ESTABLISHMENT OF OKADAIC ACID RESISTANT CELL CLONES USING CDNA LIBRARY EXPRESSION CLONING

*Tone Sandal, Johan Lillehaug, Stein Ove Døskeland, Ragnhild Ahlgren

*Department of Anatomy and Cell Biology, University of Bergen, Årstadveien 19, N-5009 Bergen, Norway
• tone.sandal@iac.uib.no

INTRODUCTION. The broad action of the phosphatases inhibited by this universal apoptogen has made it difficult to pinpoint the relevant molecular targets, and mechanisms involved in OA induced apoptosis(1,2,3,4). Retroviral cDNA-library expression cloning technique (5) report the isolation of more than twenty OA resistant cell clones, and characterisation of two of them reveals that the approach can produce OA resistant cell clones to be used as important tools to investigate mechanisms of apoptosis induced by serine/threonine phosphatase inhibitors.

METHODS. A Jurkat T-cell cDNA library, constructed in the retrovirus vector pBabeMN, was applied to produce recombinant retrovirus, according to previously described procedures (6) Mouse embryonic fibroblasts, C3H/10T1/2-cl-8. The bidirectionally oriented cDNA library, containing 1 x 106 independent cDNA inserts, was a kind gift from Dr. Nolan, Stanford University. After three rounds of screening, with lethal doses of OA, on infected cells, surviving OA resistant cell clones were selected. Isolation of cDNA from selected cell clones was performed by RT-PCR, characterised by sequencing and their antiapoptotic effect confirmed by recloning and retransfection into parental cells.

RESULTS. Twenty-one OA-resistant C3H10T1/2-cl-8 cell clones were selected after infection with a retroviral JurkatT-cell cDNA expression library (Fig. 1). Those clones with highest OA-resistance upon re-challenge (OAR-1, OAR-2) had retrovirally introduced, short cDNAs corresponding to two human genes with unknown function (Fig. 2). The oar1-cDNA sequence, in the antisense orientation, is nearly identical to a human DNA sequence (AP000795) localised on human chromosome 11 (11q14), recently submitted to the database. It is therefore most probable that oar1 acts as antisense RNA against mRNA coded by AP000795 or a similar gene not yet reported.
Oar2 is nearly identical to human DNA sequences within a gene (AK002158) with unknown function. Oar2 appears to have one region (nucleotides 226-376) that is identical to the predicted last part of the reading frame of AK0021258. The rest of oar2 corresponds, in antisense direction, to sequences within the 3'UTR of AK0021258. There is a potential large open reading frame including part of the antisense sequence and the whole of the sense sequence.

Reintroduction of oar-1 or oar-2 into wild-type cells reproduced the OA-resistant phenotype. OAR-1,2 were cross-resistant to other phosphatase inhibitors, but fully susceptible to staurosporin, indicating selective resistance to phosphatase inhibitors. The defect(s) in OAR1,2 appear to be downstream of phosphatases, since the 2-D phosphoprotein pattern was similarly affected by OA in wt cells and OAR1,2. We found that OA induced cell death was partially protected by the caspase inhibitor zVAD-fmk in both wt, OAR1 and OAR2, and neither clone was protected against microinjected cytochrome c, indicating that the defects are upstream of or independent of zVAD sensitive caspases and the apoptosome (apaf-1/caspase 9/3).
DISCUSSION. Isolated cDNAs responsible for Oa resistant affect in OAR1,2 did not point to any known protein, but rather to two recently sequenced genes (AP000795 and AK0021258) of completely unknown function. The elucidation of the importance of the two genes for apoptosis and of how the cDNAs of OAR1,2 work, will with high probability shed new light on the mechanisms of OA-induced apoptosis, and is a major undertaking in our laboratory. The novel cell clones will further be used to elucidate the significance for apoptosis of oar1,2, and their link to altered protein phosphorylation.

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REFERENCES.
1. Cohen, P.T. (1997) Trends Biochem. Sci. 22(7), 245-251
2. Honkanen, R.E. et al. (1994) Toxicon 32(3), 339-350
3. Boe, R. et al. (1991) Exp. Cell Res. 195(1), 237-246
4. Gjertsen, B.T. and Døskeland, S.O. (1995) Biochim. Biophys. Acta 1269(2), 187-199
5. Hitoshi, Y. et al. (1998) Immunity 8(4), 461-471
6. Pear, W.S. et al. (1993) Proc. Natl. Acad. Sci. U S A 90(18), 8392-8396
