Short Communication

FREE-FLOW ELECTROPHORESIS OF AN ASCITES
MAST-CELL TUMOUR

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Detailed biochemical and physiological characterization of malignant cells  
and host cells from the same tumour requires the separation of large quantities of  
each type of cell in a highly purified, viable state. Free-flow electrophoresis  
offers the advantage over many other techniques for the separation of cells from  
tumours (Pretlow & Pretlow, 1980) that 10^8–10^9 cells/h can be separated. This  
technique has been successfully applied to many blood and lymphoid cells in a viable  
state (reviewed by Pretlow & Pretlow, 1979). Many malignant or transformed cells  
have electrophoretic mobilities different from those of normal cells under similar  
conditions (reviewed by Pretlow & Pretlow, 1979). We wanted to test whether the  
electrophoretic mobilities of malignant and host cells were sufficiently different to  
permit large-scale separations by free-flow electrophoresis. The Furth mastocytoma  
(Furth et al., 1957) was chosen for our test system, because the neoplastic cells contain  
metachromatic granules that aid in their identification, and the separated malignant cells can be tested for  
their tumorigenicity in syngeneic hosts.

For each experiment the mastocytoma was grown as an ascitic tumour for 6 days  
in 3- to 4-month-old male C57L × AF1 (hereafter called LAF1) mice (The Jackson  
Laboratory, Bar Harbor, Me., U.S.A.) after the i.p. transplantation of 3·6 × 10^6  
ascites tumour cells into each mouse. The tumours were harvested by washing the  
peritoneal cavities repeatedly with 8 successive 3 ml aliquots of 10% foetal calf serum  
in Joklik’s modification of minimum essential medium (Gibco, Grand Island,  
N.Y., U.S.A.) at 4°C. The cells were centrifuged at 97 g for 8 min at 4°C, washed  
twice in electrophoretic separation buffer, and diluted with buffer after the third  
centrifugation, so that the starting suspension contained 13 × 10^6 cells/ml. Cells  
were filtered through a single layer of Nitex (TETKO, Inc., Elmsford, N.Y.,  
U.S.A.) with a pore diameter of 48 μm just before their introduction into the  
electrophoretic apparatus.

Cells were separated in an FF5 free-flow electrophoretic apparatus (Biomedical  
Instruments, Inc., New York, N.Y., U.S.A.)—a modification of the apparatus  
described by Hannig (1969)—that has a chamber width of 10 cm and a chamber  
gap of 0·7 mm. The buffers for electrophoretic separation and for the electrode  
compartment have been described by Zeiller & Hannig (1971) and used by us  
previously (Kreisberg et al., 1977). The sample was introduced at the rate of  
2 ml/h and the separation buffer at the

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rate of 380 ml/h. Electrophoresis of cells was carried out at 7-6°C in an electric field of 90 V/cm, with a current of 180 mA.

The separated cells were collected in 90 tubes at 4°C and counted with haematocytometer chambers. Slides were prepared with the Cyto centrifuge (Shandon Southern Instruments, Inc., Sewickley, Pa., U.S.A.) and stained with Wright's stain. Differential cell counts were performed on at least 500 cells from each fraction and from the starting suspension.

We obtained an average of 13-9 ± 6-3 × 10^6 cells/mouse (range 5-3–23-8 × 10^6). This starting suspension of ascites cells contained 49-7 ± 1-2% neoplastic mast cells, 28-7 ± 10-3% lymphocytes, 16-7 ± 7-9% macrophages, 2-5 ± 0-1% red blood cells, and 2-6 ± 1-1% other nucleated cells. Two examples of electrophoretic separation of the ascites mast-cell tumour are presented in Fig. 1. Previously, free-flow electrophoresis experiments have been standardized with respect to the modal population of red blood cells (Stein, 1975; Shortman et al., 1975; Pretlow & Pretlow, 1979). Since red blood cells comprised an average of only 2-5% of the cells in our starting suspensions, their modal location could not be determined precisely. These graphs have been standardized by setting the neoplastic mast-cell peak at Fraction 26.

After standardization, the peak modal fractions of lymphocytes and macrophages were within one fraction of each other in the respective graphs. Fraction 26 (Fig. 2) contained an average of 65-2 ± 23-5% neoplastic cells. The purest modal fraction for neoplastic cells (Fraction 24) contained 69-4 ± 25-7% neoplastic mast cells, 0-3 ± 0-4% red blood cells, 10-3 ± 7-3% lymphocytes, 17-3 ± 16-8% macrophages, and 2-8 ± 2-0% other nucleated cells. The populations of lymphocytes and macrophages did not form sharp peaks, but were spread over several fractions (Fig. 1). The purest population of lymphocytes was in Fraction 31 (Fig. 3) and contained an average of 59-0 ± 9-4% lymphocytes, 15-4 ± 18-3% neoplastic mast cells, 24-4 ± 10-0% macrophages, and 1-2 ± 1-1% other nucleated cells. The purest population of macrophages was in Fraction 30, which contained an average of 36-0 ± 16-5% macrophages, 24-8 ± 24-8% neoplastic mast cells, 38-3 ± 7-8% lymphocytes, and 1-1 ± 0-5% other nucleated cells. Most of the recovered neoplastic mast cells were in Fractions 18–26, whereas most of the recovered lymphocytes and macrophages had slower mobilities and were in Fractions 27–33. This is in agreement with several studies carried out with
Fig. 2.—Cells from Fraction 26 after electrophoresis. Neoplastic mast cells represent 65.2% of the cells in this fraction. (Wright's stain, original × 200.)

Fig. 3.—Cells from Fraction 31 after electrophoresis. Lymphocytes represent 59% and macrophages 24.4% of the cells in this fraction. (Wright's stain, original × 200.)
individual cells in cytopherometers (reviewed by Pretlow & Pretlow, 1979).

The various types of cell from the ascitic form of the Furth mast-cell tumour were less highly purified by electrophoresis than by velocity sedimentation (Pretlow et al., 1977; Green et al., 1980). Each type of cell exhibited heterogeneous electrophoretic mobilities, and their modal mobilities were within a few fractions of each other. Lowick et al. (1961) previously found that 2 of the 3 ascites tumours they studied had very heterogeneous electrophoretic mobilities. The electrophoretic mobilities of ascites tumour cells have been observed to vary both with the inoculum size (Hartveit et al., 1968) and with the number of days after transplantation (Hartveit et al., 1968; Mayhew, 1968). The absolute number and the proportion of neoplastic cells and host cells also vary with the inoculum size and the number of days after transplantation (Stewart et al., 1972; Norman & Cornelius, 1978). The inoculum size and number of days of tumour growth were kept constant in our experiments.

Theoretical and practical aspects of cell electrophoresis have been thoroughly discussed (Hannig et al., 1975; Zeiller et al., 1975; Pretlow & Pretlow, 1979). Although cellular aggregation is a possible cause for the overlap of fractions, it does not appear to be implicated here. Less than 10% of the cells in any fraction were aggregated, as observed in haemacytometer chambers after electrophoresis.

The total recovery of cells after electrophoresis was generally >65%; average recoveries of individual types of cells varied between 48 and 68%. These recoveries are in the same range as reported by us previously (Kreisberg et al., 1977) and somewhat less than reported by Stein (1975) for human blood cells.

Viabilities of cells, as assessed by the exclusion of trypan blue, was >95%, both before and after electrophoresis. Serial dilutions of cells purified by free-flow electrophoresis were transplanted into mice. As few as 3 cells from Fraction 26 gave rise to an ascites tumour within a month after transplantation. Thirty-one or fewer cells from Fraction 31 failed to produce tumours, even 4-5 months after transplantation; >60 cells from Fraction 31 did produce tumours within a month. Since this fraction of lymphocytes was adulterated with 15% neoplastic cells, 60 or more cells from this fraction included 9 or more neoplastic cells.

Although this first attempt, to our knowledge, to separate malignant and host cells by free-flow electrophoresis achieved only partial purification of cells from this tumour, the mobilities of malignant and stromal cells from other types of tumour may differ more markedly and warrant further investigation. Free-flow electrophoresis differs from the electrophoretic techniques used more commonly for work with cells, in that a large number of cells can be electrophoresed, and recovery is sufficient for preparative biochemical applications. The malignant cells are viable and can form tumours after electrophoresis.

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