PZR Coordinates Shp2 Noonan and LEOPARD Syndrome Signaling in Zebrafish and Mice

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Noonan syndrome (NS) is an autosomal dominant disorder caused by activating mutations in the \textit{PTPN11} gene encoding Shp2, which manifests in congenital heart disease, short stature, and facial dysmorphia. The complexity of Shp2 signaling is exemplified by the observation that LEOPARD syndrome (LS) patients possess inactivating \textit{PTPN11} mutations yet exhibit similar symptoms to NS. Here, we identify “protein zero-related” (PZR), a transmembrane glycoprotein that interfaces with the extracellular matrix to promote cell migration, as a major hyper-tyrosyl-phosphorylated protein in mouse and zebrafish models of NS and LS. PZR hyper-tyrosyl phosphorylation is facilitated in a phosphatase-independent manner by enhanced Src recruitment to NS and LS Shp2. In zebrafish, PZR overexpression recapitulated NS and LS phenotypes. PZR was required for zebrafish gastrulation in a manner dependent upon PZR tyrosyl phosphorylation. Hence, we identify PZR as an NS and LS target. Enhanced PZR-mediated membrane recruitment of Shp2 serves as a common mechanism to direct overlapping pathophysiological characteristics of these \textit{PTPN11} mutations.

The Src homology 2 (SH2) domain-containing protein tyrosine phosphatase 2 (Shp2) is an ubiquitously expressed non-transmembrane protein tyrosine phosphatase (PTP) encoded by the \textit{PTPN11} gene (1). Shp2 comprises two SH2 domains, a PTP domain and a C-terminal tail that contains a proline-rich region flanked by two tyrosine phosphorylation sites. The SH2 domains of Shp2 serve to direct protein-protein interactions with its upstream phosphotyrosyl target in a sequence-specific context, and the SH2 domains participate in maintaining a closed conformation by establishing contacts with the PTP domain (2). The phosphatase domain of Shp2 catalyzes substrate-specific dephosphorylation and is tightly regulated. In the basal state, the catalytic activity of Shp2 is suppressed by Shp2 assuming a “closed” conformation maintained through the NH2-terminal SH2 (N-SH2) domain forming surface contacts with the PTP domain, which occludes the catalytic site. SH2 domain engagement disrupts this “closed” conformation, leading to allosteric modulation and exposure of the catalytic site in an “open” state that is now substrate accessible (2). Engagement of the SH2 domains provides an elegant molecular switch mechanism to activate the phosphatase that allows for targeted substrate dephosphorylation within a precise microenvironment. As such, the repertoire of Shp2-binding proteins plays a major role in dictating Shp2 signaling complexity and specificity. Consistent with this, interference with Shp2’s SH2 domains abrogates both physiological and pathophysiological signaling (3). Thus, the integrity of the SH2 domains of Shp2 plays a critical role in its ability to signal.

A wealth of biochemical, cellular, and genetic evidence in mice, \textit{Drosophila melanogaster}, and zebrafish has established that Shp2 plays a positive signaling role downstream of numerous receptor tyrosine kinases, cytokine receptors, G protein-coupled receptors, and integrins (1). The repertoire of Shp2-interacting proteins in combination with a wide variety of substrates (4) endows this phosphatase with the capability of engaging a complex array of signaling molecules and pathways, such as the Ras/mitogen-activated protein kinase (MAPK), phosphoinositol 3-kinase (PI3K)/Akt, JAK/STAT, RhoA/Rac, and Src family kinases (SFKs). Shp2 has been shown to promote a number of cellular functions, including cell growth, cell survival, and differentiation, hence affecting organismal functions, such as development in mice (5, 6), \textit{Drosophila} (7), and zebrafish (8). The highly conserved nature of Shp2 signaling in lower organisms has provided powerful model systems for the genetic dissection of Shp2’s function in development. The realization that Shp2 plays an important role in human development arose through the discovery that germ line mutations in the \textit{PTPN11} gene coding for Shp2 cause two similar types of autosomal dominant disorders (9, 10). More than 50% of Noonan syndrome (NS) (NIM163950) cases are caused by mutations in \textit{PTPN11} that result in increased Shp2 catalysis. In contrast, as much as 90% of LEOPARD syndrome (LS) (NIM151100) mutations are caused by inactivating \textit{PTPN11} mutations that give rise to impaired Shp2 activity.

NS and LS are autosomal dominant disorders that manifest in...
short stature, ocular hypertelorism, cardiac defects, webbed neck, and increased incidence of mental retardation (11–13). NS and LS comprise part of a larger series of disorders that have thus far been characterized by mutations in components of the Ras/MAPK pathway and are collectively referred to as “RASopathies” (14). As such, NS and LS exhibit overlapping clinical presentations, such as proportionate short stature, facial dysmorphism, musculoskeletal anomalies, and, to a lesser extent, congenital heart disease. Despite the similarities between NS and LS, these “RASopathies” are caused by opposite effects on Shp2 activity (9, 15–18). These observations suggest that NS and LS may utilize common signaling components in the pathogenesis of these developmental disorders. Interestingly, one common feature of both NS and LS mutants is that both forms of Shp2 exhibit an increased propensity to adopt an “open” conformation, leading to increased affinity to upstream target binding through the SH2 domains and target substrates (2, 8, 19, 20). Hence, the “open” conformation of NS and LS disease mutants may underlie the molecular pathogenesis of these syndromes rather than altered substrate dephosphorylation by the PTP domain. Despite these highly plausible explanations, it remains unclear how activating and inactivating mutants of Shp2 liberate overlapping disease outcomes. Moreover, the target or targets for both NS and LS that must presumably be shared in order for them to elicit overlapping clinical outcomes are unknown.

PZR ("protein zero related") is a transmembrane glycoprotein with an extracellular immunoglobulin (Ig) domain and an intracellular domain containing two immunoreceptor tyrosine-based inhibitory motifs (ITIMs) (21–23). The ITIMs, when tyrosyl phosphorylated, serve as Shp2 binding sites (21). We had identified previously that NS-associated Shp2 mutants induce PZR hyper-tyrosyl phosphorylation and recruit increased Shp2 (25). Moreover, in Ptpn11Y61G/+ mouse embryos, PZR is hyper-tyrosyl phosphorylated, demonstrating that it is aberrantly regulated in NS mice during development. Interestingly, PZR is upregulated during blastocyst formation (24). Moreover, in mouse embryo fibroblasts derived from NS mice, adhesion-dependent extracellular signal-regulated kinase 1/2 (ERK1/2) activation is abrogated upon loss of PZR expression (25).

To gain further insight into how NS and LS might liberate similar phenotypes, we performed an unbiased phosphotyrosine proteomics screen in a mouse model of NS and a zebrafish model of NS and LS. We identified PZR as a major hyper-tyrosyl-phosphorylated protein in both NS and LS in mice and zebrafish. The role of PZR in development remains unclear, and the pathophysiological significance of PZR-Shp2 interactions in NS and LS is also unknown. NS and LS are linked to defects in gastrulation cell movements, and Shp2 is essential for gastrulation movements in mice and zebrafish (11, 26). We show that PZR-Shp2 interactions are required for gastrulation in zebrafish. These results define a functional role for PZR in an intact organism. Moreover, they suggest that PZR is a common target that contributes to altered development in cases of both NS and LS.

MATERIALS AND METHODS

DNA constructs and cloning. Ptpn11 was cloned previously (11). Mpl1 was cloned by nested PCR from zebrafish embryo cDNA (from bud stage to 48 h postfertilization [hpf]) with Phusion DNA polymerase (Finnzymes, Vantaa, Finland) using fwPZRout (5′-GGCGGTATCTTACGAAATCTAAA-3′), rwPZRout (5′-CTCAGGTTGGCGAGATG-3′), fwPZRstart (5′-GGCGGTATCTGAAATACGGGT-3′), and rwPZRstop (5′-GGACCGCTTCTAGTTCGATG-3′). The PCR product was cloned into pc52+ using BamHI and EcoRI. PZRM ITM Y236F, Y258F, and Y236F Y258F mutants were made using site-directed mutagenesis with the following primers (boldfacing indicates introduced mutations in the primer sequence): Fw_Y= F236_PZR (5′-GGACCGCTTCTAGTTCGATG-3′), Rw_Y= F236_PZR (5′-GGACCGCTTCTAGTTCGATG-3′), Y258F and Y236F Y258F mutants were made using site-directed mutagenesis with the following primers (boldfacing indicates introduced mutations in the primer sequence): Fw_Y= F236_PZR (5′-GGACCGCTTCTAGTTCGATG-3′), Rw_Y= F236_PZR (5′-GGACCGCTTCTAGTTCGATG-3′), Y258F and Y236F Y258F mutants were made using site-directed mutagenesis with the following primers (boldfacing indicates introduced mutations in the primer sequence): Fw_Y= F236_PZR (5′-GGACCGCTTCTAGTTCGATG-3′), Rw_Y= F236_PZR (5′-GGACCGCTTCTAGTTCGATG-3′), Y258F and Y236F Y258F mutants were made using site-directed mutagenesis with the following primers (boldfacing indicates introduced mutations in the primer sequence): Fw_Y= F236_PZR (5′-GGACCGCTTCTAGTTCGATG-3′), Rw_Y= F236_PZR (5′-GGACCGCTTCTAGTTCGATG-3′), Y258F and Y236F Y258F mutants were made using site-directed mutagenesis with the following primers (boldfacing indicates introduced mutations in the primer sequence): Fw_Y= F236_PZR (5′-GGACCGCTTCTAGTTCGATG-3′), Rw_Y= F236_PZR (5′-GGACCGCTTCTAGTTCGATG-3′), Y258F and Y236F Y258F mutants were made using site-directed mutagenesis with the following primers (boldfacing indicates introduced mutations in the primer sequence): Fw_Y= F236_PZR (5′-GGACCGCTTCTAGTTCGATG-3′), Rw_Y= F236_PZR (5′-GGACCGCTTCTAGTTCGATG-3′).

Antibodies, chemicals, cell lines, and expression reagents. Rabbit monoclonal phospho-PTP121 (Y241) and rabbit monoclonal phospho-PTP121 (Y263) antibodies were generated in collaboration with Cell Signaling. Mouse monoclonal Src antibody, rabbit polyclonal Src antibody, rabbit polyclonal phospho-ERK1/2 (T202 Y204), mouse monoclonal ERK1/2 antibody, rabbit polyclonal phospho-Akt (S473) antibodies and mouse monoclonal Akt antibodies were purchased from Cell Signaling. Rabbit polyclonal Shp2 antibodies and rabbit polyclonal ERK1/2 antibody were purchased from Santa Cruz Biotechnology. Mouse monoclonal Shp2 antibody was purchased from BD Bioscience. Rabbit polyclonal PZR (105-6) was generously provided by Z. J. Zhao. Src family kinase inhibitors PP2 and SU6656 were purchased from Calbiochem. HEK-293, NIH 3T3, SYF (Src−/−, Yes−/−, Fyn−/−) mouse embryonic fibroblasts (MEFs), and Src−/− (Src-overexpressing SYF) cells were purchased from ATCC and grown in growth medium (Dulbecco’s modified Eagle’s medium [DMEM] supplemented with 1% penicillin-streptomycin and 10% fetal bovine serum) in a 5% CO2 incubator at 37°C. Replication-deficient adenoviral (Ad) constructs harboring wild-type Shp2 (Ad-Shp2 WT), the E76A gain-of-function Shp2 mutant (Ad-Shp2 E76A), and green fluorescent protein (GFP) (Ad-GFP) were prepared as previously described (25, 28). NIH 3T3 and SYF cells were infected with adenovirus at a dosage of 50 multiplicities of infection (MOI). The pCMV vectors containing Src WT and the K295R/Y527F dominant-negative Src mutant (SrcK295R/Y527F) have been described in previously (29). The pIRE5-GFP plasmids encoding the Shp2 WT, gain-of-function/Noonan syndrome mutants of Shp2 (Shp2Y279C and Shp2T468M), and LEOPARD syndrome mutants of Shp2 (Shp2Y279C, and Shp2T468M) have been described previously (15, 25). DNA transfection into HEK-293 and SYF cells was performed using Lipofectamine 2000 according to the manufacturer’s protocol. Mouse antiphosphotyrosine antibody 4G10 (05-321) was from Merck Millipore, rabbit anti-GFP (TAP016) was from Acris, and mouse anti-HA.11 clone 16B12 was from Covance.

Animal handling. Ptpn11D61G/+ mice were provided by Benjamin Neel (University of Toronto, Toronto, Ontario, Canada) and were genotyped as described previously (30). Briefly, Ptpn11D61G/+ male mice were crossed with WT C57BL/6 × SV129 female mice, and their offspring were genotyped by PCR and digestion with AgeI for the D61G allele. Animal handling was approved by the Yale University Institutional Animal Care and Use Committee. Tissues from Ptpn11Y529F/+ mice were provided by Maria Kontaridis (Harvard University, Boston, MA).

In situ hybridization. Zebrafish were kept and embryos were staged as described by Westerfield (31). In situ hybridizations were performed as described by Thiese et al. (32). The nkb2p, chd, gsc, nlta, d1ch3, hgg, kayak20, and myod probes were described previously (11, 33). Antisense and sense Mpl21 probes, respectively, were synthesized by T7 and SP6 mRNA polymerase (Promega) from pCPS2+ PZR.

MO RNA and injections. Zebrafish were kept and embryos were injected at the 1-cell stage. PZR morpholinol oligonucleotide (MO) 5′-CAG AGACCCCTTACCTGTGCTGGGCAGCGC-3′, Shp2 MO 5′-GGTGGGACCA CTTGGGGATGTCAT-3′, nacre MO 5′-CATGGTCAACTAATGTTGCT-3′, and standard p53 MO were from Genetools (Philomath,
OR). 5′-capped sense synthetic mRNA was synthesized with SP6 or T7 mRNA polymerase using the Message Machine kit from Ambion. The amount of RNA that was injected at the 1-cell stage was optimized for each synthetic RNA. MOs were titrated down to give a specific phenotype without inducing increased apoptosis as coinjection of 1 nmol p33 MO did not attenuate the phenotype. Likewise, mRNA was titrated down to either induce a phenotype or rescue an MO-induced phenotype.

**Immunoprecipitation and immunoblotting.** Cells were lysed on ice in lysis buffer (20 mM Tris-HCl, 150 mM NaCl, 1 mM CaCl2, 1 mM MgCl2, 1% Nonidet P-40, 1 mM Na3VO4, 10 mM NaF, 1 mM benzamidine, 1 mM phenylmethylsulfonyl fluoride [PMSF], 1 μM phosphatase A, 5 μg/ml aprotinin, 5 μg/ml leupeptin). Tissues from Noonan syndrome (NS) and LEOPARD syndrome (LS) mice were lysed by homogenization with lysis buffer. Cell or tissue lysates were incubated at 4°C for 30 min and clarified by centrifugation at 14,000 rpm for 10 min. The protein concentration was determined using the bicinchoninic acid (BCA) reagent according to the manufacturer’s instructions (Pierce). For immunoprecipitations, 500 μg of lysate was incubated with 1 μg of the indicated antibodies at 4°C overnight. Immune complexes were collected on either protein A- or protein G-Sepharose beads for 4 h at 4°C, washed three times with the same lysis buffer, and then heated to 95°C in sample buffer for 5 min. Total cell or tissue lysates and immune complexes were subjected to SDS-PAGE and immunoblotting. The sites of antibody binding were visualized using enhanced chemiluminescence detection or the Odyssey imaging system.

Transfected cells were lysed in radioimmunoprecipitation assay (RIPA) buffer (150 mM NaCl, 20 mM Tris [pH 8.0], 10 mM Na3HPO4, 5 mM EDTA, 1 mM Na-orthovanadate, 10% glycerol, 1% Nonidet P-40, 1% sodium deoxycholate, 0.1% SDS, 2.5 mM sodium fluoride, 5 μM β-glycerophosphate, 1 μg/ml aprotinin, 1 μg/ml leupeptin) for 15 min on ice at 1 ml per 10-cm-diameter dish. Samples were centrifuged for 30 min at 14,000 rpm at 4°C. The samples were precleared using protein A-agarose beads for 1 h, and an aliquot of whole-cell lysate was immunoprecipitated with 1 μg antibody against HA for approximately 1 h on ice. Samples were precipitated using 25 μl protein A-agarose slurry for approximately 1 h at 4°C. Samples were boiled in 2× Laemmli buffer, and SDS-PAGE was performed.

For zebrafish, bud-stage embryos were snap-frozen in liquid nitrogen and stored at −80°C. Subsequently embryos were ground in cell lysis buffer containing 50 mM Tris (pH 7.5), 150 mM NaCl, 1 mM EDTA, 1 mM sodium orthovanadate, 1% Nonidet P-40, 0.1% sodium deoxycholate, 2.5 mM sodium fluoride, 5 mM β-glycerophosphate, 1 μg/ml aprotinin, and 1 μg/ml leupeptin. Samples were then boiled in Laemmli sample buffer containing 4% SDS and β-mercaptoethanol. Proteins equivalent to 4 embryos per lane were subjected to SDS-PAGE and blotted onto a polyvinylidene difluoride (PVDF) membrane. After blotting, membranes were stained using Coomassie blue to verify equal loading. 293T cells were maintained in Dulbecco’s modified Eagle’s medium (DMEM) or Ham’s F-12 medium containing 10% penicillin-streptomycin and 7.5% fetal calf serum (FCS), respectively. For serine starvation experiments, nonessential amino acids were added. 293T and COS7 cells were passed twice a week at 1:20 and 1:7 before transfection. Transfection of zebrafish PZFR was performed using polyethyleneimine (PEI) from Sigma with 20 μg DNA. After transfection overnight, medium was replaced with serum and harvested 40 h after transfection.

**MS analysis.** The PhosphoScan method was performed as previously described (34–37). Wild-type and Shp2 mutant (Noonan syndrome) mouse hearts were homogenized, sonicated, and centrifuged to remove cellular debris. Total protein for each tissue was normalized using the ProteinPlus Coomassie reagent (Pierce), and proteins were reduced, alkylated, and digested overnight using trypsin (Worthington). The resulting peptides were separated from nonpeptide material by solid-phase extraction with Sep-Pak classic C18 cartridges (Waters). Lyophilized peptides were redissolved, and phosphopeptides were enriched by immunoaffinity purification using py-100 phosphotyrosine antibody (9411; Cell Signaling Technology). Peptides were eluted with 0.15% trifluoroacetic acid (TFA) and concentrated with C18 spin tips immediately prior to liquid chromatography-mass spectrometry (LC-MS) analysis.

Duplicate injections of each sample were run to generate analytical replicates and increase the number of tandem MS (MS/MS) identifications from each sample. Peptides were loaded directly onto a 10-cm by 75-μm PicoFrit capillary column packed with Magic C18 AQ reversed-phase resin. The column was developed with a 45-min linear gradient of acetonitrile in 0.125% formic acid delivered at 280 nL/min. Tandem mass spectra were collected with an LTQ-Orbitrap XL mass spectrometer running XCalibur using a 10 method, a dynamic exclusion repeat count of 1, and a repeat duration of 30 s. MS spectra were collected in the Orbitrap component of the mass spectrometer, and MS/MS spectra were collected in the LTQ portion. MS/MS spectra were processed using SEQUEST and the Core platform (Gygi Lab, Harvard University) (38–41). Searches were performed against the mouse NCBI database, with reverse decay databases included for all searches to estimate false-positive rates. Peptide assignments were obtained using a 0.98-precision cutoff in the linear discriminant analysis module of Core. Cysteine carboxymidomethylation was specified as a static modification, and methionine oxidation and serine, threonine, and tyrosine phosphorylation were allowed. Results were further narrowed using mass accuracy (±5-ppm) filters and the presence of a phosphotyrosine in the peptide. Label-free quantitation was performed using Progenesis v4.1 (Nonlinear Dynamics). Peptide abundance data were manually reviewed in Progenesis for all peptides with at least a 2.0-fold change to ensure accuracy of results.

**Global phosphotyrosine differential proteomic analysis in the zebrafish heart.** Heart tissue was lysed in urea buffer (20 mM HEPES [pH 8.0], 9 M urea, 1 mM sodium vanadate, 2.5 mM sodium pyrophosphate, 1 mM β-glycerophosphate). The lysate was sonicated and clarified by centrifugation. Tissue lysate was reduced by dithiothreitol (DTT) and alkylated with iodoacetamide. Samples were then diluted 4 times with 20 mM HEPES to reduce the urea concentration to 2 M and digested by trypsin overnight at room temperature with gentle shaking. Phosphotyrosine-containing peptides were isolated using immobilized phosphotyrosine-specific antibody PY100 (PhosphoScan kit; Cell Signaling Technology). Following immunopurification, phosphotyrosine-containing peptides were analyzed by LC-MS/MS as described previously (37).

**RESULTS**

PZR is a major hyper-tyrosyl-phosphorylated protein in NS. Since NS and LS exhibit overlapping clinical outcomes, we set out to identify target proteins that are affected by both syndromes. We hypothesized that targets similarly altered in their levels of tyrosyl phosphorylation in both NS and LS likely represent molecules of convergence in these diseases. Given that NS and LS mutants exhibit opposite catalytic activities, we reasoned that a phosphatase-independent mechanism likely represents a common pathway for these two syndromes. Therefore, we focused on the identification of hyper-tyrosyl-phosphorylated proteins in a mouse model of NS rather than hypo-tyrosyl-phosphorylated ones, since the latter likely represents phosphatase-dependent Shp2 substrates.

We performed a phosphotyrosyl proteomics screen in mice harboring a knock-in of an NS-associated Shp2 mutant (30). Mice containing an Asp61 to Gly61 (D61G) mutation within the NH2-terminal SH2 domain of Shp2 (Shp2<sup>D61G</sup>) generate an NS mutation found in humans (9). Heterozygous progeny of these mice (Ptpn11<sup>D61G/+</sup>) recapitulate several NS features observed in humans, such as cardiovascular and skeletal abnormalities (30). Since both NS and LS patients exhibit significant clinical defects in the heart, we focused on this tissue. We performed a global survey of the levels of tyrosyl-phosphorylated proteins in the hearts of...
Proteomic analysis of differentially tyrosyl-phosphorylated proteins in hearts of Ptpn11D61G/+ mice. (A) Classification of hypo- and hyper-tyrosyl-phosphorylated proteins in the hearts of Ptpn11D61G/+ mice. (B) Log2-transformed values for the ratio of each phosphotyrosine-containing peptide in wild-type and Ptpn11D61G/+ mouse hearts. (C) Heat map of differentially tyrosyl-phosphorylated peptides (the site of phosphorylation is identified by MS in parentheses). (D) Extracted ion chromatogram and peptide sequence of PZR-containing tyrosine 242 (upper panels) and tyrosine 264 (lower panels) by differential proteomics. (E) Amino acid sequences of the PZR C terminus in different vertebrates. Consensus sequences for ITIM (S/I/V/LXYXXI/V/L) are indicated in boldface, and tyrosine residues are marked red with the appropriate amino acid numbering. Sequences are shown for the PZR C terminus from Homo sapiens, Mus musculus, Rattus norvegicus, Bos taurus, Canis lupus familiaris, Danio rerio, Gallus gallus.

5-day-old wild-type and Ptpn11D61G/+ mice were used in an immunofluorescence phosphoproteomics approach (34). The relative abundance of tyrosyl-phosphorylated proteins was assessed by quantifying the frequency of phosphopeptides derived from wild-type and Ptpn11D61G/+ mice (34). We identified an approximately equal repertoire of differentially hypo- and hyper-tyrosyl-phosphorylated peptides in the hearts of Ptpn11D61G/+ mice (Fig. 1A and B). The limited number of differentially tyrosyl-phosphorylated peptides suggests that the NS mutant induces a discrete influence on global tyrosyl phosphorylation levels in the heart (Fig. 1B). The highest induced hyper-tyrosyl-phosphorylated peptide represented Y264, which is present in the transmembrane glycoprotein “protein zero-related” (PZR) and was induced by 6.7-fold com-
pared to that in the wild-type controls (Fig. 1C and D). The sequence surrounding Y264 was identified to be part of the immunoreceptor tyrosine inhibitory motif (ITIM) of PZR (23). PZR contains two ITIMs, both of which are highly conserved among vertebrates and when phosphorylated bind directly to the SH2 domains of Shp2 (23). Notably, the other tyrosine-containing ITIM on PZR represented by Y242 was also detected to be hyper-tyrosyl phosphorylated by 2.2-fold (Fig. 1C and D). For clarity, we will use the human amino acid numbering of PZR (Fig. 1E). (Mouse Y242 and Y264 correspond to human Y241 and Y263, respectively.) Previously, we found that PZR was hyper-tyrosyl phosphorylated during embryogenesis in Ptpn11D61G/H11001 mice (25). These results identify Y242 and Y264 as sites of PZR hyper-tyrosyl phosphorylation in this mouse model of NS.

Zebrafish have been shown to recapitulate both NS and LS phenotypes (11). Therefore, phosphotyrosyl proteomics screens were performed in developing (26 days postfertilization) embryos to identify differentially tyrosyl-phosphorylated peptides between zebrafish embryos expressing either NS or LS mutations compared with controls. When injected with the Shp2D61G NS mutant, peptides representing the ITIMs of zebrafish PZR containing Y241 and Y263 were identified to be among the highest-induced phosphotyrosyl-containing peptides (7.7- and 4.7-fold increases for Y241 and Y263, respectively, compared with controls). In addition, LS-injected embryos also were found to contain PZR hyper-tyrosyl-phosphorylated peptides at both Y241 and Y263, representing increases of 3.4- and 3.1-fold relative to wild-type controls. These results identify PZR as a major hyper-tyrosyl-phosphorylated target in zebrafish embryos expressing either NS or LS mutations.

**PZR is expressed ubiquitously and functions in the Shp2 pathway in zebrafish.** Our data demonstrate that PZR tyrosyl...
Phosphorylation is aberrantly regulated in NS mouse embryos and postdevelopmentally in NS hearts (25) (Fig. 1) and during development of NS/LS-expressing zebrafish embryos. These results suggest that altered PZR tyrosyl phosphorylation may play a role in NS and LS pathogenesis. To further examine the role of PZR during zebrafish development, we determined the expression of mpzl1, the gene encoding zebrafish PZR. Zebrafish PZR was cloned and was used to generate an mpzl1-specific probe. In situ hybridization of mpzl1 expression was apparent from the 8-cell stage onwards, indicating maternal expression. mpzl1 expression was ubiquitous until the bud stage, and at later stages, mpzl1 expression appeared to be enhanced anteriorly (Fig. 2). The specificity of the antisense mpzl1 probe was confirmed by the lack of signal of the sense control probe (Fig. 2). Thus, PZR is maternally contributed and expressed at embryonic stages during and after zebrafish gastrulation.

**FIG 4** PZR knockdown is specific and does not affect cell specification. (A) Zebrafish embryos were injected at the 1-cell stage with 2 ng PZR MO alone or coinjected with p53 morpholino oligonucleotide (MO) and photographed at 2 dpf. (B) Zebrafish embryos were injected at the 1-cell stage with 2 ng PZR MO1, PZR MO2, or control MO and fixed at the shield stage (bmp2b, chd, gsc, and ntl), 70% epiboly (cyc), and bud stage (six3) and probed for cell fate markers by in situ hybridization.

PZR knockdown phenocopies NS and LS expression in zebrafish embryos. Next, PZR was knocked down in order to investigate the role of PZR in zebrafish signaling. The specificity of the PZR MO to block PZR splicing was confirmed (Fig. 3A to C). Knockdown of PZR but not a control MO induced a phenotype characterized by a reduced body axis length (4.9% ± 1.1% reduction compared with the control) and heart edema, and this phenotype was confirmed using a second nonoverlapping MO (Fig. 3D and E). Interestingly, similar phenotypes are observed upon Shp2 knockdown in zebrafish (42). Coinjection of a moderate dose of PZR mRNA (25 pg) rescued the PZR MO-induced defects, whereas a higher dose of PZR mRNA (100 pg) contributed and expressed at embryonic stages during and after zebrafish gastrulation.

**FIG 5** Genetic interaction between Shp2 and PZR. Embryos were injected at the 1-cell stage with Shp2 MO (0.5 pg) or PZR MO (0.5 pg), a combination thereof, or Shp2 MO (1.0 pg) and PZR MO (2.0 pg). Embryos were imaged at 4 dpf. The stacked histogram in the left panel indicates quantitation of the phenotypes shown in representative photographs in the right panel.
failed to rescue (Fig. 3D). To quantify the craniofacial defects in PZR knockdowns, embryos were fixed at 4 days postfertilization (dpf), and the cartilage was stained with alcian blue. The cartilage was greatly malformed, and the angle of the ceratothyroid displayed a significant increase, supporting the notion that PZR knockdown induced craniofacial defects, reminiscent of Shp2 knockdown (11) (Fig. 3F).

The use of knockdown approaches by itself may lead to non-specific activation of p53 and subsequent apoptosis, thereby giving rise to developmental defects (43). To rule out this possibility, we performed a co-knockdown of p53 and PZR, which did not attenuate the phenotype (Fig. 4A). We subjected PZR knockdown embryos to in situ hybridization using a panel of cell fate markers at the shield stage (gsc, bmp2b, chd, and mtl), 70% epiboly (cyc), and bud stage (six3). All markers were expressed, indicating that cell specification was not affected (Fig. 4B). Hence, PZR knockdown induced defects similar to those observed in both NS and LS Shp2-expressing embryos.

To investigate the genetic interaction between Shp2 and PZR, we partially knocked down Shp2 and PZR separately and together. If PZR is essential for Shp2 signaling, partial loss of both Shp2 and PZR should induce developmental defects that are more severe than those induced by loss of each alone (42). Knocking down Shp2 and PZR at suboptimal levels did not induce developmental defects (Fig. 6A). C2C12 cells were cotransfected with empty vector or activated glutathione S-transferase (GST)–Shp2E76A and either empty vector (vector), wild-type human PZR (WT), or PZR mutated at tyrosine 242 (Y241F), tyrosine 264 (Y263F), or both (2YF). Cell lysates were immunoblotted with anti-pPZR (Y241 or Y263), -PZR, or -Shp2 antibodies. (B) HEK-293 cells were cotransfected with empty vector (Vec) or activated Shp2E76A and either empty vector (vector), wild-type zebrafish PZR (WT), or PZR mutated at tyrosine 236 (Y241F), tyrosine 258 (Y263F), or both (2YF). Cell lysates were immunoblotted with anti-pPZR(Y241 or Y263), -PZR, or -Shp2 antibodies. ERK1/2 was used as a loading control. (C) HUVECs were infected with adenoviruses expressing either GFP as a control, wild-type Shp2, or Shp2E76A. Cell lysates were immunoblotted with anti-pPZR(Y241 or Y263), anti-total PZR, and anti-Shp2 antibodies. (D) HEK-293 cells were transiently transfected with empty vector, wild-type Shp2 (WT), or the indicated Shp2 mutants (activated Shp2, E76A; NS mutant, N308D; or LS mutants, Y279C and T468M). Cell lysates were immunoblotted with anti-pPZR(Y241 or Y263), -PZR, and -Shp2 antibodies. ERK1/2 was used as a loading control. (E) HEK-293T cells were transfected with HA-tagged zebrafish PZR with empty vector, wild-type Shp2, or Shp2E76A, or Shp2E76A. Cell lysates were immunoprecipitated with anti-HA antibodies, and immune complexes were immunoblotted with anti-Shp2 and anti-HA antibodies. Whole-cell lysates (WCL) were blotted with anti-pPZR(Y241 and Y263), -Shp2, and -HA antibodies.
defects (Fig. 5). However, partial knockdown of PZR and Shp2 together resulted in severe developmental defects at 4 dpf (Fig. 5). The embryos were scored in three categories (wild type [WT], phenotype, and severe), and it was evident that coknockdown of Shp2 and PZR greatly enhanced the percentage of embryos displaying a severe defect phenotype (Fig. 5). PZR knockdown also induced wider-set eyes, reminiscent of the phenotype that we and others reported in Shp2 knockdowns (11, 44). Together, these results are consistent with the notion that PZR and Shp2 function in a conserved signaling pathway in zebrafish.

NS- and LS-associated Shp2 mutants induce PZR Y241 and Y263 hyper-tyrosyl phosphorylation. We investigated the characteristics of PZR hyper-tyrosyl phosphorylation in further detail. To accomplish this, antibodies against tyrosyl-phosphorylated PZR at Y241 (pY241-PZR) and Y263 (pY263-PZR) were generated. When cells were cotransfected with PZR, along with an activating mutant of Shp2 (Shp2E76A) that induces PZR hyper-tyrosyl phosphorylation (25), both anti-pY241 and anti-pY263-PZR antibodies detected tyrosyl-phosphorylated PZR upon coexpression of Shp2E76A (Fig. 6A). Similarly, mutation of Y241 to F241 in PZR also resulted in the failure of both anti-pY241 and anti-pY263-PZR antibodies to detect tyrosyl-phosphorylated PZR.
PZR (Fig. 6A). The explanation for these results is that Y241 is required for Y263 phosphorylation, and conversely, Y263 is required for Y241 phosphorylation. Similar observations have been made by others (21, 45). Importantly, mutation of Y241 and Y263 to nonphosphorylatable residues abolished the immunoreactivity of both phospho-specific PZR antibodies (Fig. 6A). These antibodies also recognized Y241 and Y263 in zebrafish PZR (Fig. 6B). We tested whether activating Shp2 mutants induced either Y241 PZR or Y263 PZR tyrosyl phosphorylation in other cell types. To this end, the wild type and activating Shp2E76A mutants were introduced into human umbilical vein endothelial cells (HUVECs). These results indicated that pY241-PZR and pY263-PZR antibodies recognized endogenous tyrosyl-phosphorylated PZR at Y241 and Y263, respectively (Fig. 6C). The LS mutants from humans and zebrafish when expressed in 293 cells were capable of inducing PZR hyper-tyrosyl phosphorylation at both Y241 and Y263 (Fig. 6D and E). As reported previously (25), increased zebrafish PZR tyrosyl phosphorylation correlated with enhanced PZR/Shp2 binding (Fig. 6E). Hence, both NS and LS mutations in humans and zebrafish enhance PZR tyrosyl phosphorylation.

**PZR hyper-tyrosyl phosphorylation in NS and LS mice.** We determined whether PZR was hyper-tyrosyl phosphorylated in NS mice. Hearts isolated from 8-week-old wild-type and Ptpn11D61G/+ mice showed a significant increase in pY263-PZR tyrosyl phosphorylation compared with wild-type controls (Fig. 7A). In addition, PZR hyper-tyrosyl phosphorylation was observed in the cortex of Ptpn11D61G/+ mice (Fig. 7B), as well as in the kidney, liver, and spleen of these mice (Fig. 8A to C).

![Figure 8](http://mcb.asm.org/)  
**FIG 8** PZR tyrosyl phosphorylation in Ptpn11D61G/+ mice. Liver (A), kidney (B), and spleen (C) were isolated from 5-week-old wild-type and Ptpn11D61G/+ mice. Tissue lysates were immunoblotted with anti-pPZR(Y263) and -total PZR antibodies. Densitometric analysis of the phosphorylation levels of tyrosine 263 in PZR was performed, and the results represent the means ± SEM from 5 mice per genotype. **, P < 0.01; ***, P < 0.001 (WT versus NS).

Given that in cultured cells LS induces PZR hyper-tyrosyl phosphorylation (Fig. 6D and E), we tested whether PZR is hyper-tyrosyl phosphorylated using Ptpn11Y279C/+ knock-in mutant mice, which represented a human LS mutant (46). Hearts from Ptpn11Y279C/+ mice were isolated, and although to a much lesser extent than Ptpn11D61G/+ mice, PZR was hyper-tyrosyl phosphorylated on Y263 (Fig. 7C). In addition, PZR was robustly hyper-tyrosyl phosphorylated in the cortex of Ptpn11Y279C/+ mice (Fig. 7D). In agreement with these data, we also observed that when zebrafish embryos were injected to express either the NS or LS mutants, PZR was also hyper-tyrosyl phosphorylated at Y241 and Y263 (Fig. 7E). No apparent differences were observed in the basal levels of Shp2, phosho-ERK1/2, or phospho-Akt between NS or LS mice (Fig. 9A to D). Collectively, these results show that both NS and LS mutants induce PZR hyper-tyrosyl phosphorylation within the ITIM Shp2 binding motif of PZR in mice and zebrafish.

**NS and LS mutants induce PZR hyper-tyrosyl phosphorylation by enhancing the Src/Shp2/PZR complex.** In order to investigate the mechanisms underlying how both NS and LS mutants induce PZR hyper-tyrosyl phosphorylation, we examined the dependency of NS and LS mutants to induce PZR hyper-tyrosyl phosphorylation through the SFKs. ITIMs serve as potent Src family kinase (SFK) substrates, and SFKs phosphorylate ITIMs within PZR (47). Pharmacological inhibition of the SFKs by SU6656 impaired both NS- and LS-induced PZR hyper-tyrosyl phosphorylation at pY241 and pY263 (Fig. 10A). Similar results were also observed with the activated E76A mutant with both SU6656 and the broader-specificity tyrosine kinase inhibitor PP2 (Fig. 10B).
Next, we determined the ability of the activating Shp2 mutant to induce PZR hyper-tyrosyl phosphorylation in fibroblasts lacking the expression of Src/Yes/Fyn (SYF cells). Here, we found that an activating Shp2 mutant failed to induce PZR hyper-tyrosyl phosphorylation at Y241 and Y263 (Fig. 11A). Consistent with this, a kinase-dead mutant of c-Src blocked the ability of the activated Shp2 mutant to induce PZR hyper-tyrosyl phosphorylation at these specific sites (Fig. 11B). Both human and zebrafish PZR served as c-Src substrates in vitro (Fig. 11C and D), and coexpression of a constitutively activated mutant of c-Src promoted tyrosyl phosphorylation of zebrafish PZR in cells (Fig. 11E). Hence, c-Src phosphorylates human and zebrafish PZR at the Shp2 binding sites, demonstrating that the SFKs likely mediate PZR hyper-tyrosyl phosphorylation.

It has been previously suggested that the "open" conformations of both NS and LS mutants exhibit properties that increase their propensity to interact with other signaling proteins. Based upon our observations that SFKs may play a role, we tested whether the SFKs are NS and/or LS binding targets. To test this, c-Src was immunoprecipitated from cells expressing either wild-type Shp2 or NS or LS mutants, and these immune complexes were immunoblotted to detect the presence of Shp2. We found that in both NS- and LS-expressing cells, the amount of NS/LS Shp2 present in c-Src complexes was significantly greater than that in the wild-type Shp2-expressing cells (Fig. 12A).

To determine whether Src is present in PZR complexes and whether this is enhanced by the "open" conformation of mutated Shp2, we immunoprecipitated Src and assessed the extent of PZR interactions. We found that when Src was immunoprecipitated, PZR and Shp2 were present in the complex (Fig. 12B). However, Src failed to coimmunoprecipitate with the form of PZR that was mutated in both of the ITIMs but still exhibited enhanced Shp2 binding to the "open" conformation Shp2 mutants (Fig. 12B). Wild-type PZR, as expected, was found to exist in a complex with Shp2, and this was further enhanced when either the activating or LS Shp2-containing mutation was expressed (Fig. 12B). In contrast, double-ITIM-containing mutant Shp2 failed to form a complex (Fig. 12B). These results suggest that the "open" conformation of Shp2 mutants exhibits an increased propensity to bind c-Src and that the likely explanation for PZR-induced hyper-tyrosyl phosphorylation is causally related to the ability of Src to increase its proximity to PZR through Shp2 binding.
Convergence and extension defects induced by NS and LS are recapitulated by PZR in a phosphotyrosyl-dependent manner. The similarities between the Shp2 (42) and PZR morphant phenotypes (Fig. 3 and 5) indicate common defects during development. Previous results have shown that Shp2 knockdown and NS and LS PTPN11 mutant overexpression lead to convergence and extension (C/E) cell movement defects during gastrulation, resulting in shorter embryos with craniofacial defects and cardiac edema (11). To test whether PZR knockdown also induces C/E defects, embryos were injected with PZR, Shp2, or control MO, or LS mRNA as a positive control. Embryos were fixed at the 8- to 10-somite stage, subjected to in situ hybridization, and stained for krox20 and myoD markers for rhombomeres 3 and 5 and the somites, respectively. Loss of PZR led to shorter and wider embryos than controls at the same stage, and these phenotypes were similar to Shp2 knockdown and LS-Shp2-injected embryos (Fig. 13A). These results indicate that PZR is essential for C/E cell movements, analogous to those of both NS and LS mutants during zebrafish gastrulation.

To assess whether PZR tyrosyl phosphorylation sites were important for C/E phenotypes, we overexpressed wild-type PZR and PZR tyrosyl phosphorylation site-deficient mutants and performed in situ hybridization at the 8- to 10-somite stage using krox20 and myoD markers. Overexpression of wild-type PZR resulted in shorter and wider embryos (Fig. 13B). When mRNA encoding double ITIM tyrosine-to-phenylalanine mutants (Y241F and Y263F [2YF]) and as a control mRNA encoding GFP were injected into zebrafish, the C/E phenotype was inhibited (Fig. 13B). As expected, PZR tyrosyl phosphorylation and Shp2 binding to zebrafish PZR are inhibited in the double ITIM Y241F/Y263F mutant (Fig. 13C). These data indicate that tyrosyl phosphorylation of the ITIMs are required for PZR signaling for the promotion of the C/E phenotype during early development in zebrafish.

**DISCUSSION**

Here, we identified using nonbiased proteomic approaches PZR as a major hyper-tyrosyl-phosphorylated protein in both NS and LS models of mice and zebrafish. Remarkably, disruption of PZR function resulted in phenotypes similar to that observed in NS or LS mutant-expressing zebrafish. Mechanistically, NS and LS mutants induce PZR tyrosyl phosphorylation in an Src-dependent manner, and these mutants acquire an enhanced Src interaction compared with wild-type Shp2. PZR is required for zebrafish gastrulation, and tyrosyl phosphorylation is necessary to promote convergence and extension movements. Thus, NS and LS mutants promiscuously promote PZR hyper-tyrosyl phosphorylation, leading to dysfunctional gastrulation. Collectively, these results suggest that PZR is a novel target for both NS- and LS-associated Shp2 mutants.

The extracellular matrix serves to integrate dynamically processes that govern cell fate, tissue maintenance, and homeostasis to outcomes that dictate organ architecture (48). Growth factor receptors, chemokines, and G-protein-coupled receptors are either directly or indirectly influenced by the extracellular matrix (ECM) to control cell proliferation, differentiation, and cell survival. The “RASopathies” represent mutated components of the SOS/Ras/Raf/Erk pathway, and there has been much focus on the role this cascade plays in NS and LS as it relates to cell proliferation and cell survival (10). PZR has been shown to function in the control of cell adhesion and migration (24, 25, 45, 47, 49). Therefore, our findings imply that aberrant cell migration and adhesion likely play important roles in the pathogenesis of both NS and LS. The development of PZR mutant mice will be necessary to test these conclusions more comprehensively.

Germ line mutations in Ptpn11 that result in either increased or decreased protein tyrosine phosphatase activity lead to developmental defects in zebrafish, mice, and humans. Our findings that PZR is a target for Ptpn11 mutations that exhibit either increased

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**FIG 10** Effect of Src family kinases on PZR Y241 and Y263 phosphorylation. (A) HEK-293 cells were transiently transfected with the indicated Shp2 mutants and treated with either dimethyl sulfoxide (DMSO) as a control or 5 µM SU6656. (B) NIH 3T3 cells were infected with the adenoviruses expressing either GFP as a control or a constitutively active Shp2E76A, in the presence of DMSO, PP2, or SU6656 at the indicated concentration. Cell lysates were immunoblotted with anti-Shp2, pSrc(Y416), Src, pPZR(Y241 or Y263), and total PZR antibodies. ERK1/2 was used as a loading control.
or decreased catalytic activity argue for a central role for PZR in ECM-mediated signaling for both NS and LS mutations. Using a zebrafish model system, we found that PZR is required for development. PZR RNA was apparent at early stages in zebrafish embryos, suggesting a maternal contribution, and either knockdown or overexpression of PZR in zebrafish resulted in a gastrulation defect. PZR knockdown in zebrafish resulted in similar defects to Shp2 and NS- and LS-like phenotypes. In addition, partial knockdown of PZR and Shp2 together resulted in even more severe developmental defects. Together, this genetically links PZR to a pathway in which both NS and LS mutations participate. Importantly, we found that the loss of PZR in zebrafish led to convergence and extension defects, which is consistent with deterioration in the fidelity of cell migration. Intriguingly, overexpression of PZR induced gastrulation defects similar to those of both overexpression and reduced expression of Shp2, reported previously.

**FIG 11** Src family kinases mediate NS/LS-induced PZR hyper-tyrosyl phosphorylation and increase NS/LS Src binding. (A) SYF cells (Src−/− Fyn−/− Yes−/− MEFs) and Src−/− cells (SYF cells expressing wild-type Src) were infected with adenoviruses expressing either GFP or Shp2E76A. (B) SYF cells were transiently transfected with wild-type c-Src or kinase-dead c-SrcK295R/Y527F (KR/YF) and infected with adenoviruses expressing either GFP, wild-type Shp2, or Shp2E76A. Cell lysates were immunoblotted with anti-Shp2, pSrc (Y416), pPZR (Y241 or Y263), PZR, and Src antibodies. ERK1/2 was used as a loading control. (C and D) HEK-293 cells were transfected with Flag-tagged human PZR (C) or HA-tagged zebrafish PZR (D). The cell lysates were immunoprecipitated with indicated antibody. The immunoprecipitates were subjected to in vitro Src kinase assay with Src recombinant protein. The reaction products were immunoblotted with anti-pPZR (Y241 or Y263) antibodies. (E) HEK-293 cells were cotransfected with constitutively active Src mutant and either HA-tagged wild-type zebrafish PZR (WT), PZR mutated at tyrosine 236 (Y241F), tyrosine 258 (Y263F), or both (2YF). Cell lysates were immunoblotted with anti-pPZR (Y241 or Y263) or -HA antibodies.
These results suggest a tight balance and interplay exist between PZR and Shp2 signaling, consistent with the biochemical evidence demonstrating that Shp2 binds directly to PZR within its ITIMs (11). Collectively, these observations argue that PZR-mediated tyrosyl phosphorylation in zebrafish and mice serves as a mechanism to promote NS and LS signaling.

What are the consequences of increased PZR tyrosyl phosphorylation? PZR acts as an adhesion-responsive glycoprotein where, upon engagement to the extracellular matrix, it becomes rapidly tyrosyl phosphorylated within its ITIMs. Upon tyrosyl phosphorylation, Shp2 is recruited to PZR, and it has been shown that this event is responsible for promoting the promigratory and cell adhesive effects in a variety of cell types (24, 25, 45, 47, 49). Given the importance of cell migration and cell adhesion in developmental processes, such as gastrulation, it is conceivable that aberrant PZR tyrosyl phosphorylation induced by both NS and LS mutants disrupts this process. In mice, PZR is expressed during early development, and we have demonstrated that PZR is hyper-tyrosyl phosphorylated in the developing embryo of Ptpn11D61G/H11006 mice (25). These observations support the notion that altered PZR tyrosyl phosphorylation and hence downstream signaling are associated with developmental defects. The significance of PZR tyrosyl phosphorylation was demonstrated in experiments in which overexpression of the ITIM mutants of PZR failed to induce gastrulation defects. These results define PZR tyrosyl phosphorylation as critical for development.

FIG 12 Enhanced Src complex formation with NS/LS-associated Shp2 mutants. (A) HEK-293 cells were transiently transfected either with the Shp2 WT and the indicated Shp2 mutants. Cell lysates were immunoprecipitated (IP) with anti-c-Src antibodies, and immune complexes were immunoblotted (IB) with anti-Shp2 and -Src antibodies. The graph represents the means ± SEM of the densitometric analysis from three independent experiments. Statistical significance was derived using a Dunnett’s test comparing Shp2 mutants with the WT. *, P < 0.05; **, P < 0.01. (B) HEK-293 cells were cotransfected with the Shp2 WT or E76A or Y279C mutant and either empty vector (vector), WT human PZR, or the PZR 2YF mutant. Cell lysates were immunoblotted with anti-pPZR(Y241 or Y263) or -Shp2 antibodies. ERK1/2 was used as a loading control. Immune complexes were immunoblotted with anti-Src, -Shp2, and -pPZR antibodies.

FIG 13 PZR tyrosyl phosphorylation is necessary for the zebrafish convergence and extension phenotype. (A) Embryos were injected with Shp2 MO, PZR MO, or LS mRNA at the 1-cell stage and fixed at the 8- to 10-somite stage. In situ hybridization was performed using krox20 and myoD staining for rhombomeres 3 and 5 and somites, respectively. Quantification of the width of the rhombomeres (red) and the length of the first 8 somites (light blue) is shown schematically. krox20/myoD ratios are plotted compared to those of noninjected controls (NIC). Data represent means ± SEM. **, P < 0.01; *** P < 0.001. (B) Embryos were injected at the 1-cell stage with mRNA encoding GFP, wild-type (WT) PZR, or PZR with the indicated mutations. Ratios of krox20/MyoD (% of NIC) were calculated and compared to those of NIC. Data represent means ± SEM. **, P < 0.01; *** P < 0.001. (C) HEK-293 cells were cotransfected with either the HA-tagged zebrafish PZR WT or 2YF mutant and constitutively active Src. Cell lysates were immunoprecipitated with HA antibody and blotted with anti-pY, -Shp2, and -HA antibodies.
A provocative finding from our study is that these results implicate the actions of a tyrosine kinase that promotes PZR tyrosyl phosphorylation by both NS and LS mutants. The observation that PZR is hyper-tyrosyl phosphorylated in both NS and LS most likely indicates that the ability of the kinase to phosphorylate PZR occurs in an Shp2 catalytically independent manner. We had shown previously that enhanced PZR tyrosyl phosphorylation caused by NS mutants does not occur as a result of SH2 domain protection (25). Rather, the data here support an Src family kinase caused by NS mutants does not occur as a result of SH2 domain protection. The open conformation of NS and LS-induced PZR hyper-tyrosyl phosphorylation (25). In the study by Eminaga et al., we immunoprecipitated Shp2 and analyzed tyrosyl phosphorylation of Shp2-associated proteins. The Shp2-associated tyrosyl-phosphorylated complex that migrated with a molecular mass of ~40 kDa was found to contain at least PZR. It is likely that in this p40 complex, “site-specific” PZR tyrosyl phosphorylation was obscured by other comigrating tyrosyl-phosphorylated proteins. Whereas, in the experiments presented here, PZR tyrosyl phosphorylation is assessed directly using site-specific phospho antibodies.

We found that both NS and LS mutants contain increased levels of c-Src complexes compared with wild-type Shp2. In addition, we showed increased PZR in Src complexes from activating and LS-associated Shp2 mutants, and this was abolished in cells expressing the PZR tyrosyl phosphorylation-deficient mutant. The nature of this increased association with c-Src correlates with the common underlying property that both NS and LS mutants exhibit an “open” conformation (19, 50). Other groups have suggested that the “open” conformation of NS and LS may represent an important molecular mechanism for the pathogenesis of these mutant forms of Shp2 (19, 20, 24, 50). The open conformation of NS and LS mutants exposes the PTP and SH2 domains; presumably, this presents binding surfaces for Src that are otherwise inaccessible. Indeed, it has been shown previously that Shp2 binds to Src through its SH3 domains (51, 52). Our findings support this observation and extend it by providing a mechanism that reconciles the behavior of both NS and LS in their capacity to promote similar phenotypic outcomes.

The results from our proteomic screen are consistent with activation of an SFK-mediated pathway as we identified the Src substrate, focal adhesion kinase (FAK), and PECAM as hyper-tyrosyl phosphorylated in the hearts of NS mice. JNK was increased in its level of tyrosyl phosphorylation at one of its activating phospho-sites in NS hearts. Interestingly, Marin et al. showed that JNK and FAK were hyper-tyrosyl phosphorylated in LS mice, suggesting a catalytic-independent mechanism for the activation of these molecules through the effects of enhanced Src function (46). Whether the PZR/Shp2/Src pathway accounts for the entirety of either the NS- or LS-like phenotype will need to be determined. However, Jopling et al. have shown that Shp2 lies upstream of the SFKs in zebrafish development, and the Shp2/SFK pathway is required for convergence and extension cell movements (11). An integrated interpretation of our results supports a model in which we invoke the actions of enhanced Src-mediated PZR tyrosyl phosphorylation that results in altered development. An important aspect of Shp2-mediated signaling is that it is required to be appropriately localized to its upstream target. Disruption of the SH2 domains of Shp2 results in the abrogation of its ability to signal (1). Therefore, enhanced PZR tyrosyl phosphorylation results in increased recruitment of Shp2 to PZR to promote further PZR tyrosyl phosphorylation and presumably other Src substrates that are in close proximity (Fig. 14). It should also be noted that Src couples to Ras and in some cases does so through Shp2 (53). It is formally possible that both NS and LS engage Ras via this mechanism. However, the contribution of Ras signaling specifically for the manifestation of NS- or LS-related pathologies has yet to be determined. Collectively, these observations go some way toward providing an explanation as to how both the NS and LS mutants, which have enhanced and inactive PTP activities, respectively, result in overlapping phenotypes.

In summary, we have identified PZR as a proximal target of both NS and LS mutants. PZR functions to regulate cell adhesion and cell migration. Therefore, in NS and LS the augmentation of PZR’s interaction with Shp2 when it is hyper-tyrosyl phosphorylated presumably disrupts key developmental events that rely on the proper delivery of migratory cues. This mechanism occurs through enhanced binding of NS and LS mutants to Src. The implications of our findings clinically are that they suggest that the magnitude of PZR tyrosyl phosphorylation may reflect the “openness” of Ptpn11 mutants and hence the severity of NS- and LS-related diseases. As such, it would be instructive to assess the clinical genotype-phenotype relationship in the context of PZR tyrosyl phosphorylation in NS and LS patients.

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