Binding of SH2 Domains of Phospholipase C_{γ} 1, GAP, and Src to Activated Growth Factor Receptors

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Phospholipase $C_{\gamma}1$ (PLC_{γ}) and p21^{ras} guanosine triphosphatase (GTPase) activating protein (GAP) bind to and are phosphorylated by activated growth factor receptors. Both PLC_{γ}l and GAP contain two adjacent copies of the noncatalytic Src homology 2 (SH2) domain. The SH2 domains of PLC_{γ}l synthesized individually in bacteria formed high affinity complexes with the epidermal growth factor (EGF)- or platelet derived growth factor (PDGF)-receptors in cell lysates, and bound synergistically to activated receptors when expressed together as one bacterial protein. In vitro complex formation was dependent on prior growth factor stimulation and was competed by intracellular PLC_{γ}l. Similar results were obtained for binding of GAP SH2 domains to the PDGF-receptors. The isolated SH2 domains of other signaling proteins, such as p60^{src} and Crk, also bound activated PDGF-receptors in vitro. SH2 domains, therefore, provide a common mechanism by which enzymatically diverse regulatory proteins can physically associate with the same activated receptors and thereby couple growth factor stimulation to intracellular signal transduction pathways.

variety of polypeptide hormones that elicit cell growth and differentiation bind to cellsurface receptors with intracellular protein-tyrosine kinase domains (1). Growth factors apparently activate their receptors by inducing receptor dimerization and subsequent autophosphorylation on tyrosine, evoking a catalytically active receptor capable of phosphorylating cellular substrates (1 - 3). Activated EGF- and PDGF-receptors (EGF-R; PDGF-R) complex with a set of cytoplasmic proteins that directly regulate intracellular signal transduction pathways. These include the v1 isoform phosphoinositide-specific the phospholipase C (PLC) (4, 5), p21^{ras} GTPase activating prorein (GAP) (6, 7), phosphatidyl inositol (PI) 3'-kinase (8), and $p74^{raf}$ (9). These results suggest that critical targets for receptor tyrosine phosphorylation are selected from the pool of potential substrates by their ability to physically complex with the receptor. A simple mechanism to accomplish these interactions would be the provision of cytoplasmic ligands with a common structural domain that autophosphorylated recognizes receptors.

The proteins that associate with activated growth factor receptors have quite distinct enzymatic properties and are structurally unrelated within their catalytic domains. However, PLC_{γ}I (*10*) and GAP (*11*) each contain two adjacent copies of a noncatalytic domain of ~100 amino acids, called the Src homology (SH) region 2 (*12*) (Fig. 1). The SH2

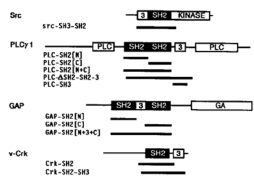
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first identified domain was in nonreceptor protein tyrosine kinases like Src and Fps, by its apparent ability to interact with the kinase domain and phosphorylated substrates (13 - 15). An SH2 sequence has also been identified in the v-Crk oncoprotein, which complexes with several tyrosine phosphorylated proteins in crk-transformed cells (16). Most SH2-containing proteins also contain a motif, SH3, which is found independently in several cytoskeletal proteins and may mediate interactions with the cytoskeleton (12, 16, 17). The SH2 domains have been implicated in protein-protein interactions that involve protein-tyrosine kinases and their substrates (13, 15). This raises the possibility that enzymes such as PLC_y and GAP associate directly with activated tyrosine kinase receptors by virtue of their SH2 domains (18).

To test this hypothesis, restriction sites were introduced into the complementary DNA (cDNA) for bovine PLC_vl, which allowed the precise excision of the NH,terminal and COOH-terminal SH2 domains (SH2[N] and SH2[C]), either alone or together (Fig. 1) (19). The individual SH2 domains, or the two SH2 domains together (SH2[N + C]) were introduced into a bacterial expression vector (pATH) and expressed as TrpE fusion proteins in Escherichia coli. These proteins were isolated from bacterial lysates-by immunoprecipitation with antibodies to TrpE (anti-TrpE) attached to Sepharose beads (20). The immobilized bacterial proteins were then incubated with lysates of either Rat-1 cells that expressed the human EGF-R, which had been stimulated with EGF or Rat-2 cells that expressed the PDGF-R, which had been stimulated with PDGF. The immunoprecipitates were recovered, washed extensively, and analyzed for associated phosphotyrosine (P.Tyr)containing proteins by immunoblotting with antibodies to P.Tyr (anti-P.Tyr) (Fig. 2). The TrpE-PLC-SH2[N] fusion

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protein complexed specifically with a 180-kilodalton (kD) P.Tyr-containing protein in lysates of EGF-stimulated cells. Immunoblotting of duplicate samples with antibodies to the EGF-R confirmed that this protein was the EGF-R and showed that its in vitro association with the PLC_{γ}l SH2[N] domain was EGF-dependent (Fig. 2). The $PLC_{\gamma}l$ SH2[N] domain was more efficient than the SH2[C] domain in its ability to bind the EGF-R. Interestingly, the fusion protein that contained both NH2- and COOH-terminal SH2 domains bound two to fourfold more EGF-R in EGFstimulated cell lysates than could be accounted for by the two individual SH2 domains. The PLC_vl SH2 domains therefore functioned synergistically in binding to the activated EGF-R. Very similar results were obtained for interactions of the PLC_vl SH2 domains with the PDGF-R (Fig. 2). The PLC_yl SH2[N] domain bound the PDGF-R in lysates of cells treated with the BB homodimeric form of PDGF but not in lysates of unstimulated cells. As observed for the EGF-R, the PLC_vl SH2[C] domain alone was inefficient in binding activated PDGF-R, but bound synergistically with the SH2[N] domain



when both domains were expressed as one bacterial protein (Fig. 2).

Within the SH2 domain, there are motifs that are articularly highly conserved. For example the NH2terminal tryptophan is invariant, and most SH2 domains start with the consensus W(Y,F)(H,F)GK (15, 21). A likely possibility is that these residues have been conserved because they are important in the interactions of SH2containing proteins with activated growth factor receptors. We therefore expressed a TrpE fusion protein that contained both PLC_vl SH2 domains, with the exception that the first four residues of SH2[N] (W-F-H-G) were deleted (PLC \triangle SH2-SH2-3). This fusion protein showed a modest ability to bind activated EGF- or PDGF-R (Fig. 2, lanes 5 and 10) that was equivalent to the SH2[C] domain alone, indicating that the removal of the four residues weakened binding activity.

Because GAP also associates with the PDGF-R, we undertook similar experiments using bacterial GAP SH2 sequences (see Fig. 1). The GAP SH2[N] domain bound the PDGF-R in a lysate of PDGF-stimulated cells (Fig. 3), but not in unstimulated cells (22). The GAP SH2[C] domain exhibited much weaker PDGF-R-binding activity. However, the two SH2 domains together (GAP-SH2[N + 3 + C]) bound the receptor threefold more efficiently than expected from their individual binding activities (Fig. 3, lanes 4 to 6 and 13 to 15). GAP contains an SH3 domain, which intervenes between the two SH2 elements and might contribute to binding to receptors. This seems unlikely, because the PLC_vl SH3 domain. expressed in isolation as a TrpE fusion protein, did not associate with the PDGF-R (Fig. 3).

Only a minor fraction of activated PDGF-R complexes with $PLC_{\gamma}l$ in vivo. We genetically modified a Rat-2 cell line

Fig. 1. Locations of SH2 and SH3 domains. The solid bars beneath the proteins indicate the regions expressed as bacterial TrpE fusion proteins for in vitro binding experiments (*32*). Abbreviations: 3, SH3 domain; PLC, catalytic regions of PLC₇I; GA, GTPase activating region of GAP; kinase, Srcttyrosine knGc domain.

to overexpress $PLC_{\gamma}l$ by tenfold as compared with the endogenous enzyme (Rat-2 PLC_{γ}l) (23). There is a proportionate increase in the amount of PDGF-R precipitated with antibodies to PLC_v1 (anti-PLC_yl) after PDGF stimulation of Rat-2 PLC_v1 cells, in comparison with parental Rat-2 cells (23). If bacterial PLC_{γ}1 SH2 domains bound to the same site(s) on the PDGFas did cellular $PLC_{\gamma}l$, R then overexpression of PLC_y1 should block binding of bacterial PLC_vl SH2 domains to activated PDGF-R in vitro. Consistent with this prediction, when the Rat-2 PLC_vl cell line was stimulated with PDGF, lysed, and incubated with immobilized PLC_yl-SH2[N] or PLC_yl SH2[N + C], only one-third as much PDGF-R associated with the bacterial protein, compared with the parental PDGF-stimulated Rat-2 cells (Fig. 4). Binding of TrpE-GAP-SH2 fusion protein to the PDGF-R was also reduced by overexpression of endogenous PLC_vl, suggesting that $PLC_{\gamma}1$ and GAP compete for sites on the activated PDGF-R.

Src-like tyrosine kinases and v-Crk also contain SH2 domains, which may bind activated receptors. Consistent with this prediction, bacterial fusion proteins that contained the SH2 domains of p60^{src} or P47^{gag-crk} bound PDGF-R in lysates of PDGF-stimulated Rat-2 cells (Fig. 3). p60^{src} is a substrate for the PDGF-R (24), and recent evidence suggests that Src-like kinases are physically associated with activated PDGF-R in vivo (25). Our data imply that this interaction involves the Src SH2 domain. Whether the normal homolog of v-Crk complexes with growth factor receptors in vivo remains to be established.

These results indicate that individual PLC_{γ}l or GAP SH2 domains, isolated as bacterial fusion proteins, can form stable complexes with activated growth factor receptors. The 100 amino acids of the PLC_{γ}l and GAP SH2 domains therefore

contain sufficient structural information to form independent, high affinity binding sites for receptors. The ability of SH2 domains to form such complexes when fused to a heterologous TrpE polypeptide suggests that SH2 domains may act in vivo to physically couple enzymatically diverse signaling proteins such as PLC_{γ} and GAP to growth factor receptors. Sequences that are structurally and functionally similar to the catalytic domains of $PLC_{\gamma}l$ and GAP have been identified in proteins that do not have SH2 domains. Several distinct isofoms of PLC have been isolated, including PLC_{β} and PLC_{δ} which share enzymatic sequences with PLC_{γ} , but lack the central PLC_{γ} SH2 and SH3 domains (26). Only the PLC_{ν}l isofom has been shown to complex with growth factor receptors (4) or to stimulate PI turnover in vivo in response to PDGF, and, indeed, deletion of the entire PLC_yl Src homology region abolishes the in vivo association of PLC_vl with the PDGF-R (27). Similarly, the IRA gene products. which stimulate Ras GTPase activity in

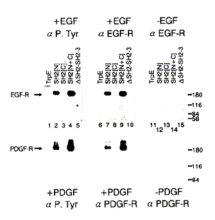


Fig. 2. SH2 domains of PLC_vl synthesized in bacteria bind synergistically in vitro to activated EGF- and PDGF-receptors. (A) Immobilized parental TrpE or the indicated TrpE-PLC,1 bacterial fusion proteins were incubated with lysates of Rat-1 cells that overexpressed the human EGF-R (RlhER), which had been serumstarved for 48 hours (lanes 11 to 15) or stimulated for 5 min at 37°C with 80 nM EGF (lanes 1 to 10). Complexes were washed, resolved on 8.25% SDSpolvacrvlamide gels. and analyzed hv immunoblotting with either $anti(\alpha)$ -P.Tvr (lanes 1) to 5) or anti-EGF-R (lanes 6 to 15) followed by I^{125} -labeled protein A. Autoradiography was for 18 hours. (B) Immobilized TrpE or TrpE- PLC_xI fusion proteins, as in (A), were incubated with lysates from Rat-2 cells that were serum-starved for 48 hours (lanes 11 to 15) or stimulated for 5 min at 37°C with 75 nM BB-PDGF (lanes 1 to 10). Samples were resolved on 6% SDSpolyacrylamide gels and analyzed by immunoblotting with either anti-P.Tyr (lanes 1 to 5) or anti-PDGF-R (lanes 6 to 15).

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yeast, contain a region related to the COOH-terminus of mammalian GAP, but do not possess SH2 domains (28). The ability of bacterial SH2 domains to mimic in vitro the interactions of native PLC_{γ} or GAP with activated growth factor receptors provides direct evidence that these sequences are sufficient for receptor-binding. These data strongly suggest that during the evolution of cellular signaling mechanisms, the acquisition of SH2 domains conferred On PLC_{γ} and GAP the capacity to interact with transmembrane tyrosine

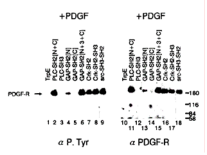


Fig. 3. Binding of TrpE fusion proteins that contain the GAP, Src, or Crk SH2 domains to PDGF-R in lysates of PDGF-stimulated Rat-2 cells. Serumstarved Rat-2 cells were stimulated for 5 min at 37°C with 75 nM BB-PDGF, lysed, and mixed with the indicated immobilized TrpE bacterial fusion proteins, Complexes were washed, resolved on 7.5% SDS-polyacrylamide gels and analyzed by immunoblotting with anti-P.Tyr (8 hour exposure; lanes 1 to 9) or with anti-PDGF-R (18 hour exposure; lanes 10 to 18).

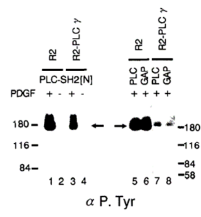


Fig. 4. Inhibition of in vitro binding of both PLC₄I and GAP SH2 domains to the activated PDGF-R in Rat-2 cells that overexpress pLC4, Rat-2 cells (lanes 1, 2, 5, and 6) or a Rat-2 cell line that overexpressed PLC₄I by tenfold (R2- PLC₅ lanes 3, 4, 7, 8) were stimulated with PDGF (lanes 1, 3, and 5-8) or maintained without PDGF (lanes 2 and 4). Cell lysates were mixed with immobilized TrpE-PLC-SH2[N] (lanes 1 to 4), TrpE-PLC-SH2[N + C] (lanes 5 and 7), or TrpE-GAP-SH2[N + 3 + C) (lanes 6 to 8). Samples were washed, separated by gel electrophoresis, and immunoblotted with anti-PTyr. Similar results were obtained by blotting with anti-PDGF-R.

kinases, and thereby to couple growth factor stimulation to PI turnover and the Ras pathway.

The high affinity association of PLC_vl and GAP with the EGF-R or PDGF-R is dependent on prior growth factor stimulation, and requires receptor tyrosine activity (5, 6). A simple explanation might be that receptor autophosphorylation elicits high affinity SH2 binding (29). The SH domains of GAP, Src, Abl, and Crk bind several proteins other than growth factor receptors (16, 29, 30), such as GAPassociated p62 (18, 29), whose common feature is tyrosine phosphorylation. In addition, autophosphorylation is required for efficient complex formation in vivo between the PDGF-R and GAP (6). It is feasible that tyrosine phosphorylation of residues in SH2-binding sites increases the affinity for SH2 domains. We note the presence of invariant positively charge d residues in SH2, which might contribute to such an interaction (15). Alternatively, or in addition, growth factor stimulation and subsequent receptor autophosphorylation induce a conformational change that forms an SH2-binding site. The affinity of PLC_yl and GAP for activated receptors was increased synergistically by the juxtaposition of two SH2 domains, There are several potential explanations for this synergistic effect. Growth factorbinding induces receptor dimerization, and the elevated binding activity of two linked SH2 domains may result from a cooperative interaction with the activated. dimerized receptor. Alternatively, a receptor monomer might have multiple SH2-binding sites, or the combined SH2 domains might bind more strongly to a single site. The ability of overexpressed $PLC_{\nu}1$ to inhibit the in vitro binding of both PLC_vl and GAP SH2 domains to PDGF-R suggests that these bind to similar sites. The EGF-R P phosphorylates $PLC_{\gamma}l$ at two tyrosines in vivo, one of which is adjacent to SH2[C] (31). Tyrosine phosphorylation of PLC_yl may decrease the affinity of SH2binding, as the phosphorylated enzyme is apparently released from the receptor (5).

In summary, SH2 domains mediate the high affinity interactions of $PLC_{\gamma}I$ and GAP with activated growth factor receptors. The use of a specialized noncatalytic domain to direct complex formation between protein kinases and their presumptive targets is

unprecedented. It is possible that a function of tyrosine phosphorylation is to regulate heteromeric protein-protein interactions.

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- 19 Restriction sites were introduced on either side of SH2 coding sequences in the cDNA's for bovine PLC_vl and human GAP with oligonucleotide-directed mutagenesis [T. A. Kunkel, J. D. Roberts, R. A. Zakour, Methods Enzvmol. 154,367 (1987)]. For each individual SH2 domain an Sph I site was created at the 5' end and an Nhe I site at the 3' end. These Sph I-Nhe I fragments were cloned into a pATH bacterial trpE expression vector whose multiple cloning site had been modified to contain unique Sph I and Nhc I sites. For fusions that contained both SH2 domains, the Sph I site of the NH2-terminal End of page in published document 981

SH2 domain and the Nhe I site of the COOH-

terminal SH2 domain were used for the excision. Src and Crk fusion proteins utilized natural restriction sites. The resulting fusion proteins contained the NH₂-terminal 323 amino acids of TrpE and retained the desired reading frame for PLC₇I or GAP.

20. Cultures of E. coli RRI with pATH expression plasmids were grown, induced, and lysed as described (29). The TrpE fusion proteins were recovered from the supernatants by immunoprecipitation with polyclonal anti-TrpE antiserum immobilized on protein A-Sepharose beads. Immune complexes were washed (29), aliquoted, flash-frozen, and stored at -70°C until mixed with mammalian cell lysates. Starved or growth factorstimulated rat fibroblasts (-5 x 106) were lysed in 2 ml of lysis buffer (50 mM Hepes, pH 7.0, 150 mM NaCI, 10% glycerol, 1% Triton X-100, 1.5 mM MgCI,, 1 mM EGTA, 100 mM NaF, 10 mM sodium pyrophosphate, 1 mM Na3V0,, 1 mM PMSF, 10 ~ g l m l aprotinin, 10 µg/ml leupeptin). Clarified mammalian cell lysate (1 ml) was mixed with immobilized bacterial fusion protein by gentle inversion for 90 min at 4°C. Complexes were recovered by centrifugation, washed three times with HNTG buffer (20 mM Hepes pH 7.0, 150 mM NaCI, 0.1% Triton X-100, 10% glycerol, 1 mM Na3V0,), and analyzed by immunoblotting with anti-P.Tyr or anti-receptor as described (6, 15, 18, 29). To ensure that the different TrpE fusion proteins were present in similar amounts in the immune complexes incubated with the

mammalian cell lysates, duplicate samples for anti-P.Tyr and anti-EGF-R immunoblotting were probed with an anti-TrpE monoclonal antibody. Equivalent amounts of the various TrpE fusion proteins were detected.

- Abbreviations for the amino acid residues are: A, Ala; C, Cys; D, Asp; E, Glu; F, Phe; G, Gly; H, His; I, Ile; K, Lys; L, Leu; M, Met; N, Asn; P, Pro; Q, Gln; R, Arg; S, Ser; T, Thr; V, Val; W, Trp; and Y, Tyr.
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- 32. Bacterial fusion proteins contained the following amino acid residues: Src-SH3-SH2, p60^{1-src} 87 to 291; PLC-SH2[N], bovine PLC₁] 547 to 659; PLC-SH2[C], PLC₁] 663 to 752; PLC-SH2[N + C], PLC₂I, 547 to 752; PLC ASH2- SH2-3, PLC₂I, 555 to 951; PLC-SH3, PLC₂I, 781 to 855; GAP-SH2[N], human GAP 178 to 277; GAP-SH2[C], GAP 348 to 444; GAPSH2[N + 3 + C], GAP 178 to 444; Crk-SH2, p47^{gag-crk} 205 to 386; Crk-SH2-SH3, p47^{gag-crk} 220 to 427.
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