In human acute myeloid leukaemia (AML), patients who fail to respond to first induction treatment as well as relapsing patients generally have a short life expectancy. It is generally assumed that this poor prognosis is due in part to chemoresistance. Alternate schedules of chemotherapy could overcome this resistance to conventional therapy, and 50% or more patients treated after failure of induction treatment could reach a subsequent remission (Herzig et al., 1983). On the other hand, a second remission could be obtained in 41% of relapsing patients with a reinduction treatment identical to the one given for the first remission induction (Peterson & Bloomfield, 1981).

The selection or the emergence of a leukaemic clone resistant to chemotherapy is therefore one of the probable explanations of induction treatment failure and relapse in acute myeloid leukaemia. The hypothesis of selection of a resistant subpopulation was commonly observed in experimental systems (Skipper et al., 1978), but was rarely explored in AML (McCulloch et al., 1981).

The drug sensitivity of clonogenic leukaemic cells (CFU-L) can be tested by in vitro assays (Preissler, 1980; Marie et al., 1987), with good in vitro–in vivo correlations, with few exceptions (McCulloch et al., 1982). We have used a leukaemic clonogenic assay giving 87% successful leukaemic growth (Marie et al., 1982; Marie, 1987) to explore modifications of the in vitro sensitivity of bone marrow CFU-L to daunorubicin (DNR) and cytosine arabinoside (ara-C) during evolution of AML in 23 adult patients treated with conventional doses of ara-C and anthracycline.

Patients and methods

Twelve patients (mean age: 55±11 years old, range: 26–77) with AML (7 M2, 5 M4; 72.5±14% bone marrow leukaemic cells before treatment) showed resistance to treatment induction including, in all cases, a combination of anthracycline (adriamycin or DNR) and ara-C at conventional dose (100 to 200 mg m⁻² × 7 days) according to ‘AML5,6,7,8’ protocols of the European Organization for Research on Treatment of Cancer (EORTC). Resistance to chemotherapy was documented 21 days following the end of the induction treatment by bone marrow aspirate showing 58±20% leukaemic cells. The interval between the two in vitro studies (before and after treatment) was 6 weeks±4 (3–18 weeks). Salvage therapy (anthracycline+ara-C, 2 cases; high doses of ara-C (HiDAC)+amsacrine, 4 cases; amsacrine+VP16213, 2 cases) was successful in 3 cases (nos. 6, 9, 10) and failed in the 5 other patients. Two patients were treated with a phase II protocol (low dose ara-C or aclacinomycine) without success, and two patients did not receive subsequent chemotherapy.

Eleven patients (mean age: 55±13 years old, range: 22–69) with AML (2 M1, 3 M2, 3 M3; 74%±18 bone marrow leukaemic cells) were tested at diagnosis and at first relapse. These patients entered into CR after induction treatment including anthracycline (adriamycin: 2 cases; DNR: 9 cases) and ara-C at conventional dose (‘AML5.6’ of ‘EORTC’). A consolidation course with the same drugs was administered in all cases. Maintenance treatment was alternation of amsacrine+HiDAC/amsacrine+5 azacytidine (6 courses) in 4 cases, DNR+ara-C (6 courses) in 3 cases, 6 mercaptopurine+methotrexate orally and DNR+VCR (× 6) in 2 cases, 6 thioguanine+ara-C+immunotherapy in one case, and HiDAC+amsacrine (‘intensive consolidation’ × 2) in one case. Relapse (74±16% bone marrow leukaemic cells) occurred after an average of 13±7 months (range: 5–30 months) following attainment of CR. Six patients achieved as second CR with DNR+ara-C (3 cases) or HiDAC+amsacrine (3 cases), 5 patients failed (2 deaths during treatment and 3 resistant cases).

Methods

Cell preparation

Mononuclear cell suspensions were obtained from marrow aspirate by centrifugation using density 1077 MSL (Eurobio Lab.).

T lymphocytes were depleted by a second centrifugation after sheep erythrocyte rosette formation as described by Minden et al. (1979).

CFU-L assay

Blast colony formation. The technique has been described previously (Marie et al., 1983). Briefly, 2 × 10⁴ T-depleted cells in 0.1 ml alpha medium were plated with methyl cellulose (0.8%), 20% foetal calf serum (Flow Laboratories) and 10% PHA-LCM in 1 ml microwells (Titertek Lab.). Eight to 10 microwells were plated and incubated in a moist atmosphere with 6% carbon dioxide.

Aggregates >20 cells were counted at day 7 (plating efficiency 1, or PEI), and several colonies were pooled for May-Grünwald Giemsa staining and detection of T cells if an unusual aspect of the colonies was noted.

Drug exposure

According to in vivo drug pharmacokinetics during AML.
protocols, the cells were exposed briefly to anthracycline and continuously to ara-C: T-depleted cells (3 × 10⁴) were incubated in alpha medium containing 10% FCS for 30 min with 10⁻⁷, 10⁻⁶ and 10⁻⁵ M daunorubicin (DNR). The cells were pelleted, washed twice in large alpha medium excess and plated as before.

For ara-C, continuous exposure to 10⁻⁷, 10⁻⁶ and 10⁻⁵ M were tested, the drug being added just before plating.

A minimum of 4 microwells was counted at day 7 for each drug. Results were expressed as (i) number of surviving colonies, percentage of inhibition of CFU-L compared to controls without drugs (a minimum of 15 colonies per well in controls was required for results to be recorded); and (ii) dose inhibiting 50% of the CFU-L for ara-C (D₁₅₀) and 70% of the CFU-L for DNR (D₁₇₀), according to our previous data (Marie et al., 1987).

**Statistics**

The Mann-Whitney test was used to compare quantitative data, and linear regression analysis for comparison of results of replicate studies in each patient.

**Results**

**Resisting patients** (Table I)

The in vitro CFU-L drug sensitivity showed few variations when tested before treatment and at time of leukaemic regrowth: the level of CFU-L inhibition to DNR (30 min exposure, Figure 1) to 10⁻⁴ M (55.4 ± 14% and 56.6 ± 36% respectively; r = 0.8, P = 0.006) and to 10⁻³ M (76 ± 31 and 79 ± 39, r = 0.98, P = 0.005) did not change. A significant increase (≥ 1 log) in the D₁₇₀ (DNR) was observed only in 3/12 patients.

The CFU-L inhibition after continuous exposure to 10⁻¹ M ara-C was less stable than for DNR (Figure 2) in each patient (89 ± 14% at induction and 79 ± 21% at leukaemic regrowth; r = 0.57, P = 0.06). An increase of D₁₇₀ (ara-C) was observed in 3/12 patients.

Four patients were treated with a ‘salvage’ protocol including HiDAC and AMSA, and a CR obtained in 3 cases, despite in vitro CFU-L resistance to 10⁻² M ara-C (equivalent to conventional doses) in one case (pl 10).

**Relapsing patients** (Table II)

The PEI remained stable in 5 patients, increased in 5 and decreased in one.

The CFU-L drug sensitivity showed variations when we compared CFU-L inhibition by ara-C and DNR at diagnosis and at relapse. In the same patient, repeat measurements failed to correlate (r = 0.4 for 10⁻⁴ M DNR, r = 0.3 for 10⁻³ M DNR; r = 0.2 for 10⁻⁰ M ara-C, r = 0.4 for 10⁻¹ M ara-C).

CFU-L sensitivity to DNR (Figure 3) remained stable (3 pts) or decreased (6 pts) in the 9 patients who received DNR during maintenance treatment; it remained stable in the 3 other patients who did not. The D₁₇₀ to DNR increased in 4/10 evaluable patients.

The CFU-L inhibition by 10⁻³ M ara-C (Figure 4) decreased dramatically (−45 ± 11%) in 5 patients despite the absence of ara-C in the maintenance regimen in 2 of them; it remained stable in 3 patients and increased in 2 patients, although all these 5 patients received ara-C during their remission. The D₁₇₀ for this drug increased in 5/11 patients.

The treatment of relapse was successful using DNR + ara-C or HiDara-C + AMSA in 6/11 cases, but only one (no. 20) had a longer second remission than the first one. Among patients showing in vitro CFU-L resistance to at least one drug at time of relapse, only 2 out of 6 entered into remission (in one case after HiDAC + AMSA, one after standard doses of DNR-ara-C), whereas all the 4 evaluable patients with CFU-L still sensitive to drug entered into CR.
Table 1 Two determinations of CFU-L first plating efficiency (PEI), in vitro sensitivity to ara-C and DNR, suicide index, treatment received and clinical results of these treatments in patients with primary clinical drug resistance.

| Pts | PEI | Ara-C 10^-7 M | 10^-8 M | 10^-9 M | DNR 10^-7 M | 10^-8 M | 10^-9 M | ^3H thymidine | Treatment received |
|-----|-----|----------------|---------|---------|-------------|---------|---------|--------------|-------------------|
| 1   | 21±3 | nd             | 0       | 0       | 25±7        | 25±6   | 8±2     | 26±5         | TAD = E2          |
| +18 | 15±2 | nd             | 21±4    | nd      | 7±1         | (95)    | nd      | 17±1         | Low dose ara-C = no change |
| 2   | 15±7 | nd             | 0       | 0       | nd          | nd      | nd      | nd           | AML5 = E2         |
| +5  | 21±4 | nd             | 5±4     | 5±4     | nd          | nd      | nd      | AML5 = E3     |
| 3   | 50±3 | nd             | 6±1     | 0       | 41±2        | 25±8   | 19±2    | 32±6         | TAD = E2          |
| +3  | 46±6 | nd             | 31±1    | 21±1    | 28±2        | 26±6   | 10±2    | 42±2         | Acylaminocynine = E2 |
| 4   | 62±5 | 24±2           | 20±2    | 15±5    | 12±2        | 5±2    | 0       | nd           | AML6 = E1         |
| w   | 59±3 | 20±2           | 21±2    | 18±2    | 37±1        | 22±12  | 0       | 52±2         | (14)              |
| 5   | 32±7 | nd             | 0       | 0       | 26±3        | 27±6   | 30±2    | 11±5         | AML5 = E1, TAD = E2 |
| w   | 15±1 | 5±1            | 10 (70) | 100 (100) | 16±1     | 16±8   | 15±2    | 8±6          | HiDAC + AMSA = CR |
| 7   | 62±12| 51±12          | 31±6    | 29      | 14±8        | 3±3    | 0       | nd           | HiDAC + AMSA = CR |
| w   | 70±8 | 49±3           | 32±2    | 20±5    | 11±3        | 21±5   | 9±3     | 37±4         | AML7 = E2         |
| 6   | 47±9 | 18±3           | 28±1    | 2±1     | 49±3        | 35±3   | 2±1     | 20±5         | (14)              |
| w   | 110±13| nd          | 64±12   | nd      | 27±6        | nd     | 94±20   | AML6 = E2     |
| 8   | 73±14| nd             | 25±4    | nd      | 18±2        | nd     | 73±14   | AML6 = E2     |
| 9   | 40±6 | 16±4           | 12±1    | 8±3     | 14±4        | 1±1    | 0       | nd           | HiDAC + AMSA = CR |
| w   | 21±4 | 3±1            | 1±5     | 0       | 10±5        | 0      | 0       | nd           | HiDAC + AMSA = CR |
| 10  | 54±1 | nd             | 56±4    | nd      | 65±7        | 45±12  | 6±3     | 44±4         | AML6 = E2         |
| w   | 55±11| nd             | 3±10    | nd      | 38±10       | nd     | 47±15   | HiDAC + AMSA = CR |
| 11  | 181±6| nd             | 50±39   | nd      | 0           | 0      | nd      | 144±7        | AML7 = E2         |
| 12  | 83±6 | nd             | 0       | nd      | 20±1        | nd     | 49±5    | AML8 = E2     |
| w   | 19±2 | nd             | 0       | 0       | 6±2         | nd     | 10±2    | VP16 + AMSA = E2 |
| +3  | 19±2 | nd             | 0       | 0       | 6±2         | nd     | 10±2    | VP16 + AMSA = E2 |

w: weeks; PEI: number of colonies/1 x 10^6 cells plated. CFU-L growth after continuous exposure to 10^-9, 10^-8 and 10^-9 M of ara-C or to 30 min exposure to 10^-5, 10^-6 and 10^-5 M of DNR. Percentage of CFU-L inhibition is given in brackets.

Treatments: TAD: 6-thioguanin, 200 mg m^-2 x 7, ara-C 200 mg m^-2 x 7, DNR 60 mg m^-2 x 3; AML5 of EORTC: adriamycin 50 mg m^-2 x 1, vincristin 1 mg m^-2, Ara-C 160 mg m^-2 x 7. AML6,7,8 of EORTC: DNR 45 mg m^-2 d1-d3 (30 mg m^-2 in AML7), vincristin 1.4 mg m^-2 d2 (AML6,7), ara-C 200 mg m^-2 d1-d7. HiDAC + AMSA: ara-C 1 or 2 mg m^-2 x 12 + AMSA 120 mg m^-2 x 3. VP16 + AMSA = VP16: 100 mg m^-2 x 5 + AMSA 100 mg m^-2 x 5. E1: Complete resistance with persistence of circulating leukaemic cells without aplasia; E2: failure with leukaemic regrowth; E3: prolonged aplasia with leukaemic regrowth; CR: complete remission.
| Pts | PE1 | Ara-C  | DNR   |
|-----|-----|--------|-------|
|     | 10^{-7} M | 10^{-6} M | 10^{-5} M | 10^{-7} M | 10^{-6} M | 10^{-5} M | Treatment received |
| 13  | 22±7 | 65±5   | 21±2   | 18±7   | 14±1   | 0       | 0       | AML6, AMSA + HiDAC/ |
|     | (0)  | (5)    | (18)   | (36)   | (100)  | (100)   |         | AMSA + 5 AZA        |
| +6  | 29±6 | 56±4   | 28±4   | 19±4   | 32±4   | 18±5    | 5±1     | HiDAC + AMSA = E2   |
| m   | (0)  | (33)   |         | (0)    | (37)   | (82)    |         |                   |
| 14  | 16±2 | 6±3    | 3±1    | 5±2    | 5±3    | 1±1     | 0       | AML5, DNR + VCR,   |
|     | (62) | (81)   | (68)   | (68)   | (94)   | (100)   |         | 6MP + PU           |
| +5  | 82±14| nd     | 60±1   | 63±3   | 60±13  | 59±6    | 26±4    | AMSA + HiDAC = E2  |
| m   | (27) | (22)   |         | (27)   | (28)   | (68)    |         |                   |
| 15  | 40±9 | 42±4   | 27±2   | 35±2   | 0      | 0       | 0       | AML6, AMSA + HiDAC/ |
|     | (0)  | (31)   | (13)   |         |         |         |         | AMSA + 5 AZA        |
| +6  | 53   | nd     | 47     | 56     | 60     | 40      | 24      | AML6 = E2          |
| m   | (11) | (0)    | (24)   | (25)   | (55)   |         |         |                   |
| 16  | 66±15| 42±10  | 16±3   | 1±1    | 65±3   | 30±9    | nd      | AML6, DNR + Ara-C  |
|     | (36) | (76)   | (98)   |         | (54)   |         |         |                   |
| +16 | 73±9 | 27±1   | 26±6   | 27±1   | 33±5   | 25±4    | 6±1     | AML6 = CR         |
| m   | (63) | (64)   | (63)   | (55)   | (66)   | (92)    |         |                   |
| 17  | 158±30| 27±4   | 18±5   | 14±6   | 63±6   | 23±5    | nd      | AML5, 6TG + ara-C+  |
|     | (82) | (89)   | (91)   | (61)   | (85)   |         |         | Immunotherapy      |
| +30 | 72±9 | 31±13  | 38±4   | 34±2   | 21±6   | 21±5    | 0       | AML6 = E4         |
| m   | (57) | (48)   | (53)   | (71)   | (71)   | (100)   |         |                   |
| 18  | 38±5 | 28±4   | 14±2   | 12±3   | 25±1   | 34±1    | 0       | AML6, DNR + 6MP + MTX|
|     | (26) | (61)   | (68)   | (34)   | (100)  | (100)   |         |                   |
| +11 | 46±2 | 39±2   | 26±6   | 26±5   | 18±6   | 31±5    | 0       | HiDAC + AMSA = CR  |
| m   | (15) | (44)   | (43)   | (61)   | (33)   | (100)   |         |                   |
| 19  | 24±4 | nd     | 13     | 6      | nd     | 13      | nd      | AML6, AMSA + HiDAC/ |
|     | (46) | (75)   |         | (46)   | (55)   | (100)   |         | AMSA + 5 AZA        |
| +10 | 114±3| 121±19 | 57±15  | 39±2   | 95±9   | 51±2    | 0       | HiDAC + AMSA = E4  |
| m   | (0)  | (50)   | (66)   | (16)   | (55)   | (100)   |         |                   |
| 20  | 46±5 | 60±6   | 58±8   | 27±1   | 22±1   | 0       | 0       | AML6, DNR + ara-C  |
|     | (0)  | (41)   | (0)    | (52)   | (100)  | (100)   |         |                   |
| +20 | 92±8 | 42±5   | 27±3   | 9±6    | 23±3   | 11±3    | 0       | HiDAC + AMSA = CR  |
| m   | (54) | (71)   | (90)   | (75)   | (88)   | (100)   |         |                   |
| 21  | 40±13| 48±11  | 19±1   | 12±4   | 55±9   | 29±4    | 20±5    | AML6, DNR + ara-C  |
|     | (0)  | (53)   | (70)   | (37)   | (27)   | (50)    |         |                   |
| +14 | 51±2 | nd     | nd     | 52±4   | 16±2   | 1±1     | 0       | AML6 = CR         |
| m   | (0)  | (69)   | (97)   | (0)    | (69)   |         |         |                   |
| 22  | 56±22| 25     | 16     | 0      | nd     | 13±2    | 0       | AML6, HIDAC + AMSA |
|     | (55) | (71)   | (100)  | (22)   | (100)  | (100)   |         |                   |
| +10 | 81±9 | nd     | 48±4   | 82±10  | 43±3   | 0       | 0       | AML6 = CR         |
| m   | (41) | (0)    | (69)   | (0)    | (47)   | (100)   |         |                   |
| 23  | 23±4 | nd     | 42±14  | 7±3    | nd     | nd      | nd      | AML6, AMSA + HiDAC/ |
|     | (0)  | (69)   | (11)   | (34)   | (34)   |         |         | AMSA + 5 AZA        |
| +17 | 54±3 | nd     | 48±3   | 35±4   | nd     |         |         | AML8 = CR         |
| m   | (11) | (34)   |         |         |         |         |         |                   |

m: months; PE1: number of colonies/10^4 cells plated. CFU-L growth after continuous exposure to 10^{-7}, 10^{-6} and 10^{-5} of ara-C or to 30 min exposure to 10^{-7}, 10^{-6} and 10^{-5} of DNR. Percentage of CFU-L inhibition is given in brackets.

_Treatments:_ AML5 of EORTC: adriamycin 50 mg m^{-2} x 1, vincristin 1 mg m^{-2}, ara-C 160 mg m^{-2} x 7; consolidation: _idem_, maintenance: either purinethol + 6-mercaptopurine (+ DNR + vincristin x 9) + androgen or 6-thioguanin + ara-C immunotherapy. AML6,8 of EORTC: DNR 45 mg m^{-2} d1-d3, vincristin 1.4mg m^{-2} d2 (AML6), ara-C 200 mg m^{-2} d1-d7; consolidation in AML6: _idem_ but DNR d1 only, Maintenance for AML6: 6 courses of either ara-C or ara-C + HiDAC/AMS + 5 AZA. AMSA + HiDAC/AMS + HiDAC/AMS + HiDAC/AMS + 5 AZA. AMSA + HiDAC/AMS 120 mg m^{-2} x 5 + ara-C 3 mg m^{-2} x 4. HiDAC + AMSA + ara-C 1 or 2 mg m^{-2} x 12 + AMSA 120 mg m^{-2} x 3; E2: failure with leukemic regrowth; E4: toxic death; CR: complete remission.

In our patients, _in vitro_ CFU-L sensitivity to 10^{-6} M DNR was higher in patients entered into CR (73±22%) than in resistant patients (55±41%), but without statistical significance.

**Discussion**

Repeated study of CFU-L drug sensitivity in 23 patients treated for AML showed different results according to the time of the second study.

In treatment failures, the sensitivity to DNR concentrations used for _in vitro_ _in vivo_ correlations did not change significantly when tested before and after chemotherapy. In these resistant patients, the interval between the _in vitro_ studies was short (6 weeks), a time probably insufficient to develop a leukemic clone with properties different than those observed before therapy. These results confirm the preliminary report of McCulloch (1982), who have repeatedly assessed the self renewal capacity and drug sensitivity of circulating CFU-L. These parameters showed little variation, except evidence of developing drug resistance in a few patients (one relapse and one primary resistance). These data suggested that drug sensitivity and self renewal
capacity are heritable characteristics in leukemic clones, but the small number of patients (7) does not permit definitive conclusions.

Treatment failure was predicted by the first in vitro CFU-L sensitivities to DNR but not to ara-C. This superior predictive value of DNR sensitivity was also found in a larger series (Marie et al., 1987).

In patients achieving a complete remission, the emergence of a leukemic clone with different properties from that observed at diagnosis was noted at relapse in a majority of cases: the sensitivity to \(10^{-5}\text{m} \text{DNR} \text{ or } 10^{-3}\text{m} \text{ ara-C} \text{ decreased in 5 patients, increased in one and remained stable in the five others.} \) This could be related to other modifications observed in the study of leukemic populations in relapse viz.: additional chromosomal abnormalities (Pui et al., 1986), or change in surface phenotype (Borella et al., 1979; Laufer et al., 1982; Pui et al., 1986; Stass et al., 1984), suggestive of clonal evolution. Schwarzmieier et al. (1984), using a short term test with \(^3\text{H} \text{ uridine incorporation into leucocytes,} \) found an in vitro acquired drug resistance in 2 patients tested repeatedly.

The increase of CFU-L drug resistance observed in several relapsing patients could explain the poor prognosis of AML patients in relapse. The reduction of leukemic cells during induction treatment and the long time between the two in vitro studies in these patients are in favour of the expansion of a subclone either existing from the beginning of the disease, or selected by mutations in the residual leukemic clone.

In the great majority of cases, the second CR is shorter than the first one, like in our present study, reflecting a relative chemoresistance, with only moderate cyto reduction during reinduction treatment. Our comparative observations in resistant and relapsing patients are surprising, if one takes into account the good correlation normally observed between in vitro sensitivity of clonogenic cells to DNR + ara-C. Clinical responses (Zittoun et al., 1987): one could except, on the contrary, the emergence of in vitro resistant clones after failure of induction treatment and persistence of sensitive clones in most relapsing patients. This could be explained either by the inability of our clonogenic assay to measure leukemic stem cell properties, or by other resistance mechanisms involving cell kinetic (Raza et al., 1987) or pharmacological factors (Plunkett et al. 1985). The properties of more primitive leukemic stem cells could be studied in liquid cultures (Nara & McCulloch, 1985), and could perhaps better account for the clinical evolution of the disease.

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