Cyclophilin A promotes cell migration via the Abl-Crk signaling pathway

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Summary

Cyclophilin A (CypA) is over-expressed in a number of human cancer types, but the mechanisms by which CypA promotes oncogenic properties of cells are not understood. Here we demonstrate that CypA binds to and prevents the CrkII adaptor protein from switching to the inhibited state. CrkII is involved in cell motility and invasion by mediating signaling through its SH2 and SH3 domains. CrkII Tyr221 phosphorylation by the Abl or EGFR kinases induces an inhibited state of CrkII, by means of an intramolecular SH2-pTyr221 interaction, causing signaling interruption. We show that the CrkII phosphorylation site constitutes a binging site for CypA. Recruitment of CypA sterically restricts the accessibility of Tyr221 to kinases, thereby suppressing CrkII phosphorylation and promoting the active state. Structural, biophysical, and in vivo data show that CypA augments CrkII-mediated signaling. A strong stimulation of cell migration is observed in cancer cells wherein both CypA and CrkII are greatly up-regulated.
INTRODUCTION

Cyclophilin A (CypA) is a highly abundant protein, accounting for up to ~0.6% of the total cytosolic protein content\(^1\). CypA is involved in a growing number of biological processes, including protein folding, signal transduction, viral infection, trafficking, receptor assembly, immune response, and transcription regulation\(^2\). Although several proteins have been identified to interact with CypA\(^3\)–\(^6\), the underlying mechanism of the CypA action and the physiological implications of the interactions remain in most cases unknown. CypA exhibits peptidyl–prolyl cis–trans isomerase (PPIase) activity by catalyzing cis–trans isomerization of peptide bonds preceding proline residues\(^7\). CypA can in principle act as an enzyme or a binding partner\(^8\) in mediating the biological processes.

Over the recent years a number of studies have consistently demonstrated that CypA is significantly up-regulated in various cancers and cancer lines\(^9\),\(^10\). The mRNA expression profile of different tumors shows an elevated level of CypA mRNA\(^11\) and proteomic studies of cells treated with anticancer drugs revealed a decrease in the level of CypA\(^10\). CypA has been directly linked to cell proliferation and tumor growth and treatment of cells with cyclosporin (CsA) or the downregulation of CypA by small interfering RNA (si-RNA) reduces the proliferative effects of CypA\(^12\),\(^13\). Although there is ample evidence to suggest that the presence of CypA confers a proliferative and anti-apoptotic effect, the molecular mechanisms of CypA function in oncogenesis remain elusive.

The Crk (CT-10 Regulation of Kinase) family of adaptor proteins are ubiquitously expressed in tissues and mediate the formation of protein complexes elicited by various extracellular stimuli, including growth and differentiation factors\(^1\),\(^4\). Crk proteins are overexpressed in many human cancers including lung and breast cancer, glioblastoma and synovial sarcoma\(^15\)–\(^19\). The Crk expression levels positively correlate with the disease stage and poor survival outcome\(^20\). Crk knockdown results in attenuated invasion and migration of cancer cell lines\(^21\). Crk proteins stimulate the activity of Abl\(^22\), a kinase whose fusion to Bcr causes chronic myelogenous leukemia\(^23\), and interact strongly with the epidermal growth factor receptor (EGFR) kinase thereby mediating oncogenic EGFR signaling\(^24\).

Cellular Crk (CrkII; 304 amino acids) consists of an SH2 domain, an N-terminal SH3 (SH3\(^{N}\)) domain and a C-terminal SH3 domain (SH3\(^{C}\)) (Fig. 1a). Previous NMR studies showed that a proline residue (Pro237, human numbering) located at the SH3\(^{C}\) domain of chicken CrkII undergoes cis–trans isomerization\(^25\),\(^26\). The isomerization process regulated the intramolecular interaction between the two SH3 domains and the rate of the interconversion was shown to be accelerated catalytically by CypA\(^25\),\(^26\). In contrast, Pro237 does not exhibit isomerization in human or other mammalian CrkII proteins\(^27\) because of variations in the amino acids flanking Pro237\(^28\). The SH3\(^{N}\) and SH3\(^{C}\) domains are tethered by a 50 residue–long linker, which contains a tyrosine residue (Tyr221) that becomes phosphorylated in vivo by Abl\(^29\) and EGFR kinases\(^30\). Tyr221 phosphorylation results in intramolecular association between the SH2 domain and this region of the linker (pY221-A-Q-P)\(^27\),\(^31\) (Fig. 1b). This intramolecular association engages the SH2 domain and prevents its binding to focal adhesion scaffold proteins such as paxillin and p130CAS (Fig. 1a and 1b), thereby reducing drastically cell migration and motility\(^32\). This negative regulatory
mechanism is absent in CrkI, an alternatively spliced isoform of CrkII that is only 204 residues long. Thus, CrkI adopts a constitutively active form and for this reason CrkI is associated with more aggressive phenotypes.

Here we show that the amino acid sequence flanking Tyr221 in CrkII, a sequence that is universally conserved in all species (Fig. 1c), constitutes a recognition site for CypA. CypA is recruited by CrkII Pro220, which undergoes cis–trans isomerization, and complex formation causes modulation of the level of CrkII phosphorylation by the Abi and EGFR kinases. This specific interaction between CypA and CrkII occurs both in vitro and in vivo. Suppression of Tyr221 phosphorylation leads to CrkII remaining in the active state wherein the SH2 domain is available to interact with paxillin or p130CAS thereby stimulating cell migration.

RESULTS

Proline isomerization at the CrkII phosphorylation site

Tyr221, the primary phosphorylation site in CrkII (Fig. 1a), is universally conserved among all CrkII proteins (Fig. 1c). Interestingly, the preceding residue, Pro220, is also universally conserved (Fig. 1c). NMR characterization of human CrkII shows that the amide signals of the residues flanking Pro220 (Glu217-Gln223) are duplicated (Supplementary Results, Supplementary Fig. 1a), indicating the presence of two conformations in slow exchange. Characteristic NOE correlations between Gly219 and Pro220, as well as \(^{13}C_β\) and \(^{13}C_γ\) chemical shift analysis of Pro220 (Supplementary Fig. 1b), indicate that the Gly219–Pro220 prolyl bond exists both in the cis and trans conformation (Supplementary Fig. 1a and c). We conclude that the source of the conformational heterogeneity in the phosphorylation site of CrkII is the presence of cis–trans isomerization about the Gly220–Pro221 prolyl bond.

The effect of Pro220 cis–trans isomerization is local, with only 7 residues (Glu217-Gln223) showing duplicate peaks. This is expected since Pro220 is located in an unstructured region of the linker. A synthetic peptide encompassing the Pro216-Ser225 region exhibits cis–trans isomerization at Pro220 demonstrating that this process is an intrinsic property of this sequence (Supplementary Fig. 2a, b). The sequence of the region around Pro220 is absolutely conserved among all CrkII proteins (Fig. 1c) and thus this isomerization process appears to be evolutionarily conserved.

CypA is recruited at the CrkII phosphorylation site

The presence of a heterogeneous proline residue undergoing cis–trans isomerization provides a potential binding site for PPIase enzymes. In fact, the Gly219–Pro220 motif in CrkII presents a preferred recognition site for CypA. To determine whether CypA interacts with this region in CrkII, we titrated CypA (human) to \(^{15}N\)-labeled CrkII (human) (Fig. 1d). Increased amounts of CypA resulted in specific chemical shift perturbations, with the residues close to the Pro220 binding site being most affected (Fig. 1e). Weaker effect was observed for several other residues in CrkII, presumably because of minor rearrangements in its assembly and domain interactions. The reverse titration, where unlabeled CrkII was titrated to \(^{15}N\)-labeled CypA, further corroborated the specific interaction between the two proteins.
proteins (Supplementary Fig 3a). Chemical shift and intensity perturbation analysis shows that CrkII binds to the catalytic site of CypA (Fig. 1f). Indeed, addition of CsA, an analog that binds with high affinity to the CypA catalytic site, abolishes binding between CypA and CrkII (Supplementary Fig. 3b, c). NMR characterization of the interaction between CypA and CrkII showed that the dissociation constant ($K_d$) of the complex is $\sim 15\pm 4\mu M$ (Supplementary Fig. 3d), an affinity typical of CypA–ligand interactions. Substitution of Pro220 by Ala in CrkII abrogates binding to CypA (Supplementary Fig. 3e) further corroborating the observation that the heterogeneous Pro220 provides the only CypA-binding site in human CrkII (Supplementary Fig. 2b). Taken together, the data demonstrate that CypA and CrkII form a stoichiometric and specific complex.

**Structural basis for the interaction of CypA with CrkII**

To understand the basis for the specific interaction between CypA and CrkII, we determined the solution structure of CypA in complex with a peptide encompassing the CypA-binding region of CrkII (Pro216-Ser225) (Supplementary Table 1). NMR analysis demonstrates that the chemical shift effect induced on the binding site of CypA by the peptide is very similar to the effect induced by full-length CrkII suggesting that the contacts between the CrkII peptide or full-length CrkII and CypA are essentially identical (Supplementary Fig. 3a).

The structural data show that the Gly219–Pro220 region of CrkII binds deep into the active site cleft of CypA (Fig. 2a), in an overall structural arrangement that is reminiscent of the binding of the Gly–Pro motif of the HIV-capsid protein (CA) to CypA (Supplementary Fig. 4b). Pro220 fits snugly into a hydrophobic pocket formed by CypA residues Phe60, Met61, Ala101, Phe113, and Leu122 (Fig. 2b). Several hydrogen bonds between polar side chains in CypA and the backbone of CrkII are also present (Fig. 2b). The side chain of Tyr221 is exposed to the solvent (Fig. 2a) and appears to form no contacts with CypA. The complex is suitably arranged for the catalysis of the cis–trans isomerization in CrkII by CypA. Indeed, $^{15}$N ZZ-exchange NMR spectroscopy shows that the addition of catalytic amounts of CypA accelerates the cis–trans isomerization process (Supplementary Fig. 3f). Thus, CypA uses its catalytic site to interact specifically with the CrkII linker region that encompasses the tyrosine-phosphorylation site.

**CypA and CrkII form a specific complex in the cell**

To further corroborate that CrkII is indeed a CypA substrate, we tested their interaction in live cells using fluorescence resonance energy transfer (FRET) microscopy imaging. We transiently transfected HeLa cells with CrkII tagged with yellow fluorescent protein (CrkII-YFP) and CypA tagged with a cyan fluorescent protein (CypA-CFP) and monitored the expression levels and distribution of the two proteins. Signal from the fluorescent tags indicated that both proteins reside in the cytoplasm and for CrkII-YFP the distribution agrees with previously published results (Supplementary Fig. 5a). In the cells cotransfected with both plasmids, maximum FRET was observed near the membrane ($\sim 70\pm 3.3\%$) (Fig. 2c and Supplementary Fig. 5b). Addition of CsA resulted in a dramatic decrease in FRET efficiency to $\sim 15\%$ (Fig. 2d). Thus, the FRET imaging data demonstrate that CrkII and CypA form a specific complex in vivo mediated by the CypA catalytic site.
**CypA inhibits CrkII phosphorylation in vitro**

When CypA is bound to CrkII, the backbone of the residues flanking the phosphorylation site is buried in the catalytic cleft (Fig. 2a). Therefore the CypA-CrkII complex could prevent phosphorylation of CrkII (replace with original). To test the hypothesis that CrkII–CypA complex formation results in suppression of phosphorylation of CrkII, we used a kinase assay to measure the efficiency of CrkII Tyr221 phosphorylation by Abl in the presence and absence of CypA. The data show that the level of CrkII Tyr221 phosphorylation is significantly lower in the presence of CypA (up to a factor of ~12) (Fig. 3a and Supplementary Fig. 6). As expected, phosphorylation increases with time because of the moderate affinity and the dynamic nature of the CypA–CrkII complex.

**CypA-mediated CrkII phosphorylation suppression in vivo**

Next we investigated whether CypA regulates CrkII Tyr221 phosphorylation in cells. To test the effect of CypA on CrkII phosphorylation, we prepared 293T cells where CypA was knocked-down using short hairpin RNA (shRNA). Consistent with the in vitro kinase assay, transient CypA knockdown (~50% efficiency) enhanced phosphorylation of CrkII at Tyr221 ~3 fold (Fig. 3b). Phosphorylation of CrkII was achieved by co-expression of CrkII and Abl in 293T cells. To determine the effect that increased amounts of CypA, as is the case in several cancer lines, have on CrkII phosphorylation, we used various approaches to ectopically overexpress CypA. However, all our efforts were unsuccessful and we could not achieve overexpression to a level greater than ~20% of the endogenous CypA level. For this reason, in an effort to increase the population of the CypA–CrkII complex in the cell we sought to design a CrkII variant with higher affinity for CypA. Analysis of the CypA–CrkII structure suggested that substitution of Pro218 (Supplementary Fig. 7a) by a bulkier hydrophobic residue would optimize juxtaposition of the two proteins. Molecular modeling indicated that a CrkII variant with a Phe residue in place of Pro218 (CrkII P218F) maximized the surface complementarity in the complex with CypA (Supplementary Fig. 7b). Indeed, the CrkII P218F variant binds CypA with a ~20-fold higher affinity than wild-type CrkII (Supplementary Fig. 7c). Interestingly, when 293T cells were transfected with CrkII P218F a significant decrease in Tyr221 phosphorylation was observed when compared with cells transfected with wild type CrkII (Fig. 3c).

**CypA suppresses CrkII phosphorylation in Cancer Cells**

CypA is overexpressed in many cancers. We screened several cancer cell lines and found that the MDA-MB-468 breast-cancer cell line showed one of the highest levels of CypA expression among them (Supplementary Fig. 8a). Incidentally this particular cell line overexpresses EGFR, which also phosphorylates CrkII at Tyr221, a reaction that can be stimulated by the addition of EGF. CypA knock-down MDA-MB-468 cells displayed up to ~10-fold higher phosphorylation at CrkII Tyr221 when compared to cells that were treated with a scrambled shRNA (at 1min) (Fig. 4a). In order to further investigate this inhibitory property of CypA, we treated serum starved MDA-MB-468 cells with CsA. As expected, the addition of CsA significantly increases phosphorylation at CrkII Tyr221 because of the disruption of the CypA–CrkII complex (Fig. 4b).
CypA binding to CrkII increases paxillin-CrkII complex

To determine the effect of CypA-mediated suppression of CrkII Tyr211 phosphorylation on CrkII association with signaling proteins, we probed changes in the interaction between CrkII and paxillin. The paxillin–CrkII complex forms when the activation of integrin receptor, in the context of cell adhesion and cell migration, results in the phosphorylation of paxillin creating a docking site for the CrkII SH2 domain. Examination of the complex formation between paxillin and CrkII in CypA knock-down MDA-MB-468 cells revealed a significant decrease (~10-fold) of paxillin association with CrkII when compared to MDA-MB-468 cells treated with scrambled shRNA. (Fig. 4c). In addition the CypA knockdown cells were passaged until the CypA levels were restored and the complex formation between CrkII and paxillin were reexamined. As expected the association of paxillin and CrkII increases to that of the control cell line. (Supplementary Fig. 8b). Thus, the specific interaction of CypA with CrkII results in a significant decrease in CrkII Tyr221 phosphorylation, which leads to an increase in the amount of CrkII bound to paxillin or other proteins interacting with the SH2 domain of CrkII such as p130CAS.

CypA–CrkII complex stimulates cell migration

To determine whether the CypA-effected suppression in CrkII phosphorylation has an impact on the physiology of the cell, we investigated the migratory properties of cells as a function of the CypA-CrkII interaction. Crk has been shown to enhance the migration of several patient-derived cancer cell lines including breast cancer. Phosphorylation of Tyr221 has been shown to inhibit Crk-mediated motility in breast cancer cells. Migration is a complex process that involves an intricate balance between several protein complexes that eventually leads to a reorganization of the cytoskeleton. We argued that a straightforward comparison of the migratory properties of cells in the presence and absence of CypA would provide direct evidence of whether the CypA–CrkII complex and the subsequent suppression of Tyr221 phosphorylation could significantly perturb the dynamic cell motility equilibrium.

We tested the effect of the interaction between CrkII and CypA on cell motility since CrkII is a key molecule in regulating cell motility. We prepared three cell lines of MDA-MB-468 cells (i) overexpressed wild type CrkII (CrkII); (ii) overexpressed CrkII(P218F); and, (iii) CrkII knock-down (CrkII-KD). A control cell line expressing EYFP was used in parallel for comparison. To obtain quantitative information about the cell migration properties, we used the xCELLigence system (Roche) that measures migration by detecting the electrical impedance using microelectronic biosensor technology. We followed the migration of MDA-MB-468 cells collecting data at 15 minute intervals. Both CrkII and CrkII(P218F) greatly enhanced cell motility compared to the control cells while CrkII-KD cells had no significant motility, in agreement with the crucial role of CrkII in cell motility (Supplementary Fig. 9a). Migration in CsA treated cells were markedly lower at the 4 hour time point. (Supplementary Fig. 9b). In a separate set of experiments, we also quantified the effects of CsA at earlier time points and increased the data collection frequency from 15 minutes to 15 seconds. Addition of CsA caused a significant decrease in migration (Fig. 5a). Taken together, the results indicate that formation of the CypA-CrkII complex stimulates the
migratory properties of the cells and that disruption of this complex by CsA results in suppression of cell migration.

To further investigate the effect of CypA-CrkII interaction on cell migration, we used a wound healing assay (Fig. 5b and Supplementary Fig. 9c). Using the control as a basis for the comparison, the CrkII-KD cells, even at the 24-hr time point displayed the least migratory tendency including a lack of cell spreading and absence of lamellipodia (Fig. 5c and Supplementary Fig. 9c, d). In contrast, overexpression of CrkII promoted cell spreading and numerous lamellipodia were observed at the leading edges of the culture (Supplementary Fig. 9d). The cell lines overexpressing CrkII and CrkII\textsuperscript{P218F} displayed increased cell proliferation and migration (Fig. 5c and Supplementary Fig. 9e) especially at the edges of the monoculture (Fig. 5b and Supplementary Fig. 9c). The CrkII\textsuperscript{P218F} mutant presented a more aggressive migratory phenotype and greater cell-spreading tendency when compared to the cells overexpressing CrkII (Fig. 5b, c and Supplementary Fig. 9c). Treatment of either CrkII or CrkP\textsuperscript{218F} cells with CsA resulted in significant decrease in migration (Fig. 5b,c and Supplementary Fig. 9e), further establishing CrkII Tyr221 phosphorylation as a key event in cell migration. This role is consistent with the observation that expression of Crk\textsuperscript{Y221F} nullifies Ephrin-B2 induced inhibition of cell migration in breast cancer cells\textsuperscript{39}. Taken together, these results suggest that the CypA-CrkII interaction delays phosphorylation of CrkII at Tyr221 thereby decreasing Crk-mediated cell motility.

Finally we followed the distribution of paxillin using fluorescent microscopy. Paxillin is integral to the formation of focal contacts and is recruited to nascent focal adhesions. It is also important for the focal adhesion turnover, which is essential for cell migration\textsuperscript{42,43}. We reasoned that the increased migratory behavior we observed in the CrkII\textsuperscript{P218F} cells would translate to an increase in the number of focal contacts that are observed. Using GFP-Paxillin as a marker for focal adhesion we stably reconstituted CrkII or CrkII\textsuperscript{P218F} plasmid in Crk(−/−) MEFs and observed focal adhesion formation at early time points after seeding (15 min, 30 min and 60 min) (Supplementary Fig. 10a). At the 60-min time point the CrkII\textsuperscript{P218F} cell line had significantly more focal contacts (by ~50%) than the cells expressing CrkII (Fig. 5d and Supplementary Fig. 10b). Treatment of CsA caused a marked decrease on focal contacts in the cells expressing CrkII or CrkII\textsuperscript{P218F}. Interestingly, we also observed that by 60 mins, a greater number of the CrkII\textsuperscript{P218F} cells were spread out with robust paxillin signal from the leading edges (Supplementary Fig. 10a). In the Crk knockout cells paxillin was diffused throughout the cell with few focal contacts. Taken together, these results support the coimmunoprecipitation data that the increased affinity for CrkII to CypA has a direct effect on the complex formation between CrkII and paxillin and as a consequence the distribution of paxillin in the cell.

**Abl-mediated CrkL phosphorylation is not affected by CypA**

Crk-like (CrkL), encoded by a distinct gene, exhibits high sequence identity (~60%) to CrkII (Supplementary Fig. 11a). Similarly to CrkII, CrkL also features a highly conserved Tyr residue (Tyr207) that becomes phosphorylated by Abl\textsuperscript{44}. CrkL is a preferred substrate of the Bcr-Abl kinase and its tyrosine-phosphorylation is used as a diagnostic tool for Philadelphia-positive leukemia\textsuperscript{45}. Notably, the residue preceding Tyr207 in CrkL is Ala, as opposed to
Pro in CrkII (Fig. 1c), and thus, CrkL does not undergo cis–trans isomerization as shown by NMR analysis. (Supplementary Fig. 11b). As a result, CypA is not recruited to this region in CrkL and no binding was detected between the two proteins (Supplementary Fig. 11c). Kinase assay performed with Abl kinase and CrkL as a substrate also show no inhibition of phosphorylation in the presence of CypA (Supplementary Fig. 11d). This provides another important difference between CrkII and CrkL in addition to the structural differences identified previously. Thus, whereas CypA regulates CrkII phosphorylation by Abl, it does not regulate CrkL phosphorylation. This may be one of the factors that contribute to the preferential phosphorylation of CrkL by Abl.

**DISCUSSION**

PPIases have been implicated in an ever-growing number of cellular processes. Their ubiquitous presence in cells suggests that their full repertoire remains to be explored. Recent studies have identified CypA as one of the proteins upregulated in many human cancers. However, the molecular mechanisms underlying CypA action and the signaling pathways through which it exerts its function are poorly understood. Here we show that CypA binds to the CrkII adaptor protein and inhibits a key phosphorylation process that shuts down signaling by CrkII. As a result, CypA augments CrkII-mediated signaling giving rise to enhanced cell motility and invasion (Fig. 6).

Integrin-mediated cell adhesion and migration are important processes both in normal cell activities as well as in pathological conditions, such as tumorigenesis and metastasis. Tyrosine phosphorylation of the paxillin or p130CAS scaffold proteins, which are important players in the formation of focal adhesions, creates binding sites for the SH2 domain of Crk proteins. Crk proteins bind to Rho-GTPase exchange factors DOCK180, SOS1 and C3G via their SH3N domain enabling efficient localized activation of small GTPases, such as Rac1 or Rap1 at the cell membrane. These events induce cell migration through actin cytoskeleton remodeling, pseudopodia extension and focal adhesion turnover. Because the Crk family of proteins form a bridge between the focal adhesion proteins and the GTPases, dysregulation of Crk signaling either by mutation or expression levels negatively regulates cell migration. Phosphorylation of CrkII Tyr221 by Abl or EGFR kinases is a key event that induces intra-molecular binding of pTyr221 to the SH2 domain giving rise to an autoinhibited form of CrkII. In this form, CrkII cannot associate with paxillin or p130CAS scaffolding proteins causing the signaling cascade to be interrupted and cell movement to decrease (Fig. 6).

Interestingly, it has been shown that the SH3N domain of CrkII, which binds to Rho-GTPase exchange factors, is not accessible for binding in the phosphorylated form of CrkII. Thus, suppression of Tyr221 phosphorylation not only extends the lifetime of the paxillin-CrkII complex but it also allows CrkII to continue mediating signaling through its SH3N domain.

The Pro residue preceding Tyr221 is conserved in all CrkII proteins (Fig. 1c). Using NMR spectroscopy we demonstrate that Pro220 undergoes cis–trans isomerization and creates a binding site for CypA. The structure of the CypA–CrkII complex shows that the CrkII
region encompassing Tyr221 is buried in the binding cleft of CypA and thus cannot be accessed by kinases (Fig. 2a). As a result, CypA binding to CrkII suppresses Tyr221 phosphorylation and CrkII remains in the active form that mediates cell motility (Fig. 6).

In principle, CypA can function as a binding partner (in stoichiometric amounts) or as an enzyme (in catalytic, sub-stoichiometric amounts). Pro220 is located in an unstructured region of the 50-residue long linker that tethers the two SH3 domains. The NMR data show that Pro220 isomerization has a very local and limited effect since only about seven residues exhibit duplicated peaks. Furthermore, the NMR data showed that both the cis and trans conformers of the Pro220–Tyr221 prolyl bond in CrkII are phosphorylated by Abl and both the cis and trans conformers of pTyr221 bind to the SH2 domain (Supplementary Fig. 12). Thus, there seems to be no conformer-specific binding selectivity. Moreover, the effect of catalytic amounts of CypA on inhibiting CrkII phosphorylation is negligible further corroborating that CypA exerts its inhibitory effect as a binding partner and not as an enzyme. Because the affinity of CrkII for CypA is relative low (K$_d$ ~15 µM), the effect of CypA on suppressing Tyr221 phosphorylation becomes significant and noticeable only when the proteins are present in significant amounts. This is especially the case in cancer cells wherein both CypA and CrkII are vastly upregulated.

Binding of CypA to CrkII is expected to have a similar, albeit likely diminished, phenotype to the CrkII$^{Y221F}$ mutant, which cannot be phosphorylated. Indeed, motility of cells expressing CrkII$^{Y221F}$ is not inhibited downstream of extracellular stimuli that induce Tyr221 phosphorylation such as Ephrin B2$^3$. Moreover, dephosphorylation of pCrkII by phosphatase PTB1B enhances motility of cells expressing CrkII$^4$. CrkI, an alternatively spliced Crk isoform that consists only of the first 204 residues of CrkII, and thus lacks the Tyr221 phosphorylation site, remains always in the active form (Fig. 6) and is associated with a particularly aggressive phenotype$^1$. The lack in CrkI of the segment encompassing Pro220 explains why CrkI does not associate with CypA$^5$.

In contrast to CrkII, CrkL showed no interaction with CypA. CrkL is missing the critical heterogeneous Pro residue next to the Tyr phosphorylation site (Tyr207 in human CrkL; Fig. 1c) and is unable to recruit CypA. This highlights a potential mechanism by which CypA can influence signaling by discriminating between two similar proteins and may account, to some extent, for the differential behavior of the CrkII and CrkL proteins.

**ONLINE METHODS**

**Protein preparation and purification**

The following constructs were used for the experiments: human CrkII (1–304), human CrkII linker-SH3$^C$ (193–304), chicken CrkII (1–305), chicken CrkII linker-SH3$^C$ (193–305), human CrkL (1–303), human CrkL linker-SH3$^C$ (191–304) and human CypA (1–165). Crk and CrkL proteins were cloned into pet42a using NcoI and XhoI restriction sites. A TEV protease cleavage site was introduced between the histidine tag and the protein. Isotopically labeled samples for NMR studies were prepared by growing the cells in M9 minimal media supplemented with 1 g l$^{-1}$ of $^{15}$NH$_4$Cl and 2 g l$^{-1}$ of $^{13}$C$_6$ glucose. All CrkII and CypA constructs were grown at 37°C, and protein synthesis was induced by the addition of 0.25
mM IPTG at OD$_{600}$ ~0.4. Cells were lysed by sonication and the cytosolic fraction was separated by centrifugation at 50,000 × g. The lysate was loaded onto Ni-NTA agarose resin (GE) equilibrated with Tris buffer and 1 M NaCl, pH 8. Elution was performed with 400 mM imidazole and after TEV cleavage the sample was concentrated and applied to a Superdex 75 size – exclusion column (GE). For NMR studies the samples were dialyzed in NMR buffer (50mM KPi, pH 6.5, 150 mM NaCl and 1 mM β-mercaptoethanol) and concentrated using Amicon cell units (Millipore). All samples were monomeric in solution at concentrations used for the NMR studies (0.3mM–0.5mM) as confirmed by multi angle light scattering (MALS).

**Kinase assay**

Kinase assay was carried out in the NMR buffer with either recombinant Crk or CrkL supplemented with 5 mM MgCl$_2$ and 5 mM ATP. The kinase reaction was started by addition of enzymatic quantity of recombinant Abl kinase and the reaction was stopped using 2× Laemmli buffer. The samples were analyzed using western blotting with the indicated antibody. Quantitation was performed using imageJ.

**FRET assay and imaging**

CFP-CypA and YFP-CrkII were constructed by inserting the respective genes into pcDNA3.1 already containing either the CFP or YFP genes. Each plasmid was sequenced for verification. Fluorescence imaging to measure FRET was conducted 24 hrs post-transfection using the Nikon TE2000 Fluorescence Microscope equipped with a motorized barrier filter wheel. Images were acquired using the CFP channel (excitation: 430 nm, emission: 470/24nm), YFP channel (excitation: 500 nm, emission: 535/30nm) and CFP-YFP channel (excitation 430 nm, emission: 535/30nm). Cells transfected with only CFP-CypA and only YFP-CrkII were used to calibrate the FRET measurements for spectral bleed-through. The NIS Elements Imaging Software was used to measure the FRET efficiency using the sensitized emission method.

**Cell culture and plasmid transfection**

HeLa cell lines were seeded in 24-well polystyrene-coated culture plates in Dulbecco’s modified eagle medium (DMEM). The medium was supplemented with 10% fetal bovine serum (FBS), 100 U/ml penicillin, 100 µg/ml streptomycin, and 2 mM L-glutamine. Cells were seeded to achieve 70% confluence at the time of transfection. The cells are transfected with the fusion plasmids using the commercial available transfection reagent X-tremeGENE (Roche). Briefly, equal amounts of the CFP-CypA and YFP-CrkII plasmids were mixed together and complexed with the recommended volume of XtremeGENE for 20 min. The complexes were then added dropwise to the cells in serum-free OptiMEM media. After 6 hr the cells were washed two times with fresh cell culture medium. For the inhibition studies, CsA was dissolved in DMSO and added to the cell culture medium to achieve a final concentration of 25 µM CsA and 0.1% DMSO. CsA was added 30 min prior to conducting fluorescence imaging.
Isothermal Titration Calorimetry

All calorimetric titrations were performed on an iTC200 microcalorimeter (GE). Protein samples were extensively dialyzed against the ITC buffer containing 50 mM KPi, pH 6.8, 150 mM NaCl and 1 mM TCEP at 8 °C. The sample cell was typically filled with ~40 µM of CypA and the injection syringe with ~400 µM of Crk proteins or peptides. Crk peptide solutions were prepared by dissolving the peptide in the flow through of the last exchange buffer. Each titration typically consisted of a preliminary injection followed by 7–8 subsequent injections. Data for the preliminary injection, which are affected by diffusion of the solution from and into the injection syringe during the initial equilibration period, were discarded. The data were analyzed with Origin 7.0.

Reagents

The following antibodies were used: anti-pY221 CrkII (Sigma SAB4503813), anti-CrkII (Sigma SAB1405658), anti-actin (Sigma A2066), anti-Flag (Sigma F1804) and anti-Abl (Sigma SAB4501043). Protein G agarose beads were purchased from Santa Cruz Biotechnology, HRP-conjugated goat anti-mouse and goat anti-rabbit were purchased from Jackson Immunologicals and Type I Collagen was purchased from BD. Anti-Crk (RF-51) anti-sera was prepared as described previously. Transfections were performed with the XtremeGene HP reagent according to the manufacturers’ protocol (Roche). Human EGF was purchased from Life Technologies.

Cell culture and growth factor stimulation

MCF10A, MDA-MB-468, MCF7, MDA-MB-231, T47D and 293T cells were obtained from ATCC. Cells were cultured in DMEM with 10% Fetal Bovine Serum (FBS) and 1% Penicillin/Streptomycin. Cells were starved in DMEM containing 0.5% FBS over-night and then stimulated with EGF at 100 ng ml⁻¹. Where indicated, cells were pre-treated before stimulation with CsA (Cell Signaling Technology).

Generation of stable cell lines

For stable over-expression, EYFP or Crk proteins in the pMSCV vector backbone, pCL-Eco and pMD2.G (encoding the VSV glycoprotein) were transiently transfected into the BOSC23 packaging cell line. Viral supernatants were collected 48 hours post-transfection and transferred to MDA-MB-468 cells in the presence of Polybrene (8 µg ml⁻¹). Stable polyclonal populations were selected in the presence of puromycin (1 µg ml⁻¹). For stable knockdowns, scrambled shRNA, CypA shRNA or Crk shRNA in the pLKO.1 vector backbone, psPAX2 and pMD2.G were co-transfected into HEK 293T cells. Viral supernatants were collected 60 hours post-transfection, virus was concentrated by ultracentrifugation and transferred to MDA-MB-468 cells. Stable polyclonal populations were selected in the presence of puromycin (1 µg/ml).

Cell migration assays using xCelligence

CIM plates (Acea Biosciences) were coated on the underside with 10 µg/ml collagen. 40,000 viable cells were seeded into the top chamber while EGF was added only to the bottom chamber. In this system, the bottom of the transwell insert is lined with electrodes that send
a pulse of current periodically to record impedance. As cells move through the membrane and make contact with the electrode, they contribute to impedance which is used to generate a cell index (y-axis) that is directly proportional to the number of migrating cells. Data are plotted as Delta cell index versus time where delta cell index refers to cell index after background subtraction from each well.

**Wound healing migration assay**

Cell migration for the four MDA-MB-468 cell lines (control EYFP, CrkII\textsuperscript{P218F}, CrkII and knock-down CrkII) was evaluated by a scratched wound healing assay. All four cell lines were seeded in 6-well polystyrene-coated culture plates in Dulbecco’s modified eagle medium (DMEM). The medium was supplemented with 10% fetal bovine serum (FBS), 100 U ml\textsuperscript{−1} penicillin, 100 µg ml\textsuperscript{−1} streptomycin, and 2 mM L-glutamine. Cells were grown to confluence and then wounded using a pipette tip. Two wounds were made for each sample, and all were imaged at 0 hr, 6 hr and 24 hrs at the same position to compare the degree of migration. For the inhibition studies, CsA was dissolved in DMSO and added to the cell culture medium to achieve a final concentration of 25 µM CsA and 0.1% DMSO. CsA was added immediately after wounding.

**Western blotting and immunoprecipitation**

Samples were analyzed by SDS PAGE and then transferred to a PVDF membrane (Millipore). After blocking, the membrane was incubated with primary and secondary antibodies according to the manufacturer’s protocols. Antibody dilutions used were Anti-CypA: 1:1000, anti-CrkII: 1:5000, anti-CrkII-pY221: 1:2000, anti-Flag: 1:1000, anti-Abl: 1:1000, anti-actin: 1:1000. Binding was detected by chemiluminescence and then quantified using ImageJ software. Immunoprecipitation was performed as described previously\textsuperscript{52}.

**Image acquisition and processing**

Images were taken on an inverted wide field Zeiss microscope equipped with a Plan-Apo 63x oil immersion objective with NA 1.4 using Axiovision 4.8.2 software (Carl Zeiss, Inc). For each field binning was set to 2x2 using a CoolSNAP HQ2 (Photometrics) camera and 60 z-stacks with a step size of 0.240 µm. Deconvolution was iterative with autolinear normalization for all images. Focal contacts were quantified using ImageJ. (Rasband, W.S)

**Statistics**

Bar graphs represent mean ±SEM. Data were analyzed using Excel. Statistical significance was tested using two-tailed t-tests.

**NMR spectroscopy**

NMR experiments were conducted at 25 or 32 °C on Bruker 700 MHz spectrometer equipped with a TCI cryoprobe. Spectra were processed with NMRPipe\textsuperscript{53} and visualized using Sparky. Complete backbone and sidechain assignment for the [U-\textsuperscript{13}C,\textsuperscript{15}N]-CypA was achieved using standard 3D triple-resonance, 3D-\textsuperscript{1}H-\textsuperscript{13}C -TOCSY, and COSY experiments. Assignments were extended, verified and distance restraints were extracted from 3D \textsuperscript{15}N,\textsuperscript{13}C-edited NOESY spectra acquired with 120 ms mixing time. The \textsuperscript{1}H assignment
for the CypA-bound CrkII peptide and intermolecular protein-protein NOE contacts were obtained using a double half-filtered (3D F1-$^{13}$C/$^{15}$N-filtered, F3-$^{13}$C-edited) NOESY experiment. $^{1}$H-$^{15}$N ZZ-heteronuclear exchange NMR spectra of $^{15}$N-CrkII in the absence and presence of catalytic amounts (2–10%) of CypA were performed as detailed before$^{25}$.

**Structure calculation and refinement**

Structures for CypA–CrkII complex were calculated using CYANA 3.0$^{54}$. Initial restraints for the backbone torsion angles $\Phi$ and $\Psi$ were obtained from chemical shifts using TALOS-N$^{55}$. Resonance assignments, NOESY peak lists from all NOESY experiments, and TALOS-derived dihedral angles were used as CYANA input. Structural refinement was performed using restrained molecular dynamics in explicit water with CNS 1.3$^{54}$. The ensemble of 20 lowest energy structures and restraints for CypA–CrkII complex have been deposited in PDB (PDB ID 2MS4). Chemical shifts were deposited in BMRB (BMRB ID 25104). The summary of NMR restraints and structure refinement statistics is presented in Supplementary Table 1. HADDOCK 2.1$^{56}$ was used to model the CypA-CrkIIP218F structure.

**Supplementary Material**

Refer to Web version on PubMed Central for supplementary material.

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Figure 1. Binding between CypA and CrkII
(a) Domain organization of human CrkII. Tyr221 and the Pro220 residues are highlighted.
(b) Phosphorylation of Tyr221 in CrkII results in intramolecular interaction between
pTyr221 and the SH2 domain of CrkII. (c) Sequence alignment of the region Pro216-Pro231
of CrkII from different species. (d) Overlaid $^1$H-$^{15}$N HSQC NMR spectra of labeled CrkII
(blue) and equimolar unlabeled CypA (orange). (e) Chemical shift mapping of the effect of
CypA binding to CrkII. A stretch of ~6 residues flanking Pro220 and encompassing Tyr221
is most affected. CrkII is shown in a ribbon representation (PDB ID 2EYZ). Tyr221 and
Pro220 side chains are shown as sticks. (f) Chemical shift mapping of the effect of CrkII
binding to CypA. CypA residues strongly affected by CrkII binding are colored green. CypA
is shown as a solvent-accessible surface.
Figure 2. Specific interaction between CypA and CrkII in vitro and in vivo
(a) Structure of CypA in complex with the CrkII peptide consisting of residues Ser216-Pro225. CypA is shown in a solvent-accessible surface representation whereas CrkII is shown as a ball-and-stick model. CrkII residues Glu217-Gln223 are shown (CrkII residues Pro216, Pro224 and Pro225 do not interact with CypA). The backbone of the CrkII region interacting with CypA is buried inside the catalytic cleft of CypA. Two views related by a 90° rotation about the x axis are shown. CrkII is in the trans conformation. (b) Interacting interface between CypA and CrkII. CypA is shown in light blue ribbon and the side chains
of the residues interacting with CrkII are shown in ball-and-stick (green). CrkII is shown in yellow ball-and-stick. The magenta lines denote inter-molecular polar contacts (hydrogen bonds and/or salt bridges). (c) FRET analysis of HeLa cells transiently co-transfected with CrkII-YFP and CypA-CFP. The FRET efficiency is color coded from 0–100% on the left bar. Images were acquired on live cells after confirming comparable expression of both proteins. An average FRET efficiency of 70% is observed between the two fluorophores. Scale bar: 10 µm. (d) FRET analysis of HeLa cells transiently co-transfected with CrkII-YFP and CypA-CFP and treated with 25 µM of CsA before imaging. The average FRET efficiency drops to ~15% indicating that the complex is disrupted in the cell. Scale bar: 10 µm.
Figure 3. CypA attenuates CrkII Tyr221 phosphorylation in vitro and in vivo

(a) Effect of CypA on CrkII Tyr221 phosphorylation by Abl kinase was investigated using an in vitro kinase assay (left). The intensity of the pTyr221 bands were quantified and plotted (right). Full gels are shown in Supplementary Fig. 13. Without CypA (orange) and with CypA (green)

(b) Effect of CypA on CrkII Tyr221 phosphorylation in 293T cells was investigated by stably expressing scrambled shRNA or CypA shRNA and transiently transfected with Abl kinase and CrkII plasmids for 48 hours. Lysates were prepared and analyzed by western blotting (top) with the antibodies indicated. Intensity of bands were quantified and plotted (bottom) Color code is as displayed in the figure panel. Full gels are shown in Supplementary Fig. 13.

(c) Effect of CrkII Tyr221 phosphorylation on CrkII P218F in 293T cells transfected with the plasmids or treatment indicated for 48 hours followed by immunoprecipitation of CrkII from lysate. Quantification of CrkII Tyr221 phosphorylation normalized to total CrkII (bottom) was performed after western blotting (top). Full gels are shown in Supplementary Fig. 13. For all graphs data represents the results from 3 independent experiments. (±SEM)
Figure 4. CypA attenuates CrkII Tyr221 phosphorylation cancer cell line MDA-MB-468

(a) Effect of CypA on CrkII Tyr221 phosphorylation was measured upon EGF stimulation in MDA-MB-468 cells. Lysates were analyzed by western blotting with the antibodies indicated. The intensity of the bands (top) was quantified and pTyr221 CrkII phosphorylation levels were normalized to overall CrkII expression and plotted (bottom). Full gels are shown in Supplementary Fig. 13. Scrambled shRNA (blue) and CypA-KD (magenta).

(b) Effect of CsA on CrkII Tyr221 phosphorylation was investigated in MDA-MB-468 cells pretreated with DMSO or CsA and stimulated with EGF for the times...
indicated, and lysates analyzed for CrkII Tyr221 phosphorylation. The intensity of the bands (top) was quantified and pY221 CrkII phosphorylation levels were normalized to overall CrkII expression and plotted (bottom). Full gels are shown in Supplementary Fig. 13. Without CsA (blue) and with CsA (magenta).

(c) Effect of CypA on paxillin-CrkII complex formation was investigated in MDA-MB-468 cells stably expressing scrambled shRNA or CypA shRNA transiently transfected with flag-paxillin and stimulated with EGF for 1 and 2 minutes. Immunoprecipitation of CrkII from lysates was analyzed with anti-flag antibody. The intensity of the bands (top) representing paxillin were quantified, normalized against the paxillin level in untreated cell line expressing CypA shRNA and plotted (bottom). Scrambled shRNA (blue) and CypA-KD (magenta). The two bands of paxillin represent differentially phosphorylated forms of paxillin. All experiments were analyzed by western blotting. Full gels are shown in Supplementary Fig. 13. For all graphs data represents the results from 3 independent experiments. (±SEM)
Figure 5. Effect of the CypA–CrkII complex on cell migration
(a) MDA-MB-468 cells stably expressing EYFP or CrkII were serum starved and motility was analyzed towards EGF (25 ng ml\(^{-1}\)) in real-time using xCelligence technology (Roche). Delta Cell Index is plotted on the y-axis as an average of duplicate samples versus time on the x-axis (b) Wound healing assay of wild type CrkII (left) and CrkII\(^{P218F}\) (right) in the presence and absence of CsA. Cells were cultured to near confluence, scratched using a pipette and imaged after 24 hours. Scale bars: 500µm (c) Summary of the measurement of cell migration from the wound healing assay in panel b and Supplementary Fig. 9c was quantified for each condition indicated. Data represents mean of 3 independent experiments ±SEM. Two-tailed t-test, **** p≤0.001. (d) Focal contacts based on signal from GFP-Paxillin were quantified for each cell line and treatment using Image J and plotted. Light grey without CsA, dark grey with CsA. (Supplementary Fig. 10a, b). Data represents a mean of 6 cells ±SEM (n=6; two-tailed t-test **p and, §§≤0.05. § and **** ≤0.001.

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Figure 6. Effect of CypA binding to CrkII in integrin signaling
(i) Integrin activation elicits paxillin/p130CAS phosphorylation by tyrosine kinases (TK), and, as a result, Crk proteins are recruited. (ii) Abl-induced phosphorylation of CrkII forces its dissociation from paxillin/p130CAS and thus results in signaling suppression. (iii) Binding of CypA to Pro220 sterically inhibits CrkII Tyr221 phosphorylation and CrkII remains in the active form. (iv) GEFs (for example, DOCK180 and C3G) associate with CrkII via its SH3N domain, giving rise to efficient localized activation of small GTPases at the membrane. (v) CrkI is a short alternatively spliced form of Crk that lacks the phosphorylation site and thus forms the active state constitutively.