevent induction mortality in this most curable blood cancer.

Introduction

Patients belonging to the Jehovah’s Witness congregation refuse blood transfusions resulting in specific challenges to the health care team especially in diseases with good outcomes. The belief is based on their interpretation of the scripture that forbids eating of blood; these passages have been interpreted as precluding transfusion of blood.1 Acute promyelocytic leukemia (APL) commonly presents with profound cytopenias and an aggressive strategy of blood product transfusion is the mainstay of supportive care during induction. This results in both practical as well as ethical problems during leukemic induction. Witnesses have been treated without blood product support during induction period of acute leukemias including APL induction and have also successfully received autologous and allogeneic transplants.2-4 Excellent drugs are available to treat APL; specifically those that are mildly suppressive thereby limiting the need for transfusion support. This is an ideal leukemia where these agents along with growth factors developed for different blood lines; and supportive care could be judiciously used. This will eliminate the need for transfusions during treatment. However the consistent problem in APL across the general population is a high early death rate of approximately 30% during induction.5,6 The main causes of death during induction are bleeding, differentiation syndrome, infection and multi-organ failure. We treated three Witnesses with APL and herein report our experience. We also reviewed the literature on APL induction and identified a scant amount of data. It is likely that patients with negative outcomes were not reported resulting in limited available data. From reviewing the APL literature in Witnesses (similar to APL in general), patients who survived induction achieved hematologic and molecular remission and in all likelihood cured. As such preventing induction mortality in this patient population is the most critical aspect of management.

Materials and Methods

We performed a Pubmed search to include acute promyelocytic leukemia and Jehovah’s Witness. Published literature on leukemias, cancers, supportive care in Jehovah’s Witness patients was reviewed.5,7-11 Our own experience in treating three APL Jehovah Witness patients is also documented.

Case Report #1

A 71 year old female with hypertension and diabetes presented in 2013 with a white blood cell (WBC) count of 10×10⁹/L, Hemoglobin (Hb) of 6.6 g/dL, platelets 7×10⁹/L, fibrinogen 107 mg/dL and markedly elevated D-dimers. After confirmation of the diagnosis of APL, Al-trans retinoic acid (ATRA) at 45 mg/m² and arsenic trioxide (ATO) 0.15 mg/kg were started. Patient accepted cryoprecipitate but refused packed red cells and platelets. Erythropoietin at 40,000 units once a day was given as well as dexamethasone 10 mg IV twice a day. On day 4 of admission, the patients Hb decreased to 3.6 g/dL, became progressively weaker, unresponsive and died on day 7.

Case Report #2

A 56 year old Hispanic female presented with flu-like symptoms and was treated with oseltamivir in late 2013. Two weeks later patient was found to have a DVT and heparin therapy was started. The patient was also pancytopenic with a WBC of 0.8×10⁹/L, Hb 11.8 g/dL, platelets 47×10⁹/L, INR 1.7, Fibrinogen 132 mg/dL, D-dimers 5463 (normal <295); a diagnosis of APL was made. ATRA at 45 mg/M² per day was started, an IVC filter placed and heparin was stopped. The patient accepted cryoprecipitate but no other blood products. Arsenic Trioxide was started on day 3 at 0.25 mg/kg twice a week. Blood draws were restricted to once a day initially and subsequently twice a week. On day 10, ATO was held for a rising white count of 10×10⁹/L and darboepoetin started for Hb of 8.8 g/dL. Hydroxyurea was started for a WBC of 39.8×10⁹/L on day 16 and ATO restarted on day 21 with a 30% dose reduction three times a week. Day 24 Hb dropped
to 7.9 g/dL with angina; darbopoeitin was switched to erythropoietin (EPO) with oral iron supplementation. On day 37 WBC increased $2.6 \times 10^9$/L, Hb 10.2 g/dL and platelets $183 \times 10^9$/L and she was discharged. The patient completed 4 cycles of ATO and ATRA based consolidation and is in molecular remission one year post treatment.

**Case Report #3**

A 59 year old white female was admitted in mid-2014 with a 2 week history of aches and pains, shortness of breath, fevers and headache. A CT scan of the head showed multiple acute infarcts. The WBC was $38.6 \times 10^9$/L, Hb 7.4 g/dL and platelets $31 \times 10^9$/L. A diagnosis of APL was made and ATRA and ATO were started. The patient had multiple comorbid conditions to include diabetes, hypertension, CAD, multiple pulmonary edema but subsequently recovered. A 10 year old treated with ATRA and cytarabine had a nadir Hb of 4.4 g/dL but responded to EPO started on day 10. A 62 year old patient treated with ATRA and ATO with EPO support from the start did not develop symptomatic anemia and had an uncomplicated induction and is in remission 10 years post treatment.

### Table 1. Induction of acute promyelocytic leukemia patients belonging to the Jehovah’s witness congregation.

<table>
<thead>
<tr>
<th>Year (ref)</th>
<th>Age/sex</th>
<th>Risk status</th>
<th>Induction</th>
<th>Initial Hb/Nadir Hb</th>
<th>Supportive care</th>
<th>Complications</th>
<th>F/U months</th>
<th>Death</th>
</tr>
</thead>
<tbody>
<tr>
<td>1990 (10)</td>
<td>18/F</td>
<td>Intermediate</td>
<td>Chemotherapy</td>
<td>7.5/4.6</td>
<td>Tranexamic acid, Aprotinin, norethisterone</td>
<td>Anemia</td>
<td>Yes</td>
<td></td>
</tr>
<tr>
<td>1997 (19)</td>
<td>10/M</td>
<td>Intermediate</td>
<td>ATRA higher dose/ Ara-C</td>
<td>8.0/NR</td>
<td>EPO</td>
<td>Anemia</td>
<td>9</td>
<td>No</td>
</tr>
<tr>
<td>1998 (12)</td>
<td>28/F</td>
<td>High</td>
<td>Low dose ATRA/Ara-C</td>
<td>6.9/4.4</td>
<td>EPO</td>
<td>Anemia</td>
<td>Yes</td>
<td></td>
</tr>
<tr>
<td>1999</td>
<td>17/F</td>
<td>High</td>
<td>ATRA/DA</td>
<td>7.0/5.0</td>
<td>EPO</td>
<td>Anemia</td>
<td>120</td>
<td>No</td>
</tr>
<tr>
<td>2000 (14)</td>
<td>62/F</td>
<td>Intermediate</td>
<td>Low dose ATRA</td>
<td>8.6</td>
<td>EPO</td>
<td>Anemia</td>
<td>Yes</td>
<td></td>
</tr>
<tr>
<td>2004 (4)</td>
<td>39/M</td>
<td>Intermediate</td>
<td>ATRA, Dauno, Ara-C</td>
<td>6.6/3.3</td>
<td>EPO</td>
<td>Anemia</td>
<td>32</td>
<td>No</td>
</tr>
<tr>
<td>2007</td>
<td>39/M</td>
<td>Low</td>
<td>Low dose ATRA/ATO</td>
<td>11.8/7.9</td>
<td>Darbopoeitin, EPO</td>
<td>Anemia, Angina</td>
<td>12</td>
<td>No</td>
</tr>
<tr>
<td>2013</td>
<td>71/F</td>
<td>High</td>
<td>ATRA/ATO</td>
<td>7.4/6.7</td>
<td>EPO</td>
<td>Stroke</td>
<td>Yes</td>
<td></td>
</tr>
<tr>
<td>2014</td>
<td>59/F</td>
<td>Low</td>
<td>ATRA</td>
<td>10.5/5.0</td>
<td>EPO</td>
<td>Pregnancy, Anemia</td>
<td>Yes</td>
<td></td>
</tr>
<tr>
<td>2015 (15)</td>
<td>28/F</td>
<td>High</td>
<td>ATRA</td>
<td>8.6</td>
<td>EPO</td>
<td>Anemia</td>
<td>No</td>
<td></td>
</tr>
</tbody>
</table>

### Table 2. Acute promyelocytic leukemia induction suggestions for the Jehovah’s witness patient.

**Work up**

Use pediatric tubes. CBC, CMP, D-dimers, PT, PTT, fibrinogen at admission. Fibrinogen and CBC three times a week. Chest X-ray. Echocardiogram. Bone marrow examination (aspirate, biopsy, flow, cytogenetics, FISH for PML-RAR alpha, PCR for PML-RARalpha). PICC line; NO central lines or invasive procedures (bronchoscopy, spinal tap). Day 14 marrow is NOT necessary. APL is a medical emergency - start ATRA asap.

**Supportive care**

Allopurinol 300 mg daily. Antibiotic prophylaxis - levofloxacin 500 mg daily or similar antibiotic. Antifungal prophylaxis - posaconazole 300 mg daily, voriconazole 200 mg 2 daily or miconafungin 50 mg daily or a similar drug. Acrlovir 400 mg 2 daily or valacyclovir 1000 mg daily.

**Treatment of coagulopathy**

Keep fibrinogen above 150. Use cryoprecipitate if needed and agreeable. Amicar during the first 2 to 3 weeks if there is bleeding or profound thrombocytopenia

**Prevention/treatment of APL differentiation syndrome**

Daily weights - bedside scales only; Keep I/O matched – use diuretics for fluid retention or weight gain. Prednisone 0.5 mg/kg for 14 days and taper if no evidence of DS. If WBC >10,000 at presentation or clinical evidence of DS, dexamethasone 10 mg twice daily.

**Treatment**

ATRA 25 mg/m² for 14 days. Increase to 45 mg/m² after 14 days if stable and no complications. ATO to be incorporated during consolidation.

**Results**

We identified a total of 12 patients (9 published and 3 from our practice) with APL that have been treated (Table 1). Seven were female, 4 male and one not known. Age range was 10 to 62 years; 2 high risk, 4 intermediate risk, 1 low risk and 5 unknown. 5 of the 12 patients died during induction. Three of the 5 deaths were high risk. Two patients were pregnant; one died along with fetal demise. All 5 patients that died had severe anemia reported as complication but without evidence of bleeding. This anemia appears to be secondary to myelosuppression. Four of the 7 surviving patients developed symptomatic anemia (3.7 to 7.9 g/dL); EPO was started in 3 of these patients after developing symptoms with a good response. A pregnant woman treated with ATRA alone developed fetal distress with a nadir Hb of 3.7 g/dL and had EPO started on day 10; the patient responded achieved remission and had an uneventful delivery. One patient treated with ATRA and ATO without aggressive EPO support from initiation dropped his Hb to 5.2 g/dL, had a myocardial infarction and pulmonary edema but subsequently recovered.
Discussion

A thorough discussion should be had with the patient regarding the nature of the disease, the high rate of curability, acceptable and unacceptable blood fractions. While the family members are generally a part of the discussion, the treating Oncologist/Hematologist should privately discuss with the patient his/her wishes in a life threatening circumstance to avoid peer pressure.5 Patients in this reported series have been treated from 1990 to 2014. Over that time, therapeutic agents available for APL as well as supportive care have changed dramatically. The most effective agents are not markedly myelosuppressive but caution should be used to tide the patient through induction.

Supportive care

In 5 of 12 patients that died, there was symptomatic anemia with hemoglobin values as low as 3.6 g/dL without evidence of bleeding and appears to be myelosuppression. This certainly was the reason for myelosuppression in our patients. Of the remaining seven, 4 patients developed symptomatic anemia that prompted use of EPO later in the course of treatment with response but substantial morbidity. Six of the 12 patients were treated with ATRA only or ATRA/ATO. Five of them developed symptomatic anemia; the one patient who did not develop symptomatic anemia was treated with EPO from day 1. From these observations it is apparent that the major cause of morbidity and mortality in Witnesses is non-hemorrhagic anemia and the goal from the onset should be to improve or prevent development of anemia. General supportive measures such as folic acid and iron therapy can be started at diagnosis. In one study intravenous iron was superior to oral iron supplementation for pre-operative stimulation of hemoglobin synthesis using EPO.17 Early institution of erythropoietin to increase or prevent a drop in hemoglobin will be beneficial and continued to maintain it above 10. This is a consideration even if ATRA/ATO induction is being used as they are known to cause cytopenias. The dose of EPO used in leukemia or APL induction has been variable in the limited available literature. A reasonable dose of erythropoietin is 200 units/kg and is adjusted based on the severity of the anemia.

Patients with profound thrombocytopenia and clinical evidence of bleeding, epsilon aminocaproic acid (Amicar) has been used with success. Since APL is also a pro-coagulant condition, thrombosis is quite frequent. If amicar is used, careful monitoring for resolution of DIC should be undertaken after starting differentiating agents as it resolves quickly; at which point amicar can be discontinued.18

Induction

Using single agent ATRA might be the safest and most logical approach. Using both ATRA and ATO in combination causes leukocytosis in >50% of patients. This may require use of hydroxyurea or anthraclycines to decrease the white count thereby causing unwanted myelosuppression. Also a full dose of ATRA can cause leukocytosis and predispose to DS. Single agent ATRA at a lower dose (25/M2-pediatric dose) followed by increasing to full dose after 14 days if there are no complications may be better tolerated. ATRA can be continued until hematologic remission is achieved. Prophylactic steroids to prevent DS at least during the first 2 weeks would be helpful to avoid unnecessary complications. It may not be necessary to use ATO during induction but can be effectively incorporated during consolidation. The goal during induction in these patients is to reverse the DIC, prevent early deaths and achieve hematologic remission without blood product support.

Conclusions

Jehovah’s Witnesses with APL are a unique population due to their refusal to accept blood product support. The biggest hurdle for cure is early death during induction. Patients who survive induction achieve hematologic and molecular remission and in all likelihood are cured. Anemia is the most frequent cause of morbidity and mortality. We suggest aggressive use of erythropoietic growth factors and supportive care to treat anemia. Single agent ATRA during induction followed by incorporation of ATO during consolidation may be the most logical approach to minimize complications during induction. Based on our own experience and published literature, we outline suggestions regarding strategies that may be practical especially during induction to prevent early deaths (Table 2).

Suggested outline: recommendations

The following recommendations are proposed: i) discuss with the patient regarding APL and the excellent outcome; ii) acceptable/unacceptable blood products; iii) blood draws in pediatric tubes; iv) folic acid and intravenous iron; v) erythropoietin 200 u/kg three times a week to start at the beginning of induction; vi) epsilon aminocaproic acid (Amicar) in patients with platelets <10,000 or clinical evidence of bleeding; can be discontinued after 14 to 21 days; vii) prednisone 0.5 mg/kg for 14 days and taper if no evidence of differentiation syndrome; viii) ATRA 25 mg/M2 for 14 days; increase to 45 mg/M2 if no complications (in patients >65 and without comorbid conditions) till hematologic remission; ix) avoid ATO during induction; can be incorporated during consolidation.

References