Establishment of a murine leukaemia cell line resistant to the growth-inhibitory effect of bryostatin 1

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Summary Bryostatin 1 is a novel macrocyclic lactone activator of protein kinase C (PKC) which has clinical potential as an anti-cancer agent. The mechanism of action of this agent is unknown, but protein kinase C has been implicated. In order to investigate this possibility, we have developed P388 sublines resistant to bryostatin 1 by continuous challenge of the parent cell line with increasing incremental concentrations of the drug over 4 months. Cell lines were established at monthly intervals yielding four sublines: P388 BR A, which were removed at 1 month; P388 BR B, obtained after 2 months; P388 BR C, obtained after 3 months; and P388 BR D, which were established after 4 months. All four P388 BR sublines show an equal degree of resistance to the growth inhibitory effects of bryostatin 1, with a relative resistance ratio (RR) IC50 of approximately 4.000. The ability of the cytosol of cells to phosphorylate PKC-specific substrate is decreased by 41% for BR A, 57% for BR B 80% for BR C and 94% for BR D compared with the parental cell line, even when grown in the absence of bryostatin 1 for up to 4 weeks. Similar decreases are seen for cytosolic phosphor bryostatin ester binding and whole-cell PKC isoenzyme expression. All four P388 BR sublines show high and equal levels of cross-resistance to the PKC activatory phorbol ester, phorbol 12-myristate 13-acetate (PMA). There is no loss of resistance to either bryostatin 1 or PMA up to 3 months after termination of exposure of the sublines to bryostatin 1. There was no significant degree of cross-resistance to daunorubicin in the bryostatin 1-resistant cell lines. P388 BR A, B, C or D, when compared with the parental cell line, P388.

Bryostatin 1 is the prototype of a family of naturally occurring activators of protein kinase C (PKC) (Berkow & Kraft, 1985; Smith et al., 1985: Kraft et al., 1986; Fields et al., 1988). It is a macrocyclic lactone (see Figure 1) isolated from the marine invertebrate Bugula neritina, a member of the phylum Ectop trocta (Pettit et al., 1982). Bryostatin 1 exerts a wide range of biological effects, including antineoplastic activity, induction of differentiation, haemopoietic stimulation, platelet aggregation, and immunoenhancing activity. Significant antineoplastic activity has been demonstrated against leukemias, lymphomas, ovarian sarcoma, reticulum cell sarcoma and melanoma in a variety of murine and human cell lines and also in vivo against murine tumours (NCI Antitumour Screening Program; Pettit et al., 1982; Dale & Gescher, 1989; Hornung et al., 1992). Bryostatin 1 therefore has potential as an anti-cancer agent and it is currently undergoing phase I and phase II clinical trials in patients with malignant disease at the Christie Hospital, as part of the Cancer Research Campaign clinical trials initiative.

Bryostatin 1 shares many of its biological effects with phorbol esters, which are potent activators of PKC (Castagna et al., 1982). In several biological systems, however, bryostatin 1 behaves differently from phorbol esters. Unlike phorbol esters, bryostatin 1 does not induce differentiation in human bronchial epithelium (Jetten et al., 1989) or primary mouse epidermal cells (Sako et al., 1987). Furthermore, bryostatin 1 and phorbol esters show differential effects on the hydrolysis of phosphatidylylthanolamine (Kiss et al., 1991). Furthermore, bryostatin 1 is inactive as a complete tumour promoter or carcinogen and acts to inhibit the tumour-promoting properties of phorbol esters (Hennings et al., 1987). Some of these differences may be attributed to the diversity of responses to PKC activation mediated by the multiple isoforms that constitute the PKC family (α, β, γ, ε, ζ, η, etc.) and their tissue-selective distribution (Nishizuka, 1988; Parker et al., 1989; Gescher, 1992). Differences in the nature of the binding to the PKC receptor may provide an alternative mechanism for generating heterogeneity (Konig et al., 1985; Nakadate & Blumberg, 1987; Nelsestuen & Bazzz, 1991; Blumberg & Pettit, 1992).

The substantial involvement of protein kinase C in cellular signal transduction gives it a central position in the regulation of growth and differentiation. It is, therefore, of great importance to identify and understand the mechanisms that account for the multiplicity of response within the protein kinase C pathway (Azzo et al., 1992). In order to investigate the role of PKC in the anti-tumour effect of bryostatin 1, we have developed murine leukaemia sublines exhibiting resistance to the growth-inhibitory effects of the drug. These have been compared with the parental line in terms of their PKC activities and isoenzyme expression in order to determine whether alterations in this signalling is associated with decreased response to this agent.

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Figure 1 Bryostatin 1.
Materials and methods

Drugs and chemicals

Phorbo1 12-myristate 13-acetate (PMA) used in the growth inhibition assays and dimethylsulphoxide (DMSO) were obtained from Sigma, UK. ['H]Phorbol dibutyrate ([3H]PDBu) was purchased from New England Nuclear, UK, and Ethanol AR was supplied by BDH. PMA used in the PKC assay, phosphatidylserine, Triton X-100 mixed micelles, PKC substrate [Ac-MBP (4–14)], PKC inhibitor [PKC (19–36)]. calcium chloride, Tris, leupeptin, aprotonin, β-mercaptoethanol and sodium chloride were obtained from Gibco BRL, UK. Specific anti-peptide antisera for PKC isoforms (α, β1, βII, γ, ε) were a kind gift from W.F. Heath, Lilly Research Laboratories. RPMI medium and serum were also obtained from Gibco BRL.

Cell culture

P388 is an immortal murine acute leukaemia cell line. P388 PR8/22 is a P388 cell line with acquired resistance to daunorubicin (DrN), which was previously developed from the parental cell line, P388, by incremental challenge with DrN in vitro (McGown et al., 1983). Both cell lines were grown as a suspension culture in RPMI medium supplemented with 10% horse serum. P388 PR8/22 was routinely maintained in the presence of DrN (0.1 μg ml−1) except during experimental procedures. All cell lines were cryopreserved free and replaced from frozen stock at 3 month intervals.

All cell lines were grown in the absence of bryostatin 1 for 4 weeks before experimentation.

Cell size was measured using a Coulter Channelizer 256.

Establishment P388 sublines resistant to bryostatin 1-induced growth inhibition

We established P388 sublines resistant to bryostatin 1-induced growth inhibition by continuous exposure to increasing concentrations of the drug in vitro for over 4 months. Briefly, exponentially growing P388 cells were collected and adjusted to 10⁶ cells ml⁻¹. Ten millilitres of this cell suspension was plated in 50 cm² flasks and incubated for 7 days at 37°C in a humidified atmosphere containing 5% carbon dioxide. Bryostatin 1 was dissolved in 100% ethanol and added at a final concentration of 0.5% ethanol to the cell suspension each week after the cells had been readjusted to 10⁶ cells ml⁻¹ in fresh medium. The cell suspension was incubated at 37°C. Bryostatin 1 was dissolved in dimethylsulphoxide at a concentration of 0.5 nM and by gradual weekly increments was increased over 4 months to 1,000 nM. Cells were removed from bryostatin 1 challenge at monthly intervals (P388/BR/A, removed at 1 month and up to 50 nM bryostatin 1 challenge; P388/BR/B, 2 months and 100 nM; P388/BR/C, 3 months and 500 nM; and P388/BR/D, 4 months and 1,000 nM), adjusted to 10⁶ cells ml⁻¹ and frozen in triplicate at −27°C. On removal from the freezer, P388/BR/B, C and D were cultured in bryostatin 1-free medium for at least 4 weeks before each experiment.

Growth inhibition studies

Each drug (bryostatin 1, PMA or DrN) was tested for its ability to inhibit the growth of each cell line (P388, P388 PR8/22 or P388/BR/A, B, C or D) when exposed continuously to 1 ml wells for 5 days. Bryostatin 1 was dissolved in ethanol, PMA in DMSO and DrN in water. Exponentially growing cells were adjusted to 10⁶ cells ml⁻¹ and plated in 2 ml wells. Various concentrations of drug were added to appropriate wells with the final concentration of diluent 0.5% (v/v). Cells were incubated for 5 days. Cell number was determined in each well using a Coulter counter and a growth curve was constructed. Each experiment was performed in quadruplicate and carried out three times. The IC₅₀ was defined as the drug concentration necessary to achieve a 50% inhibition of cell growth when compared with the control. Relative resistance was defined as:

Relative resistance = IC₅₀ for resistant subline

IC₅₀ for parental cell line

Whole-cell protein kinase C assay

Assay of PKC was based on measurement of the phosphorylation of an acetylated synthetic peptide based on myelin basic protein, Ac-MBP (4–14; Yasuda et al., 1990). PKC specificity was confirmed by using the PKC pseudosubstrate inhibitor peptide PKC (19–36), which inhibits PKC-catalysed substrate phosphorylation potently (Yasuda et al., 1990).

Cells (approximately 1 x 10⁶ per assay) were rapidly sedimented, washed with ice-cold phosphate-buffered saline and resuspended in ice-cold extraction buffer (20 mM Tris, pH 7.5, 0.5 mM EDTA, 0.5 mM EGTA, 0.5% Triton X-100 and 25 μg ml⁻¹ each aprotonin and leupeptin) and sonicated. PKC was partially purified by binding to DEAE (diethylaminoethyl) cellulose, which was equilibrated with ice-cold wash buffer (20 mM Tris, pH 7.5, 0.5 mM EDTA and 0.5 mM EGTA). PKC was collected as a single fraction with ice-cold elution buffer (20 mM Tris, pH 7.5, 0.5 mM EDTA, 0.5 mM EGTA, 10 mM β-mercaptoethanol and 0.2 M sodium chloride). Enzyme activity was assayed in triplicate after a 5 min incubation at 30°C of a reaction mixture containing 20 mM Tris, pH 7.5, 20 mM magnesium chloride, 1 mM calcium chloride, 20 μM ATP, 0.2 μCi of [γ-³²P]ATP, 50 μM Ac-MBP (4–14), 0.25 mM EDTA, 0.25 mM EGTA, 5 mM β-mercaptoethanol and 0.1 M sodium chloride with either 10 μM PMA, 0.28 mg ml⁻¹ phosphatidylserine and Triton X-100 mixed micelles or 20 μM PKC inhibitor. PKC (19–36). Reactions were terminated by spotting aliquots of the reaction mixture onto individual phosphocellulose discs, which were washed with 1% (v/v) phosphoric acid and water to remove non-incorporated γ-ATP. Each phosphocellulose disc was placed in a scintillation vial with scintillation fluid (10 ml) for measurement of radioactivity in a Beckman (USA) LS 1801 scintillation counter. Conditions were adjusted to ensure that the reaction was linear with respect to time of incubation and concentration of cells.

Phorbo1 ester binding assay

Formation of mixed micelles from a semipurified extract of the cytosolic fraction and measurement of phorbo1 ester receptor binding was performed as described by Hannun and Bell (1987) using [³²P]HPDBu as ligand. Non-specific binding was <10% of total binding.

PKC isoenzyme expression

The expression of PKC isoenzymes α, βI, βII, γ and ε was studied. Cells in exponential growth were pelleted by centrifugation and washed twice with ice-cold PBS. Pellets were lysed in 300 μl of SDS reducing buffer [0.0625 M Tris–HCl, glycerol 10% (v/v), SDS 2% (w/v), 2-β-mercaptoethanol 5% (v/v) 1.25 x 10⁻³ M (w/v) Tris, pH 7.4). The lysate was heated to 95°C for 10 min. An aliquot of lysate equivalent to 35 μg of protein was subjected to 10% SDS–PAGE and electrotransferred to nitrocellulose paper with a current of 60 mA for 1 h. The nitrocellulose membrane was stored overnight at 4°C in blocking buffer (5 g of casein milk powder in 100 ml or TBS/Tween [1:211,1-1 Tris, 8.18 g 1-1 sodium chloride, 0.001% (v/v) Tween 20, pH 7.4]. The nitrocellulose membrane was exposed for 2 h to monoclonal antibodies against PKCα, βI, βII, γ and ε. The nitrocellulose membrane was washed five times with blocking buffer and then stained for 1 h with anti-rabbit Ig peroxidase conjugate diluted 1:1,000 in blocking buffer. The nitrocellulose was then further washed five times with TBS/Tween. The immunoreactive bands were then visualised using the ECL.
Western blotting detection system (Amersham, UK) according to the manufacturer’s instructions.

Results

Cell growth

P388 cells were exposed to bryostatin 1 for up to 4 months. Bryostatin 1 was removed from the medium after 1, 2, 3 or 4 months and the cell lines obtained were designated P388/BR/A, B, C and D respectively. After removal of the drug the cells grew at the same rate as the parental cell line. Significant differences were not observed in either cell size or protein content between the parent cell line and the sublines which had been exposed to bryostatin 1. The general characteristics of the parent line (P388) and cell lines P388/BR/A, P388 BR B, P388 BR C and P388 BR D are summarised in Table 1.

Growth inhibition

Growth-inhibitory effects were assessed using a 5 day continuous exposure assay. The growth of P388/BR/A, B, C and D cells was not inhibited when they were cultured with 100 nM bryostatin 1, however growth inhibition was seen at 500 nM. At 1 μM proliferation of all lines was arrested by 52.6% in the case of P388/BR/A cells, 65.1% for P388/BR B, 59.9% for P388/BR C cells and 48.1% for P388/BR D (see Figure 2). All sublines showed a similar and very high relative resistance (RR) to bryostatin 1 (approximately 4,000-fold). This high level of resistance developed after 1 month of incubation with bryostatin 1 at doses of up to 50 nM. Further incubations for 3 months at doses of up to 1 μM did not increase the degree of resistance. Similarly, all four P388 BR sublines also showed a high and equal level of cross-resistance to phorbol 12-myristate 13-acetate (PMA), with an RR of approximately 4,000 also (Figure 3). Resistance to either bryostatin 1 or PMA was not lost up to 3 months after removal of the cells from the bryostatin 1 challenge. The IC₅₀ of DnR for P388 (6 nM) was very similar to that seen in the bryostatin 1-resistant cell lines P388/BR/A (9.5 nM), P388/BR/B (10.2 nM), P388/BR/C (9.8 nM) and P388/BR/D (10.5 nM) (Figure 4). The IC₅₀ of bryostatin 1 for P388 (0.25 nM) was almost the same as that in the daunorubicin-resistant cell line, P388 PR8 22 (0.30 nM) (Figure 5).

Whole-cell PKC activity

Whole-cell PKC activity was measured in the parent cell line. DnR-resistant subline and bryostatin 1-resistant sublines (Figure 6). Whole-cell PKC activity in response to PMA was reduced to 59 ± 5% for P388 BR A, 43 ± 3% for BR B, 20 ± 6% for BR C and 6 ± 2% for BR D compared with P388 cells (= 100%). Enzyme activity was not significantly altered in the daunorubicin-resistant cell line. P388 PR8 22 (110% ± 10%).

Phorbol receptor assay

Cytosolic phorbol ester receptors were reduced in the bryostatin 1-resistant sublines to similar degrees as whole-cell

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**Table 1** Morphological characteristics of parent cell line (P388) and bryostatin 1-resistant sublines (P388 BR A, B, C and D)

|       | BR A | BR B | BR C | BR D |
|-------|------|------|------|------|
| Doubling time (h) | 14.6 (± 1.2) | 14.8 (± 1.3) | 15.1 (± 1.0) | 14.9 (± 0.8) |
| Cell volume (μl) | 537.7 (± 10.4) | 568.8 (± 17.7) | 576.4 (± 9.4) | 586.5 (± 0.0) |

P388, parent murine leukaemia cell line; BR A, BR B, BR C and BR D, bryostatin 1-resistant P388 sublines. Errors are standard errors of the mean.
PKC activity: P388 BR A (53.9 ± 1.4% of P388 control), P388 BR B (34.5 ± 1.55%), P388 BR C (15.2 ± 1.6% of P388 controls) and P388 BR D (10.5 ± 2.8%) compared to P388 controls (100%o) (Figure 7). Cytosolic phorbol ester receptor levels are essentially unchanged in the daunorubicin-resistant subline. P388 PPR 22 (97.8 ± 5.3%) compared to P388 controls (P388 PR 22 > BR A > BR B > BR C > BR D). Indeed, no PKC-x was detected in the BR D line by this method. These results mirror closely the changes seen in whole-cell PKC phosphorylating activity and cytosolic PKC receptors. PKC-x levels in P388 PR 22 were similar to those found in P388 wild-type cells.

Discussion

In common with phorbol esters, brystatin 1 is a very potent activator of PKC. Bryostatin 1 induces only a subset of the characteristic biological responses seen for phorbol esters.
For those responses which bryostatin 1 fails to induce it often blocks the same responses to the phorbol ester if the two compounds are present together (Sako et al., 1987; Jetten et al., 1989). The mechanisms which give rise to this diversity of response following PKC activation are currently unclear, but two possibilities which may be involved are (1) differential stimulation of PKC isoenzymes by bryostatin 1 and PMA and (2) differential binding of the two agents to PKC.

The murine leukaemia cell line, P388, is very sensitive to the growth-inhibitory effects of bryostatin 1 (IC₅₀ = 0.25 nM) if exposed for 5 days to the drug. We have established P388 sublines with high levels of resistance to bryostatin 1 by continuous and increasing incremental challenge with this drug over 4 months. Each subline was found to possess relative resistance of approximately 4,000 against bryostatin 1. A similar high level of cross-resistance has also been found to the phorbol ester, PMA. The reduced activation of PKC by PMA observed in these sublines – 41% and 94% reduction for P388 BR A and P388 BR D respectively compared with parental cells – shows that the continuation of drug exposure continued to cause an alteration in PKC content of the sublines. Similar changes were observed for cytoplasmic phorbol ester receptor levels. It is not clear whether it is time of exposure or increase in drug concentration which caused decreased PKC activity. What is clearly evident is that down-regulation of PKC as assayed by the methods described is not linearly related to the decrease in sensitivity to bryostatin 1. However, it is possible that loss of a specific PKC isoenzyme was responsible for conferring bryostatin 1 resistance. It must be noted that the PKC levels in this study are not necessarily indicative of the activity of each PKC isoform.

Activity was measured using a specific peptide inhibitor based on the pseudosubstrate region common to the α, β and γ isoenzymes of PKC. The pseudosubstrate regions for the δ, ε, ζ isoenzymes diverge (Parker et al., 1989) and therefore the inhibitor is probably not recognised by these enzymes. Similarly, the affinity of the substrate is isoform dependent (Yasuda et al., 1990). Therefore, although PKC activity was reduced in these cells, it is not possible from the data to draw conclusions as to specific PKC isoforms. An analysis of these isoenzymes was necessary to address this issue. The expression of five individual PKC isoenzymes (α, β1, β2, γ and ε) was examined. Only α was detectable in the parental cell line, P388 (Figure 8). The enzyme showed a gradual decrease in expression with increasing duration and concentration of bryostatin 1 exposure reflecting the reductions observed in whole-cell PKC activity and cytoplasmic phorbol ester receptor levels. No increased expression in β1, β2, γ or ε was seen in the resistant cell lines. This result suggests that bryostatin 1 resistance is not associated with selective PKC down-regulation, although we cannot exclude the possibility that other PKC isoenzymes are involved. Nevertheless, the result is strongly suggestive that down-regulation of a cellular signal transduction pathway other than those involving PKC-α, -β, -γ or -ε is important in mediating bryostatin 1 resistance and that down-regulation of PKC is either just part of the resistance mechanism or perhaps only an associated finding.

It is of interest that the P388 BR D subline, with 94% loss of whole-cell PKC activity, continued to grow at the same rate in serum supplemented with medium as the parent cell line. It seems, therefore, that only 6% of the PKC activity as measured by these techniques need be associated with normal rates of cell proliferation.

An increased activity of PKC has been reported in certain multidrug-resistant (MDR) cell lines (Palayoor et al., 1987). We observed a small increase in whole-cell PKC activity when the daunorubicin-resistant cell line, P388 PR8 22, was compared with the parent cell line, P388. However this increase was not significant.

We could not demonstrate any significant degree of cross-resistance towards daunorubicin in the bryostatin 1-resistant cell lines. Equally, bryostatin 1 demonstrates similar growth-inhibitory effects on both P388 and the daunorubicin-resistant subline. We conclude that bryostatin 1 is not one of the 'mdr' class of agents. This is in agreement with Haiit and De Rosa (1991), who have shown that a subclone of HL-60 human promyelocytic leukaemia with acquired resistance to phorbol esters showed no change in sensitivity to doxorubicin.

In conclusion, this work has shown that the relationship between levels of PKC activity and sensitivity towards bryostatin 1 or phorbol esters is a complex one. However, continuous exposure of cells to bryostatin 1 results in the rapid development of a stable cell line with decreased total PKC activity. The importance of this residual PKC activity to cell proliferation has yet to be determined.

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Figure 8 PKC-α isoenzyme expression in parent cell line P388, daunorubicin-resistant subline (P388 PR8/22) and bryostatin 1-resistant sublines (P388 BR A, B, C and D).
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