Gas1 Is Related to the Glial Cell-derived Neurotrophic Factor Family Receptors α and Regulates Ret Signaling*

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The growth arrest-specific gene 1 (Gas1) protein has been proposed to function during development as an inhibitor of growth and a mediator of cell death and is also re-expressed in adult neurons during excitotoxic insult. Here we have demonstrated that the Gas1 protein shows high structural similarity to the glial cell-derived neurotrophic factor (GDNF) family receptors α, which mediate GDNF responses through the receptor tyrosine kinase Ret. We found that Gas1 binds Ret in a ligand-independent manner and sequesters Ret in lipid rafts. Signaling downstream of Ret is thus modified through a mechanism that involves the adaptor protein Shc as well as ERK, eventually blocking Akt activation. Consequently, when Gas1 is induced, Ret-mediated GDNF-dependent survival effects are compromised.

Gas1 is a glycosylphosphatidylinositol (GPI) anchored protein, expressed at growth arrest and down-regulated under proliferative conditions (1, 2). During development, Gas1 is widely expressed in the nervous system (3) and is associated with growth inhibition and cell death (4). In Gas1−/− mice the cerebellum is smaller than in wild type (5), indicating an unexpected role for Gas1 in proliferation. Moreover, Gas1 is induced by Wnt and interacts directly with sonic hedgehog (SHH), antagonizing SHH patterning function (6). In the adult, Gas1 is expressed during physiological apoptosis in some tissues (7), although not in brain (3). Aberrant Gas1 expression was nonetheless found in adult neurons during excitotoxicity, and its expression in neuroblastoma cells is associated with proapoptotic effects (8). Together, this suggests that Gas1 may have different functions within distinct cell contexts and that its mechanism of action depends on its spatiotemporal expression. No mechanisms have yet been proposed by which Gas1 transmits signals that affect decisions about cell proliferation, growth arrest/differentiation, or cell death.

The GFRα (GDNF family receptors α) are four GPI-anchored proteins that serve as a link between the neurotrophic factors GFL (GDNF family ligand) and the transmembrane receptor tyrosine kinase Ret (reviewed in Ref. 9). The GFL consist of four proteins, GDNF, neurturin, artemin, and persephin, representing an important class of soluble mediators of neuronal survival, neurite growth, and differentiation. GFL are critical regulators of neurodevelopment and support the survival of midbrain dopaminergic and spinal motor neurons (reviewed in Ref. 10). These effects are mediated through the multicomponent receptor system consisting of GFRα and Ret (11, 12). Activated Ret binds many different adaptor proteins to activate divergent downstream signaling pathways. Recruitment of Ret to lipid rafts seems to be important for the choice of adaptor molecule (13, 14), although the mechanisms are not yet fully understood (reviewed in Ref. 15).

To study the Gas1 mechanism of action, we undertook detailed sequence analysis and found a structural relationship between Gas1 and the GFRα. Based on this finding, we analyzed Gas1 involvement in GFRα-Ret receptor complex formation and downstream effects. Our results suggest that Gas1 is related to the GFRα and may have a regulatory function in Ret signaling. In a ligand-independent manner, Gas1 sequesters Ret to lipid rafts, mediates Shc and ERK recruitment to this membrane domain, and blocks Akt activation without affecting ERK activation. This indicates that the proapoptotic function of Gas1 could be associated to its interaction with Ret and to inhibition of a GDNF-dependent survival pathway.

**EXPERIMENTAL PROCEDURES**

Sequence Analysis—For sequence analysis, we related distant protein families via intermediate searches (16) using global hidden Markov model profiles (17). To improve profile quality, we searched against EST (expressed sequence tag) data bases. Alignment was produced with T-Coffee software (18) and was slightly refined manually; it is viewed with the Belvu program (www.cgb.ki.se/cgb/groups/sonnhammer/Belvu.html). Secondary structure was predicted using PhD (19). The location of predicted signal peptides and transmembrane regions is based on Signal-P and TMHMM2 programs, respectively (20, 21).

Phylogenetic Analysis—To determine the phylogenetic distribution of the domains in homologous proteins, the domains were considered as individual entries. We used standard methods based on progressive alignment (22), generating Neighbor-Joining trees (23) to establish topologies in 10,000 bootstrap replicates. As the sequences were small and divergent, we used a probabilistic approach using MrBayes (24), running for 900,000 generations and four independent Markov chains. We sampled 25,000 trees and generated a consensus. Trees were drawn using the TreeView tool (25). For clarity, only the probabilistic unrooted tree is shown (Fig. 2).

Molecular Modeling—Fold recognition analyses were performed using the 3D-Jury metaserver (26). The model was based on the published crystal structure from domain 3 of the GFRα1 receptor protein.
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GAS1 Is Related to the GDNF Family Receptors α—Multiple alignment of the GAS1 protein primary sequence from several species identified two cysteine-rich repeats, residues 48–147 and 166–243 in the human sequence, in which the relative positions of the cysteines were highly conserved (Fig. 1). Searches with the global profile HHMmer (hidden Markov model) of the cysteine-rich repeat region of GAS1 proteins three times with lyase buffer and solubilized in sample buffer. Anti-Gas1 has been described (8). Antibodies against Ret, Src, GFRA1, FRS2, ERK-2, thyroglobulin-stimulating hormone receptor (TSHR), and actin were from Santa Cruz Biotechnology, phosphotyrosine (4G10), Shc, and phospho-Shc were from Upstate Biotechnologies; Akt, phospho-Akt, and cAMP-response element-binding protein (CREB) were from Cell Signaling. TR was from Zymed Laboratories Inc., Myc, (9E10) was from BD Biosciences, and phospho-ERK was from Sigma. Blots were developed with chemiluminescence (Super Signal West Dura; Pierce or ECL Advance, GE Healthcare) and quantified using NIH Image software. Fold induction was calculated by ratio of phosphorylated form to total protein, and the ratio in basal condition was set to 1.0. Statistical analysis was performed using Student’s t-test or one-way analysis of variance followed by Tukey’s post hoc test for the significance of differences.

Flotation Gradient and Detergent-resistant Membranes—For analysis of detergent-insoluble complexes in flotation gradients, 1.5 × 10^6 cells were cooled on ice, washed with phosphate-buffered saline, and lysed in 350 μl of TNE buffer (50 mM Tris-HCl, pH 7.4, 150 mM NaCl, 5 mM EDTA) with 0.3% Triton X-100 (Calbiochem) as described (32). Cells were extracted (20 min, on ice) and the extract subsequently brought to 35% (v/v) Optiprep (Reactiva). The lysate (250 μl) was overlaid sequentially with 3.5 ml of 30% (v/v) Optiprep and 200 μl of TNE with detergent in SW60 tubes. After centrifugation (4 h, 170,000 × g, 4°C), six 600-μl fractions (1 to 6) were collected from the gradient (top to bottom) and precipitated with trichloroacetic acid (all steps performed in a 4°C cold room, on ice). Normalized protein amounts for each fraction were analyzed by SDS-PAGE and immunoblotting. As controls for the gradients, we used anti-Src (present in the raft-associated fraction one) and anti-transferrin receptor (present in the nonraft-associated fraction six). Detergent-resistant membrane fractions were isolated as described (13).

Flow Cytometry Analysis of Propidium Iodide Incorporation—SH-SY5Y cells were plated and infected after 12 h in culture. At 24 h postinfection, medium was replaced with serum-free medium; at 48 h, cells were treated with GDNF until 72 h postinfection. Cells were collected, prepared for cytometry using the Coulter DNA-Prep Reagents kit (Beckman–Coulter), and analyzed in a Coulter Epics XL-MCL Flow Cytometer (Beckman–Coulter). Ten thousand events were counted on the scatter gate. Expo32 was used as statistical analysis software. Experiments were repeated twice in quadruplicates and analyzed by one-way analysis of variance followed by Bonferroni test for the significance of differences.

RESULTS

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Cell Culture, Transfections, and Treatments—Human embryonic kidney (HEK)293T and mouse neuroblastoma Neuro2a (N2a) cells were cultured in Dulbecco’s modified Eagle’s medium (DMEM), human neuroectodermic cells SK-N-MC in DMEM/F12 Glutamax, and human neuroblastoma SH-SY5Y in RPMI; all media were supplemented with 10% fetal bovine serum, Glutamax-I, and penicillin/streptomycin. To select for cells SK-N-MC in DMEM/F12 Glutamax, and human neuroblastoma Neuro2a (N2a) cells were cultured respectively. The PCR product was cloned into pGEM-TEasy, sequenced, and the ratio in basal condition was set to 1.0. Statistical analysis was performed using Student’s t-test or one-way analysis of variance followed by Tukey’s post hoc test for the significance of differences.

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found GFRα with an E-value of 0.049 (sptrembl_id:Q6UXV0, residues 131–210). Reciprocally, the profile of the GFRα repeat domain detected Gas1 proteins with an E-value of 0.55 (swissprot_id:GAS1_HUMAN, residues 166–243). None of these profile HMMer searches retrieved any other related sequences. The human sequence sptrembl_id:Q6UXV0, GRAL(33), was automatically identified as a new GFRα member by Pfam (www.sanger.ac.uk/Software/Pfam/). In addition, secondary structure prediction for the Gas1 protein (Fig. 1) showed good agreement with the crystal structure of GFRα1 domain 3 (27).

We used sequence alignment of the repeated domain common to the GFRα and the GAS1 proteins for phylogenetic analysis. Taking into account both the short length and the high divergence of the alignment, low clade confidence values at deep branches of the tree are predicted. Nonetheless, the topology was consistent and sufficient to establish a relation among the GFRα domains (Fig. 2A). In addition to the sequence similarity, we found an overall resemblance in the domain architecture of GFRα and GAS1 proteins (Fig. 2B). We confirmed a signal peptide located at the N terminus of both GAS1 and GFRα1 and a GPI anchor at the C terminus of GFRα (11, 12) and GAS1 proteins (2). Mammalian GRAL has a C-terminal transmembrane domain.

To determine whether fold recognition analysis generated consistent results, we submitted the two GAS1 cysteine-rich domains (swiss-
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Because Gas1 does not affect GFRα1-Ret receptor complexes, we analyzed whether Gas1 influences Ret autophosphorylation and activation. We used the N2a neuroblastoma cell line, which expresses Ret and very low GFRα1 levels (11). Gas1 expression in serum-deprived N2a cells did not result in Ret tyrosine phosphorylation, although Gas1 coimmunoprecipitated with RetMyc in an ligand-independent manner.

Gas1 is widely expressed in the nervous system during mouse embryonic development (4) at times when Ret expression has also been reported, e.g. in brain at embryonic day 14 (E14) (35). To demonstrate that the Gas1-Ret interaction can occur in vivo, we used brain extract from E14 mice for immunoprecipitation with an anti-Ret antibody. Immunoprecipitated Gas1 was detected by Western blot (Fig. 4D, upper panel). To confirm this interaction, Gas1 was immunoprecipitated from E14 mouse brain extract and Ret was detected by Western blot (Fig. 4D, lower panel). These data indicate that Gas1 could interact with Ret in developing mouse brain.

**Gas1 Does Not Bind to GDNF**—As the Gas1-Ret interaction is ligand independent, we tested whether Gas1 interferes with the phosphorylation-competent complex formed by GDNF, GFRα1, and Ret in cross-linking experiments with radiolabeled GDNF in transfected HEK293T cells. GFRα1 bound 125I-GDNF, forming a complex of ~85 kDa (Fig. 5, upper panel) as described (11, 12), whereas Gas1 did not bind to 125I-GDNF (lane 2). In cells transfected with RetMyc alone or together with Gas1, we observed no binding to 125I-GDNF (lanes 4, 5). Cotransfection of Gas1 with RetMyc resulted in the formation of two complexes, at ~85 and 200 kDa, corresponding to the 125I-GDNF-GFRα1 and 125I-GDNF-GFRα1-RetMyc complexes, respectively (lane 6). Cotransfection of Gas1 with GFRα1 and RetMyc showed the same banding pattern as for GFRα1 and RetMyc alone, with no change in the level of the complex at 200 kDa (lane 7). The results indicate that neither Gas1 alone nor the Gas1-RetMyc complex is able to bind 125I-GDNF and that Gas1 does not interfere with GDNF-GFRα1 binding to Ret.

**Gas1 Modifies Ret Downstream Signaling**—Because Gas1 did not affect GFRα1-Ret receptor complex formation, we analyzed whether Gas1 influenced Ret autophosphorylation and activation. We used the N2a neuroblastoma cell line, which expresses Ret and very low GFRα1 levels (11). Gas1 expression in serum-deprived N2a cells did not result in Ret tyrosine phosphorylation, although Gas1 coimmunoprecipitated...
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with Ret (Fig. 6A); cells transfected with GFRα1 and stimulated with GDNF showed robust Ret phosphorylation (Fig. 6A) as described (11). To determine whether Gas1 is involved in the regulation of Ret downstream signaling, we transfected N2a cells with GFRα1 with or without Gas1. We analyzed the phosphorylation of Akt and ERK, two major effectors downstream of Ret, after serum deprivation and GDNF stimulation. To permit comparison after GDNF stimulation, we verified similar GFRα1 expression in all samples in each experiment. Coexpression with Gas1 did not modify GFRα1-GDNF-induced Ret autophosphorylation (Fig. 6B). The presence of Gas1 severely decreased GDNF-induced Akt phosphorylation from 2.9- to 1.4-fold compared with control cells (p < 0.01, n = 4) (Fig. 6B), whereas ERK phosphorylation increased from 2.0- to 4.5-fold (p < 0.01, n = 4). Time course analysis of Ret, Akt, and ERK phosphorylation showed that in the presence of Gas1, Akt phosphorylation was impaired at all times analyzed up to 2 h, whereas ERK phosphorylation took place earlier and lasted longer (not shown).

We analyzed the influence of endogenously expressed Gas1 on Ret signaling using the SH-SY5Y human neuroblastoma cell line, which expresses Ret and GFRα1 and responds to GDNF (36). Gas1 was expressed at very low levels in the presence of serum, whereas Gas1 mRNA was strongly induced after 16 h of serum deprivation (Fig. 6C). This correlated with increased Gas1 protein levels detected by immunoprecipitation (Fig. 6C). We used this cell model to analyze Ret signaling with or without Gas1. SH-SY5Y cells were serum deprived for 3 h (absence of Gas1) or for 24 h (presence of Gas1). Cells were subsequently stimulated with GDNF, and phosphorylation of Ret, Akt, and ERK was analyzed. Concurring with the results in transfected N2a cells (Fig. 6, A and B), in Gas1-expressing cells Ret phosphorylation was independent of Gas1 expression (Fig. 6D), whereas Akt phosphorylation decreased appreciably from 2.0- to 0.9-fold (p < 0.05, n = 3) and ERK phosphorylation increased from 1.9- to 3.2-fold (p < 0.01, n = 3) (Fig. 6D). As serum deprivation can affect other pathways, however, these...
data do not allow us to conclude that Gas1 is uniquely responsible for inhibition of Akt activation and increased ERK activation after GDNF stimulation.

To confirm the influence of Gas1 on Ret signaling, we analyzed GDNF-induced Ret, Akt, and ERK phosphorylation in SH-SY5Y cells in which Gas1 induction after serum deprivation was knocked down by lentivirus-mediated expression of an antisense Gas1 mRNA. Ret phosphorylation was similar in GFP-infected cells (Gas1-expressing cells) and in cells infected with antisense Gas1 (knockdown) (Fig. 6E). Gas1-expressing cells (GFP-infected) did not activate Akt after GDNF stimulation (1.3-fold) whereas cells infected with antisense Gas1 (knockdown) were able to do so (2.2-fold, p<0.001, n = 4) (Fig. 6E). ERK phosphorylation was similar both in Gas1-expressing cells and in Gas1-knockdown cells, 3.2- and 3.3-fold, respectively (Fig. 6E). These results indicate that endogenously induced Gas1 cannot activate Ret but modifies Ret downstream signaling induced by GDNF-GFRα1 so that Akt phosphorylation is reduced. The relevance of Gas1 in ERK phosphorylation is nonetheless unclear. Overexpressed Gas1 mediates an increase in ERK phosphorylation in response to GDNF. ERK is also strongly activated after 24 h of serum deprivation and GDNF stimulation as compared with cells serum deprived for 3 h. This increase was not reversed by antisense Gas1 mRNA, however, suggesting a minor role for Gas1 in ERK activation. Alternatively, serum deprivation may generate sufficient oxidative stress to strongly activate ERK after GDNF stimulation, also in the Gas1 knockdown cells. Taken together, our results show that Gas1 is a regulator of Ret downstream signaling, specifically blocking Akt activation.

Gas1 Recruits Ret to Lipid Rafts—The location of phosphorylated Ret in the cell membrane is an important determinant for the activation of distinct downstream signaling pathways (14). GFRα1 is constitutively located inside lipid rafts through its GPI anchor, whereas in unstimulated cells Ret is found mainly outside rafts. After GDNF stimulation, GFRα1-GDNF complex formation induces transient Ret recruitment to lipid rafts (13) and activated Ret is in equilibrium between detergent-resistant membranes and the soluble membrane fraction (14). To determine Gas1 location in the plasma membrane, we used N2a cells and SK-S3, a clone derived from the human neuroectodermic SK-N-MC cell line. SK-N-MC cells express endogenous GFRα1 and GFRα2 (37), and the SK-S3 clone stably expresses Ret9, the short isoform of Ret (38). We transiently transfected N2a and SK-S3 cells with Gas1 and sepa-
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FIGURE 7. Ret is recruited to lipid rafts in the presence of Gas1. A, N2a or SK-S3 cells were transfected with Gas1 + GFRα1 or Gas1, respectively, and serum deprived. Flotation gradients were prepared and fractions analyzed by immunoblotting. B, Ret recruitment to rafts was analyzed in SK-S3 cells. In empty vector-transfected cells, Ret is recruited only in the presence of GDNF (upper and lower left panels). Gas1 promotes strong Ret recruitment to the first fraction with or without ligand (upper and lower right panels). C, detergent-resistant and soluble membrane preparations from transfected N2a cells were used for the analysis of Ret phosphorylation. Cells expressing Gas1 showed higher levels of Ret phosphorylation in the detergent-resistant fraction (upper panel) with a corresponding reduction in the soluble fraction (lower panel).

Gas1 Mediates the Recruitment of Activated Shc and ERK2 to Lipid Rafts—Ret can activate the PI3K/Akt pathway through various adaptors (15). To explain Gas1-mediated impairment of Akt activation, we analyzed activation of the adaptor FRS2, which transduces Ret signaling in rafts and is proposed to activate the PI3K/Akt pathway (39). We also analyzed the Shc adaptor, which transduces Ret signaling outside rafts (14) and activates the PI3K/Akt pathway (40). We immunoprecipitated FRS2 from N2a cell extracts and analyzed tyrosine phosphorylation by immunoblotting. GDNF stimulation induced a strong tyrosine-phosphorylated band at 75 kDa (3.5-fold), corresponding to activated FRS2. In the presence of Gas1 and GDNF, the level of activated FRS2 was higher (5.2-fold, p < 0.05, n = 3) (Fig. 8A, left), in agreement with the Ret increase in rafts. Gas1 thus increases GDNF-induced FRS2 activation, which concurs with the fact that Gas1 is changing the raft/non-raft equilibrium of Ret. It cannot, however, explain the reduction in Akt phosphorylation.

In cells transfected with GFRα1 and stimulated with GDNF, Shc phosphorylation increased 1.8-fold (Fig. 8A, right) as described (40). Strikingly, in cells transfected with GFRα1 and Gas1, Shc was strongly activated, 3.5-fold (p < 0.01, n = 3) in the absence of GDNF. Addition of GDNF resulted in rapid Shc dephosphorylation to basal levels (1.2-fold) (Fig. 8A, right). The strong, Gas1-mediated Shc activation in the absence of GDNF did not result in activation of Akt or ERK, as shown in Fig. 6B. GDNF addition thus reversed Shc phosphorylation, suggesting that Shc is involved in the mechanism that blocks Akt activation in the presence of Gas1.

Gas1 recruits non-phosphorylated Ret to lipid rafts (see Fig. 7), which may contribute to the modified function of adaptor proteins. We thus analyzed recruitment to lipid rafts of Shc, as well as of ERK, a major downstream effector that is implicated in negative loops that regulate adaptor signaling (41, 42). In the absence of Gas1 and prior to GDNF stimulation, we detected no Shc or phospho-ERK recruitment to lipid rafts (Fig. 8B, upper left). In unstimulated cells transfected with Gas1, we observed strong Shc recruitment and a single band of activated ERK in lipid rafts (Fig. 8B, upper right). An ERK2-specific antibody identified the single phospho-ERK band as ERK2 (Fig. 8B).

Gas1 Blocks GDNF-induced Survival—Because Gas1 blocks GDNF-induced Akt phosphorylation, we studied the effect of endogenous Gas1 expression on GDNF-mediated cell survival after serum deprivation. We quantified cell viability after knock down of endogenous Gas1 expression induced by serum deprivation, using a lentivirus expressing
antisense Gas1 mRNA (LV-ASgas1). We used a lentiviral vector coding for GFP (LV-GFP) as a negative control and to monitor equivalence of infectivity between experiments as determined by fluorescence-activated cell sorter. Infection with lentiviral particles had no effect on SH-SY5Y neuroblastoma cell viability compared with uninfected cells (Fig. 9 and not shown). In LV-GFP-infected cells, serum deprivation-induced cell death was only slightly modified by GDNF treatment (100 ng/ml) (Fig. 9). Knock down of endogenously induced Gas1 in LV-AS-gas1-infected cells improved cell survival only slightly. Treatment with the same GDNF concentration in Gas1 knockdown cells, however, significantly rescued neuroblastoma cells, reducing the percentage of cell death by half (Fig. 9). These data suggest that endogenous Gas1 expression counteracts the GDNF-activated survival pathway by providing a scenario in which Gas1 and Ret interaction has clear physiological consequences.

**DISCUSSION**

Sequence alignments, secondary structure predictions, and threading analyses yielded a template suitable for building a feasible model of GAS1. The GFRα1 structure is maintained by cysteine bridges (27), and although the conservation in the alignment was low, Gas1 proteins conserve these critical residues. The E-values obtained in HMMer searches showed a high degree of reliability, and based on the biological evidence presented here, we conclude that Gas1 is related to the GDNF family receptors α. The phylogenetic analysis did not provide high confidence branches; this is expected due to the short and highly divergent nature of the sequences, but tree topology was consistent when analyses were independent. This suggests that Gas1 was separated from GFRα very early in metazoan evolution and developed behavior different from that of GFRα. The ability of Gas1 to interact with Ret in the absence of ligand reflects this early divergence, suggesting that Gas1 could have functions or partners different from those described for the GFRα.

Our data suggest that Gas1 is a regulator of Ret signaling. The temporal expression patterns of these two proteins at defined developmental stages suggests a physiological function for the Gas1-Ret interaction. Gas1 and Ret are both expressed in midbrain, hindbrain, and spinal cord at E10–10.5 (4, 35). At E14–14.5, Gas1 and Ret are present in hindbrain, spinal cord, dorsal root ganglia, lung bronchioles, and gonads, although coexpression has not been analyzed in the different cell types in these tissues (4, 35). Another putative site for a Gas1-Ret interaction is the mouse embryonic retina. Ret is expressed strongly at E13.5 in neural retina and in the retinal pigmented epithelium, a layer in which Gas1 transcripts have been detected from E12.5 to E14.5 (43). Gas1-deficient mice show transdifferentiation of

![Image](https://example.com/figure8.png)

**FIGURE 8.** Gas1 modifies the profile of adaptor proteins in lipid rafts. A, transfected N2a cells were serum deprived and stimulated with GDNF. After immunoprecipitation with an anti-FRS2 antibody, FRS2 phosphorylation (left panel) was analyzed by immunoblotting with a phosphotyrosine-specific antibody. Phosphorylation of Shc (right panel) was analyzed by immunoblotting using a specific anti-phospho-Shc antibody. B, flotation gradients prepared from transfected N2a cells, serum deprived and GDNF stimulated, were analyzed by immunoblotting. In cells transfected with GFRα1 alone, recruitment of nonphosphorylated ERK2 was detected neither before nor after GDNF stimulation (upper and lower left panels). Cells transfected with Gas1 and GFRα1 showed Shc and activated ERK2 recruitment to lipid rafts prior to GDNF stimulation (upper right). After GDNF stimulation, the pattern of raft recruitment was similar to that in GFRα1-only-transfected cells (lower right).

![Image](https://example.com/figure9.png)

**FIGURE 9.** Gas1 blocks GDNF-induced survival. SH-SY5Y cells were infected, serum deprived to induce Gas1, and then GDNF treated. Results are the mean of two experiments performed in quadruplicate. **, p < 0.01.
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...the retinal pigmented epithelium to neural retina, (43), which may indicate a Gas1 association to differentiation. Taken together, the expression data and the present results suggest that Gas1 may have an influence on Ret signaling during development.

Through the tyrosine kinase Ret, the GDNF family ligands are able to activate many different signaling pathways (15). Akt and ERK are two main effectors of Ret signaling, and both proteins are usually activated together. Because Akt is associated with survival and proliferation (37), whereas ERK is implicated in differentiation and neurite outgrowth (37, 44), there must be a coordinating mechanism that permits them to mediate the different biological activities of Ret. Ectopic Gas1 expression in N2a cells suggests that Gas1 negatively regulates GDNF-induced Akt signaling whereas ERK activation is increased. Transient knock down of endogenous Gas1 in SH-SY5Y cells confirmed the ability of Gas1 to block GDNF-induced Akt phosphorylation. The role of endogenous Gas1 as a positive regulator of ERK activation has not been completely resolved, however, because induced levels of activated ERK, which are resistant to Gas1 knock down, could be due to activation of parallel signaling pathways during serum deprivation.

In the absence of Gas1, Ret is located mainly outside rafts; following GDNF stimulation, Ret is recruited to the rafts and is tyrosine phosphorylated. Activated Ret is in equilibrium between rafts and soluble membrane, where Ret associates with the adaptors FRS2 and Shc, respectively (14). Ret downstream signaling from outside rafts seems to be less efficient, according to results using the GFRα1 (13) and Gas1 TM constructs (not shown). FRS2 and Shc both activate Akt and MAPK, eventually resulting in survival/differentiation. These two pathways are activated by the formation of different, independent complexes. One complex involves Grb2-Gab1/2 and leads to recruitment of p85, which activates the PI3K/Akt pathway; the other, mediated by Grb2-Sos, leads to Ras/MAPK activation (39, 40).

Gas1 is located in lipid rafts and recruits Ret to these microdomains, resulting in a higher level of activated FRS2. A consequent increase in the activation of Akt and ERK would be predicted. Remarkably, the Akt pathway was inhibited whereas the ERK pathway was not. To explain these modifications, it is important to consider the Gas1-induced changes that occur before GDNF stimulation; these are Ret recruitment to rafts, which promotes Ret interaction with FRS2, and the recruitment of activated Shc and ERK2. The biological significance of these latter modifications is presently unknown, but they can be mechanistically linked to the negative regulation of the PI3K/Akt pathway.

Signals through Ret activate PI3K/Akt through many different adaptors, which must be blocked by Gas1 to inhibit the PI3K/Akt pathway. One candidate mediator of this inhibition is the scaffold adaptor protein Gab1/2, which links the adaptors and the PI3K regulatory subunit p85 (15). Gab1/2 can be inhibited or activated by ERK via phosphorylation at different threonine residues. Phosphorylation of Gab1 at Thr477 enhances recruitment, whereas phosphorylation at a still unidentified threonine residue correlates with inhibition of PI3K activation (42). We observed ERK2 inhibition of Gab1/2 in the presence of Gas1 before stimulation; this could agree with existing data (39, 40), as the block in PI3K activation via Grb2-Gab1/2-p85 does not impair Grb2-Sos-Ras formation and activation of MAPK. In addition to Gab1/2, another protein(s) must participate in mediating the Gas1 block of Akt activation. Among these, we might hypothesize a protein phosphatase that rapidly inactivates Shc after GDNF stimulation in the presence of Gas1. The nature of this phosphatase, as well as of other mediators of Gas1-induced Akt inactivation, is presently unknown.

Gas1 is proapoptotic during development (4) and participates in excitotoxic neuronal death in adult brain (8). Here we have shown that GDNF-induced Akt phosphorylation is blocked by Gas1. GDNF-induced neuron survival and proliferation is dependent on PI3K/Akt activation and not on the Ras/ERK pathway (40, 37). Suppression of the Akt phosphorylation blockade by antisense Gas1 knock down concurs with the improved survival effect of GDNF in serum-deprived SH-SY5Y cells. Our data show that Gas1 also modifies GDNF-induced ERK activation. ERK phosphorylation is associated to differentiation (37, 44), and it is tempting to speculate that Gas1 could function as a switch between proliferation and differentiation during neuron development. In the adult brain, ERK activation has also been associated with a proapoptotic function, because in vivo inhibition of ERK activation after ischemia promotes cell survival (45). Emerging data relating ERK activation with neuron degeneration and apoptosis were recently reviewed (46). A putative dual mechanism for the proapoptotic function of Gas1 in adult brain could thus be linked to Akt blockage as well as to ERK activation.

Gas1 regulates Ret signaling in a ligand-independent manner, and our data show that SHH does not affect the Ret-Gas1 interaction. Recently, however, SHH was described as repressing GDNF-induced differentiation and migration of neural crest cells (47). Gas1 interacts with SHH (6) and with Ret; for this reason, an interaction involving SHH and Ret cannot be excluded. Gas1 blocks the cell cycle in fibroblasts (1), a cell type that does not express Ret, indicating that other transmembrane receptor(s) could also mediate the biological effects of Gas1 in different cell systems. GFRα1 is known to interact with p140 N-CAM (30), for instance, suggesting that cell adhesion molecules (CAM) could be potential partners for Gas1 biological effects. Nevertheless, because of the early divergence between GFRα and Gas1 during evolution, this putative interaction between cell adhesion molecules and Gas1 must be demonstrated experimentally. The interaction between Gas1 and Ret provides a new framework to understand the functions described for Gas1 during development and neurodegeneration.

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