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Expression of $V_{1\alpha}$ and GRP receptors leads to cellular transformation and increased sensitivity to substance-P analogue-induced growth inhibition

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Small-cell lung cancer (SCLC) is a particularly aggressive cancer, which metastasises early. Despite initial sensitivity to radio- and chemo-therapy, it invariably relapses, so that the 2-year survival remains less than 5%. Neuropeptides particularly arginine vasopressin (AVP) and gastrin-releasing peptide (GRP) act as autocrine and paracrine growth factors and the expression of these and their receptors are a hallmark of the disease. Substance-P analogues including [D-Arg$^1$,D-Phe$^5$,D-Trp$^7,9$,Leu$^{11}$]-substance-P (SP-D) and [Arg$^8$,D-Trp$^{7,9}$N$^\text{Me}$Phe$^3$]-substance-P (6–11) (SP-G) inhibit the growth of SCLC cells by modulating neuropeptide signalling. We show that GRP and $V_{1\alpha}$ receptors expression leads to the development of a transformed phenotype. Addition of neuropeptide provides some protection from etoposide-induced cytotoxicity. Receptor expression also leads to an increased sensitivity to substance-P analogue-induced growth inhibition. We show that SP-D and SP-G act as biased agonists at GRP and $V_{1\alpha}$ receptors causing blockade of G$q$-mediated Ca$^{2+}$ release while directing signalling to activate ERK via a pertussis toxin-sensitive pathway. This is the first description of biased agonism at $V_{1\alpha}$ receptors. This unique pharmacology governs the antiproliferative properties of these agents and highlights their potential therapeutic potential for the treatment of SCLC and particularly in tumours, which have developed resistance to chemotherapy.

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Lung cancer is the most common fatal malignancy in the developed world. Small-cell lung cancer (SCLC), which constitutes 25% of the total, is a particularly aggressive form of lung cancer. It metastasises early and over 90% of patients have widespread metastasis at presentation precluding curative surgery. Despite initial sensitivity to radio- and chemo-therapy, SCLC almost invariably relapses, so that the 2-year survival remains less than 5% (Smyth et al, 1986). Small-cell lung cancer is a paradigm for neuropeptide-driven tumourigenesis. Small-cell lung cancer cells proliferate in response to a range of neuropeptide growth factors and in many cases these neuropeptides are involved in autocrine and paracrine growth loops, which fuel unrestrained proliferation (Moody et al, 1983; Sethi and Rozengurt, 1991; Sethi et al, 1992; North, 2000). This is thought to be part of a special process of oncogenic transformation called Selective Tumour gene Expression of Peptides essential for Survival (STEPS) (North, 2000). The expression of neuropeptides and their receptors in tumour cells may render them sensitive to pharmacological therapeutic intervention and may be of diagnostic importance for the early detection of lung cancer and for the selection of appropriate treatment. Much of the recent studies in SCLC have focused on the roles of the specific mitogens gastrin-releasing peptide (GRP, the mammalian homologue of bombesin) and arginine vasopressin (AVP).

Gastrin-releasing peptide receptors are frequently aberrantly expressed in human neuroendocrine tumours of the breast, prostate, lung and colon (Carroll et al, 1999; Jensen et al, 2001) where they have both mitogenic and morphogenic roles and confer a survival advantage in proliferating cancer cells. It has been suggested that the detection of these markers in the peripheral blood of patients may be useful as early markers of SCLC. Scintigraphy with the labelled peptide ligand $99m$Tc-bombesin/GRP has been shown to detect prostate cancer and invasion of pelvic lymph nodes and should be applicable to other endocrine tumours particularly SCLC (De Vincentiis et al, 2004). Monoclonal antibodies have been developed against circulating bombesin and one such antibody 2A11 has been shown to inhibit the growth of SCLC in vitro and as xenografts in nude mice (Chaudhry et al, 1999); however, it has limited efficacy in human trials.

The presence of AVP and $V_{1\alpha}$ receptors on SCLC tumours has been extensively studied and the potential presence of an autocrine growth loop has been established. Small-cell lung cancer patients frequently display symptoms of inappropriate antidiuretic hormone secretion such as hyponatremia and urinary hyperosmolality (Johnson et al, 1997). Independent studies have shown the expression of $V_{1\alpha}$ receptors in five out of five SCLC lines and
zero out of four non-small-cell lung cancer (NSCLC) lines (Ocejo-Garcia et al, 2001) while we showed expression of V1A receptors in four out of four SCLC lines (Waters et al, 2003). Expression of V1A receptors and vasopressin is the most useful diagnostic tool for differentiating SCLC from NSCLC and other cancers (Coulson et al, 2003). Therapy that targets V1A receptors may be of potential benefit for SCLC. Small-cell lung cancer cells can also express the AVP gene as provasopressin, which remains attached to the cell membrane and conceivably contributes to the autocrine-driven mitogenesis (Friedmann et al, 1994). Antibodies recognising this cell surface antigen have been developed as a potential diagnostic and therapeutic tool that targets SCLC tumours in vivo (Keegan et al, 2002).

However, the main focus of research in this drug development strategy has been in the development of broad-spectrum neuropeptide antagonists. Synthetic analogues of substance-P for example, [Arg^i, d-Trp^7, N^mePhe^i]-substance-P (6–11) (SP-G) were initially identified as antagonists of substance-P-mediated cellular effects and were subsequently found to also antagonise the cellular effects of bombesin (Jensen et al, 1984). SP-G and its analogue SP-D inhibit calcium mobilisation stimulated by bombesin and vasopressin in SCLC cells (Woll and Rozengurt, 1988; Langdon et al, 1992; Sethi et al, 1992). They also inhibit mitogenesis by the same neuropeptides in both Swiss 3T3 cells and SCLC cells (Woll and Rozengurt, 1988; Sethi et al, 1992; Seckl et al, 1995) In addition to the in vitro growth inhibitory effects of substance-P analogues, these compounds inhibit the growth of tumours in xenograft models in nude mice (Langdon et al, 1992) and are more effective than specific neuropeptide receptor antagonists. SP-G has recently completed a phase I clinical trial where it shows minimal toxicity and therapeutic plasma concentrations (Clive et al, 2001). However, its mechanism of action is still being investigated.

Our previous work has shown that SP-G and SP-D do not act as simple competitive antagonists of GRP receptors but rather act as biased agonists, inhibiting GRP-stimulated PLC activation via Gq, while directly stimulating JNK and ERK via Gi and Gq, respectively (Tallet et al, 1996; Jarpe et al, 1998; MacKinnon et al, 1999, 2001). The prolonged stimulation of JNK and ERK coupled with an inhibition of intracellular $Ca^{2+}$ is fundamental for the antiproliferative and proapoptotic effects of these agents (MacKinnon et al, 2001, Waters et al, 2003). It is not known whether these compounds interact with vasopressin receptors in the same fashion.

It was the objective of the present study to examine the effect of GRP receptors and V1AT expression, the two most common neuropeptide receptors present on SCLC, on growth and transformation of the epithelial cell line, CHO-K1. We show that expression of these receptors leads to an increase in basal and anchorage-independent growth in low serum, and an increased sensitivity to substance-P analogues. As well as directing GRP receptor signalling, substance-P analogues also act as biased agonists at V1A vasopressin receptors. These combined effects will be useful for SCLC therapy and particularly in targeting well-differentiated tumours, which have developed resistance to chemotherapy.

**METHODS**

**Materials**

CHO-K1 cells were purchased for the European Cell Culture Collection; Dulbecco’s modified Eagle’s medium (DMEM), GRP, bombesin and monoclonal antibody to dihydropyridylated ERK 1 and 2 (M 8159) were from Sigma (Poole, UK); [d-Arg^i, d-Phe^i, d- Trp^7, Leu^i]-substance-P (SP-D) and [d-Arg^i, d-Trp^7, N^mePhe^i]-substance-P (6–11) (SP-G) were synthesised by Cancer Research UK (London, UK). The human V1A receptor construct in pcDNA3.1 was a kind gift from M Thibonnier (Case Western Reserve University School of Medicine, OH, US). The human GRP receptor in pcDNA3.1 was from J Battey (Albert Einstein College of Medicine, NY, USA). [123I]-Arginine vasopressin (2000 Ci mmol$^{-1}$) and [125I]-GRP were from Amersham International (Amersham, UK).

**Cell culture and transfection**

CHO-K1 cells were maintained in DMEM supplemented with 10% (v/v) foetal bovine serum (heat-inactivated at 37°C for 1 h), 50 $\mu$m -1 penicillin, 50 $\mu$m -1 streptomycin and 5 $\mu$m -1 -glutamine in a humidified atmosphere of 5% CO$_2$:95% air at 37°C. CHO-K1 cells were transfected with full-length human GRP receptor or human V1A receptor using lipofectamine plus (Invitrogen) as per the manufacturer’s instructions. Stable cell cultures were maintained in the presence of 400 $\mu$g ml$^{-1}$ G418 sulphate.

**Receptor binding**

Receptor binding was carried out in confluent and quiescent cultures of CHO-K1-V1AT and GRP cells in a binding medium containing DMEM, 1 mg ml$^{-1}$ bovine serum albumin and radioligand (1.0 nM GRP containing 2ng [125I]-GRP or 0.6 nM AVP containing 45 nCi [125I]-AVP. Incubations were carried out at 37°C for 30 min in the presence of inhibitors as indicated. Nonspecific binding was defined in the presence of 1 $\mu$m GRP or AVP, respectively. The reaction was stopped on ice and unbound ligand was removed by washing $\times$3 with ice-cold phosphate-buffered saline. After solubilisation in 0.1% NaOH, 2% Na$_2$CO$_3$ containing 1% SDS, bound ligand was estimated by liquid scintillation counting. The binding parameters $K_d$ and $B_{max}$ were calculated from competition binding isotherms with unlabelled ligand (DeBlasi et al, 1989). The $IC_{50}$ (concentration of drug displacing 50% specific binding) was converted to the inhibitory constant ($K_i$), where $K_i = IC_{50} / (1 + [ligand]/K_d)$ (Cheng and Prusoff, 1973).

**Growth assays**

**Liquid growth** Exponentially growing CHO-K1 cells were trypsinised and suspended in DMEM with 5% FCS at a density of $5 \times 10^7$ cells per plate in the presence or absence of mediators in triplicate. Cells were grown for 1–9 days and cell number determined using a Coulter Counter (model Z1, Coulter).

**MTT assay** In some assays, MTT (3-[4,5-dimethylthiazol-2-yl]-2,5-diphenyl tetrazolium bromide) formazan production (Sigma) was used to measure proliferation as per the manufacturer’s instructions.

**Clonogenic assay** CHO-K1 (2x10$^5$) viable cells were mixed with DMEM containing 0.3% agarose in the presence or absence of mediators and layered over a solid base of 0.5% agarose in DMEM in 35 mm plastic dishes. The cultures were incubated at 37°C for 1–10 days, and then stained with 1 mg ml$^{-1}$ nitro-blue tetrazolium (NBT, Sigma) overnight at 37°C. Colonies from 10 separate fields were counted using a microscope with a $\times$4 objective. Cloning efficiency is calculated as the percentage of original number of seeded cells forming colonies of > 6 cells.

**Aggregation assay** CHO-K1 cells were suspended in DMEM in the presence of 5% FCS and seeded into low adhesion tissue culture plates on top of a layer (1 ml) of 0.5% agar. Under these conditions, the cells did not adhere. Cells were maintained in culture for 7 days briefly trypsinised to disaggregate clusters and viable cells counted.

**Determination of intracellular Ca$^{2+}$ concentration**

CHO-K1 cells expressing the GRP or V1A receptor were grown to confluence on 10 cm plates and quiesced overnight in DMEM
containing 0.1% FCS. Cells were trypsinised and loaded with fura-2-tetraacetoxymethylester AME (FURA-2-AM, 1 μM) in calcium-free Hank’s balanced salt solution for 10 min at 37°C. The cells were pelleted and resuspended in 2 ml of Hank’s balanced salt solution containing 1.8 mM CaCl₂. Fluorescence was recorded in a fluorescence spectrophotometer (Perkin Elmer). Alternate dual wavelength excitation at 380 and 410 nm allowed ratiometric analysis of bound and unbound FURA-2AM when measured at 505 nm. [Ca²⁺] was calculated according to the equation 

\[ [Ca^{2+}] = \frac{K(F - F_{min})}{(F_{max} - F)} \]

where \( F \) is the ratio of the unknown sample, \( F_{max} \) is the ratio after the addition of 0.1% Triton X-100 and \( F_{min} \) is the ratio after Ca²⁺ chelation with 10 mM EGTA. \( K \) is the dissociation constant for Fura-2, which is 224 nM.

**Western blotting**

Quiescent cell cultures in six-well plates were treated as described in figure legends and lysed at 4°C in 0.25 ml lysis buffer containing: 25 mM HEPES pH 7.4, 0.3 mM NaCl, 1.5 mM MgCl₂, 0.2 mM EDTA, 0.5% Triton X-100, 20 mM β-glycerophosphate, 0.5 mM dithiothreitol, 1 mM sodium orthovanadate and protease inhibitors (Boehringer Mannheim, Sussex, UK; prepared as per the manufacturer’s instructions). Lysates were clarified by centrifugation, equilibrated for protein using BCA protein assay reagent (Perbio Science, Cheshire, UK) and denatured by boiling (5 min) in SDS-PAGE loading buffer. In all, 20 μl lysate/lane was resolved on 12% SDS-PAGE gels and electroblotted onto nitrocellulose membranes.

**RESULTS**

**Receptor binding**

To assess receptor expression, stable cultures of CHO-K1-GRP and CHO-K1-V1A cells were incubated with the appropriate radioligand, and receptor number and affinity were measured from competition binding curves with unlabelled ligand. Figure 1 shows that GRP receptors were expressed with \( K_d = 2.55 ± 0.84 \) nM and \( B_{max} \) of 1151 ± 326 sites/cell (n = 4). V1A receptors were expressed at a slightly lower level with a \( B_{max} = 450 ± 80 \) sites/cell and \( K_d = 2.98 ± 0.71 \) nM (n = 4, Figure 1B). The substance-P analogues SP-G and SP-D inhibited \([^{125}I]-\)GRP binding with affinities in the micromolar range (SP-G \( K_i = 19.4 ± 6.3 \) μM (n = 4), SP-D \( K_i = 0.64 ± 0.05 \) μM (n = 4, Figure 1)). Both analogues also inhibited V1A receptor binding (SP-G \( K_i = 3.50 ± 0.82 \) μM (n = 4), SP-D \( K_i = 8.58 ± 1.47 \) μM (n = 4)). This data show that SP-G is relatively (six-fold) more selective for the V1A receptor whereas SP-D is GRP receptor selective (13-fold).

**Figure 1**  
Receptor binding. Gastrin-releasing peptide (A, C) or V1A (B, D) receptor expressing CHO cells were incubated with \([^{125}I]-\)GRP or \([^{125}I]-\)AVP for 30 min at 37°C as described in Methods in the presence of various concentrations of unlabelled GRP (A), unlabelled AVP (B), SP-D or SP-G (C, D). Results are expressed as % total binding and represent the mean ± s.e.m. of four experiments performed in triplicate.
Effects of V1A and GRP receptor expression on cell growth in liquid culture and semisolid media

Figure 2A shows that in liquid culture, in the presence of 5% FCS, the GRP and the V1A expressing cells proliferated at a similar rate to untransfected CHO, but on day 6 after seeding, the GRP cells had reached a density of 4.5 × 10^5 cells per 100 mm dish. The cells did not appear to be contact inhibited and exhibited a rounded morphology. Cells over grew one another and significant numbers of cells were observed to be detached and growing in clusters (Figure 2B). Trypan blue exclusion revealed that the cells were > 95% viable. A similar, though less pronounced, effect was observed with the V1A-transfected cells, which reached a density of 3.1 × 10^6 cells per 100 mm dish. The parental cell line exhibited normal contact inhibition and had flat fibroblast morphology and did not show significant detachment. The end point density of these cells was 2.1 × 10^6 cells per 100 mm dish. These observations suggested that transfection of GRP and V1A receptors caused CHO-K1 cells to lose contact inhibition and adopt a more transformed phenotype. These findings led us to examine their growth in liquid culture and semi-solid media, which is widely regarded as being a hallmark of the transformed phenotype. Cells were trypsinised and suspended in 0.3% agarose in DMEM. Figure 2C shows that at day 9 after seeding, clusters of cells (> 6 cells) were visible with MTT staining. The cloning efficiency was significantly higher in the GRP- and V1A receptor-expressing cells (14.1%, 7% and 1.5%, P < 0.05, ANOVA) and the V1A receptor-expressing cells (8.5 ± 1.9%, P < 0.05, ANOVA) than the vector control cells in the presence of 5% FCS (5.5 ± 0.02%). In a separate series of experiments, the effects of SP-D and SP-G were examined (Figure 3A). At 30 μM, both analogues significantly reduced colony formation in the GRP and V1A receptor expressing cells but had no effect on the cloning efficiency of vector control CHO-K1 cells. The effect of receptor expression was also examined in an aggregation assay where cells are prevented from adhering and form aggregates in suspension. After 6 days, cells are briefly trypsinised and viable cells counted. These experiments showed that expression of the GRPr and V1A receptors increased anchorage-independent growth compared to wild-type cells or vector control cells. Similarly SP-G and SP-D were able to inhibit cell growth in receptor expressing cells (Figure 3B). Together, these data suggest that expression of the GRP and V1A receptors induce a more transformed phenotype in CHO cells and induce sensitivity to substance-P analogue-induced growth inhibition.

Chemosensitivity

The response to etoposide in control and receptor-transfected cells was measured by MTT accumulation. Figure 4 shows that after 48 h in culture in the absence of serum, etoposide produced a dose-dependent inhibition of proliferation in all cell types (IC50 = 12.4 ± 3.1, 8.1 ± 2.3 and 14.2 ± 4.0 μg/ml in CHO-WT, CHO-GRPr and CHO-V1Ar cells, respectively). Incubation with 50 nm of either bombesin or AVP added to CHO-GRPr or CHO-V1Ar produced a small but significant protection from etoposide, which was not observed when both neuropeptides were added to wild-type cells (IC50 = 13.0 and 26.9 μg/ml in control and AVP-treated V1Ar-expressing cells, respectively; and 6.30 and 12.7 μg/ml 1 bombesin-treated GRPr-expressing cells, respectively). At 40 μg/ml etoposide, vasopressin-treated V1Ar-expressing cells accumulated 92% more MTT than control cells.

Figure 2 Effect of neuropeptide receptor expression on cell growth liquid growth (A) CHO-K1 cells expressing vector, the GRP or V1A receptor were plated at a density of 1 × 10^4 cells/100 mm tissue culture dish and incubated at 37° C. Cells were harvested at various times and counted. Results represent the mean ± s.e.m. of three experiments performed in duplicate (*significantly different from vector control cells P < 0.05, ANOVA). (B) Representative images showing morphological features of transfected cells are shown. The rounded cobblestone features and higher cell density are evident in neuropeptide receptor expressing cells compared to the elongated contact inhibited monolayer exhibited by vector transfected controls. (C) Cholangial growth vector (open bars), GRP receptor (black bars) and V1A receptor (grey bars) transfected cells were plated at 1 × 10^4 cells well−1 in 0.3% agar in DMEM containing 1% (C) or 5% (D) FCS. At various time points, cells were stained with MTT and colonies counted at ×10 magnification. Results represent the mean ± s.e.m. of three experiments performed in duplicate (*significantly different from vector control cells P < 0.05, ANOVA).
The mobilization of calcium from intracellular stores is one of the earliest events triggered by neuropeptide receptor activation of Giq leading to PLC activation and subsequent generation of IP₃. In untransfected CHO-K1 cells, neither GRP nor AVP produced a significant change in [Ca²⁺]ᵢ (data not shown). In the CHO-GRP cells, GRP produced a concentration-dependent increase in [Ca²⁺]ᵢ (EC₅₀ = 2.00 ± 0.4 nM, n = 5, Figure 5A). SP-D and SP-G produced no change in [Ca²⁺]ᵢ on their own but inhibited GRP-induced [Ca²⁺]ᵢ elevation. Figure 5A and B show that SP-D and SP-G inhibited GRP-induced [Ca²⁺]ᵢ, with resultant pA₂ values of 7.21 and 5.72 for SP-D and SP-G respectively. In V₁A-expressing cells, AVP increased calcium mobilisation with an EC₅₀ = 3.1 ± 0.60 nM (n = 4, Figure 5C and D). As in the GRP-expressing cells, SP-D and SP-G did not mobilise intracellular calcium directly but inhibited the response to AVP (Figure 5C and D). SP-G was more efficacious in inhibiting the vasopressin response than the GRP response. PA₂ values for AVP antagonism were 7.23 and 6.53 for SP-D and SP-G, respectively. Together, these data suggest that when stably transfected into CHO cells the GRP and the V₁A receptor effectively couple to Giq to increase intracellular calcium and that the substance-P analogues are effective antagonists of this response.

**ERK activation**

AVP and bombesin have been shown to activate extracellular regulated protein kinases (ERK 1 and 2) in a variety of cell types expressing the receptors. Much of these data suggest a Giq- and PKC-dependent mechanism although these observations may be cell type-dependent (Della Rocca et al, 1999; Sinnett-Smith et al, 2000). Previous studies have shown that substance-P analogues reversibly inhibit ERK activation by neuropeptides in Swiss 3T3 cells; however, our studies suggest that in rat-1 fibroblasts and in human small-cell lung carcinoma cells (SCLC), in the absence of bombesin, substance-P analogues can activate ERK and c-jun kinase (JNK) in a GRP receptor-dependent manner (MacKinnon et al, 2001). In this study, ERK activation was measured in CHO cells expressing GRP or V₁A receptors in response to the appropriate neuropeptide or substance-P analogue. Activation was estimated in immunoblots from stimulated lysates using a

(Edward et al, 1995). In GRPR-expressing cells, bombesin treatment caused an increase of 52% MTT accumulation compared to untreated cells (P<0.01). These results suggest that neuropeptide receptor stimulation may contribute to an increase in chemoresistance.

**Figure 3** Effect of SP-D and SP-G on clonal growth. (A) Cells were grown in 0.3% agar with 1% FCS for 8 days in the presence or absence (open bar) of 30 μM SP-D (black bar) or SP-G (grey bar). Results represent the mean ± s.e.m. of three experiments performed in duplicate (*significantly different from wild-type controls P<0.005, ANOVA). Aggregation assay. (B) Wild-type, GRPR-transfected and V₁A-R-transfected CHO-K1 cells were plated into low adhesion tissue culture plates on top of a layer of 0.5% agar in DMEM containing 5% FCS at a density of 5 x 10⁴ ml⁻¹ in the presence of varying concentrations of SP-G. Cells were maintained in culture for 7 days, briefly trypsinised to disaggregate clusters and viable cells counted by propidium iodide exclusion.

**Figure 4** Effect of neuropeptide on chemosensitivity. Wild-type CHO-K1 cells (left) and cells expressing the V₁A (middle) or GRP (right) receptor were plated at a density of 1x⁴ cells per well of a 96-well tissue culture plate in DMEM with 10% FCS and incubated overnight 37°C. Cells were then incubated in serum-free media containing etoposide as indicated and in the absence (filled squares) or presence of either 50 nM AVP (open squares) or 50 nM bombesin (open circles) or both neuropeptides (wild-type cells) for 48 h at 37°C. Cell viability was assessed by MTT staining. Results are expressed as % viability in the absence of neuropeptide and are mean ± s.e.m. of four independent experiments (*significantly different from untreated etoposide control, P<0.05 ANOVA).
phosphorylation state-specific antibody. Lysates from quiesced cells stimulated for 10 min with either neuropeptide or SP-D or SP-G were separated on 12% gels and blotted for pERK2 or total ERK1/2. Figure 6A shows that bombesin stimulated ERK phosphorylation in GRP-expressing CHO cells. Quantification of five separate experiments showed that bombesin produced a maximal 3.5-fold increase in ERK phosphorylation at 1 mM with an EC50 of 0.56 nM. In V1A-expressing cells, AVP produced an increase in ERK phosphorylation of 4.2-fold with an EC50 of 0.72 nM, which correlated with its affinity for V1A receptors measured in binding studies and for the stimulation of intracellular calcium. Neither AVP nor GRP produced any stimulation of ERK phosphorylation in untransfected CHO cells (data not shown).

SP-D and SP-G stimulated ERK activation in GRP receptor expressing CHO cells (Figure 6B). Activation of ERK was evident at 3 µM for SP-D, but the maximal stimulation was less than that observed with bombesin (2.8-fold for SP-D and 2.1-fold for SP-G at 50 µM). The substance-P analogues also stimulated ERK in V1A-expressing cells. However, in these cells SP-G was more efficacious than SP-D giving a 4.3-fold stimulation at 50 µM compared to a 2-fold stimulation by the same concentration of SP-D. The observed increase in ERK phosphorylation was not due to a change in total ERK expression, as ERK1/2 immunoreactivity did not change with drug treatment. These data show that at concentrations, which block neuropeptide-stimulated calcium release, substance-P analogues are agonists for ERK activation in GRP and V1A expressing CHO cells.

Mechanisms of ERK phosphorylation

The mechanisms of GRP- and V1A-receptor-mediated ERK activation are not fully understood, although it has been shown in some cell types that activation is mediated by Gqi-induced PKC activation and subsequent activation of raf. Other GPCRs have been shown to activate ERK in a Gi and ras-dependent mechanism, which may or may not involve activation of PI3 K (Della Rocca et al., 1999). To examine the possible recruitment of Gs, CHO cells were incubated overnight in quiescent media containing 100 ng ml⁻¹ pertussis toxin. Figure 7 shows that in GRP receptor expressing cells, PTX produced a small but nonsignificant inhibition of bombesin-stimulated ERK activation whereas the response to 10 µM SP-D was completely abolished (P < 0.01, ANOVA). In V1A-expressing cells, the response to AVP was not inhibited by PTX pretreatment but the response to SP-G (20 µM) was completely inhibited (P < 0.01, ANOVA). In both cell types, ERK activation by 1 µM LPA was completely inhibited by PTX treatment (data not shown). This differential sensitivity to pertussis toxin indicates that dual Gi/Gq-coupled mechanisms activate the ERK 1/2 cascade via GRP and V1A receptors expressed in CHO cells.

DISCUSSION

This study utilised a model CHO cell system for neuropeptide receptor expression as these cells have an acceptable null background for these receptors. The results from this work shows that (1) GRP and V1A receptor expression leads to the development of a transformed phenotype in CHO-K1 cells. (2) Receptor expressing cells showed some increased resistance to the chemotherapeutic agent etoposide and increased sensitivity to the substance-P analogues, SP-D and SP-G. (3) SP-D and SP-G act as biased agonists at GRP and V1A receptors. This is the first demonstration of biased agonism at vasopressin receptors. (4) This pharmacological activity is crucial for the antiproliferative effect of these agents, which may be of particular benefit in more differentiated cancers that have developed resistance to chemotherapy.

Introduction of the neuropeptide receptors into CHO cells lead to an increase in proliferation and an increase in the ability of cells to grow as colonies in semisolid medium. The cells also showed a greater propensity to grow as nonadherent clusters. Colony growth was also evident in low serum suggesting that expression of these receptors allows for anchorage- and serum-independent growth consistent with transformation. Moreover, it suggests that these
Neuropeptide receptors induce transformation

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Figure 6 (A) ERK phosphorylation by neuropeptides. Quiescent GRP (left) or V1A (right) receptor expressing CHO cells were stimulated for 5 min with neuropeptide at the indicated concentrations. Aliquots of cell lysate were resolved by SDS–PAGE and Western blots probed with monoclonal anti-pERK1/2 antibody (upper panel) or polyclonal anti-ERK2 antibody (lower panel). Representative Western blots are shown. Bar graphs represent the mean optical density (mean ± s.e.m.) of four separate experiments. (B) ERK phosphorylation by substance-P analogues. Quiescent GRP (left) or V1A (right) receptor expressing CHO cells were stimulated for 5 min with SP-D (filled bars) or SP-G (open bars) at the indicated concentrations. Aliquots of cell lysate were resolved by SDS–PAGE and Western blots probed with monoclonal anti-pERK1/2 antibody (upper panel) or polyclonal anti-ERK2 antibody (lower panel). Representative Western blots are shown. Bar graphs represent the mean optical density (mean ± s.e.m.) of four separate experiments.

receptors show some constitutive activity, as wild-type receptor expression alone was sufficient to increase transformation in the absence of exogenously added neuropeptide. The ability to observe constitutive GPCR activity is not only determined by the intrinsic molecular properties of the GPCR studied but also by the specific cellular background. Apparent constitutive activity of a wild-type GPCR may vary substantially in various systems. Differences in GPCR expression level, GPCR desensitisation, G-protein complement and GPCR/G-protein stoichiometry can all contribute to such differences (Seifert and Wenzel-Seifert, 2002) This could explain why GRP receptors expressed in NCM460 cells, a nonmalignant human colon epithelial cell line, are constitutively active (Ferris et al, 1997) but are not in other cell types (Benya et al, 1994). Our findings are unlikely to be due to an overexpression artefact as the receptors in this study were expressed at levels not greater than those measured from a variety of other cancers (100–50 000 sites/cell, Jensen et al, 2001) So at these moderate levels, wild-type receptor was sufficient to activate downstream signals that promote cell survival under anchorage-independent conditions. The cells were, however, still able to respond to agonist, which resulted in a small but significant increase in resistance to the cytotoxic effects of etoposide. It is unclear how significant this change in sensitivity relates to the development of chemoresistance in vivo, but highlights an important area for future investigation.

Our previous work shows that sensitivity to AVP and GRP increases during the progression to chemoresistance in a set of SCLC cell lines (Waters et al, 2003). This would be analogous to the development of autocrine and paracrine growth loops in tumour development in vivo, with tumour cells becoming more dependent on neuropeptide growth factors for survival and subsequently developing chemoresistance. The mechanisms governing the survival advantage gained in neuropeptide receptor expressing
cells are poorly understood. During cancer progression, increased expression of GRP receptors leads to tumour cell differentiation, increased motility and adhesion to extracellular matrix via an increased activation of focal adhesion kinase (FAK) (Glover et al., 2004). Recent reports have shown that GRP receptors can activate insulin-like growth factor-1 receptors (IGF-IR) leading to Akt activation in prostate cancer cells (Sumitomo et al., 2001), a mechanism which may underlie bombesin-mediated cell survival. In another study, human chorionic gonadotrophin (hCG) increased ovarian cell survival by up-regulating IGF-1 (Kuroda et al., 1998). Conversely, specific bombesin antagonists have been shown to inhibit growth and decrease EGF receptor expression in SCLC cell lines (Koppan et al., 1998) and breast cancer cells (Bajo et al., 2002). Bombesin antagonists in combination with GIHRH antagonists have also been shown to down-regulate IGF-1, IGF-II, IGF receptors, GRP and EGF receptors in SCLC xenografts (Kanashiro et al., 2003). There is therefore potential for neuropeptides to regulate growth factor signalling pathways that control survival.

The present study shows that expression of GRP and V_1A receptors in CHO cells also leads to an increased sensitivity to substance-P analogues. This was demonstrated by an inhibition of clonal growth by substance-P analogues only in receptor-transfected cells. We previously showed that substance-P analogues are more effective at inhibiting growth of chemoresistant tumour cells expressing high GRP receptor levels and can sensitize cells to etoposide-mediated cell death (Waters et al., 2003). Moreover, diverse tumour types expressing GRP receptors have a greater sensitivity to SP-D (Figs. et al., 2003). Together, these data suggest that the increase in neuropeptide receptor expression that occurs as cells become more differentiated may contribute to the development of chemoresistance and may be exploited as a target for drug therapy. The potential role of neuropeptide receptors in the development of chemoresistance is an interesting finding that requires further study.

The mechanism of action of substance-P analogues was studied in receptor-transfected cells. Upon neuropeptide receptor transfection, the cells responded to the appropriate neuropeptide as expected for these predominately G_i-coupled receptors. The substance-P analogues behaved as antagonists for Ca^2+ release with potencies that reflected their affinities in receptor binding experiments, with SP-G showing some selectivity for V_1A receptors. This may be useful in tailoring therapy to a specific

tumour phenotype in as much as SP-G may be more beneficial in V_1A receptor expressing tumours while SP-D may be better for tumours, which express both GRP and V_1A receptors. Both neuropeptides and the substance-P analogues activated ERK in CHO cells expressing the GRP receptor and the V_1A receptor. No ERK activation was observed in untransfected CHO cells suggesting that the activation is receptor mediated. These observations suggest that substance-P analogues act as dual efficacy receptor agonists directing receptor signalling via the GRP and the V_1A receptor towards ERK activation while blocking liberation of intracellular calcium. Moreover, the substance-P analogues activate ERK via G_q as the response is pertussis toxin sensitive and more prolonged than that observed with neuropeptide (data not shown). These data support a multitrack signalling complex leading from neuropeptide receptor to ERK activation; where one track is mediated by G_q and most likely involves PKC, whereas another feeds into a G_i-mediated pathway (Wetzker and Bohmer, 2003). Although by far the majority of evidence suggests a preference for interacting with G_q some previous work has shown V_1A receptors coupling to G_{q3} in quiescent fibroblasts and in hepatocytes (Strakova et al., 1997; Abel et al., 2000) There is also evidence that GRP receptors couple to G_i in pancreatic acinar cells (Profrock et al., 1992), and that G_i mediates bombesin-stimulated chemotaxis in monocytes (Djannati et al., 2003). Our data suggest that in the presence of substance-P analogues, the neuropeptide receptors exist in a conformation that is unable to activate G_q but which is able to interact with G_i and activate ERK. These analogues may act at a common region of the GPCRs perhaps at the receptor/ G-protein interface, which favours coupling to G_i. Although activation of ERK is normally associated with a mitogenic response as is seen with the neuropeptides, deregulated ERK activation has also been shown to cause disruption of the cell cycle, growth arrest and apoptosis. In SCLC-activated raf-1, the upstream regulator of ERK, activation while blocking liberation of intracellular calcium. Moreover, the substance-P analogues activate ERK via G_q as the response is pertussis toxin sensitive and more prolonged than that observed with neuropeptide (data not shown). These data support a multitrack signalling complex leading from neuropeptide receptor to ERK activation; where one track is mediated by G_q and most likely involves PKC, whereas another feeds into a G_i-mediated pathway (Wetzker and Bohmer, 2003). Although by far the majority of evidence suggests a preference for interacting with G_q some previous work has shown V_1A receptors coupling to G_{q3} in quiescent fibroblasts and in hepatocytes (Strakova et al., 1997; Abel et al., 2000) There is also evidence that GRP receptors couple to G_i in pancreatic acinar cells (Profrock et al., 1992), and that G_i mediates bombesin-stimulated chemotaxis in monocytes (Djannati et al., 2003). Our data suggest that in the presence of substance-P analogues, the neuropeptide receptors exist in a conformation that is unable to activate G_q but which is able to interact with G_i and activate ERK. These analogues may act at a common region of the GPCRs perhaps at the receptor/ G-protein interface, which favours coupling to G_i. Although activation of ERK is normally associated with a mitogenic response as is seen with the neuropeptides, deregulated ERK activation has also been shown to cause disruption of the cell cycle, growth arrest and apoptosis. In SCLC-activated raf-1, the upstream regulator of ERK causes growth arrest (Ravi et al., 1998). It is our hypothesis that deregulated ERK coupled with a decrease in intracellular Ca^{2+} are important factors in the growth inhibitory/apoptosis inducing activity of substance-P analogues.

In summary, the results presented here show that neuropeptide receptor expression leads to the development of a transformed phenotype in CHO epithelial cells. These cells display increased sensitivity to substance-P analogues that parallels the situation observed in SCLC cells, which develop chemoresistance in vivo. These analogues activate GRP and V_1A receptors in such a way as
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To cause activation of only a subset of possible downstream signals. They are competitive antagonists for PI hydrolysis mediated via Gαq but have agonist activity that results in activation of ERK via Gi. This presents a novel pharmacological mechanism whereby substance-P analogues utilise the normally mitogenic neuropeptide receptors to transduce an antiproliferative and apoptogenic cell signal. Moreover, the concentration of SPG used in this study was detected in patient plasma following administration with no noticeable toxicity (Clive et al, 2001), suggesting therapeutic doses can be achieved clinically. Development of these agents may provide a unique opportunity to treat SCLC and may be of particular benefit for tumours that have developed resistance to chemotherapy.

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