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Susceptibility to apoptosis is differentially regulated by c-myc and mutated Ha-ras oncoproteins and is associated with endonuclease availability

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Summary Oncogenes and oncosuppressors can deregulate cell replication in tumours, and recently have been shown to influence the probability of apoptosis. The effects of human c-myc and mutated (T24) Ha-ras oncoproteins on susceptibility to apoptosis were investigated by introducing them into immortalised rat fibroblasts. The resulting family of transfectants showed closely similar measures of proliferation, but widely divergent rates of apoptosis, differing by up to fifteen-fold, that correlated inversely with population expansion rates in vitro. T24-ras transfectants with moderate or high p21 expression showed reduced apoptosis, and this was reversed by pharmacological inhibition of membrane localisation of p21 expression by mervinolin. In contrast, c-myc stimulated apoptosis, and this was further enhanced by serum deprivation. Inducibility of effector proteins represents one possible mechanism of genetic control of the susceptibility to apoptosis, and its investigation showed that c-myc was associated with expression by viable cells of latent calcium/magnesium sensitive endonuclease activity characteristic of apoptosis. In contrast, T24-oncogene activity was not detected in viable cells of a T24-ras transfectant expressing high levels of p21. Thus, there appeared to be differential regulation of susceptibility to apoptosis, positively by c-myc and negatively by activated ras, and this was associated with availability of endonuclease activity. Genetic modulation of apoptosis in human neoplasms is likely to influence net growth rate, retention of cells acquiring new mutations and response to certain chemotherapeutic agents.

Recently it has become clear that oncogene and oncosuppressor gene activity can influence the probability of cell death as well as that of cell replication. Expression of c-myc appears to be associated with a cellular state in which DNA replication occurs provided suitable growth factors are present, but from which cells die by the process of apoptosis should these growth factors be withdrawn (Evans et al., 1992). Hence, somewhat paradoxically, induction of c-myc expression in factor starved cells leads to death. The oncogene bel-2 has a major effect in protecting cells from apoptosis, including the apoptosis induced by c-myc and a variety of other stimuli, physiological and otherwise (Fanidi et al., 1992; Bissonnette et al., 1992). Activated v-abl oncogene has also been shown to rescue cells from apoptosis (Evans et al., 1993). In contrast, expression of the p53 oncosuppressor gene has been reported to initiate apoptosis in epithelial, lymphoid and myeloid cells (Yonish-Rouach et al., 1991; Shaw et al., 1992; Clarke et al., 1993). Furthermore, mice that are homozygous for a targeted deletion of the retinoblastoma oncosuppressor gene (Rb-1) die in utero, showing a striking increase in apoptosis in particular locations within the developing nervous system (Clarke et al., 1992; Lee et al., 1992; Jacks et al., 1992). No mechanism has been proposed whereby these important genes exert their control over programmed cell death.

In this paper we provide evidence for the regulation of susceptibility to apoptosis by ras as well as myc oncoproteins, and indicate a possible mechanism for its control. We have constructed a family of cell lines by independent transfections of the human c-myc and mutated (T24) Ha-ras oncoproteins into a common parental rat fibroblast. We show that, under conditions in vitro in which the members of the family are closely similar in terms of proliferation, apoptotic rates differ over a fifteen-fold range. In confirmation of previous independent reports by ourselves and others, we show that high apoptotic rates are associated with c-myc expression and we present new data that expression of the Ha-ras oncogene has the opposite effect. These differences in apoptotic rates are associated with differences in the cellular content of an endogenous endonuclease, considered to be one of the effector elements of apoptosis (Wyllie, 1980; Arends et al., 1990; Wyllie et al., 1992). The results suggest that one action of these oncogenes is to influence the availability of apoptosis effector proteins, in a manner analogous to their action on proteins critical for cell replication.

Materials and methods

Cell lines

The parent cell line, the Fischer rat lung fibroblast 208F (Quade, 1979), was transfected with human c-myc linked to the Moloney virus LTR and a Hygromycin resistance marker (pHRMCGM1) using electroporation. Selection with Hygromycin B (HmB) was followed by picking single colonies as monoclonal cell lines (M7 and M8). The presence of exogenous DNA was confirmed by Southern analysis and PCR using c-myc specific primers. Human myc RNA expression was confirmed by reverse transcription-PCR using exon connection primers and RNA dot blot analysis (data not shown). A third c-myc transfectant was constructed independently by calcium phosphate transfection of pMCGM1 and neomycin selection (Spandidos & Wilkie, 1984). These were compared with transfectants of 208F cells containing the mutated T24-ras oncogene with a codon 12 valine substitution (Santos et al., 1982). T1 was generated by calcium phosphate transfection of the high expression plasmid pHOST1, in which the T24-ras is linked to the SV40 enhancer and a neomycin resistance gene (Spandidos & Wilkie, 1984). T2 and T3 contain pHOST1 modified to include a Hygromycin B resistance gene. Clones of these two transfectants were selected with HmB following electroporation. The parental line, 208F, was also transfected with non-oncogene-containing vector alone (pHOST5) and similarly selected by drug resistance, the resulting transfectants were non-transformed and appeared morphologically identical to the parental 208F cells. The presence of exogenous DNA was confirmed by Southern hybridisation analysis and PCR using Ha-ras specific primers (data not shown) (Bos et al., 1987), and enhanced expression of p21 confirmed by immunocytochemistry as described below.

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Cells were seeded into flasks in quadruplicate. 20 ml of Glasgow’s Modification of Eagles Medium (GMEM) and 10% Heat Inactivated Newborn Calf Serum (HINCS) was added and the flasks maintained at 37°C in a 5% CO₂ atmosphere for 48 h prior to analysis. The number of cells added was sufficiently low to avoid the establishment of a confluent monolayer by 48 h, as confluent monolayers tend to demonstrate increased apoptosis (Perotti et al., 1990). Cells were initially grown in the presence of GMEM/10% HINCS. At approximately 50% cell confluence the media was replaced by 20 ml of (i) fresh GMEM supplemented with 10% HINCS, (ii) fresh GMEM with 0.01% HINCS, (iii) GMEM without serum, or (iv) serum-free GMEM with added 25 μM mepulinolin, and thereafter the cultures were maintained at 37°C for 48 h. Mepulinolin (lovastatin) (kindly donated by Dr Alberts) was converted to its sodium salt prior to use as described by Kita et al. (1980).

Detection of ras product

Cell monolayers were harvested and resuspended in 2 ml of GMEM with 10% HINCS. 0.25 ml of cell suspension was added to sterilised slides, placed in sterile multiplate dishes (Lum Scientific Corporation). The cells were allowed to adhere (6–10 h) before being flooded in GMEM with 10% HINCS. For growth in a humidified 5% CO₂ atmosphere at 37°C the slides were washed in PBS and fixed for 4 min at room temperature using PLPD, a fixative known to be optimal for subsequent detection of p21ras with antibody Y13-259 (Going et al., 1988). Slides were washed in PBS and allowed to dry before storage at −20°C.

Relative levels of p21ras protein expressed by each cell line were measured in immunocytochemical preparations (Williams et al., 1985; Going et al., 1988) using the monoclonal antibody Y13-259 that specifically recognises p21ras proteins (Furth et al., 1982). The antibody was serially diluted in 20% newborn goat serum (NGS) to concentrations ranging from 1:500 to 1:20,000, and each dilution was pipetted on to a separate cell preparation for incubation for 30 min at room temperature, washed and detected by biotinylated goat anti-rat antibody diluted 1:50 in 20% NGS incubated for 30 min at room temperature. This was followed by three drops of avidin-biotin complex containing biotinylated peroxidase (ABCComplex; Dako) for 30 min, two washes, and the reaction was visualised with excess 3,3-diaminobenzidine tetrahydrochloride (DAB) solution for exactly 4 min. Samples with brown reaction product deposited on the membrane or in the cytoplasm of greater than 5% of cells were scored positive.

Cell turnover parameters

A proportion of fibroblasts growing in culture die by releasing contacts with their neighbours and the substrate, contracting in size, rounding up and floating as cellular bodies in the culture medium. This process was confirmed as apoptosis by fluorescence and electron microscopy of the cell bodies harvested from the media, which showed the characteristic morphology of apoptosis. (Wyllie et al., 1980; Wyllie, 1987; Arends & Wyllie, 1991) and gel electrophoretic analysis of extracted DNA, which demonstrated the typical 'chromatin ladder' representing oligonucleosomal fragments due to DNA cleavage by endogenous endonuclease (data not shown).

Apoptosis was measured as the number of apoptotic bodies accumulated over 2 days by a subconfluent monolayer. 75 cm² flasks were seeded with 1–8 × 10⁵ cells to generate, after 48 h, monolayers of 50%–80% confluence. The overlying medium was collected, including a further PBS rinse that was swirled over the monolayer cells, and together these were centrifuged at 3000 rpm for 10 min. Cell bodies released from the monolayer into the media – Released Cell Bodies (RCB) – were resuspended in a known volume of PBS and the total numbers of RCB were counted by haemocytometer. Twenty μl of this sample was mixed with an equal volume of 10 μg ml acridine orange on a glass slide and viewed under UV light. One to two hundred cellular bodies were counted and identified as viable, apoptotic or necrotic – on the basis of their characteristic morphological appearances (Kerr et al., 1972; Wyllie et al., 1980; Arends & Wyllie, 1991). This was used to calculate the proportion of apoptotic bodies (%A) and viable (%V) cells comprising the RCB. The monolayer was carefully harvested, ensuring complete removal of residual cells from the substratum by phase contrast microscopy, and counted by Coulter Counter (Model ZM; Coulter Electronics) and haemocytometer, to enumerate the total number of monolayer cells (MC). The apoptotic index (AI) was calculated as a measure of the production of apoptotic bodies per 100 attached cells over 48 h, using the following equation:

Apoptotic Index = [%A × RCB × 100]/(%V × RCB + MC)

The rate of population expansion (PE) was assessed as the ratio of the mean monolayer-cell number at 48 h to that at 24 h, in triplicate subconfluent experiments, whilst the cells were in maximal growth phase. This was calculated as a single measure of the proportional increase in cell number over one day, in order to compare different cell lines.

Cell proliferation was analysed for each cell line, using nuclei prepared for flow cytometric measurement of DNA content (Vindelov et al., 1983), harvested from monolayers growing in log phase. The computer software program, PCA 1 (Coulter Electronics), was used to determine the proportion of cells in each phase of the cell cycle. The alternative program SFITS (Coulter Electronics) was also used and gave very similar data. The growth fraction (GF) was calculated as the proportion of cells in S plus G2/M phases of the cell cycle.

Endogenous endonuclease activity

To obtain a semi-quantitative assessment of the nuclear endonuclease activity associated with viable cells in each cell line, subconfluent monolayers were prepared and fed with either GMEM/10% HINCS or GMEM/0.5% HINCS, for 24 h and again for 4 h prior to harvest. At harvest, monolayers were rinsed twice with PBS to remove any apoptotic cells in the supernatant, and the monolayers were stripped with trypsin and EDTA as before. Nuclei were prepared by isotonic lysis and centrifugation through a glycerol gradient (Arends et al., 1990). Approximately 10⁷ nuclei were resuspended in 200 μl incubation buffer containing 100 mM Tris-HCl, 2 mM CaCl₂, with or without MgCl₂ at 2 mM, and incubated at 37°C for 18 h. Previous studies in this laboratory (data not shown) have suggested that these conditions are likely to produce maximal demonstration of endonuclease activity with preference for internucleosomal cleavage. 2 mM zinc ions were also added to the incubation buffer in some experiments to inhibit nuclease activity (Cohen & Duke, 1984). DNA from these incubations was phenol/chloroform extracted, ethanol precipitated and analysed by 2% agarose gel electrophoresis, to reveal chromatin cleavage by the endogenous endonuclease activity. Prior to incubation, more than 99% of the nuclei were found to be non-apoptotic as judged by acridine orange staining and UV fluorescence microscopy.

Chromatin cleavage into the characteristic oligonucleosomal 'ladder' pattern of apoptosis was observed. To obtain a quantitative measure of the extent of digestion to lower order nucleosomal fragments, photographs of the ethidium bromide stained gels were scanned by laser densitometer (Ultrascan XL, enhanced laser densitometer, LBK). For each cell line a 'Digestion Ratio' (DR) was calculated. The DR numerator was the amount of DNA fluorescence in the lower order oligonucleosomes that comprised the mono-and di-nucleosomes together with the lower half of the tri-nucleosomal DNA band (an arbitrary cut off chosen for its accurate reproducibility on the densitometry graph), whereas the DR denominator was the total DNA fluorescence in the electrophoretic track, including the high molecular weight DNA smear.
Results

Myc and ras transfectants show similar measures of proliferation but divergent apoptotic rates

The six cell lines resulting from transfection of 208F with c-myc and T24-ras DNA all showed closely similar proliferation rates, analysed as Growth Fractions (S + G2/M phases of the cell cycle) during growth in log phase in 10% serum, and there were no statistically significant differences by ANOVA or Kruskal-Wallis tests (Table I). In contrast, the apoptotic indices differed from each other by more than fifteen-fold under these conditions (Figures 1 and 2a). The three c-myc transfectants showed significantly higher levels than the control 208F (P < 0.004 in all cases), and apoptosis was further increased by serum deprivation (0.01% serum) (P < 0.004 in all cases) (Figure 1). Two of the three T24-ras transfectants, T1 and T2, showed significantly reduced apoptosis compared to the control 208F (P = 0.004 and P = 0.005 respectively) (Figure 2a). T3, the third of the T24-ras transfectants, showed an apoptotic index not significantly different from the parental line. As expected, these large differences in apoptosis had great effects on the overall population expansion rates of the monolayers as a whole, and these two parameters showed a strong inverse correlation (r = -0.77) for the six oncogene transfectants (Table I).

High expression of mutated ras is associated with reduced apoptosis

We sought to establish the role of ras in modifying apoptosis by two classes of experiments. In the first, the three ras

| Table I Cell turnover parameters |
|----------------------------------|
| **Cell line** | **Growth fraction** | **Population expansion** | **Apoptotic index** |
|----------------|---------------------|--------------------------|---------------------|
| 208F           | 41.7 (± 2.98)       | 1.53                     | 6.98 (± 1.75)       |
| M1             | 44.3 (± 2.17)       | 2.89                     | 26.5 (± 7.97)       |
| M7             | 41.8 (± 3.37)       | 1.54                     | 30.2 (± 11.1)       |
| M8             | 45.5 (± 2.96)       | 1.20                     | 15.4 (± 4.44)       |
| T1             | 46.2 (± 1.17)       | 3.56                     | 1.7 (± 0.14)        |
| T2             | 45.9 (± 1.62)       | 3.67                     | 2.2 (± 0.12)        |
| T3             | 41.0 (± 1.32)       | 1.77                     | 8.4 (± 3.07)        |

Growth Fraction data (S + G2/M phases of the cell cycle analysed flow cytometrically using the PARA 1 program) are means (± s.e.m.) of 9-10 experiments and show no statistically significant differences by ANOVA or Kruskal-Wallis tests. Population Expansion (PE) data are means of triplicate experiments. Apoptotic Index (AI) data (at 10% serum) are means (± s.e.m.) of 6-21 experiments and were log transformed to normalise the distributions for statistical analysis. The means of the log AI values for the six oncogene transfectants correlate inversely (r = -0.77) with their PE values.

Figure 2 a, The 2 T24-ras-transfectants, T1 and T2 show less apoptosis than the control 208F cells, when grown at 10% serum (P = 0.004 and P = 0.005 respectively). Under conditions of serum deprivation, T1 shows a lower apoptotic index than 208F (P = 0.006), whereas T2 and T3 demonstrate higher rates (P = 0.004 and P < 0.00001 respectively). The addition of 25 µM mevinolin to serum-deprived cultures resulted in a considerable increase in the apoptotic index for all 3 T24-ras transfectants compared to 208F (P < 0.00001 in all cases) (all comparisons by Student's t-test, based on eight experiments). No significant change in the apoptotic index of 208F was observed. b, Apoptotic indices for ras transfectants and 208F cells grown at 10% serum are compared with relative levels of p21 expression (arbitrary units). High and intermediate levels of p21 expression (T1 and T2) are associated with significantly lower rates of apoptosis than that of the control, whereas a low level of p21 expression (T3) is associated with an apoptotic index similar to 208F. Apoptotic indices correlated inversely (r = -0.72) with p21 expression.

transfected lines were compared in terms of apoptotic rate and relative levels of expression of p21measured semi-quantitatively as the dilution of antibody Y13-259 at which membrane staining of monolayer-cultured cells was no longer evident. Whereas expression of p21 in T3 was approximately two-fold greater than in the parental line, T2 exhibited four-fold greater expression, and T1 ten-fold greater expression by this method (Figure 2b). These semi-quantitative assessments of p21 protein expression correlated well with estimates of the relative copy number of the transfected genes obtained from Southern hybridisation and densitometric analysis of the three cell lines (data not shown). The levels of p21 detected showed an inverse correlation with apoptotic indices (r = -0.72). Growing in log phase in 10% serum, apoptosis was lowest in the cell line with the highest copy number and p21 expression (T1) and highest in T3 in which expression and copy number were lowest. T2 showed an intermediate pattern both in p21 expression and apoptosis. Even in serum-deprived conditions, T1 showed low levels of apoptosis and retained high expression of p21. In contrast, both T2 and T3 showed elevated apoptotic rates under these conditions with reduction in p21 expression to undetectable levels (data not shown).
In a second series of experiments the isoprenylation inhibitor mevinolin was used in an attempt to inhibit the biological activity of the p21sup"ras protein by blocking its membrane attachment. Mevinolin, applied to serum-deprived cultures at a concentration known to inhibit p21sup"ras function (Hancock et al., 1989; Schaefer et al., 1989; De Clue et al., 1991), had little effect on the apoptotic rate of the parental 208F cell line. In contrast, there was a profound increase in apoptotic rates in all three T24-ras transfected lines, compared with apoptosis in its absence (P <0.001 in all cases) (Figure 2a). The parental line 208F showed no significant increase in apoptosis following treatment with mevinolin indicating that low level endogenous ras within this line is presumably not the prime regulator of apoptosis.

**Cellular content of endogenous endonuclease varies in proportion to susceptibility to apoptosis**

Previous experiments have shown that lymphoma cells about to enter apoptosis accumulate an endogenous endonuclease which can be activated by calcium and magnesium ions (Wyllie et al., 1986b and 1992). Elsewhere we have described such cells as 'primed' for apoptosis, to distinguish them from cells lacking this component of the effector pathway of apoptosis (Arends & Wyllie, 1991). It was therefore of interest to compare the nuclear content of this calcium/magnesium sensitive endonuclease (or endonucleases) in log phase cultures of the transfected cell lines. Parallel incubations of nuclear preparations from the parental and transfected cell lines revealed several consistent differences. First, incubation of the nuclei from many of the cell lines for several hours frequently produced internucleosomal cleavage of chromatin which was optimum in the presence of calcium plus magnesium ions, but could be inhibited by zinc ions (Figure 3). In every case, more than 99% of the nuclei showed a normal morphology (i.e. without the stigmata of apoptosis) prior to incubation. The three myc transfectant lines (M1, M7 and M8) consistently showed the highest nuclear content of the enzyme, as evident by the intensity of ethidium bromide staining of the chromatin ladder from equivalent cell numbers. The high p21sup"ras expressing cell line T1 consistently showed only minimal evidence of internucleosomal cleavage, even after overnight incubation with calcium and magnesium ions. The two T24-ras transfecants with lower levels of p21sup"ras expression (T2 and T3) generated fragmented chromatin to a varying degree, which was most conspicuous in cells harvested after several hours incubation in low serum (Figure 4). At high serum concentration there was some cleavage of chromatin, mostly to large oligonucleosomal fragments, but at low serum a greater proportion of digested chromatin appeared as small oligonucleosomes. This was confirmed by densitometric measurement of DNA bands and calculation of 'Digestion Ratios' (DR). Compared with serum supplemented cells, the DR increased in assays of serum-deprived T2 (from 27% to 45%) and T3 (from 53% to 61%), indicating more complete digestion to smaller oligonucleosomal chains by the endonuclease activity present in populations of serum deprived cells. The calcium and magnesium ion sensitivity of the endonuclease activity was confirmed in T2 and T3 nuclei prepared from cells grown at both serum concentrations. In contrast, T1 showed no significant endonuclease activity at low serum, whereas M8 demonstrated marked nuclelease activity at high serum that was further increased by serum deprivation (Figure 4).

**Discussion**

*C-myc is associated with a high turnover state*

The data show that cell lines of common rat fibroblast parentage can vary fifteen-fold in rates of apoptosis, whilst retaining similar cell proliferation kinetics. The high apoptotic indices in the c-myc transfecants confirm the observation made by ourselves and others that constitutive c-myc activation is associated with initiation of apoptosis, particularly in cells in which cell-cycle progression is inhibited by a variety of circumstances including reduced serum growth factor support (Wyllie et al., 1987 and 1992; Askew et al., 1991; Bertrand et al., 1991; Evan et al., 1992). However, there is incomplete data regarding the regulation of apoptosis by endogenous myc in normal fibroblasts. These conclusions may not apply to all cell types as some appear to die by apoptosis following the disappearance of myc (Yuh & Thompson, 1989). The c-myc transfecants reported here appear to be in a 'high turnover' state whilst growing in log phase, in which cell proliferation and death by apoptosis co-exist, so that the overall population expansion is substantially slower than expected from consideration of the cell
proliferation rate alone. There was no other obvious means of exit from the proliferating pool in these cultures (such as necrosis or differentiation). Thus, the rate of apoptosis appears to be the major regulator of the rate of population expansion for these cell lines growing in culture. Furthermore, high turnover states of this sort are almost universal in tumours growing in vivo in which deregulation of c-myc is also a common event (Field & Spandidos, 1990), supporting an important role for apoptosis in determining net tumour growth in vivo, although c-myc may not be the only cellular protein of significance in this regard.

Mutated Ha-ras is associated with suppression of apoptosis

Of the mutated Ha-ras transfectants, two lines showed suppressed apoptosis relative to the control. In experiments of this design, in which ras expression is not selectively reversible, it is not possible to be completely certain that the differences in apoptotic rate are due exclusively to expression of the transfected oncogene rather than incidental cellular changes arising during transfection and selection. We show here, however, that inhibition of apoptosis is proportional to expression of functional p21ras protein in the three independent transfectants, following serum withdrawal and immediately after blockade of p21ras processing by mevinolin in dosages previously shown to be effective in inhibition of isoprenylation (Hancock et al., 1989; Schaffer et al., 1989; DeClue et al., 1991). Strikingly, this pharmacological blockade induced similarly high apoptotic rates in all the three T24-ras transfectants although there was no immediate toxic effect in the parental 208F fibroblast. Inhibition of ras expression has been reported to precede, and perhaps be a prerequisite for apoptosis of cultured choleoreukemic cells (Servomaa & Rytoma, 1988). In a mouse mast cell line transfected with human activated Ha-ras oncogene in a regulable construct, ras activation was also associated with rapid growth, whilst reduced ras expression was accompanied by cell death (Andrejauskas & Moroni, 1989).

Susceptibility to apoptosis reflects availability of endonuclease in viable cells

We have argued elsewhere that entry to apoptosis requires two cellular events: 'priming', in which the effector proteins for apoptosis accumulate within the cell, and 'triggering', in which these proteins are activated (Arends & Wyllie, 1991; Dive & Wyllie, 1993). By implication, cells might exist in an unprimed state in which apoptosis would be impossible, at least temporarily, until the effector proteins accumulated. Certain cell types such as cortical thymocytes appear sensitive to apoptosis induced by a wide variety of disparate agents, ligands for the T cell receptor, dexamethasone, toxic agents such as TCD-Dioxin, or steroid hormones (Van Haelst, 1967; Umansky et al., 1981; Wyllie & Morris, 1982; Wyllie et al., 1984; McConkey et al., 1988; Yamada & Ohayama, 1988; Smith et al., 1989; Walker et al., 1991). These stimuli have clearly different initial effects upon the thymocyte, as shown by the fact that calcium signalling (McConkey et al., 1989a and 1989b) and p53 dependence (Clarke et al., 1993) are features of some but not all of them. Nonetheless, apoptosis is the final common event, complete with chromatin condensation and activation of an endogenous endonuclease which appears to be present constitutively in these cells (Wyllie, 1980; Arends et al., 1990). Similarly, divergent stimuli such as c-myc expression together with growth arrest or exposure to etoposide can induce apoptosis in cultured fibroblasts (Evan et al., 1992; Fanidi et al., 1992). However, fibroblasts and lymphocytes can be rendered insensitive to apoptosis in response to a similar variety of stimuli by suitable gene expression, such as ras as shown here and bcl-2 (Vaux et al., 1988; Nunez et al., 1990; Hockenbery et al., 1990; Evan et al., 1992; Fanidi et al., 1992; Bissonnette et al., 1992; Wyllie et al., 1992). The mechanism whereby these changes in susceptibility to apoptosis are engendered is not known, but one of the possibilities is that elements within the apoptosis effector pathway may themselves be regulable.

This proposition is difficult to test as no element of the effector pathway has been definitively identified and purified. In this paper, we have examined the availability of a nuclear calcium/magnesium sensitive endonuclease (or endonucleases), whose activity is apparently responsible for the chromatin cleavage of apoptosis and is induced prior to the onset of chromatin condensation (Wyllie et al., 1986a, 1986b, 1992 and 1993; Arends et al., 1990; Walker et al., 1991). Endonuclease activation appears to be a mid to late event in apoptosis, as in the T cell line death in other circumstances (Hengartner et al., 1992) and may provide a useful indication of the cells' ability to undergo the process as a whole. Here we demonstrate that abundance of readily activated endonuclease in viable fibroblast nuclei is associated with susceptibility of these cells to undergo apoptosis, being high in c-myc transfectants, apparently absent in the T24-ras transfectant, T1, with the lowest apoptotic rate and the highest expression of p21ras oncprotein, and substantially increased in serum-deprived cultures of the intermediate and low p21ras expressing lines, T2 and T3, in which apoptosis is also increased. Therefore, these experiments provide evidence for a mechanism whereby oncogenes may regulate apoptosis, even in the absence of definitive information concerning any of the effector molecules. Hopefully, when these effectors are isolated, it will be possible to test more precisely the hypothesis advanced here.

The role of oncogene modulation of apoptosis in tumour growth

Finally, the question arises as to the role of ras and myc oncogenes in the growth of authentic human tumours. The data reported here show that differences in myc and ras oncogene expression, are not limited to effects upon cell
proliferation, but can profoundly influence overall population expansion through modifying apoptosis. Elsewhere we show (Arends et al. unpublished data) that these transfected cell lines have differences in their rates and aggressiveness of tumour growth in vivo that correspond to the phenomena described here in vitro, extending the previously published data (Wyllie et al., 1987) that showed that a T24-ras expressing cell line (T1) produced a higher primary tumour take-rate and a significantly higher proportion of test mice with metastasis at 14 days, compared with a c-myc expressing line (M1). There is also evidence from human pathology that expression of myc is associated with tumours with high cell turnover (Field & Spandidos, 1990), whereas ras expression or oncogenic activation is associated with cell population expansion in potentially premalignant tumours of the colon and breast, rather than acquisition of specific features of the malignant phenotype (Williams et al., 1985; Gomp et al., 1992). Thus, the data support the hypothesis that oncogenes such as ras and bcl-2 can suppress apoptosis (Vaux et al., 1988; Hockenbery et al., 1990; Nunez et al., 1990). Reduction of apoptosis directly causes an increase in population expansion likely to produce greater retention of cells acquiring new mutations (Wyllie, 1985), increasing the risk of malignant transformation in a population of cells in a premalignant lesion, or increasing tumour progression in an already malignant population of cells, and both of these have been associated with ras activation (Brown et al., 1986; Buchmann et al., 1991). Furthermore, genetic modulation of apoptosis, by myc, bcl-2 and p53, determine to a large extent the cellular response to certain anticancer chemotherapeutic agents such as etoposide (Walker et al., 1991; Fanidi et al., 1992; Clarke et al., 1993; Evans & Dive, 1993). Thus, the levels of apoptosis in tumours appear to influence net growth, acquisition of new properties, and response to certain cytotoxic agents, and susceptibility to apoptosis is regulated by many genes including myc and ras.

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