CLINICAL OBSERVATIONS, INTERVENTIONS, AND THERAPEUTIC TRIALS

A Randomized Comparison of All Transretinoic Acid (ATRA) Followed by Chemotherapy and ATRA Plus Chemotherapy and the Role of Maintenance Therapy in Newly Diagnosed Acute Promyelocytic Leukemia

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All transretinoic acid (ATRA) followed by daunorubicin (DNR)-AraC chemotherapy (CT) has improved the outcome of acute promyelocytic leukemia (APL) by comparison to CT alone. In a randomized trial, (1) we compared 2 induction schedules (ATRA followed by CT [ATRA−CT] and ATRA plus CT [ATRA+CT, with CT added on day 3 of ATRA treatment]) and (2) we assessed the role of maintenance treatment. Four hundred thirteen patients ≤75 years of age and with newly diagnosed APL were included. Induction treatment was stratified on white blood cell (WBC) count and age: patients ≤65 years of age and with an initial WBC count of ≤5,000/µL (n = 208) were randomized between ATRA−CT and ATRA+CT (initially randomized patients); patients with a WBC count greater than (high WBC count group, n = 163) and patients 66 to 75 years of age with a WBC count greater than 5,000/µL (elderly group, n = 42) were not initially randomized and received ATRA+CT from day 1 and ATRA−CT, respectively. All patients achieving CR received 2 additional DNR-AraC courses (only in patients 66 to 75 years of age) and were then randomized for maintenance between no treatment, intermittent ATRA (15 days every 3 months) for 2 years, or both, using a 2-by-2 factorial design. Overall, 16% in the ATRA=CT group (P = .0002) and 13% in patients randomized to intermittent ATRA and 25% in patients randomized to no ATRA (P = .02). An additive effect of continuous maintenance CT and intermittent ATRA was seen, and only 6 of the 74 patients who received both maintenance treatments had relapsed. Overall survival was improved in patients who received maintenance CT (P = .01), and there was a trend for better survival in patients who received maintenance ATRA (P = .22). Our findings strongly suggest that early addition of chemotherapy to ATRA and maintenance therapy combining continuous CT and intermittent ATRA can reduce the incidence of relapse in APL. This effect already translates into significantly better survival for maintenance treatment with continuous CT.

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ACUTE PROMYELOCYTIC leukemia (APL) is a specific type of acute myeloid leukemia (AML) characterized by the morphology of blast cells1,2, by t(15;17) translocation,3 which fuses the PML gene on chromosome 15 to the retinoic acid receptor (RAR) α gene on chromosome 17;4-5 and by specific coagulopathy.6,7 Until recently, combination chemotherapy (CT) with anthracycline-cytosine arabinoside (AraC) was the only treatment of APL, with complete remission (CR) in 65% to 80% of newly diagnosed cases.8-12 The remaining patients suffered early death, mainly from bleeding due to worsening of coagulopathy or resistance to CT. Fifty percent to 65% of the patients who achieved CR subsequently relapsed, and 30% to 40% survived at 2 years.8-12

All transretinoic acid (ATRA) can differentiate abnormal promyelocytes into mature granulocytes in APL in vitro and in vivo13-16 and induced CR in 80% to 90% of newly diagnosed13,15-17 and first-relapsing APL.18 ATRA rapidly improved coagulopathy without inducing aplasia. However, in 15% to 25% of the patients, it was associated with an ATRA syndrome that generally occurred with a rapid increase in white blood cells (WBC) and often had a fatal outcome.19-22 Furthermore, patients who achieved CR with ATRA and were maintained on ATRA alone or mild CT generally rapidly relapsed.16,18 Thus, we designed a strategy combining ATRA and intensive CT in newly diagnosed APL. CT was administered as consolidation or, in patients with rapidly increasing WBC counts, added to ATRA to prevent the ATRA syndrome.23,24

In a previous randomized study, we found that treatment with ATRA followed by CT gave better survival than CT alone in newly diagnosed APL, mainly by reducing the incidence of
relapse,25,26 and these results were confirmed by a US inter-
group randomized study.27 However, approximately 30% of the
patients treated by ATRA followed by CT still relapsed, and new
strategies were required to further reduce that number.

Results of the combination of ATRA and CT suggested that
those 2 treatments had an additive or even synergistic effect in
reducing the incidence of relapse in APL; whether this effect
was optimal by using ATRA first, followed by CT, or by
combining the 2 treatment modalities upfront was the first issue.
Another issue was the possible role of maintenance treatment in
APL. Pharmacological studies suggested that maintenance

treatment with ATRA, using an intermittent rather than continu-
ous administration, could reduce the relapse rate in APL.28 A
similar benefit had also been suggested in nonrandomized studies
for continuous low-dose maintenance CT.29,30

To address those issues, we performed a randomized study
that (1) compared 2 induction schedules (ATRA followed by CT
v ATRA plus CT) and (2) assessed the role of maintenance
treatment with intermittent ATRA and continuous low-dose CT
using a 2-by-2 factorial design.

PATIENTS AND METHODS

Eligibility Criteria

Between April 1, 1993 and December 31, 1996, 439 patients from 97
European centers (listed in the Appendix) with newly diagnosed APL
were included in this trial (APL93 trial), which had been approved by
ethical committees in all participating countries.

Inclusion criteria were as follows: (1) diagnosis of APL, based on
morphology criteria1,2; (2) age 75 years or less; and (3) written informed
consent. Diagnosis had to be subsequently confirmed by presence of
(t(15;17) or PM-L-AR group rearrangement. In the absence of t(15;17) and
if no analysis of the rearrangement could be made, review of initial
marrow slides by an independent morphologist was mandatory.

Design

Objectives of the trial were to assess the optimal timing of ATRA
treatment (before or with CT) and the role of maintenance therapy.
Induction treatment was stratified by age and initial WBC count.

Patients ≤65 years of age with a WBC count less than 5,000/µL were
randomized to receive the reference ATRA treatment of our previous
trial (APL91 trial),25 ie, 45 mg/m²/d ATRA followed by CT (ATRA−CT
group) or ATRA plus CT (ATRA+CT). In the ATRA−CT group,
patients received 45 mg/m²/d ATRA orally until CR, with a maximum
of 90 days. After CR achievement, they received a course of 60 mg/m²/d
donormycin (DNR) for 3 days and 200 mg/m²/d AraC for 7 days
(course I). However, course I was added to ATRA if the WBC count
was increased to greater than 6,000/µL, 10,000/µL, or 15,000/µL by day 5,
10, and 15 of ATRA treatment, respectively, because, from our
experience, patients were at risk of ATRA syndrome above those
thresholds.20 Patients randomized to the ATRA+CT group received
the same combination of ATRA and CT, with course I of CT starting on day
3 of ATRA treatment. This 48-hour interval before onset of CT was
based on our previous report, because it allowed correction of coagulop-
thathy.20

Patients with a WBC count greater than 5,000/µL at presentation
(irrespective of their age) and patients 66 to 75 years of age with a WBC
count ≤5,000/µL were not randomized but received ATRA plus CT
course I from day 1 (high WBC group) and the same schedule as in the
ATRA−CT group (elderly group), respectively.

Treatment of coagulopathy during the induction phase was based on
platelet support to maintain the platelet count at a level greater than
50 × 10⁹/L until the disappearance of coagulopathy. The use of heparin,
patients with confirmed diagnosis of APL. Censored criteria were analysed with the Kaplan-Meier estimate, the log-rank test, and Cox’s model. For each endpoint, treatment comparison was adjusted on a predetermined subset of prognostic parameters using Cox’s model: age, sex, WBC count, platelet count, absolute number of circulating blasts, M3 subtype (classical M3 or microgranular variant M3), and fibrinogen level. Continuous covariates were entered into native form after checking for log linear relationships. Cox’s model was used for testing interaction in the maintenance factorial design as well as to adjust for possible baseline confounders, including result of induction assignment. In the absence of interaction, it was decided to analyze separately each maintenance treatment, with the main analysis (adjusted or unadjusted for baseline covariates) being stratified on the other treatment using a Cox’s model. Relative risks were estimated with 5% confidence intervals (CI). All tests were 2-sided. The SAS software (SAS Institute, Cary, NC) was used.

RESULTS

Initial Characteristics of the Patient Population

In 413 of the 439 patients included, diagnosis of APL was confirmed by the presence of t(15;17) (n = 352), by the presence of PML-RARα transcript by reverse transcriptase-polymerase chain reaction (RT-PCR; n = 31), or by review of marrow slides (n = 30). The remaining 26 patients, in whom diagnosis of APL was not confirmed by cyto genetics, RT-PCR, or review of marrow, slides were excluded, according to the protocol. The median age of the 413 eligible patients was 46 years; 16 patients were children (≤15 years of age), 42 patients more than 65 years of age were included in the elderly group, 163 patients presenting with a WBC count greater than 5,000/µL were included in the high WBC group, and 208 patients were randomized between the ATRA—CT (109 patients) and the ATRA+CT (99 patients) groups. The ATRA—CT and ATRA+CT groups were well balanced for all pretreatment characteristics (Table 1).

Results of Induction Treatment

CR induction and early deaths. CR was obtained in 381 patients (92%; 95% exact CI, 90% to 95%), 31 (7%) patients were older age (P = .02) and in the high WBC group (P = .03). In the remaining 4 patients with moderate ATRA syndrome died from CNS bleeding (1 case), subsequent occurrence of sepsis (2 cases), and leukemic resistance (1 case), respectively. No pretreatment factor, including the WBC count, was significantly associated with the occurrence of the ATRA syndrome.

Results according to induction stratification. Seventy-five of the 381 patients who achieved CR had relapsed, 29 had died in first CR, 18 of them from sepsis after a consolidation course, and 277 remained in first CR. The CR rate and incidence of ATRA syndrome did not significantly differ between randomized patients (ATRA—CT and ATRA+CT groups) and patients included in the high WBC count and elderly groups (Table 2).

Kaplan-Meier estimates of EFS were significantly lower in the high WBC group than in randomized patients (P = .0002) and close to significance in the elderly group (P = .086; Table 2 and Fig 1). Estimates of survival were significantly lower in the elderly group (P = .02) and in the high WBC group (P = .03). The high WBC group had significantly more and earlier relapses (P = .0001) than patients with WBC counts less than 5,000/µL.

![Table 1. Initial Characteristics of the Patients Included in APL 93 Trial According to Induction Treatment](https://www.bloodjournal.org/content/early/2017/08/21/blood.2017-042745.DataSupplement)

In the ATRA—CT group, CT was added before CR achievement due to increasing WBC counts (>6,000/µL by day 5, >10,000/µL by day 10, or >15,000/µL by day 15 of ATRA treatment) in 60 of the 109 patients (55%).

**ATRA syndrome.** In the absence of biological diagnostic criteria of ATRA syndrome, diagnosis of ATRA syndrome was made on clinical grounds by the association of at least 3 of the following signs, in the absence of other causes: fever, weight gain, respiratory distress, lung infiltrates, pleural or pericardial effusion, hypotension, and renal failure. Clinical signs of ATRA syndrome occurred in 64 (15%; 95% exact CI, 12% to 19%) patients. Sixty patients received CT and 58 also received intravenous dexamethasone. In 2 of them, with WBC counts of 186,000/µL and 38,000/µL upon admission, respectively, dyspnea, hypoxia, and pulmonary infiltrates were already present, before the onset of ATRA. Worsening of those signs occurred within hours on ATRA onset. In the other patients, signs developed after 1 to 35 days (median, 7 days) of ATRA treatment. They occurred later than day 14 of ATRA treatment in 21 (33%) patients. In 11 of those 21 patients, ATRA syndrome was observed upon recovery from the phase of aplasia that followed the addition of CT (indicated because of high WBC counts at diagnosis or during treatment). Fifty-five (86%) of the 64 patients achieved CR. Five died from ATRA syndrome, including 1 of the 5 patients in whom the addition of CT was not made and 1 of the 6 patients in whom the addition of dexamethasone was not made. The remaining 4 patients with moderate ATRA syndrome died from CNS bleeding (1 case), subsequent occurrence of sepsis (2 cases), and leukemic resistance (1 case), respectively. No pretreatment factor, including the WBC count, was significantly associated with the occurrence of the ATRA syndrome.
Results of induction randomization in patients less than 65 years of age with a WBC count less than 5,000/µL. The CR rate was 95% (95% CI, 86% to 97%) in the ATRA—CT group and 94% (95% CI, 87% to 98%) in the ATRA+CT group (P = .79 by the Fisher’s test; Table 2). Relapse at 2 years was estimated at 6% ± 3% in the ATRA+CT group as compared with 16% ± 4% in the ATRA—CT group (P = .04 by the log-rank test; Fig 2A). Relative risk (RR) was estimated at .41, with a 95% CI of 0.17 to 0.99. Adjustment for age, sex, WBC count, platelet count, circulating blast count, microgranular variant M3, and fibrinogen slightly modify these estimates (RR = .38, 95% CI, 0.16 to 0.94; P = .04 by the Wald test). Kaplan-Meier estimate of EFS at 2 years was 84% ± 4% in the ATRA+CT group as compared with 77% ± 4% in the ATRA—CT group (P = .10 by the log-rank test; crude RR = .62 [95% CI, 0.35 to 1.10]; adjusted RR for baseline covariates = .62 [95% CI, 0.10 to 0.94]; P = .1 by the Wald test; Fig 2B). Two-year survival was estimated at 84% ± 4% in the ATRA+CT group and 81% ± 4% in the ATRA—CT group (P = .40 by the log-rank test, RR = .76 [95% CI, 0.4 to 1.44]; adjusted RR for baseline covariates = .67 [95% CI, 0.35 to 1.29]; P = .23 by the Wald test; Fig 2C).

Maintenance

Of the 381 patients who had achieved CR, 348 had received full consolidation CT, were alive in CR, and could therefore be randomized for maintenance. Two hundred eighty-nine were randomized for maintenance. The remaining patients were not randomized because they had received an allograft (7 cases), had an unrelated disorder or were in poor medical condition (11 patients), had experienced side effects of ATRA during induction treatment (3 cases), or because of patient refusal (20 cases) or physician omission (18 cases).

Of the 289 patients randomized for maintenance, 73 were allocated to the absence of maintenance, 72 to intermittent ATRA, 70 to continuous CT, and 74 to the association of both treatments. One patient, who was included in the last group, refused to start treatment. No imbalances in pretreatment characteristics were seen between the treatment groups (Table 3). Overall, 56 relapses occurred before the reference date of January 1, 1998. Thirty-three patients had died, including 27 after relapse and 6 in first CR. Because no interaction between continuous CT and intermittent ATRA was found by the Cox’s model whatever the endpoint, we used the statistical strategy defined in the protocol, ie, separate analysis of each treatment, stratifying on the other maintenance treatment. Finally, to assess the influence of possible confounders measured at baseline, we adjusted analyses for baseline prognostic covariates as well as for the result of the induction assignment. Results are summarized in Table 4.

Forty-one and 15 relapses were recorded in patients without and with continuous CT (P = .0003, stratified Cox model). Thirty-five and 21 relapses were recorded in patients without and with intermittent ATRA (P = .024, stratified Cox model; Table 4 and Fig 3A and B). In patients randomized to receive both maintenance treatments, only 6 relapses occurred (Fig 3C), confirming the additive effect of these 2 maintenance treatments (2-year relapse rate estimated at 7.4%). These results were slightly modified when the comparison was adjusted on the predefined set of baseline characteristics (Table 4). Similar results were obtained with EFS (62 events; 56 relapses and 6 deaths in CR; Table 4). In patients randomized to receive both maintenance treatments, only 6 events occurred (2-year EFS estimated at 93% ± 3%). Overall survival was improved in patients who received maintenance CT (P = .01, RR = .36 [95% CI, 0.17 to 0.78]; stratified Cox model). A trend towards better survival was found in patients who received maintenance ATRA (P = .22, RR = .65; stratified Cox model). In patients receiving both maintenance treatments, only 4 deaths occurred (2-year survival estimated at 93%).

Patients in the high WBC count group seemed to benefit particularly from maintenance treatment, especially from the combination of ATRA and CT; indeed, 15 of the 28 (54%) patients of that induction group who were randomized to no maintenance relapsed, as compared with 11 of the 30 (37%) patients randomized to intermittent ATRA alone, 7 of the 29 (24%) patients randomized to continuous CT alone, and only 3 of the 29 (10%) randomized to both maintenance treatments.
Differences in the maintenance treatment groups were less important in patients initially included in the ATRA—CT group: 6 of 22 (23%), 2 of 18 (11%), 1 of 21 (5%), and 2 of 19 (10%) relapses after no maintenance, ATRA alone, CT alone, and both, respectively ($P = .06$); numbers of patients were too small in the elderly group for adequate comparisons.

Side effects of ATRA treatment during maintenance included liver function test abnormalities in 9% of the patients maintained with ATRA alone and 32% of the patients maintained with ATRA plus CT. One patient had severe myositis during maintenance, which was attributed to ATRA. No ATRA syndrome was recorded during maintenance. Liver function test abnormalities were found in 36% of the patients who received CT alone for maintenance. Maintenance CT was also associated with lung pneumocystis infection in 2 cases. Cytopenias were usual after CT, often requiring dose reduction or transient discontinuation. Overall, maintenance ATRA and maintenance CT were prematurely stopped in 7 and 16 of the patients, respectively. In the 74 patients allocated to combined maintenance treatment, 3 had early discontinuation of CT and 6 had early discontinuation of both ATRA and CT. None of the 6 deaths in CR that occurred during maintenance treatment could be attributed to the drugs used for maintenance.

**DISCUSSION**

Results of the present trial showed that, even when a large number of centers are involved, CR rates greater than 90% can be achieved in newly diagnosed APL by adding CT to ATRA from the onset of treatment or when WBC counts rapidly increase. This confirmed the 91%, 95%, and 89% CR rates obtained with the combination of ATRA and CT in our previous APL 91 trial, in the Italian experience, and in the Japanese experience.25,34,35 The US intergroup study found a CR rate of only 70% in newly diagnosed APL treated with ATRA followed by CT.27 However, in the subset of patients who fulfilled all diagnostic criteria and whose response was fully evaluated, the CR rate ranged between 80% and 85%. Because the CR rates obtained with CT alone rarely exceeded 75% to 80%, it appears that combining ATRA and CT, in addition to lowering the incidence of relapse, can also somewhat increase the CR rate in APL. Improved CR rates result in part from the almost absence of cases of resistant leukemia with ATRA combined with CT. They also probably result from a low incidence of fatal cases of ATRA syndrome and from reduction of the early death rate in APL presenting with high WBC counts. Indeed, in the present report, the CR rate was 90% in the high WBC group (and 89% in patients with WBC counts >10,000/µL). By comparison, CR rates ranging between 50% and 70% have been reported in patients with WBC counts greater than 10,000/µL treated with CT alone.7-12,25

ATRA syndrome, the major side effect of ATRA treatment, occurred in 15% of our patients. A high peak value of circulating WBC and basal expression of CD13 on blast surface were predictive of the ATRA syndrome in the New York experience,36 but, in our trial, no predictive factors of the ATRA syndrome, including pretreatment WBC count, were found.37 The absence of higher incidence of ATRA syndrome that we observed in patients with high WBC counts could have resulted from the systematic addition of CT to ATRA in those patients. Addition of CT to ATRA in patients with high WBC counts at diagnosis or during ATRA treatment in APL and administration
of high dose dexamethasone at the earliest symptom also appear to have reduced the incidence of fatal ATRA syndrome to 1.2% in the present trial and to 1% to 1.5% in other large published experiences.34,35

Although the combination of ATRA and CT has improved the outcome of APL, some patients still relapse. In the present trial, we tested several approaches to reduce the occurrence of relapse, including the comparison between ATRA—CT and ATRA + CT, and the role of maintenance therapy by intermittent ATRA, continuous low-dose CT, or both.

Reduction of the occurrence of relapses in APL with the combination of ATRA and CT over CT alone, which was observed in our previous APL 91 trial, suggested an additive or even synergistic effect of both treatment modalities. Here, in patients presenting with a WBC count less than 5,000/µL, ATRA + CT gave a significantly lower incidence of relapse and, with borderline significance, better EFS than ATRA—CT. These findings therefore strongly suggest that the additive or synergistic effect between ATRA and CT is optimal when both treatments are administered simultaneously in APL patients presenting with low WBC counts. Simultaneous administration of ATRA and CT is also recommended, from our experience, in patients with high WBC counts to reduce the risk of severe ATRA syndrome. A third possible association between ATRA and CT, ie, CT followed by ATRA, was also tested by the US intergroup study.27 These investigators found a similar incidence of relapse after ATRA followed by CT and CT followed by ATRA. However, administration of ATRA upfront is generally preferred in APL, because it allows for better control of coagulopathy.

Two nonrandomized studies had suggested that maintenance with continuous low-dose CT using 6 mercaptopurine and methotrexate and using a POMP regimen, respectively, could decrease the relapse rate in APL.29,30 In the present trial, we confirmed in a randomized design that maintenance with continuous low-dose 6 mercaptopurine and methotrexate significantly reduced the incidence of relapse and significantly improved EFS and survival. These findings strongly support a role for maintenance CT in APL, even in patients who have previously received intensive consolidation CT.

We also observed that intermittent maintenance with ATRA significantly reduced the incidence of relapse. The US intergroup study had shown a beneficial role for continuous maintenance with ATRA. In patients who had received ATRA followed by CT, 10 of the 46 cases who received continuous maintenance with ATRA relapsed, as compared with 21 of the 54 cases who received no maintenance.27 Furthermore, continuous maintenance with ATRA was significantly associated with improved survival. Prolonged use of ATRA rapidly leads to a sharp decrease of its serum levels, due to hypercatabolism of the drug.38 However, this metabolic pathway is reversible a few weeks after ATRA discontinuation, and high serum levels of ATRA can then be obtained by new drug intake.28 Thus, intermittent maintenance with ATRA, as used in the present study, could prove more efficient than continuous maintenance with ATRA in reducing the incidence of relapse. However, relapses that occur in patients who receive or have recently received ATRA have been found to be highly resistant to this

### Table 3. Initial Characteristics of Patients According to Randomization for Maintenance

<table>
<thead>
<tr>
<th></th>
<th>No ATRA (n = 143)</th>
<th>ATRA (n = 146)</th>
<th>No CT (n = 145)</th>
<th>CT (n = 144)</th>
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<tr>
<td>No. of patients</td>
<td>143</td>
<td>146</td>
<td>145</td>
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<tr>
<td>Male gender</td>
<td>13% (57%)</td>
<td>14% (47%)</td>
<td>14% (51%)</td>
<td>15% (52%)</td>
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<tr>
<td>Age (yrs)</td>
<td>45 (13-74)</td>
<td>44 (1-72)</td>
<td>43 (1-72)</td>
<td>45.5 (13-74)</td>
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<tr>
<td>WBC count (10^9/L)</td>
<td>3 (0-97)</td>
<td>2.9 (0-76.8)</td>
<td>3.3 (0-76.8)</td>
<td>2.7 (0.3-96.9)</td>
</tr>
<tr>
<td>Platelets (10^9/L)</td>
<td>31 (3-249)</td>
<td>31 (4-169)</td>
<td>35 (4-249)</td>
<td>25 (3-215)</td>
</tr>
<tr>
<td>Circulating blasts (10^9/L)</td>
<td>1.3 (0-82.4)</td>
<td>1.1 (0-72.2)</td>
<td>1.5 (0-72.2)</td>
<td>1.1 (0-82.4)</td>
</tr>
<tr>
<td>Microgranular M₃</td>
<td>18 (13%)</td>
<td>20 (14%)</td>
<td>16 (11%)</td>
<td>22 (16%)</td>
</tr>
<tr>
<td>Fibrinogen (g/L)</td>
<td>1.6 (0.3-7)</td>
<td>1.5 (0.2-6.5)</td>
<td>1.5 (0.2-6.5)</td>
<td>1.6 (0.2-7)</td>
</tr>
<tr>
<td>Induction assignment</td>
<td></td>
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</tr>
<tr>
<td>(ATRA — CT)</td>
<td></td>
<td></td>
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<tr>
<td>ATRA + CT</td>
<td></td>
<td></td>
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</tr>
<tr>
<td>group (elderly group)</td>
<td>44/33/58/10</td>
<td>37/49/10</td>
<td>41/37/59/10</td>
<td>40/36/59/10</td>
</tr>
</tbody>
</table>

*Adjusted on induction assignment, age, gender, baseline WBC count, circulating blasts, platelet count, fibrinogen, and M₃ subtype (classical M₁ v microgranular M₂).
Thus, long-term follow-up will be required to determine if the beneficial effect of maintenance with ATRA on the relapse rate we observed in this study clearly translates into survival benefit, although there was already a trend for improved survival in patients who received this maintenance. Finally, continuous low-dose CT and intermittent ATRA had an additive effect in this study, and the 2-year relapse rate was estimated at only 7.4% in patients who received both maintenance treatments. Maintenance treatment and especially the combination of intermittent ATRA and continuous CT seemed to benefit particularly patients with high WBC counts.

In conclusion, our results confirm that high CR rates (>90%) can be obtained in a large multicenter study in APL by combining ATRA and CT. They strongly suggest that early addition of CT to ATRA and maintenance combining continuous CT and intermittent ATRA can reduce the incidence of relapse. Whether all of these effects translate into better survival will only be determined by longer follow-up.

**APPENDIX**

Dr P. Fenaux and Dr L. Degos served as cochairmen. Dr C. Chastang and S. Chevret-Chastang (Department of Biostatistics, Hopital St Louis, Paris, France) served as biostatisticians. The following clinical departments participated in APL93 trial.

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**Cooperative AML study group, Germany.** H. Link (Hannover), A. Ganser (Frankfurt), E. Wandt (Nurnberg), A. Breitenbach (Stuttgart), B. Brennsmoiten (Freiburg), D. Hermann (Ulm), H. Soucek (Dresden), H. Strobel (Erlangen).

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**Belgian groups.** J.L. Michaux (Bruxelles), A. Bosly (Yvoir), E. Meuws (Anvers), A. Boulet (Mons).

**Dutch groups.** P. Daenen (Groningen), P. Muus (Nijmegen).
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