Elevated phosphorylation of EGFR in NSCLC due to mutations in PTPRH

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Abstract

The role of EGFR in lung cancer is well described with numerous activating mutations that result in phosphorylation and tyrosine kinase inhibitors that target EGFR. While the role of the EGFR kinase in non-small cell lung cancer (NSCLC) is appreciated, control of EGFR signaling pathways through dephosphorylation by phosphatases is not as clear. Through whole genome sequencing we have uncovered conserved V483M Ptprh mutations in PyMT induced tumors. Profiling the downstream events of Ptprh mutant tumors revealed AKT activation, suggesting a key target of PTPRH was EGFR tyrosine 1197. Given the role of EGFR in lung cancer, we explored TCGA data which revealed that a subset of PTPRH mutant tumors shared gene expression profiles with EGFR mutant tumors, but that EGFR mutations and PTPRH mutations were mutually exclusive. Generation of a PTPRH knockout NSCLC cell line resulted in Y1197 phosphorylation of EGFR, and a rescue with expression of wild type PTPRH returned EGFR phosphorylation to parental line values while rescue with catalytically dead PTPRH did not. A dose response curve illustrated that two human NSCLC lines with naturally occurring PTPRH mutations responded to EGFR tyrosine kinase inhibition. Osimertinib treatment of these tumors resulted in a reduction of tumor volume relative to vehicle controls. PTPRH mutation resulted in nuclear pEGFR as seen in immunohistochemistry, suggesting that there may also be a role for EGFR as a transcriptional co-factor. Together these data suggest mutations in PTPRH in NSCLC is inhibitory to PTPRH function, resulting in aberrant EGFR activity and ultimately may result in clinically actionable alterations using existing therapies.

Author summary

One of the major genetic causes of lung cancer is EGFR activity. Traditionally this is caused by mutations in the EGFR receptor tyrosine kinase resulting in unchecked activity,
which ultimately results in lung cancer. A series of tyrosine kinase inhibitors have been
developed that treat these EGFR positive lung cancers, with remarkable efficacy. Here we
describe work from a mouse model that revealed mutations in PTPRH, a phosphatase that
we show dephosphorylates EGFR. We show that mutation or loss of wild type PTPRH
results in elevated EGFR activity. Searching for similar mutations in human lung cancer
revealed that 5% of all lung cancers had PTPRH mutations. Since activation of EGFR by
mutation and loss of PTPRH function would be redundant, we tested and demonstrated
that these events only occurred separately. Patient data revealed that a subset of PTPRH
mutant lung cancer did have elevated EGFR activity. Testing two tumor lines from
patients with naturally occurring PTPRH mutations revealed a sensitivity to EGFR inhibi-
tors. The broad implications of this work are that there are a large number of lung cancer
patients with PTPRH mutations that could potentially benefit from a revised treatment
based on sequencing. Currently the PTPRH mutations are not detected and these
patients are treated with chemotherapy as a standard of care while they could potentially
be more effectively treated with EGFR inhibitors. The ability to use EGFR inhibitors in
PTPRH mutant lung cancers is a new area for investigation and is the primary impact of
this research.

Introduction

Lung cancer results in the greatest number of U.S. cancer deaths in both men and women, and
5 year survival rates remain poor [1]. Lung cancer is classified into two major histological sub-
types, including small-cell (SC) and non-small cell lung cancer (NSCLC) with NSCLC
accounting for approximately 85% of cases. NSCLC is further delineated into Adenocarcin-
oma, Squamous cell carcinoma, and Large cell carcinoma subtypes [2]. 5-year survival rates
for localized NSCLC approach 63%, but with distant metastasis the 5 year survival rates drop
to 7% (American Cancer Society). Prognosis is complicated by a number of factors, including
EGFR mutation status [3].

A member of the ERBB family, EGFR plays a role in numerous cancers and functions
through pathways PI3K/AKT, Stat3, and Ras/Raf/Mek/Erk to increase cellular growth, prolif-
eration, and evasion of apoptotic signals. Ligand binding stimulates EGFR dimerization
through conformational shifts mediated by the extracellular domains [4, 5], resulting in a
switch to the active structure. Once in the active conformation, phosphorylation occurs on the
numerous tyrosine residues in the carboxy-terminal tail of EGFR [6–8]. Interestingly, specific
ligands are capable of inducing differential tyrosine phosphorylation and activation of various
downstream signaling pathways [9, 10]. Genetic mutations are also capable of inducing the
EGFR active state, and these mutations are common in multiple cancers. Common mutations
leading to constitutively active EGFR in NSCLC include a deletion in exon 19, and the L858R
point mutation [11, 12]. EGFR stimulation leads to transcription of numerous gene products,
from immediate early genes to secondary late response genes [13]. After signaling, EGFR is
internalized and returned to the cell surface or marked for degradation [14, 15]. Interestingly,
a body of literature also supports a role for EGFR in the nucleus. Indeed, EGFR has been
found to act as a transcriptional activator via direct binding to A/T-rich sequences (ATRS) in
the promoters of certain genes, such as cyclin D1 [16] and can act as a co-activator through
interactions with transcription factors such as STAT3 to recruit nuclear EGFR to the iNOS
promoter [17]. As a result, nuclear EGFR has prognostic value for a variety of cancers,
including breast and non-small cell lung cancer [18, 19]. Taken together, EGFR is extensively involved in cancer progression through a variety of mechanisms.

With the demonstrated importance of EGFR, it is not surprising that approximately 15% of NSCLC patients have tumors presenting with amplification or activating mutations in EGFR, with higher percentages in Asian patients [20]. 80% of these EGFR mutations are putative oncogenic drivers, with the vast majority of these mutations being either missense L858R mutations or a small deletion surrounding amino acid 750, potentially resulting in an increased dimerization ability [21]. Tyrosine kinase inhibitors are standard of care for NSCLC patients who have tumors presenting with these canonical EGFR activating mutations. First generation Tyrosine Kinase Inhibitors (TKIs), such as Erlotinib and Gefitinib, were designed to target the ATP binding domain of EGFR. These TKIs successfully enhanced progression free survival, however resistance mechanisms develop in patients, usually in the form of a T790M EGFR mutation which causes a structural shift and prevents binding of TKIs to the ATP binding domain [22]. Second generation TKIs, such as Afatinib, have also been developed to target the ATP binding domain, but do so in an irreversible covalent manner. However, these second generation TKIs still suffer from resistance mechanisms due to the T790M mutation. Third generation TKIs, such as osimertinib, circumvent this structural inhibition by binding to a nearby cysteine residue and have begun to see use as first line treatment as it increases survival rates [23]. Currently, 4th generation TKIs are being developed based on allosteric inhibition of EGFR to alleviate mutations associated with Osimertinib resistance. Taken together, while oncogenic mutations in EGFR are impactful, patients with these mutations have better 5-year survival outcomes due to a series of targeted tyrosine kinase inhibitors.

A critical component of EGFR activity is regulation of phosphorylation by phosphatases. A recent global screen for EGFR phosphatases revealed Protein Tyrosine Phosphatase Receptor Type H (PTPRH) as an EGFR phosphatase [24]. PTPRH, also known as Stomach Cancer-Associated Phosphatase 1 (SAP-1) is a member of the receptor like protein phosphatases. PTPRH has an extracellular region composed of several fibronectin domains, a transmembrane domain, and an intracellular phosphatase domain. The structure of PTPRH is largely conserved between humans and mice, with humans having eight fibronectin domains and mice having six [25]. In the phosphatase screening study, Yao et. al. found that PTPRH dephosphorylated EGFR, suggesting a specificity for tyrosine residue 1197.

While some phosphatases, such as PTEN [26, 27], have well defined tumor suppressive capabilities, many phosphatases are undefined roles in the context of cancer. PTPRH studies have been largely carried out in hepatocellular tumors. Within cancers of the liver, lower PTPRH expression is associated with poorly differentiated hepatocellular carcinomas (HCC) relative to higher levels in normal liver tissue. Furthermore, overexpression of PTPRH in HCC cell lines with low PTPRH expression drastically reduced cellular motility and growth rate in vitro, suggesting PTPRH has a tumor suppressive role within hepatocellular carcinoma. Overexpression of PTPRH has been noted in NSCLC, with correlative hypomethylation of PTPRH being suggested as the cause [28].

Here we have examined mutations that inactivate PTPRH, resulting in aberrant phosphorylation of EGFR using a combination of cell lines and mouse models. The role of the mutant PTPRH in NSCLC was not previously appreciated but this work illustrates that specific mutations in PTPRH may be clinically actionable using EGFR TKIs.

**Results**

In our prior work involving whole genome sequencing of MMTV-PyMT FVB mice, we uncovered a conserved V483M mutation in Pptrh that was associated with increased
phosphorylation of EGFR [29]. Examination of exome sequence from MMTV-PyMT in other background strains revealed a variety of other Ptprh mutations. Here we have sequenced a total of 67 PyMT mouse tumors to show the conserved V483M mutation occurs in 82% of tumors (Fig 1A). Analysis of publicly available data on other PyMT strains [30] revealed these mutations were conserved between primary tumors and matched pulmonary metastases, suggesting that the Ptprh mutation occurs early in tumor progression (Fig 1B). To directly test when mutations arose in tumor progression, we extracted DNA from 21 and 35 day old MMTV-PyMT mammary glands (Fig 1C) and tested the PCR amplified sequence for presence of mutations.

Fig 1. Ptprh Mutant Mouse Tumors have Increased Phosphorylation of AKT. Conserved metastasis and downstream regulation of EGFR pathways is seen in PyMT tumors with Ptprh mutations. A) Protein domain map of mouse PTPRH shows the location of conserved V483M mutated Ptprh within out PyMT FVB mice. B) Exome sequencing data of PyMT FVB mice from Kent Hunter’s lab shows Ptprh mutations are not conserved to one location. Furthermore, Ptprh mutation status is conserved between primary tumors and their matched metastasis. C) Wholemount of a day 21 MMTV-PyMT mammary gland with the hyperplastic growth on the left and the lymph node embedded in the fat pad on the right. The entire gland was used for DNA extraction and sequencing of PTPRH. D) Diagram shows the main tyrosine residues capable of being phosphorylated on the c-terminal tail of EGFR. While the diagram is not comprehensive, as signaling pathways are convoluted and undergo numerous feedback mechanisms, some of the main downstream pathways that have been characterized are shown. E) Western blotting of PyMT tumor lysates shows no increased phosphorylation of STAT3 or ERK in Ptprh mutant tumors as compared to WT tumors. F) Western blotting shows increased phosphorylation of AKT within Ptprh mutant tumors as compared to WT tumors.

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of Ptprh mutations. This revealed that 9 of the 13 day 21 mammary glands and 7 of 8 day 35 mammary glands had already accumulated Ptprh mutations. Importantly, these samples were only hyperplastic as tumors form on average at day 45 in this background. These data suggest that there is a strong selective pressure for Ptprh mutations prior to overt tumor formation.

Given that specificity of EGFR signaling is mediated by specific tyrosine residues (Fig 1C), we postulated that specific pathways would be activated based on which tyrosine site mutant Ptprh was failing to dephosphorylate. In Fig 1C the canonical EGFR tyrosine residue numbers are listed. Given confusion in the literature and available antibodies to EGFR due to the 24 amino acid signaling peptide that is cleaved, we have also included the alternate residue number. To investigate activation of downstream pathways, Ptprh wild type and mutant samples were assayed for STAT3, ERK and AKT activity. No alteration to STAT3 or ERK phosphorylation was noted with Ptprh mutation (Fig 1D). However, mutation of Ptprh was associated with increased phosphorylation of AKT (Fig 1E). These data as well as our prior work with a Y1197-EGFR antibody suggest a hypothesis that PTPRH dephosphorylates Y1197 on EGFR and that Ptprh mutation results in an inability to downregulate signaling, leading to an increase in the PI3K / AKT signaling axis in these tumors. We have also examined pJNK / JNK and found no alterations to pJNK levels.

To determine which human tumors contained PTPRH mutations, a pan-cancer search of the International Cancer Genome Consortium (ICGC) and The Cancer Genome Atlas (TCGA) data was completed (Fig 2A). Both datasets contained PTPRH mutations within several cancers, including a mutation prevalence of approximately 5% in non-small cell lung cancer (NSCLC). Given the incidence of EGFR mutations in NSCLC, and our data suggesting that mutations in mouse Ptprh resulted in increased EGFR activity, we hypothesized that PTPRH mutant human tumors would have increased EGFR signaling. To test this, we predicted EGFR activity in PTPRH mutant human tumors through a gene set enrichment analysis (GSEA) approach (Fig 2B). As shown in the lollipop plot, there are numerous PTPRH mutant NSCLC tumors with increased predicted EGFR pathway activity. These mutations were clustered within the fibronectin and phosphatase domains with localized hotspots of EGFR activity. Examining the relationship between EGFR and PTPRH mutation status from a collection of databases revealed that mutations in these two genes were mutually exclusive (p < 0.0001, n = 307, Fig 2C). Interestingly, the mutations noted in PTPRH were largely missense mutations (green squares). To examine the pathways that were activated in these tumors we used single sample Geneset Enrichment Analysis (ssGSEA) on EGFR mutant tumors (L858R), PTPRH mutant tumors, and tumors that were wild type for both EGFR and PTPRH mutations. Unsupervised clustering of the ssGSEA results revealed that a subset of tumors with PTPRH mutations clustered together with the EGFR mutant tumors, suggesting a similar pathway activity profile (Fig 2D). Identity of each pathway is listed (S1 Table). Importantly, we confirmed the identification of the PI3K / AKT pathway and used GSEA to compare PTPRH mutant tumors with high EGFR activity to wild type, revealing a significant enrichment of the PI3K / AKT signaling axis (Fig 2E).

Given these results were correlative in nature, we sought to directly test whether loss of PTPRH activity resulted in increased pY1197 EGFR. To test this, we used CRISPR to create knockouts of PTPRH in the H23 NSCLC cell line and flow sorted into individual clones. The generation of a knockout through insertion of an adenosine base pair at the cut site, leading to a frameshift and early stop, is shown for clone 2 (Fig 3A). Given a lack of functional antibodies for PTPRH, we instead sequenced PTPRH in the clonal cell lines. Importantly, we noted that several clones with an engineered knockout of PTPRH resulted in elevated levels of pY1197 EGFR (Fig 3B). To ensure specificity, we then rescued the knockout by transfecting a plasmid expressing wild type PTPRH. This restored endogenous levels of EGFR phosphorylation at
PTPRH mutations cause EGFR activity in lung cancer

A. PTPRH SNV Mutations - ICGC

B. ssGSEA Predicted EGFR Activity

C. Protein sequence with alterations

D. Heatmap of gene expression

E. Enrichment plots
Moreover, rescue with a catalytically dead version of PTPRH (D986A) failed to result in decreased phosphorylation of EGFR (Fig 3D).

Given that both the mouse and human computational predictions suggested a role for the PI3K / AKT pathway but not the ERK or STAT3 pathways, we examined the various PTPRH knockout lines for phosphorylation of these downstream pathways. As expected, no alterations were noted in the ERK or STAT3 pathways (S1 Fig). In contrast, AKT phosphorylation was noted but was variable between the knockout clones (Fig 3E). Given the potential for clonal effects, Y1197F mutations in EGFR were engineered in the PTPRH KO clone with elevated pAKT. Both heterozygous and homozygous Y1197F EGFR mutations were examined and a step-wise reduction in EGFR Y1197 phosphorylation was detected (Fig 3F). Within the two Y1197F clones we also noted a reduction in pAKT, consistent with the hypothesis that Y1197 was the target of PTPRH (Fig 3G).

Due to the potential for clonal effects that would impact analysis, pooled knockouts of PTPRH were generated in the H23 cell line. Tracking of Indels by Decomposition (TIDE) [31] analysis revealed a knockout efficacy of 45% (S2 Fig). As expected, Western blotting revealed increased phosphorylation of EGFR at Y1197 in this pooled line (Fig 4A). Loss of PTPRH was also associated with increased growth rate and proliferation in growth curves (Fig 4B) and MTT assays respectively (Fig 4C).

Given that loss of PTPRH resulted in elevated phosphorylation of EGFR, we hypothesized that these tumors would be susceptible to EGFR TKIs, demonstrating this in a proof of principle experiment in PyMT tumors [29]. To test this hypothesis in human lung cancer cell lines, dose response curves were completed for four cell lines using the EGFR TKI Osimertinib. NSCLC lines A427 (WT for EGFR and PTPRH) was a negative control while the H1975 line with classic L858R / T790D activating EGFR mutations was a positive control. Two NSCLC lines with naturally occurring PTPRH mutations were also tested including H1155 with a M188I mutation in the second fibronectin domain and H2228 with a Q887P mutation in the phosphatase domain. Both of these spontaneous mutations were predicted to have moderate EGFR activation and were not in the regions with the highest EGFR predicted activity for PTPRH mutations. As expected, the A427 negative control had no response while the H1975 positive control had a robust response. Interestingly, the two PTPRH mutant NSCLC lines had an intermediate response (Fig 4D). To test whether this would have phenotypic effects in vivo, we used a strategy of injecting H2228 cells into the flank of mice and treating with the EGFR TKI osimertinib once tumors reached 6mm (Fig 4E). The positive control responded well with tumors rapidly shrinking with TKI treatment (Fig 4F). While slower growing, the H2228 line with the PTPRH mutation had an appreciable tumor reduction at 50 mg/kg dosage (Fig 4G) (p<0.001 at endpoint by t-test), but had no effect at 25 mg/kg. The study was stopped at 14 days due to endpoint concerns. While not as robust as the EGFR mutant result, these proof of principle data demonstrate that PTPRH mutation status can induce a susceptibility for EGFR TKIs.
Fig 3. Increased 1197 p-EGFR in H23 PTPRH CRISPR KO Cells. Western blotting using lysate from PTPRH KO cells and PTPRH KO cells with transient overexpression of PTPRH plasmids demonstrates PTPRH indeed targets EGFR within human lung cancer cells. A) Electropherogram of PTPRH KO clones shows an A insertion at the CRISPR cut site. This indel was present for both clones in 3B. B) PTPRH KO CRISPR clones have increased 1197 PTEGFR. Rescue with: 1) WT PTPRH (Panel C) and 2) D986A (KD) PTPRH (Panel D).

C) PTPRH KO CRISPR clones have increased 1197 PEGFR. Total EGFR, and Beta Tubulin.

D) PTPRH KO CRISPR clones have increased 1197 PEGFR. Total EGFR, and Beta Tubulin.

E) PTPRH KO CRISPR clones have increased 1197 PEGFR. Mutate EGFR Y1197F in Clone 1. Total AKT, and Vinculin.

F) PTPRH KO CRISPR clones have increased 1197 PEGFR. Total AKT, and Vinculin.
To determine how widespread EGFR activation was within Ptprh mutant mouse tumors, and PTPRH knockout human tumors injected into the flank of mice, we employed an immunochemistry approach using an antibody specifically recognizing phosphorylation at Y1197 in EGFR. In a negative control (MMTV-PyMT) that was wild type for Ptprh mutations, we noted no appreciable staining for pY1197 EGFR (Fig 5A and 5B). In MMTV-PyMT tumors with a Ptprh mutation, there was widespread staining for pY1197 EGFR (Fig 5C). Interestingly, higher magnification revealed that this staining was predominantly nuclear (Fig 5D). This was present in multiple samples from multiple tumors, a second example is shown (Fig 5E and 5F).

Examining human NSCLC in the H23 PTPRH parental line revealed weak staining localized to the membrane (Fig 5G 10x and 5H 40x). H23 with a PTPRH knockout revealed increased pY1197 EGFR staining as compared to wild type controls (Fig 5I), but this staining was noted to be primarily located on the membrane (Fig 5J).

The activity of PTPRH is likely not limited to Y1197 of EGFR. To determine what other kinases were regulated by PTPRH, we screened a phosphorylated receptor tyrosine kinase (RTK) array with lysate from wild type H23 cells (Fig 6A) and lysate from CRISPR PTPRH KO H23 cells (Fig 6B). Several RTKs with differential phosphorylation patterns with PTPRH knockout lysate were identified, including FGFR1 with a 3.8 fold increase and IGF-1R with a 2.4 fold increase (Fig 6A–6C). Examining these RTKs in the publicly available genomic data through FGFR1 and IGF-1R ssGSEA signatures revealed predicted activation of these kinases in many of the mutations that also resulted in EGFR activation, although others were unique to each RTK (Fig 6D). To confirm predicted activation of FGFR1, Westerns were completed using cell lysates from PTPRH WT and PTPRH KO H23 cells revealing a clear increase in phosphorylation of FGFR1 within PTPRH KO cells (Fig 6E).

**Discussion**

Here we have identified conserved V483M Ptprh mutations in mouse mammary tumors from MMTV-PyMT transgenic mice. In a mixed background these mutations were conserved in matched pulmonary metastases, indicating this mutation occurs early in tumorigenesis. We demonstrated mutation of Ptprh to be impactful since PTPRH no longer dephosphorylated Y1197 of EGFR, as predicted by Yao et al [24], resulting in activation of the PI3K / AKT signaling cascade. Given the importance of EGFR activity in lung cancer, we confirmed the causal nature of PTPRH loss on EGFR activity through CRISPR mediated knockout of PPTPRH in a NSCLC line by observing increased pEGFR and pAKT. Importantly, a rescue experiment demonstrated that this was a specific event as plasmid expressed PTPRH was able to dephosphorylate EGFR while the rescue with a catalytically dead PTPRH did not. In addition, loss of PTPRH resulted in increased growth rate, potentially as a function of activation of the EGFR / PI3K / AKT signaling pathway. These results are summarized in Fig 7. This has potential to be a tumor driving event as it occurs early in tumor etiology and allows activation of a major signaling pathway to inappropriately persist. The conservation of the V483M mutation in over 80% of tumors from the genetically engineered mice in the FVB background also indicates that there is a remarkable selective pressure for EGFR pathway activity.
A pan-cancer analysis of human PTPRH mutations found numerous cancers harboring mutations, suggesting mutated PTPRH may play a role in tumor development across the spectrum of cancer types. PTPRH mutations were found in approximately 5% of NSCLC patients.

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with far higher rates in skin in multiple datasets. There is therefore a compelling reason to explore the role of PTPRH mutations in the future in skin cancer to determine if these mutations act in the same manner. Focusing on NSCLC, we noted mutations spread across the PTPRH exome. This is an interesting contrast to the conserved V645M mutation found within PyMT tumors, with potential implications for which mutations may be impactful on tumor growth. The human PTPRH mutations appear to cluster in several regions associated with activity of the targets (Fig 7) but mutations are scattered across the coding sequence. This is similar to a recent report for PTPRT in Head and Neck cancer and suggests a tumor suppressor type role [32]. While a mechanism has yet to be explored for each of the various PTPRH mutations, it is likely that the mutations are acting in a different fashion from each other. Mutations within the phosphatase domain may abrogate catalytic activity, while mutations in the fibronectin domains may prevent dimerization and binding of target substrates [25]. Since some of the phosphatase domain mutations with predicted high EGFR activity lie outside the conserved phosphatase activity HC(X3)R motif, it is also possible that these mutations impact access to the phosphatase domain and prevent recognition of substrate binding sites. Given the lack of conserved human PTPRH mutations, eventual utility of PTPRH mutation screening would need to be combined with a functional output screen. Ultimately the mutations in PTPRH and their functional impact on EGFR and response to TKIs may be paired with pEGFR or pFGFR1 status to predict response to EGFR TKI.

Our findings demonstrated increased phosphorylation of EGFR upon loss of PTPRH in NSCLC in both in vitro and in vivo models. With this causality and given that 5% of NSCLC tumors have mutations in PTPRH, and an estimated 235,000 cases of lung cancer occurring yearly within the United States, over 10,000 patients who present with PTPRH mutations could potentially benefit from EGFR targeted TKI therapy. Interestingly, the mutation types may allow for prediction of the therapeutic response. For instance, here we identified two NSCLC lines with PTPRH mutations that only had a medium level of predicted EGFR activity and treated them with the TKI Osimertinib. While both responded in vitro, the H2228 cell line also responded in vivo, with a decrease in tumor volume. The second line H1155 has a KRas mutation and this likely provides resistance to Osimertinib. In the future, cataloging PTPRH mutations with EGFR TKI response would allow for appropriate clinical action. Other potential options for treatment of PTPRH targets include dual inhibition of kinases whose signaling pathways are altered by PTPRH loss, or targeting RTKs with proteolysis targeting chimera (PROTAC) molecules, which target them for degradation. Overall, treatment of downstream targets regulated by phosphatases, rather than the phosphatases themselves, may be a viable solution, although this will require considerable characterization of the pathways affected by deregulated phosphatases. This is especially important to consider with the context dependent nature of PTP regulation, such as PTPRH deactivating EGFR.

Beyond EGFR, a kinase array showed increased phosphorylation of numerous RTKs within PTPRH knockout cell lines, including FGFR1 and IGFR1. Interestingly, increased phosphorylation of EGFR was not detected on the array. However, a closer examination of the phosphorylated antibodies used in the array revealed that EGFR Y1197 was not included. The increased
PTPRH mutations cause EGFR activity in lung cancer

**A**

WT Lysate

**B**

PTPRH KO Lysate

| RTK   | Fold Change |
|-------|-------------|
| FGFR1 | 3.81        |
| IGF-IR| 2.42        |
| CSK   | 1.89        |
| TRKB  | 1.29        |
| EphA2 | 0.20        |

**D**

Predicted Kinase Activity

**E**

MW  WT  WT Clone  PTPRH KO Clones

- pFGFR1 145 kd band
- pFGFR1 120 kd band
- totalFGFR1 145 kd band
- totalFGFR1 120 kd band
- totalFGFR1 90 kd band

Beta Tubulin
The phosphorylation of FGFR1 was further confirmed through western blotting. The activation of the FGFR1 pathway has interesting implications, both at the level of cellular pathways that may be affected, as well as potential treatment options for those with non-functional PTPRH. Moreover, the role of FGFR1 in other cancers has potential to open EGFR therapy in the other tumor types. In addition, with further preclinical work, a dual drug inhibition approach of targeting FGFR1 and EGFR may be of clinical use.

Finally, Ptprh mutant mouse tumors have increased staining of nuclear EGFR. Nuclear EGFR has been noted in times of cellular stress, as well as regenerating liver tissue. While in the nucleus, EGFR can act as a cofactor, or direct transcriptional activator by binding to the promoters of certain genes, such as cyclin D1. Increased nuclear EGFR upon loss of PTPRH activity could have profound impacts on cellular signaling pathways. The mechanism behind increased nuclear localization of EGFR with the V483M mutation is undetermined at this point but could include potential alterations to dimerization or an inability to maintain membrane localization. This is an area for future research which will be complemented by an

Fig 6. PTPRH Regulation of Receptor Tyrosine Kinases outside the ERBB Family. Ablation of PTPRH results in differential activation of numerous receptor tyrosine kinases outside of EGFR. A) A human phosphorylated RTK array shows differential phosphorylation of numerous RTKs when incubated with lysate from H23 PTPRH WT cells. B) A human phosphorylated RTK array shows differential phosphorylation of numerous RTKs when incubated with lysate from H23 PTPRH KO cells. Top five differentially phosphorylated RTKs are highlighted in red. C) Table showing the RTKs with the top 5 largest fold changes between PTPRH KO and PTPRH WT lysates. D) The lollipop plot in Fig 2B was recreated, adding predicted activation for FGFR1 and IGF-1R. Briefly, each dot on the PTPRH exome plot corresponds to a PTPRH mutant NSCLC tumor. Color-coded bars above each dot correspond to the predicted activity of EGFR, FGFR1, or IGF-1R. E) Western blotting confirms increased phosphorylation of FGFR1 at tyrosine residues 653/654, within PTPRH KO cell lysates compared to PTPRH WT lysates.

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phosphorylation of FGFR1 was further confirmed through western blotting. The activation of the FGFR1 pathway has interesting implications, both at the level of cellular pathways that may be affected, as well as potential treatment options for those with non-functional PTPRH. Moreover, the role of FGFR1 in other cancers has potential to open EGFR therapy in the other tumor types. In addition, with further preclinical work, a dual drug inhibition approach of targeting FGFR1 and EGFR may be of clinical use.

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Fig 7. Schematic of Ptprh mutant mouse tumors failing to dephosphorylate EGFR. Ordinarily, PTPRH is responsible for regulating EGFR signaling through dephosphorylation of tyrosine residues on the C-terminal tail of EGFR. We have shown PTPRH mutant mouse tumors and PTPRH KO human cells to have increased phosphorylation of EGFR, and subsequent increased activation of the PI3K/AKT pathway.
exploration of a variety of human PTPRH mutants to determine if any mutants result in
human nuclear pEGFR.

Methods

Targeted Resequencing of PyMT tumors

DNA was extracted from flash frozen tumors using lysis buffer (50 mL Tris HCl, 5 mL 500 mM
EDTA, 10 mL 10% SDS, 20 mL 5M NaCl, H2O up to 500 mL), or FFPE tissue using the Qiagen
FFPE extraction kit. The region flanking V483M was PCR amplified using the following primers;

Forward 5’ GGCCTTAGGTTCAATTGTAATAC 3’
Reverse 5’ CCTTAGCTTCCCGAGTATTTGGTT 3’

Amplified DNA was sent to GeneWiz for Sanger sequencing with the following primer 5’
TCATCCAAACTACATCTATGATCGGA 3’. Geneious software (https://www.geneious.com/)
was used for alignment to reference DNA.

Analysis of PTPRH Mutations in exome sequence data

Pre-annotated VCF files were downloaded for 64 tumors from GEO ascension number
GSE142387. Data was processed within R by reading in VCF files, then filtering to only keep
mutations within the Chr 7 bp 4548992–4604041 range (location of Ptprh in mouse genome).
These files were then converted to Annovar format, exported, and annotated using Annovar in
Linux based command. Statistical analysis was completed using a student’s t test (unequal vari-
ance, 2 tailed) between the metastasis group (mutations per met sample), and the primary
group (mutations per primary tumor).

PTPRH Mutations in human cancers

Pan-Cancer datasets from numerous sources, including TCGA and ICGC, were analyzed
through CBioPortal and the ICGC portal. Lung cancer mutation percentage were analyzed
specifically using TCGA 2016 dataset accessed through CBioPortal (https://www.cbioportal.
org/). The South Korean and U.S datasets showing discrepancy in percentage of PTPRH muta-
tions were analyzed on the ICGC portal (https://dcc.icgc.org/). Both datasets were filtered to
include only patients with exonic mutations.

All NSCLC datasets available on CBioPortal were used for mutual exclusivity analysis and
are listed below. PTPRH and EGFR SNV mutation data were downloaded and combined.
Duplicate samples were removed, and any sample with a PTPRH or EGFR mutation was con-
sidered. A 2x2 contingency table was run to determine mutual exclusivity. Datasets include;
MSK—Cancer Cell 2018, MSKCC—J Clin Oncol 2018, TRACERx—NEJM 2017, University of
Turnin, 2017, MSK—Science 2015, TCGA—Nat Genet 2016 (Pan), Broad—Cell 2012,
MSKCC—Science 2015, TCGA—Firehose Legacy, TCGA—Nature 2014, TCGA—Pan-cancer
Atlas, TSP—Nature 2008, MSKCC—Cancer Discov 2017, TCGA—Nature 2012

Demographics of PTPRH mutations

Age, overall survival, and race demographics were analyzed using the Lung Adenocarcinoma
TCGA Pan-Cancer Atlas data set downloaded from CBioPortal. Two-tailed Student’s T-Tests
assuming unequal variance were completed for PTPRH mutant VS. EGFR mutant samples, as
well as PTPRH mutant VS. WT (non-EGFR mutant) samples for age of diagnosis and overall
survival. Samples without age or OS data were excluded. Only samples with missense or trun-
cating mutations were included, and overexpression samples were excluded. Race was ana-
yzed using a 2x2 contingency table.
EGFR Activity and pathway activity predictions

TCGA pan-cancer RNA-seq dataset (downloaded from UCSC Xena) was analyzed for 
PTPRH, EGFR, FGFR1, and IGF1R mutations. This mutation list was downloaded and filtered to keep samples that had a mutation in PTPRH, EGFR, or that were wild type for PTPRH, EGFR, FGFR1, and IGF1R. Any sample with a mutation in PTPRH was kept, resulting in 53 samples. 10 samples of each of the two categories were kept; WT for PTPRH and the above three RTKs, and L858R mutant EGFR that were WT for PTPRH, FGFR1, or IGF1R. To decide which WT and EGFR samples to keep, samples from those subsequent groups were assigned a random number using the RAND() function in Excel. These numbers were then sorted from highest to lowest, keeping the top 10 samples. RSEM(log2 X+1) normalization was applied to the filtered sample list, resulting in 47 PTPRH mutant samples (WT for the kinases), 9 samples that WT for PTPRH and the three kinases, and 8 samples with EGFR mutations (WT for PTPRH, FGFR1, and IGF1R). ssGSEA was run on the samples to predict pathway activation status. Pathways for each kinase were filtered down, selecting the most relevant and robust pathway. A ranking sum score was applied to the pathway prediction data for each sample.

For GSEA analysis of PTPRH mutant tumors, the pan-cancer RNA-seq dataset was again downloaded from UCSC Xena. Twelve tumors for each of the three categories were kept; PTPRH mutant tumors predicted to have high EGFR activity, EGFR L858R mutants, and tumors that were WT for both PTPRH and EGFR. GSEA was completed using the GenePattern server.

CRISPR Knockout

Benchling was used to design the guide RNA (AGCACACACTAACATCACCG) targeting the fourth exon of PTPRH. The guide was cloned into px458 using AgeI and EcoRI. Transient transfection of px458 into H23 cells was completed using Promega’s Viafect. GFP positive cells were sorted into single cell clones into 96 well plates using FACS. Once clones had grown into a colony, they were subsequently moved to 24-well plates, then 6-well plates. DNA was harvested and sent to ACTG for sanger sequencing.

CRISPR Knock-In mutations

Guide RNA was designed in Benchling with the PAM (NGG) sequence 5 bp downstream of the desired EGFR Y1197 mutation site. The single stranded region of homology was designed in Benchling by choosing desired length for homology arms as well as the desired mutation, then taking the reverse complement of that strand. The oligo was designed with 36 bp upstream of the desired mutation site and 90 bp downstream. The desired mutation resulting in a Y1197F amino acid substitution was added. This mutation also resulted in the addition of an EcoRI cut site, which was used for downstream screening. The mutation also altered the guide RNA enough to prevent re-annealing once HR mediated repair occurred. Guide RNA was cloned into px458. H23 PTPRH KO cells were transfected using Viafect in a 6:1 ratio. 1 ug of px458 with guide, and 4 ug of ss repair template were transfected. Sorting was completed using FACS for GFP. Clones were screened using a digest for EcoRI and confirmed with sequencing.

Western blotting

Tumor lysates were harvested from flash frozen tumors by crushing with a mortar and pestle, then dissolving in TNE lysis buffer (5 mL 1 M Tris HCl pH 8, 3 mL 5M NaCl, 1 mL NP40, 400 uL .5M EDTA, 2.0 mL .5M NaF, H2O to 100 mL). Roche mini protease tablets and sodium
orthovanadate were used as protease and phosphatase inhibitors respectively. Primary antibodies were incubated overnight. Antibodies used were as follows; total EGFR (Cell Signaling D38B1), 1197 EGFR (Invitrogen PA5-37553), total AKT (Cell Signaling 11E7), p-s473 AKT (Cell Signaling D9E), total STAT3 (Cell Signaling 79D7), p-Y705 STAT3 (Cell Signaling D3A7), total FGFR1 (Cell Signaling D8E4), p-Y653/654 FGFR1 (Cell Signaling 3471s), beta tubulin (Proteintech 10094–1), vinculin (Cell Signaling E1E9V), total ERK (Cell Signaling 9102), p-ERK (Cell Signaling 4370).

Overexpression
PTPRH cDNA within plasmid PRc-CMV was kindly provided by Dr. Takashi Matozaki at Kobe University. Site directed mutagenesis was used to achieve a D986A mutant. Both WT and D986A mutant PTPRH plasmid constructs were transiently expressed in PTPRH KO cells using Viafect.

RTK Array
The manufacturer’s protocol for RayBiotech Human RTK Phosphorylation Array C1 kit was followed. Membranes were incubated with lysate from H23 WT cells or H23 PTPRH KO cells.

IHC Nuclear EGFR
Human cell lines H23 PTPRH WT or H23 PTPRH KO were injected into the left flank of nude mice. H23 cell line tumors were grown to approximately 10 mm in the largest direction prior to necropsy. Mouse PyMT tumors, and tumors grown from human H23 cells were necropsied with portions of tumor tissue preserved in formalin, and portions of tumor flash frozen for further downstream analysis. Formalin fixed paraffin embedded tumors were subjected to staining using an antibody specific for 1197 EGFR (Thermo PA5-37553).

Pooled CRISPR Knockout
Guide RNA (AGCACACACTAACATCACCG) for PTPRH was designed using Benchling. And cloned into a lentiviral Cas9 plasmid (Addgene # 52961). Viral generation was completed through transfection of 293T cells with packaging plasmid psPAX2 and envelop plasmid pMD2.G in a ratio of 3.7:1:2.5 with the Cas9 plasmid respectively. Viral supernatant was collected from 293T cells 3 days after transfection, and filtered through a .22 uM syringe filter. 1 mL of filtered viral supernatant was applied to H23 WT cells at ~30% confluency. Sanger sequencing was used to confirm knockout, and for TIDE analysis.

MTT Assay and growth curves
MTT assay kit (Roche 11465007001) instructions were followed. Graphpad was used to plot and statistically analyze results. A Welch’s two-tailed t-test yielded a p-value of .0137.

For growth curves 1.0 x 10⁵ cells were plated in triplicate within 6-well plates. On days 1–5, cells were trypsinized and cell number was read using an automated cell counter. Graphpad was used to plot results.

Dose response curves
Cells were diluted to 5.0 x 10⁴ cells per mL, and 20 uL of cell suspension was added to wells of an opaque 384 well plate. After overnight recovery, cells were subjected to a dose response curve of increasing drug concentration in half log steps. For single drug curves, osimertinib (Cayman AZD9291) range was .00003 to 30 uM. For dual drug curves, osimertinib range was
.03 to 10 uM, and either KRAS inhibitor (ARS853, Cayman) or FGFR1 inhibitor (PD166866, Cayman) range was .00003 to 30 uM. 10 mM stocks of drugs were made by diluting with DMSO, and half-log drug series were diluted fresh with complete media. Cell viability was read after 48 hours using Promega’s Cell Titer Glo. Luminescence values were normalized to non-drug treated controls, and plotted using Graphpad.

**In vivo treatment**

H2228 and H1975 cell lines were injected into the left flank of 6–12 week old nu/nu mice. After tumors reached 6mm in the largest dimension, mice were randomized into treatment groups; vehicle control, 25 mg/kg osimertinib, or 50 mg/kg osimertinib. The 50 mg/kg dose was only used for mice with H2228 tumors. Osimertinib (AZD9291 Cayman) was diluted using the following in order to achieve a final ratio: 5% DMSO, 40% polyethylene glycol, 5% tween-80, 50% H2O. Max volume of treatment was 10 uL for 1 gram of body weight. Mice were weighed on first day of treatment, and volume of drug was adjusted to achieve proper dose.

**KI67 scoring**

Slides were scored on a scale of 1–10 by three blinded reviewers independently. The mean for each tumor slide was then taken by averaging the three reviewer scores for each slide. A two-tailed student’s T-test assuming unequal variance was then completed across the osimertinib and vehicle control sample groups, using the means for each tumor slide (each group n = 4).

**Supporting information**

S1 Fig. Stat3 and Erk phosphorylation is unchanged in PTPRH KO cell lines. (PDF)
S2 Fig. H23 Cell line Pooled knockout PTPRH events using TIDE analysis. (PDF)
S1 Table. ssGSEA Genesets from Fig 2D. (XLSX)

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