Functional proteomic analysis of long-term growth factor stimulation and receptor tyrosine kinase co-activation in Swiss 3T3 fibroblasts

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Running title: Growth factor-specific regulation of protein synthesis

Key words: RTK signaling, EGF, IGF-1, PDGF, protein synthesis regulation

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Abbreviations
GF, growth factor; GFR, growth factor receptor; RTK, receptor tyrosine kinase; EGF, epidermal growth factor; IGF-1, insulin-like growth factor 1; PDGF, platelet-derived growth factor; NGF, nerve growth factor; TGF, transforming growth factor; PTM, post-translational modification; PI3K, phosphoinositide 3-kinase; AKT, serine/threonine kinase PKB; mTOR, mammalian target of rapamycin; eEF2, eukaryotic translation elongation factor 2; NAP1L1, nucleosome assembly protein-1 like-1; PRS7, 26S protease regulatory subunit 7; RACK-1, receptor for activated C kinase-1; Hsp, heat shock protein; PLC, phospholipase C; Gab, growth factor receptor-binding protein-associated binder; FRS2, fibroblast growth factor receptor substrate 2; SHP2, tyrosine-specific protein phosphatase; GAP, GTPase-activating protein; MAPKs, mitogen-activated protein kinases; Erk1/2, p38 and MEK, different members of the MAPK-family of serine/threonine protein kinases; PKC, protein kinase C; STATs, signal transducers and activators of transcription; FAK, focal adhesion kinase; MLCK, myosin light chain kinase; ROCK, Rho-associated protein kinase 1; TCP-1/CCT, T-complex protein 1; Lasp-1, LIM and SH3 domain protein 1; Sid23, actin-depolymerizing factor Destrin; CHX, cycloheximide; TSA, trichostatin A; 5aC, 5-azacitidine; SH2, Src homology 2 domain; PTB, phosphotyrosine-binding domain.
Abstract

In Swiss3T3 fibroblasts, long-term stimulation with PDGF, but not IGF-1 or EGF, results in the establishment of an elongated migratory phenotype, characterized by the formation of retractile dendritic protrusions and absence of actin stress fibers and focal adhesion complexes. To identify receptor tyrosine kinase (RTK)-specific reorganization of the Swiss 3T3 proteome during phenotypic differentiation, we compared changes in the pattern of protein synthesis and phosphorylation during long-term exposure to PDGF, IGF-1, EGF and their combinations using 2DE-based proteomics after $^{35}$S- and $^{33}$P-metabolic labeling. One hundred and five differentially regulated proteins were identified by mass spectrometry and some of these extensively validated. PDGF stimulation produced the highest overall rate of protein synthesis at any given time and induced the most sustained phospho-signaling. Simultaneous activation with two or three of the growth factors revealed both synergistic and antagonistic effects on protein synthesis and expression levels with PDGF showing dominance over both IGF-1 and EGF in generating distinct proteome compositions. Using signaling pathway inhibitors, PI3K was identified as an early site for signal diversification, with sustained activity of the PI3K/AKT pathway critical for regulating late protein synthesis and phosphorylation of target proteins and required for maintaining the PDGF-dependent motile phenotype. Several proteins were identified with novel PI3K/Akt-dependent synthesis and phosphorylations including eEF2, PRS7, RACK-1, acidic calponin, NAP1L1, Hsp73 and fascin. The data also reveal induction/suppression of key F-actin and actomyosin regulators and chaperonins that enable PDGFR to direct the assembly of a motile cytoskeleton, despite simultaneous antagonistic signaling activities. Together, the study demonstrates that long-term exposure to different growth factors results in RTK-specific regulation of relatively small sub-proteomes, and implies that the strength and longevity of RTK-specific signals are critical in defining the composition and functional activity of the resulting proteome.
Introduction

Receptor tyrosine kinases transduce key extracellular signals and trigger multiple cellular events, including proliferation, differentiation, and cytoskeletal rearrangement. A variety of intracellular signaling molecules associate with the phosphorylated tyrosine residues on the cytoplasmic tail of activated RTKs via their SH2 or PTB domains. Mutation of these docking sites results in severely disturbed cellular phenotypes in different model systems, demonstrating the physiological importance of RTK signaling (1). As every RTK recruits and activates a different combination of signaling proteins via their own tyrosine autophosphorylation sites and sites on closely associated docking proteins (e.g., Gab1, FRS2) (2, 3) it has been considered for a long time that this combinatorial recruitment determines signal specificity.

However, mutant platelet-derived growth factor receptor (PDGFR) lacking binding sites for activation of the PLCγ, PI3K, SHP2 and RasGAP pathways still retain partial ability to induce 64 out of 66 immediate early genes (IEG) induced by the ‘wild-type’ receptor (4). The authors also observed induction of almost identical sets of IEGs in response to PDGFRβ and fibroblast growth factor receptor 1 activation in murine fibroblasts, suggesting that different RTK-activated pathways exert broadly overlapping effects on IEG induction. More recently, three groups have demonstrated that specific expression of marker genes for RTK-dependent cell fate decision in Drosophila does not appear to arise from RTK-specific intracellular signaling, but does so through a combination of RTK and non-RTK-signaling that feed into pre-existing cell-specific transcription factors which are induced during earlier developmental stages (5-7). Thus, distinct responses to RTKs in different cells can be explained by differences in the cells’ developmental histories. These results also suggest that RTKs utilize general signaling pathways or networks with some redundancy, and that the specificity provided by combinatorial recruitment of particular signaling proteins is not that stringent. However, these studies only deal with transcriptional responses to RTK activation, and gene expression changes alone are unlikely to determine biological outcome. For this reason, global analyses of protein synthesis, expression level, activity status and degradation rates are necessary to understand how signals from different RTKs are diversified to provide specific biological outcomes.

The early signaling events associated with RTK activation have been intensively studied and a detailed map of the integrated signaling circuits is emerging. However, the fact that commitment to specific cellular responses such as cell cycle progression, seem to require the continuous exposure to growth factor (GF) for several hours, suggests that there must be essential signaling events occurring throughout this time period (8, 9). The importance of this
sustained versus transient signaling in defining biological outcome is illustrated by the finding that NGF triggers sustained MAPK activation and differentiation of PC12 cells, whereas EGF triggers transient MAPK activation and cell proliferation (10). Moreover, EGF receptor overexpression in these cells leads to sustained MAPK activation, resulting in their differentiation, further implicating that biological outcome is dependent upon quantitative modulations of the strength and duration of the inducing signals. Cross-talk between pathways may also act synergistically to enhance an internal pathway, as shown for MAPK activation in response to attachment of cells to the extracellular matrix during GF treatment (11). Finally, pathways may also antagonize one another as found for interferon-γ, which inhibits TGF-β signaling by inducing the expression of the inhibitory Smad7 protein (12).

Changes in fibroblast morphology in response to short-term growth factor exposure are well characterized. However, these changes are normally transient responses occurring during short-term stimulation, and cytoskeletal changes triggered by long-term stimulation of RTKs have not been well studied. For example, while short-term PDGF treatment induces membrane ruffling and lamellipodial extensions in fibroblasts through activation of the small GTPases Cdc42 and Rac (13, 14), the establishment of a motile cytoskeleton with retractile dendritic protrusions requires exposure to PDGF for 12 hours (15).

Here, a 2DE-based proteomic analysis was undertaken to identify GF-specific proteome modifications in Swiss 3T3 cells that define distinct phenotypic outcome, and to investigate the effects of receptor co-activation. As our main objective was to describe the changes in the fibroblasts proteome that underlie the establishment of an elongated, migratory phenotype during long-term PDGF-stimulation, PDGF was used at a concentration of 50 ng/ml, which according to earlier reports should ensure optimal rates of proliferation and cytoskeletal reorganization (16-18). The concentration of EGF and IGF-1 were then adjusted to instigate a similar mitogenic effect in the murine fibroblasts. At the used concentration of 100 ng/ml EGF and IGF-1 both promoted the establishment of a well-anchored, round, immotile phenotype. Long-term treatment with higher concentrations of one or both of these growth factors did not change the phenotypic outcome.

The rate of protein synthesis was monitored by biosynthetic labeling with [³⁵S]-methionine and -cysteine, changes in protein abundance were monitored by densitometry analysis of fluorescent dye-stained gels and immunoblotting, RTK-induced phospho-signaling was examined by a combination of [³³P]-orthophosphate labeling, phosphatase treatment and immunoblotting and GF-treatment in the presence of kinase inhibitors was used to investigate
the signaling pathways/networks that control RTK-specific proteome regulation. The study finally investigates the specific, dominant changes in proteome activities that enable PDGF to control the organization of contractile actomyosin structures during the assembly of a migratory cytoskeleton, despite simultaneous antagonistic signals from other RTKs.
Materials and Methods

Cell culture, metabolic labeling, and preparation of whole cell lysates

Swiss 3T3 cells were maintained in Dulbecco’s modified Eagle’s medium (DMEM) (GIBCO) supplemented with 10% fetal bovine serum (FBS), 100 U/mL of penicillin, and 100 μg/mL of streptomycin in 5% CO₂ at 37 °C. For biosynthetic labeling, sub-confluent cells were serum-starved in DMEM supplemented with 0.1% FBS for 24 h, and subsequently stimulated with PDGF (50 ng/mL), IGF-1 (100 ng/mL), EGF (100 ng/mL) (PeproTech) or their combinations for 6, 12, and 18 hr prior to lysis. Cells were labeled with [³⁵S]-Met/Cys (0.2 mCi/mL of Promix; Amersham Pharmacia) during the last 3 hr of each stimulation condition. In some experiments, cells were cultured in the presence of various inhibitors (10 μM LY294002, 50 μM PD98059, 20 nM PD184352, 50 nM rapamycin, 2 μM SU6656, 1 μM ML-7, 3 μM Y-27632, 1 μM PS-341, 50 μM lactacystin, 10 ng/ml actinomycin D, and 10 μg/ml cycloheximide), during the entire time-course, whilst in others, cells were pre-treated with inhibitors for 30 min prior to metabolic labeling for 3 hr. For [³²P]-orthophosphate labeling, synchronized cells grown to 70% confluence were washed twice in phosphate-free DMEM containing 0.1% FBS, incubated for one hour in the same media, and labeled with 0.1 mCi/mL [³²P]-orthophosphate for 6 hr prior to short-term GF stimulations. In long-term stimulation experiments (6-18 hr), the cells were either incubated with [³²P]-orthophosphate during the entire stimulation period, or the radioisotope was added to the culture medium after 6 or 12 hr.

Cells were routinely lysed in a solution containing 8 M urea, 2 M thiourea, 4% CHAPS, 65 mM dithiothreitol (DTT), protease inhibitor cocktail with EDTA (Boehringer Mannheim) and phosphatase inhibitors (1 μM okadaic acid, 5 μM α-cyano-3-phenoxybenzyl α-(4-chlorophenyl) isovalerate, and 5 μM potassium bisperoxo (1,10-phenanthroline) oxovanadate). For carrier ampholyte-based isoelectric focusing (IEF), cells were lysed in 2% NP-40, 9.6 M urea, 100 mM DTT, 2% ampholines (pH 3-10) plus the above inhibitors. Protein concentrations were determined by Bradford assay (Coomassie Plus Protein Assay Reagent; Pierce). For alkaline phosphatase (AP) treatment, cells were lysed in a solution containing 50 mM Tris-HCl (pH 8.5), 1 M urea, 2 M thiourea, 4% CHAPS, 65 mM DTT and 2% ampholines. The lysates were incubated at 37°C for 30 min with or without AP (7.5 U). Artifactual sample modification by carbamylation was minimal.

Two-dimensional gel electrophoresis (2-DE)

Isoelectric focusing with carrier ampholines was performed as previously described (19). Isoelectric focusing with in-gel reswelling of immobilized pH gradient strips (Immobiline DryStrip pH 3-10NL 18 cm, pH 6-9NL 18 cm (GE
Healthcare), or ReadyStrip IPG strips pH 3-6, 17 cm (Biorad)) was performed as described elsewhere (20). Second dimension electrophoresis was routinely performed in 9-16% gradient SDS-PAGE gels. Proteins were silver stained according to Shevchenko (21), or stained with the fluorescent dye OGT MP17 according to the manufacturer’s protocol (22). Radio-labeled proteins were electro-blotted onto nitrocellulose or PVDF membranes, and scanned by phosphoimaging on a Molecular Imager FX scanner (Biorad). The membranes were subsequently stained by Coomassie or used for immunoblotting. Immunoblots were visualized by ECL (Amersham Pharmacia). Experiments where secondary antibodies were used alone gave no signals. All experiments were repeated with samples from separately stimulated cell cultures.

**Image analysis** Gel image analysis was performed using the Melanie 3 software (Genebio). Quantitation and comparison of radio-labeled protein spot volumes and intensities were performed using a combination of Melanie 3 and the in-house written program ChiMap. The individual spot density values were calculated as the relative difference between the spot density at the given time point, and the spot density on reference gel images, and the final value given is an average of at least three independent experiments. In some cases, PDGF-stimulated gel images (18 hr) were used as reference, as we were unable to detect all the differentially regulated proteins on 35S-gel images from unstimulated cells. The in-house programme ChiClust (23) was used to identify patterns of protein spots that were similar or different between gels. Variations between spot density and/or volume on the different images are incorporated into a generic % homology value, which can be used to generate a dendrogram or relational tree of homology, as illustrated in Figure 2. All computer alignments and target selections based on densitometry values were visually crosschecked by at least two operators. Proteins for MS analysis were excised from stained gels either manually or automatically using a robotic cutter (Oxford GlycoSciences).

**In-gel digestion** Tryptic in-gel digests were essentially prepared as described earlier (24). In brief, 2D-gel pieces were destained and rehydrated with 25 mM ammonium bicarbonate (AmBic)/50% acetonitrile (ACN), and vacuum-dried in a SpeedVac centrifuge. Then, gel pieces were rehydrated with 10 mM DTT/25 mM AmBic solution and incubated for 1 hr at 50°C, followed by removal of the DTT solution and the addition of an equivalent volume of 50 mM iodoacetic acid /25 mM AmBic solution. After incubation in the dark for 1 hour at room temperature the samples were washed with 25 mM AmBic/50% ACN and vacuum-dried in a SpeedVac centrifuge. The samples were then rehydrated and digested overnight with 30 ng modified porcine trypsin (Promega) at 37°C. Peptides were extracted three times with 50% ACN/5% trifluoroacetic acid solution, dried in a SpeedVac and resuspended in 5 μL of water.
Protein identification by mass spectrometry For matrix-assisted laser desorption/ionization (MALDI) MS analysis, the digested peptide mixture was spotted onto the target plate together with a saturated aqueous solution of 2,5-dihydroxybenzoic acid (DHB). All samples were analysed on a Reflex III reflector time-of-flight (TOF) mass spectrometer (Bruker-Daltonik, Bremen, Germany) equipped with delayed extraction and a Scout 384 ion source similar to a previously reported study (25). Briefly, the instrument was externally calibrated using calibration mixture 2 from the Sequazyme kit (Applied Biosystems). Spectra were obtained by averaging 32-128 single-shot acquisitions and internal calibration was carried out using the trypsin autolysis peaks at m/z 842.51 and m/z 2211.10. Peak masses were assigned by labeling the first monoisotope. The peak mass lists were submitted to ProteinProspector’s database search routine MS-Fit for peptide mass mapping (http://prospector.ucsf.edu/prospector/mshome.htm), searching the NCBInr database version 20010107 (601,500 sequences) or 20010303 (640,428 sequences) with trypsin as the enzyme, a ±100 ppm mass tolerance, max. 2 missed cleavages and carboxymethylation of cysteine as a constant modification. Acetylation at the protein’s N-terminus, methionine oxidation, and pyroglutamic acid for N-terminal glutamine were used as variable modifications. Positive identifications were accepted if there were at least 5 peptides matched to a protein entry, there was a 20% minimum coverage of the protein sequence, the molecular weight was within ±25% of the gel-curated molecular weight and the pI was within ±10% of the gel-curated pI value.

When peptide mass mapping by MALDI MS was unsuccessful or ambiguous, LC-electrospray ionization (ESI) MS/MS analysis was performed. Briefly, the remainder of the digested peptide mixture was run on a micro- or nano-capillary HPLC. After loading the sample, peptide separation was performed on a fused silica C18 column (LC Packings, Amsterdam, Netherlands). The HPLC column was coupled to a 50 μm I.D. fused silica capillary via a zero dead volume connector, which was coupled to the standard nano-LC interface of a Q-Tof hybrid quadrupole orthogonal time-of-flight mass spectrometer (Micromass, Wythenshawe, UK). Mobile phase A was 0.1% formic acid and mobile phase B was 80% ACN/0.08% formic acid. Gradient elution was carried out from 5% B to 42% B in 90 min. Mass spectra were acquired using the MassLynx 3.3 software package with automatic precursor ion selection. Fragment ion mass lists were submitted to the database search engine Mascot 1.8 (Matrix Science, London, UK) to search the same NCBI databases described above with trypsin as the enzyme, a ±100 ppm parent ion mass tolerance, ±100 mmu fragment ion mass tolerance, max. 1 or 2 missed cleavages and the modifications described above.
Significant hits were defined when two or more peptide sequences matched a protein entry and the MASCOT score was above the significance level ($p<0.05$).

**Immunofluorescence imaging** Immunofluorescence staining was performed as described previously (15). Cells were grown on coverslips, stimulated with GFs as described above for 18 hr, and stained with anti-vinculin (Sigma), anti-RACK-1 (Transduction Laboratory), or rhodamine-phalloidin. Specimens were visualized with an LSM 510 confocal microscope using a x40 NA 1.3 Neofluar objective (Zeiss, UK) or with a Coolview 12 integrating cooled CCD camera (Photonic Science, East Sussex, U.K.) mounted over an Axiophot microscope fitted with a x63 NA 1.4 oil immersion objective (Zeiss UK). No staining was observed when secondary antibodies were applied alone. For time-lapse analysis, cells were grown on 35 mm tissue culture dishes and image data sets collected once every 4 minutes for a period of 22 hours with a KP-M1 monochrome video camera (Hitachi Denshi, London, U.K.) mounted on the side port of an Axiovert 135 microscope (Zeiss UK) and using Tempus image acquisition software (Kinetic Imaging, Merseyside, UK).
Results

**Biologic effects of growth factor stimulation** Treatment with recombinant PDGF (50 ng/mL), IGF-1 (100 ng/mL) and EGF (100 ng/mL) induced significant and roughly equivalent mitogenic activity in the murine fibroblasts as detected by \[^{3}H\]-thymidine and BrdU-incorporation, while the induction of specific receptor phosphorylation and degradation patterns (for EGFR and PDGFR) confirmed ligand-specific receptor activation (Supplementary Data; Figure S1). PDGF stimulation induced a unique morphological phenotype, which was characterized by an elongated cell shape and dendritic protrusions, and absence of actin stress fibers and focal adhesions (Figure 1A). The disruption of stress fibers and focal adhesions occurred within the first hour of PDGF stimulation, while full development of dendritic protrusions appeared after 12 hours and was sustained throughout the rest of the stimulation period (Figure 1B). Furthermore, a significant enhancement in cell motility on tissue culture dishes was observed by time-lapse microscopy following 12 hours of PDGF stimulation, and characterized by continuous extension and retraction of dendritic protrusions and changes in cell shape (data not shown). In contrast, stimulation with EGF (or IGF-1) only slightly increased cell motility over the same time period (15). Short-term stimulation with PDGF (<1 hour) has previously been reported to induce membrane ruffling and lamellipodial extensions (13), but not dendritic protrusions, which seem to be unique features of the long-term growth factor stimulation employed in this study.

Conversely, long-term EGF or IGF-1 stimulation resulted in a moderate increase in stress fiber and focal adhesion formation, compared to unstimulated cells. Actin stress fibers and focal adhesion complexes were abundant in 20% of unstimulated cells, in 26% of IGF-1-stimulated cells and in 28% of EGF-stimulated cells (Figure 1C, top). The large, flat immobile phenotype induced by EGF and IGF-1 had almost twice the cell surface area compared to the PDGF-induced phenotype (Figure 1C, bottom). Interestingly, the EGF or IGF-1-induced morphological changes were greatly influenced by cell confluence, while the PDGF-induced changes were less affected, suggesting that PDGFR signaling may override signaling induced by cell-cell contact in Swiss 3T3 cells. Co-stimulation experiments demonstrated that PDGF had a dominant influence on morphology over EGF and IGF-1, resulting in the formation of an elongated cell shape with dendritic protrusions and suppression of stress-fibers and focal adhesions (Figures 1A and C). Taken together, these data show that long-term stimulation with PDGF inhibits formation of stress fibers and focal adhesions in Swiss 3T3 cells, and promotes formation of dendritic protrusions and motility, even during simultaneous stimulation with either or both EGF and IGF-1.
The PDGF-specific morphological transformation was abrogated in fibroblasts where DNA-dependent RNA-synthesis was blocked by treatment with actinomycin D, protein translation was blocked by cycloheximide treatment, the 26S proteasome was inhibited by PS341 treatment, where the activity of Src kinases was inhibited by SU6656 treatment, and where methyltransferase and deacetylase activities were blocked by 5aC and TSA treatments, respectively (Supplementary Data; Figure S2). These findings demonstrate that the PDGF-induced reorganization of the actin cytoskeleton in Swiss 3T3 fibroblasts involves the complex regulation of gene transcription, protein synthesis, post-translational modification and degradation.

**Analysis of growth factor-regulated protein synthesis in Swiss 3T3 fibroblasts** The rate of protein synthesis was next monitored by biosynthetic labeling with $[^{35}\text{S}]$-methionine and cysteine during the last 3 hours of GF stimulation, employing 6, 12, and 18 hour stimulation periods. Radio-labeled proteins were separated by 2-DE, electro-transferred to PVDF membranes and scanned by phospho imaging (Figure 2A). As a control, protein synthesis was abolished by addition of cycloheximide. The digitized images were subjected to computer analysis to define alterations in the *de novo* synthesis of proteins.

The three growth factors induced $[^{35}\text{S}]$-incorporation into several hundred protein spots within the first 6 hours of treatment, in addition to the 631 spots which were radiolabeled in untreated cells (Figures 2A and 2B). This is assumed to represent *de novo* protein synthesis. While the majority of the detected protein spots showed similar regulation over time, 21% of the matched spots were differentially regulated by the three growth factors. PDGF stimulation induced the highest overall rate of protein synthesis at any given time, followed by IGF-1 and then EGF, as judged by the total quantity of radioisotope incorporation detected by autoradiography of gels (Figure 2A). Computer-aided global densitometry analysis showed that the number of protein spots with up-regulated synthesis reached a maximum at the 6 hour time point and that the three GFs induced almost identical numbers of new proteins by this time (Figure 2B). The more than 1,250 different spots resolved, represents a doubling of the number of isotope-labeled protein species detected in extracts from unstimulated cells (Figures 2A and B). Whilst the number of newly synthesized protein spots stayed rather constant during continuous PDGF and IGF-1 stimulations, it declined over time during EGF stimulation, resulting in the greatest qualitative difference after 18 hours (Figure 2B); by 18 hours, PDGF had increased or reduced protein synthesis (by $>2$-fold) in a higher number of matched features compared to EGF or IGF-1 (Figures 2C and D).
Comparison of protein expression levels revealed that 18 hours treatment with PDGF and EGF induced the most distinct proteome compositions (Supplementary Data; Figure S3), which were presumably sustained by the most distinct patterns of de novo protein synthesis; 18 hours stimulation with PDGF more than doubled the synthesis rate of 15.8% of matched proteins compared to EGF, which more than doubled the synthesis rates of 5.3% of proteins compared to PDGF (Figure 2C). The suppressive effects of PDGF treatment were likewise dominant; PDGF reduced the synthesis of 11.4% of matched proteins by greater than 2-fold compared to EGF, whereas EGF reduced the synthesis rates of 3.9% of matched proteins compared to PDGF (Figure 2D).

GF-specific temporal modulations and the effects of co-stimulation were reflected by hierarchical clustering analysis. For Figure 2E, the images were clustered using the density patterns of a group of 172 proteins, whose synthesis rates were found to be similarly regulated by the three GFs after 6 hours. In accordance with the gradual reduction in both the numbers of induced proteins and their synthesis rates seen in response to EGF treatment, the images from the EGF 12 and 18 hour treatments clustered in proximity to those from untreated cells (Figure 2E). Changes in protein synthesis induced by EGF were thus rapid and short-lived. The images obtained from cells treated with PDGF and IGF-1 for 18 hours clustered with images obtained from cells exposed to GF for shorter periods, indicating that the synthesis rates of the majority of the 172 reference proteins did not change significantly during long-term treatment with these two GFs. In the ChiClust analysis shown in Figure 2F, images were clustered according to similarities between all radio-labeled spots. It demonstrates the divergence over time in the patterns of newly synthesized proteins generated following PDGF treatment versus EGF or IGF-1 treatment (Figure 2F). Indeed, long-term exposure to PDGF elevated the synthesis of more than 800 proteins. Whilst the number of newly synthesized proteins also declined over time in IGF-1 treated cells, IGF-1 maintained increased de novo synthesis significantly longer than EGF. Notably, the progressive divergences in synthesis patterns between the different treatments coincided with the development of the two distinct morphological phenotypes (see above). The formation of a migratory cytoskeleton requires continuous exposure to PDGF for 12 hours (15), the time point at which PDGF treatment had produced the most unique pattern of de novo protein synthesis in 3T3 fibroblasts (Figure 1F).

Co-stimulation with two GFs resulted in both additive and synergistic increases in protein synthesis rates and the positive effects of receptor co-stimulations were generally more pronounced than the suppressive (Figure 2C and D). Notably, while the number of proteins with a more than 2-fold increase in $^{35}$S-incorporation was significantly elevated by any combination
of two GFs, co-stimulation with all three GFs had a less pronounced effect on protein synthesis and actually resulted in more proteins showing a >2-fold reduction in synthesis (Figures 2C and D). As a consequence, images from cells treated with any combination of two GFs for 18 hours clustered together, while long-term treatment with all 3 GFs resulted in a synthesis pattern more similar to those obtained following short-term treatment with individual GFs (Figures 2E and F). However, co-stimulation experiments also revealed that for some proteins, one GF could dominantly suppress an increased synthesis rate induced by another GF (see examples in Supplementary Data; Figures S4 and S5). Notably, PDGF induced the most pronounced regulatory effects in co-stimulation experiments.

**Differentially regulated proteins identified by mass spectrometric analysis** Proteins, which were dominantly induced or suppressed by a specific growth factor or combination of growth factors, and proteins that displayed differential timing of induction or suppression were targeted for identification by MS analysis, since they are likely to be of importance for GF-specific morphological differentiation. We were able to align approximately two-thirds of the differentially regulated proteins on either silver-stained or fluorescent dye-stained gels by computer alignment. MALDI-TOF MS and LC-ESI MS/MS were then used to identify 143 protein species from 107 differentially regulated protein spots with two or more proteins identified in 25 of the spots (Supplementary Data; Table S1). These represented 105 unique gene products, the majority of which belonged to the following functional groups; proteins involved in the translational machinery, chaperone function, protein degradation, vesicular transport, DNA/RNA synthesis and transport, metabolic pathways and cytoskeleton organization; the latter subset represented by more than twenty species. The synthesis rates of 33 of the identified proteins were predominantly controlled by PDGF. Examples of differentially regulated proteins are provided in Supplementary Data; Figure S5. Notably, some proteins demonstrated bi-phasic regulation over the observation period, suggesting that secondary signaling events such as autocrine loops may be initiated during long-term GF-stimulation.

**Analysis of growth factor induced phospho-signaling** In order to study the underlying GF-induced regulatory phospho-signaling pathways and whether the enhancing and suppressive effects receptor co-activation instigates on protein synthesis reflects changes in the RTK-induced signaling cascades, [³³P]-orthophosphate labeling was combined with immunoblotting, immunoprecipitation and kinase inhibitor and phosphatase treatment experiments.
Immunoblotting with phosphorylation site-specific antibodies showed that sustained stimulation with PDGF and IGF-1, but not EGF, induced phospho-activation of Akt (pS473) that lasted for at least 18 hours (Figure 3A and B). Compared with EGF and IGF-1, PDGF induced a longer lasting and dominant phospho-activation of ERK1 (pT203/pY205), ERK2 (pT183/pY185), PKCβII (pS660), PLCγ (Y783), STAT3 (Y705) and Src (pY424) (Figure 3A). Indeed, EGF only induced transient phosphorylation of ERK1/2 and PKCβII, whilst IGF-1 produced barely any detectable activation of these signalling proteins by comparison with the PDGF-induced signals. Phosphorylation of the inhibitory site of Src (Y535) was also transiently induced by PDGF similar to the Y424 activation site, and this was abrogated by pre-treatment with 2 μM of the c-Src inhibitor SU6656 (Figure 3D). Whilst both phosphorylation events may not be mutually exclusive, this data hints that distinct pools of activated and inactivated Src may exist in cells stimulated with PDGF. PDGF was also dominant over EGF in inducing transient phosphorylation of p38γ (MAP kinase 12) at its activation sites (T183/Y185) (Figure 3B). Notably, the PDGF-induced phosphorylation of p38γ and PKCβII was reduced in cells pre-treated with LY294002 (Figures 3B and C), indicating that PDGF at least in part activate p38γ and PKCβII via the PI3K/Akt pathway. Short-term PDGF treatment (15 min) also caused a transient reduction in the phosphorylation of FAK at Y428, and a transient increase in cofilin phosphorylation at S3, consistent with cofilin inactivation (Figure 3A). These changes are likely to facilitate actin remodeling and focal adhesion turnover, and may explain the PDGF-mediated reduction in stress fibers (SF) and focal adhesions (FA) observed in Swiss 3T3 fibroblasts within the first 30 min of treatment (15). Finally, PDGF induced the specific phosphorylation of c-Myc (T58/S62) at the 6 hour timepoint. These sites are known to regulate the stability and thereby the duration of c-Myc activity following mitogenic induction. Thus, the signal detected is likely to reflect a higher level of induced c-Myc protein in the PDGF stimulated cells, which is being directed for ubiquitin-mediated degradation following its phosphorylation (Figure 3A).

Both qualitative and quantitative differences were observed between the groups of phosphoproteins that co-immunoprecipitated with Src from [33P]-orthophosphate labeled 3T3 cells stimulated with EGF, IGF-1 and PDGF, and the pattern of co-isolated phosphoproteins showed temporal variations (Figure 3E). GF-stimulation in the presence of the Src-specific inhibitor SU6656 (2 μM) strongly reduced both the number and amount of co-precipitated phosphoproteins (Figure 3E), suggesting that the majority of the interacting proteins are Src kinase substrates. The results imply that the three growth factors not only generate different Src kinase activities, they also direct the kinase towards slightly different repertoires of substrates.
While the number of co-isolated phosphoproteins was reduced in immunoprecipitates from cells treated with EGF and IGF-1 for 4 hr compared to the numbers found at the 2 hr timepoint, several new phosphoproteins were co-precipitated from PDGF-stimulated cells after 4 hr (white oblique arrows in Figure 3E). This finding further supports the notion that PDGF-induces a longer-lasting activation of Src kinase than EGF and IGF-1.

Following both short and long-term GF-treatment, the highest levels of $^{33}$P-incorporation were found in cells stimulated or co-stimulated with PDGF and were distinct from those induced by long-term stimulation with EGF or IGF-1 (Supplementary Data; Figure S6). Notably, $^{33}$P-orthophosphate labeling experiments revealed that receptor co-stimulation influenced the specificity, strength and kinetics of RTK-mediated phospho-signaling (Figure 4), which might explain the enhancing and suppressive effects receptor co-activation instigates on de novo protein synthesis.

**The effect of kinase inhibitors on RTK-induced protein synthesis and phosphorylation** To further understand how these phosphorylation events are regulated and how they affect protein synthesis, cells were treated with kinase inhibitors targeting PI3K (LY294002), mTOR (Rapamycin), c-Src (SU6656), MEK (PD98059), ROCK (Y-27632) and MLCK (ML-7) prior to GF-stimulation and metabolic labeling with $^{33}$P-ortho-phosphate and/or $^{35}$S-methionine/cysteine. LY294002 treatment had the most profound effect on the PDGF-regulated phosphoproteome, as illustrated by comparison of the alterations induced by LY294002, ML-7 or rapamycin with EGF treatment (Supplementary Data; Figure S6). Consistent with this, the rate of protein synthesis in PDGF treated cells was significantly reduced in the presence of LY294002 (Figure 5A), indicating that the PI3K/Akt signaling pathway is critical for PDGF’s ability to control proteome composition in 3T3 fibroblasts. One particularly relevant target of PDGF-induced protein synthesis (that was reversed by LY294002 treatment) was identified as eukaryotic translation elongation factor 2 (eEF2) (Figure 5A). Cluster analysis showed that the images from cells treated with PDGF for 18 hours in the presence of LY294002 were more similar to those from cells treated with EGF alone (Figure 5B), whilst those treated with PDGF in the presence of the MEK inhibitor PD98059, clustered with the images from cells exposed to two growth factors for a similar length of time (Figure 5C). Our results thus identify PI3K as an early point of signal diversification in Swiss 3T3 fibroblasts and imply that lipid kinase activity is critical for the regulation of delayed protein synthesis in PDGF-activated cells.

Phospho radio-labeling with kinase inhibitor and phosphatase treatments, revealed that PDGF-specific phospho-forms of the histone chaperone nucleosome assembly protein-1 like-1
(NAP1L1) and protein chaperone HSP73 were regulated via PI3K/Akt-dependent signaling (Supplementary Data; Figure S7). For example, the abundance of the acidic HSP73 phospho-form in PDGF-stimulated cells was reduced by simultaneous LY294002 treatment, but unaffected by treatment with rapamycin, indicating that the regulatory signals are generated downstream of Akt, but upstream of mTOR kinase (Figure S7B).

**PI3K/Akt signaling suppresses the synthesis and expression of AAA-family member 26S protease regulatory subunit 7** Nine of the 117 proteins identified are involved in the regulation of protein stability, including the 26S proteasome subunits β type-3, 7, 8, 9 and 13. Whilst all three GFs induced similar fluctuations in 26S proteasome subunit β3 synthesis, long-term EGF-stimulation induced a strong, specific reduction in COP9 signalosome subunit 4 synthesis (Supplementary Data; Figure S8). Conversely, synthesis of 26S protease regulatory subunit 7 (PRS7) was unaltered by EGF treatment (Figure 6A), but dominantly suppressed by PDGF or IGF-1 co-treatment (Figures 6B and D), resulting in a decrease in protein levels, as assessed by analysis of fluorescent dye-stained gel images (Figure 6F). While treatment with the MEK inhibitor PD98059 or the TOR kinase inhibitor rapamycin had no effect on the reduced PRS7 synthesis in PDGF and/or IGF-1 stimulated cells, the suppressive effect was abrogated when GF-stimulation was performed in the presence of the PI3K inhibitor LY294002 (Figures 6C and E). These results demonstrate that RTK-induced PI3K/Akt pathway activation can suppress the synthesis and expression of one of six highly-related 26S proteasome ATPases that belong to the AAA family. Whilst primarily involved in proteasomal degradation, these ATPases have also been implicated in the regulation of transcription (26). Whether PI3K-specific PRS7 regulation exemplifies a general mechanism that enables RTKs to modify gene expression and/or protein stability by changing the abundance of AAA-family members remains to be demonstrated.

**IGF-1 and PDGF enhance the de novo synthesis of the receptor for activated C kinase-1 (RACK-1) via the PI3K/AKT pathway and increase binding of phosphoproteins to the RACK-1 signaling scaffold** Given the dissimilar regulation of PI3K and PKC phosho-signaling in EGF, IGF-1 and PDGF-treated cells, a differentially regulated protein of particular interest was the receptor for activated C kinase-1 (RACK-1/GNB2L1). *De novo* synthesis of RACK-1 was induced by PDGF or IGF-1 and any kind of co-stimulation, but was suppressed by EGF (Figure 7A and B). RACK-1 induction by PDGF was rapid, and the elevated rate of synthesis was maintained during continuous stimulation, while IGF-1 increased RACK-1
synthesis only after prolonged stimulation (Figure 7A and B). Densitometry analysis of immunoblots and fluorescence-stained gels revealed that long-term exposure to PDGF and IGF-1 increased RACK-1 expression by more than 30%, whereas EGF reduced its abundance by more than 50% (Figure 7B and C). Notably, RACK-1 synthesis was inhibited by LY294002, but unaffected by PD98059 or rapamycin co-treatment (Figure 7D). Thus, PI3K/Akt-induced signals regulate RACK-1 synthesis upstream of TOR kinase, and are independent of the MAP kinase pathway.

RACK-1 was next immunoprecipitated from \(^{33}\text{P}\)-orthophosphate labeled cells to investigate whether protein interactions at the RACK-1 scaffold remain susceptible to RTK-specific phospho-signals hours after initiation of GF-stimulation. Distinct patterns of RACK-1 interacting phospho-proteins were isolated from cells treated with the three growth factors for 2 or 4 hours (Figure 7F). The most abundant and diverse array of phosphoproteins was copurified with RACK-1 from cells stimulated with IGF-1 for 2 hours and we were able confirm the presence of phosphorylated FAK, Src and PKC\(\beta\)II in the RACK-1 immunoprecipitates (Figure 7E and F). After 4 hours stimulation the repertoire of strongly phosphorylated proteins bound to RACK-1 in IGF-1 treated cells had declined, whilst this was sustained, and in some cases increased, in the PDGF treated cells (Figure 7F). This finding is consistent with the observation that PDGF produces the most robust and long-lasting phospho-activation of key signal transducers (Figure 3). Immuno-fluorescence staining demonstrated that RACK-1 was localized to the cytosol of Swiss 3T3 fibroblasts, but was also found in the dendritic protrusions specific to long-term PDGF stimulation (Figure 7G). Several areas of dense RACK-1 staining were seen in the retractile protrusions, mainly at sites of cell-cell contact.

**PDGF-specific control of cytoskeleton regulators during the establishment of a migratory phenotype** Several F-actin and actomyosin regulators, including acidic calponin, gelsolin, tropomysin, Lasp-1, septin 2, Sid23p, zyxin, fascin and four subunits of the cytosolic chaperonin CCT were specifically regulated by PDGF. During the initial 6 hours of treatment, PDGF reduced the synthesis rates of acidic calponin, gelsolin, septin 2, septin 11, Lasp-1 and Sid23p, compared to EGF and IGF-1 (Figure 8). Moreover, the suppression of gelsolin, acidic calponin, septin, and Sid23 synthesis was dominant in co-stimulation experiments with PDGF and EGF or IGF-1 (Table S1). PDGF induced the most distinctive changes in the synthesis of the actomyosin regulators and septins after 12-18 hours of treatment (Figure 8), correlating
with development of full morphological differentiation and induction of motility in the Swiss 3T3 fibroblasts (Figure 1B).

Lasp-1, an adaptor protein involved in the regulation of specific dynamic actin-plasma membrane-associated activities (27), displayed an initial reduction in synthesis with PDGF and IGF-1 treatment, but was then dominantly increased at later timepoints with PDGF (Figure 8). Similarly, long-term PDGF treatment was dominant in increasing the synthesis rates of four TCP-1/CCT subunits; component of the cytoskeletal protein biogenesis machinery crucial for the native folding of actin and tubulins (Figure 8). These changes are again consistent with the induction of the retractile dendritic protrusions and development of a migratory cytoskeleton.

Acidic calponin was identified from two differentially regulated protein spots migrating at apparent $p\!/\!Is$ of 5.3 and 5.1, respectively. PDGF treatment led to a reduction in the synthesis of both calponin forms compared to EGF or IGF-1 treatment, whilst the intensity of the most acidic form was increased by PDGF, but not by EGF or IGF-1 treatments (Figure 9A-C). This suggests that acidic calponin becomes phosphorylated following synthesis in PDGF-stimulated cells. Treatment with LY294002 reduced the abundance of both calponin forms in $^{35}$S-labeled gel images, suggesting that calponin synthesis is in part regulated by PI3K-dependent signaling, whilst the acidic isoform was reduced by treatment with rapamycin, implicating mTOR-dependent phosphorylation (Figure 9D).

The actin-cross linking protein fascin was detected in digests from four $^{35}$S-labeled gel spots that were differentially regulated; however reliable densitometry data could only be derived from two, including the spot containing the most basic fascin form (Figure 10A and Table S1). While the synthesis of the two fascin isoforms remained unaltered in IGF-1 stimulated cells, their synthesis was slightly increased by EGF, but reduced by PDGF treatment (Figure 10A, left). Lowered levels of fascin expression in PDGF stimulated cells was confirmed by immunoblotting and densitometry analysis of fluorescent dye-stained gels (Figure 10B). In contrast, the ratio between the acidic and the basic isoform was increased dominantly by PDGF treatment and displayed a biphasic profile with elevations following 6 and 18 hours of exposure to PDGF (Figure 10A, right). 2-D immunoblotting suggested up to five isoforms of fascin were generated in response to GF treatment, with the response much lower in EGF-treated cells (Figure 10C). The PDGF and IGF-1 induced modifications and expression were attenuated by treatment with either LY294002 or PD184352 (a specific inhibitor of MEK), with LY294002 producing the greatest effect on the ratio between the putative phosphorylated isoforms. These findings suggest that fascin is phosphorylated at multiple sites via a PI3K-dependent pathway. Immuno fluorescence experiments localized fascin to the cytosol, with
high abundance in the perinuclear area of Swiss 3T3 fibroblasts and in the dendritic protrusions of PDGF-stimulated cells, where distinct areas of dense staining were observed (Figure 10D). This supports a role for fascin in the extension and retraction of these unique protrusions, whose full development coincide with the observed late increase in the ratio between the acidic and basic isoforms of fascin (Figure 10A, bottom right).

Since several of the differentially regulated proteins identified are known calcium-dependent regulators of actin polymerization and actomyosin ATPase activity (namely acidic calponin, gelsolin and tropomyosin), Ca\(^{2+}\) binding capacity was next examined using a 2-D \(^{45}\)Ca-overlay strategy. Although preliminary, this analysis revealed specific differences in the response to different GF treatments (Supplementary Data; Figure S9). Similarly, a Fura-2 binding experiment revealed different levels of free intracellular calcium in cells stimulated with the three growth factors (data not shown). It is thus tempting to speculate that the GF-specific changes in the expression of these actomyosin regulators are also coordinated through alterations in calcium-dependent signaling.
Discussion

This multiplexed proteomic study has revealed that while activation of distinct RTKs induce broadly overlapping patterns of protein synthesis, with some synergy displayed in response to co-stimulation, small subsets of the proteome show specific and differential regulation in response to individual growth factors, and that this is likely to determine specific biological outcomes of signaling from different RTKs. We observed a range of distinct regulatory patterns, the majority of which can be explained by differences in timing, strength and duration of their inducing signals, with PI3K activation representing an important point of signal diversification for PDGF versus EGF or IGF-1 signaling. The findings imply that the phenotypic outcome from GF-stimulation depends on the corresponding receptor’s ability to sustain distinct signaling activities during long-term exposure to the ligand.

Simultaneous activation of two or more RTKs was also shown to modify protein synthesis and phospho-signaling events. Co-stimulation experiments showed that for some proteins, one GF could have a dominant regulatory effect over the others and was particularly apparent in the case of PDGF-mediated modulation of the cytoskeleton regulators. Moreover, these dominant regulatory effects correlated with the timing of PDGF-specific changes to the Swiss 3T3 fibroblast actin cytoskeleton and conversion to a migratory cell type. Co-stimulation with EGF and IGF-1 also resulted in both additive and synergistic increases in protein syntheses, and correlated with similar phenotypic effects, such as cytoskeletal rearrangement.

This study has also identified several novel regulatory pathways, particularly those involving PDGF-induced PI3K-dependent signaling. For example, the apparent regulated phosphorylation and synthesis of the histone chaperone nucleosome assembly protein-1 like-1 (NAP1L1) is novel, and suggests that chromatin modulation may be required in determining specific responses to PDGF in Swiss 3T3 fibroblasts. Likewise, PDGF and IGF-1 both repressed the synthesis of PRS7, a component of the 26S proteasome. This was reflected in a reduced level of PRS7 protein expression and could be blocked by inhibition of the PI3K/Akt pathway, but was independent of MAPK activity. The synthesis of several other proteasome subunits was also differentially modulated. Whilst this may in turn feed forward to regulate protein turn-over and hence the expression and activity of other identified proteins, there is some evidence implicating these proteasomal proteins in the regulation of transcription initiation and activator destruction (26), and this may be an alternative mechanism by which specific growth factors can differentially regulate protein expression in a PI3K-dependent manner. Another protein whose expression was regulated by PI3K/Akt signaling was RACK-1, a scaffold protein involved in the recruitment and
regulation of a variety of signaling molecules (28), some of which have been demonstrated here. RACK-1 may promote cell spreading and migration through the recruitment and regulation of PKCβII, PI3K, RhoA, Src and FAK. Notably, RACK-1 also interacts with IGF-1R to modulate IGF-1-dependent integrin signaling, with RACK-1 overexpression in fibroblasts resulting in increased cell spreading, stress fibres and focal adhesions and reduced IGF-1-dependent cell proliferation (29). We speculate that the observed changes in RACK-1 synthesis may at least in part explain the prolonged activation of PKCβII, Akt and Src observed in PDGF stimulated cells (Figure 3), which in turn may promote the PDGF-dependent migratory phenotype. This is supported by previous work showing that RACK-1 regulates integrin-mediated adhesion, protrusion and cell migration via its Src-binding site (30), with PKC activation inducing the RACK-1-Src interaction (31) and leading to inhibition of Src activity (32). RACK-1 is also a ribosomal component, and has been shown to promote translation through recruitment of PKCβII and the subsequent phosphorylation of the translation initiation factor EIF6 (33, 34). Thus, the observed increase in RACK-1 expression in response to PDGF may also promote PKCβII recruitment and activation which in turn increases protein translation. This may at least in part explain the higher rate of de novo protein synthesis in PDGF-stimulated cells. Other studies have suggested that ribosomal RACK-1 can support miRNA-mediated gene repression (35) and nascent polypeptide-dependent translation arrest (36). Thus, it could be that PDGF-induced RACK-1 expression may specifically block translation of the cytoskeletal regulators identified here (Figure 8), to promote a motile phenotype.

With respect to activity suppression, the PDGF-specific, dominant increase in S-adenosylhomocysteine hydrolase (copper binding protein) synthesis (Supplementary Figure S5I), is also noteworthy. S-adenosylhomocysteine hydrolase is the only enzyme known to cleave S-adenosylhomocysteine, a product and a potent inhibitor of all S-adenosylmethionine-dependent transmethylation reactions. Increased S-adenosylhomocysteine hydrolase activity thus favors DNA/RNA methyltransferase activation through the removal of S-adenosylhomocysteine, which PDGF might exploit to silence specific genes.

As mentioned, the regulatory dominance of PDGF receptor triggering was particularly apparent with respect to the temporal induction and suppression of several regulatory cytoskeletal components that correlated with the timing of the PDGF-specific reorganization of the actomyosin cytoskeleton, the development of dendritic protrusions and the conversion into a migratory cell type. Our demonstration of PDGF-specific regulation of acidic calponin, gelsolin, Lasp-1, tropomyosin alpha-1 chain and fascin is noteworthy, as they are known to
interact functionally in regulating actin polymerization, cellular protrusions and actomyosin ATPase activity. Tropomyosin modulates the actin severing activity of gelsolin and can, in conjunction with caldesmon, inhibit the actin-bundling activity of fascin (37-40). Likewise, interaction of acidic calponin and the Rac-activated protein caldesmon has been demonstrated and is implicated in regulation of smooth muscle cell contraction (41). The activities of fascin and gelsolin have been shown to be regulated downstream of Rac and/or Cdc42 (42-44). Since PDGF is a known activator of Rac and Cdc42, the PDGF-specific regulation of these actin modulators in determining the migratory phenotype is likely to be coordinated downstream of these small GTPases.

We also provide evidence for the differential phosphorylation of acidic calponin and gelsolin in response to PDGF versus EGF or IGF-1. For example, phosphorylation of calponin at Ser175 by PKC inhibits its actin binding capacity and attenuates its ability to inhibit actomyosin ATPase (45, 46). The prolonged PKC activation and increased calponin phosphorylation observed in PDGF-stimulated cells may thus alleviate inhibition of actomyosin ATPase activity, allowing the high actin-myosin cross-bridge cycling rate necessary for cell contraction and motility. Calcium binding is also likely to play a role in regulating these actin modulators, and we provide preliminary evidence of GF-specific differences in calcium-binding capacity. Indeed, calcium is known to increase the association between calponins and caldesmon for regulation of smooth muscle cell contraction (41), with calponin also stimulating actin polymerization and bundling to stabilize actin microfilaments (47, 48).

Taken together the results imply that the morphological transformation induced by PDGF is achieved via concerted changes in the expression pattern and activity of key actomyosin regulators that become increasingly apparent during long-term GF-treatment. This suggests that once GF-specific reorganization of the actin cytoskeleton is achieved, its maintenance involves controlled expression of key cytoskeletal components through the prolonged activation of multiple signaling cascades. The observations that Src inhibition blocked the PDGF-induced motile phenotype and PDGF treatment induced a relatively sustained phospho-activation (Y424) and transient phospho-inhibition (Y535) of Src, is consistent with a recent report showing that PDGF regulates two spatially distinct pools of Src kinases which control different biological outcomes (i.e. cell growth and morphological change) in NIH 3T3 fibroblasts (49). Src kinases also facilitate the formation of focal adhesion complexes, and it is tempting to speculate that PDGF-induced signaling leads to temporary silencing of localized Src kinase pools in the proximity to FA complexes (perhaps through RACK-1 interaction), while simultaneously activating Src kinases in other compartments to promote the motile phenotype. We thus propose
an initial, rapid demolition of existing higher order structures such as actin SF and FA complexes in response to PDGF, facilitated by short-term inhibition of FAK and coflin activities, and a Src-induced, MIR-mediated ubiquitination and subsequent degradation of MRLC (15). The second more prolonged reorganization phase is characterized by specific temporal changes in the synthesis and expression of key cytoskeletal components to direct the assembly of a motile cytoskeleton, independent of EGF and IGF-1-induced antagonistic signaling activities.

Finally, the study also emphasizes that quantitative differences in the density of $^{35}$S-labeled protein spots on 2D gels do not simply reflect temporal changes in protein synthesis, but also reflect changes in post-translational modifications (PTMs) that alter the migration of newly synthesized proteins or change their stability. For this reason, biosynthetic labeling experiments must be combined with the analysis of expression levels (e.g. by fluorescence gel staining and immunoblotting), degradation rates (e.g. by CHX-chase experiments) and post-translational modifications (e.g. by PTM-specific radio-labeling procedures, immunoblotting, and selective modification of signal transduction pathway activities), before ligand-specific regulation of proteins can be correctly interpreted by 2D gel-based proteomic approaches.

Acknowledgements
This work was funded by the Ludwig Institute for Cancer Research, and by a grant from the Karen Elise Jensen foundation to Soren Naby-Hansen. Rob Stein and John Timms are supported by the UCLH/UCL Comprehensive Biomedical Research Centre.
References


**Figure legends**

**Figure 1 A.** Immunofluorescence analysis of cytoskeleton changes in Swiss 3T3 fibroblasts induced by long-term stimulation with PDGF, IGF-1, EGF and their combinations. Focal visualized by fluorescence staining with anti-vinculin antibody and rhodamine-phalloidin staining, respectively. Dendritic protrusions are indicated by arrows (middle column). Note the dominance of PDGF in co-stimulation experiments, resulting in development of an elongated cell shape with retractile dendritic protrusions and disruption of stress fibers (SF) and focal adhesion (FA) complexes. **B.** Immunofluorescence analysis of temporal changes in the actin cytoskeleton during long-term GF-treatment. Actin filaments were stained with rhodamine phalloidin. In the PDGF-treated cells, SF and FA had disappeared by 6 hr, dendritic protrusions began to appear after 12 hr and the development of an elongated, motile cytoskeleton was fully accomplished by 18 hr treatment. **C.** Changes in focal adhesion density and cell surface area in response to growth factor stimulation.

**Figure 2 A.** Global analysis of protein synthesis in Swiss 3T3 cells stimulated with PDGF, IGF-1, EGF, and their combinations. Newly synthesized $^{35}$S-methionine- and cysteine-labeled proteins were separated on broad range, non-linear (pH 3-10NL) IPG/PAGE gels. Radio-labeled proteins were detected by phosphoimaging. The elevated *de novo* synthesis in PDGF-stimulated cells was blocked by cycloheximide. Note the profound reduction in protein synthesis in cells where long-term PDGF treatment (18 hr) was performed in the presence of the PI3K inhibitor LY290042 (10 μM). **B.** Number of 2DE protein spots detected in extracts from cells treated with PDGF, IGF-1, EGF, and their combinations. Left; temporal changes in the number of $[^{35}$S]-labeled protein spots detected following 6, 12 and 18 hr stimulation. Right; numbers of radio-labeled proteins detected following 18 hr of stimulation. Numbers are the average taken from three independent experiments. **C.** Comparison of the number of matched protein spots showing a $>$2-fold increase in $^{35}$S-incorporation between different GF treatments at the 18 hr time point. Values were calculated as a percentage of total matched protein spots and compare the synthetic responses to growth factors and their combinations to those generated by individual growth factor stimulations. The analysis provides an indication of the comparative potency of each GF in stimulating or suppressing protein synthesis. **D.** Comparison of the number of matched protein spots showing a $>$2-fold reduction in $^{35}$S-incorporation between different GF treatments at 18 hr. **E.** Hierarchical clustering analysis of protein synthesis in GF-stimulated cells. 2D gel images of biosynthetic labeled proteins were
clustered according to similarity between the density patterns of a group of 172 proteins, whose synthesis rates were almost identically regulated by the three growth factors after 6 hr. In the resulting relational tree, short branches connect similar images, while increasingly long branches join gel images with diminishing similarity. F. ChiClust clustering analysis of protein synthesis during long-term GF-stimulation. The 2DE images were clustered according to similarities between all radio-labeled protein spots.

Figure 3 Analysis of growth factor induced phospho-signalling. A. Temporal changes in key signal transducer activities demonstrated by immunoblotting with phosphorylation site-specific antibodies. β-actin served as loading control. B. Immunoblot analysis of Akt and p38 phosphorylation in PDGF and EGF treated cells (top). Pre-treatment with the PI3K-inhibitor LY294002 (10 μM) abolished the PDGF-induced increase in p38 phosphorylation at the 60 min time point (bottom). C. Immunoblot analysis of PKC and JNK phosphorylation in EGF and PDGF treated cells. Both GFs induced a weak, long-lasting increase in JNK phosphorylation. While high levels of phospho-activated PKC was sustained for several hours in PDGF treated cells, less than 30% of the increase in PKC phosphorylation found in EGF treated cells at the 15 min time point were detected after 60 min treatment. Pre-treatment with LY294002 reduced the level of phosphorylated PKC in cells treated with PDGF for 60 min (bottom). D. The PDGF-induced increase in phosphorylation of c-Src at the inhibitory site (Y535) at the 15 min time point was abolished by pre-treatment with the Src-specific inhibitor SU6656 (2 μM), indicating that the modification is induced by autophosphorylation. E. 2D autoradiograms of [33P]-orthophosphate labelled proteins isolated by immunoprecipitation with an anti-c-Src antibody following 2 hr and 4 hr growth factor treatment. The white oblique downwards arrows indicates phosphoproteins that co-precipitated with Src from cells treated with PDGF for 4 hr, but were absent or weakly represented in immunoprecipitates from cells that had been exposed to the growth factor for 2 hr. The radiolabelled phosphoprotein isolates were separated by IEF/SDS-PAGE, employing broad pH-range (3-10 NL) dry strips in the first dimension.

Figure 4 A. Analysis of changes in phosphorylation patterns induced by stimulation with EGF, PDGF or both. Synchronized cells were grown to ~70% confluence and labeled with 0.1 mCi/mL [33P]-orthophosphate for 4 hr prior to GF stimulations. The radiolabelled proteins were separated by IEF/SDS-PAGE, transferred to PVDF-membranes and visualized by phosphoimaging. PDGF induced the strongest increase in 33P-incorporation into the commonly
induced phosphoprotein isoforms, indicating that PDGF induces phosphorylation of more copies of the proteins than EGF at both time points. The number of phospho-proteins induced by PDGF stimulation was also higher than the number induced by EGF. Receptor co-stimulation influenced both the specificity and strength of protein phosphorylation at the 10 and 30 min time points. Examples of PDGF-regulated phosphorylations that were influenced by co-stimulation with EGF and vice versa are indicated by the arrows. B. 2D immunoblotting with phosphotyrosine-specific antibody demonstrating a reduction in the number and abundance of tyrosine phosphorylated proteins in co-stimulated cells (black arrows), those unaffected by co-stimulation (while oblique arrow), and those increased in co-stimulated cells (white upward arrows). C. EGF and PDGF treatment also triggered dephosphorylation of proteins that were phosphorylated in unstimulated, serum-starved cells.

**Figure 5 A.** $^{35}$S-autoradiograms demonstrating kinase inhibitor-induced modulation in PDGF-regulated protein synthesis at the 18 hr time point. Newly synthesized EF2 isoforms are boxed by white rectangles. The abundance of newly synthesized in general were less affected by inhibition of MEK or TOR than by inhibition of the PI3K/Akt pathway in cells exposed to long-term PDGF treatment. Newly synthesized eEF2 isoforms are indicated by white boxes. B. Hierarchical clustering analysis of protein synthesis in cells treated with EGF or PDGF for 6, 12 and 18 hr, or cells treated with PDGF in the presence of LY290042 for 18 hr. The resulting 2DE images were clustered according to similarities between all radio-labeled protein spots. C. Clustering analysis demonstrating that cells treated with PDGF for 18 hr in the presence of the MEK inhibitor PD98059 clustered in proximity to images from cells exposed to two growth factors.

**Figure 6 A.** Graphical representation of temporal changes in 26S protease regulatory subunit 7 (PRS7) synthesis during long-term GF-stimulation. B. Enlarged area of autoradiographic images demonstrating PRS7 synthesis at the 18 hr time point. C. $^{35}$S-autoradiograms demonstrating PRS7 synthesis in cells treated with PDGF and IGF-1 for 12 hr in the presence of TOR, MEK or PI3K inhibitors. While the three inhibitors had similar effect on the synthesis of the protein indicated by a box in the upper left corner of the images, inhibition of PI3K caused a significant increase in the amount of $^{35}$S-labeled PRS7 (indicated by upward arrows). D. $^{35}$S-autoradiograms demonstrating IGF-1 dominant suppression of PRS7 synthesis in cells co-stimulated with EGF for 18 hr. E. PRS7 synthesis in cells treated with PDGF for 18 hr in the presence or absence of LY290042. F. Enlarged areas of 2D gel separated whole cell extracts
stained with the fluorescent dye OGT MP17 demonstrating PDGF-induced dominant suppression of PRS7 expression in co-stimulated cells.

**Figure 7 A.** Graphical representation of temporal variations in the synthesis of RACK-1 in response to different GFs. **B.** Enlarged area of autoradiograms demonstrating RACK-1 synthesis at the 18 hr time point (upper) and immunoblotting of RACK-1 expression following 18 hr GF-stimulation (lower). **C.** Enlarged area of fluorescence stained gels demonstrating the expression levels of RACK-1 and a glutathione-S-transferase homolog in cells treated with EGF, PDGF or both for 18 hr. The elevated expression of both proteins in response to PDGF was maintained in cells simultaneous exposed to EGF. **D.** Biosynthetic $^{35}$S-label analysis of RACK-1 synthesis in cells co-simulated with PDGF and IGF-1, and in cells simultaneous treated with different kinase inhibitors. **E.** Immunoblot analysis of immunoprecipitates and whole cell extracts. The observed co-purification of PKC$\beta$II and RACK-1 with the anti-RACK-1 antibody, verified that this antibody, in contrast to the anti-PKC antibody, could be reliably employed in immunoprecipitation experiments. **F.** 2DE autoradiograms of $[^{35}$P$]$-orthophosphate labeled phospho-proteins that co-purified with RACK-1 in immunoprecipitates from GF-stimulated 3T3 cells. The area indicated by a white box in the IGF-1 (2 hr) image shows the migration of FAK isoforms of which 4 were detected by immunoblotting with an antibody against pY428 of FAK (inserted image). The positions of co-purified PKC$\beta$II and c-Src proteins are indicated by white boxes in the PDGF (2 hr) image and were detected by immunoblotting (inserts). **G.** Immunofluorescence localization of RACK-1 in Swiss 3T3 cells stimulated with PDGF, IGF-1 or EGF for 18 hr. RACK-1 was detected using a FITC-tagged antibody, while the actin cytoskeleton was visualized by rhodamine-phalloidin staining. Arrows indicate RACK-1 localized in dendritic protrusions with dense staining at cell-cell contact sites (right images).

**Figure 8 A.** 2DE autoradiogram of $^{35}$S-labeled proteins in IGF-1 treated cells, indicating the position of differentially regulated cytoskeletal proteins and regulators. The representative autoradiogram images at the right indicate the different synthesis rates of ten cytoskeleton proteins in GF-stimulated or co-stimulated cells. **B.** Graphic illustrations of the regulation of cytoskeletal protein synthesis in EGF, IGF-1 and PDGF treated cells.
**Figure 9 A.** Graphic representations of GF-induced temporal changes in the synthesis of two acidic calponin isoforms. **B.** Representative autoradiogram images demonstrating the abundance of newly synthesized calponin forms in GF-stimulated cells. **C.** Fluorescence stained gel images demonstrating reduced expression and inverse ratio between calponin isoforms in PDGF-treated, compared to EGF-treated, 3T3 cells. **D.** Autoradiograms showing the effects of treatment with PI3K or mTOR inhibitors on PDGF-induced calponin synthesis.

**Figure 10 A.** Graphic representation of temporal variations in de novo synthesis of two fascin isoforms. The combined density of the two spots is given at the left, while changes in the ratio between the isoforms are illustrated to the right. **B.** Western blot analysis of fascin expression following GF-treatment (left) and post-electrophoretic protein staining with the fluorescent dye OGT MP17 used to examine the expression of the two fascin isoforms in EGF and PDGF stimulated cells (right). **C.** Immunoblotting of fascin isoforms in cells stimulated with PDGF, IGF-1 or EGF in the presence or absence of kinase inhibitors. **D.** Localization of fascin in PDGF, IGF-1 and EGF-stimulated cells. Cells were stained with anti-fascin antibody following 18 hr GF stimulation. Arrows show the localization of fascin in dendritic protrusions, and arrowheads indicate dense areas of staining within protrusions.
Figure 1
Figure 2
Figure 4
Figure 5
Figure 6

A Temporal changes in the de novo synthesis of PRS7

B Biosynthetic label after 18 hr stimulation period

C $^{35}$S-incorporation following 12 hr exposure to PDGF & IGF-1 plus

D Dominant suppression of PRS7 synthesis by IGF-1

E Changes in PRS7 synthesis following PI3K inhibition

F Fluorescence stained whole cell extracts following 18 hr treatment
Figure 7
Temporal changes in acidic calponin synthesis

Acidic calponin total
(isoform 1 + 2)

Acidic calponin ratio
(isoform 1/[1+2])

B
Biosynthetic label of calponin isoforms following 18 hr

PDGF  IGF-1  EGF  P+I  P+E  E+I

C
Expression of acidic calponin isoforms in whole cell extracts

EGF  PDGF

D
Acidic calponin isoforms in $^{35}$S-labeled 2D gels

PDGF  + LY294002  + rapamycin

Figure 9
Figure 10