CDK-dependent phosphorylation of PHD1 on serine 130 alters its substrate preference in cells

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ABSTRACT

PHD1 (also known as EGLN2) belongs to a family of prolyl hydroxylases (PHDs) that are involved in the control of the cellular response to hypoxia. PHD1 is also able to regulate mitotic progression through the regulation of the crucial centrosomal protein Cep192, establishing a link between the oxygen-sensing and the cell cycle machinery. Here, we demonstrate that PHD1 is phosphorylated by CDK2, CDK4 and CDK6 at S130. This phosphorylation fluctuates with the cell cycle, and can be induced through oncogenic activation. Functionally, PHD1 phosphorylation leads to increased induction of hypoxia-inducible factor (HIF) protein levels and activity during hypoxia. PHD1 phosphorylation does not alter its intrinsic enzymatic activity, but instead decreases the interaction between PHD1 and HIF1α. Interestingly, although phosphorylation of PHD1 at S130 lowers its activity towards HIF1α, this modification increases the activity of PHD1 towards Cep192. These results establish a mechanism by which cell cycle mediators, such as CDKs, temporally control the activity of PHD1, directly altering the regulation of HIF1α and Cep192.

KEY WORDS: EGLN2, HIF, Cep192, Hypoxia

INTRODUCTION

Decreased oxygen levels, or hypoxia, present a major stress to the cell. Many of the crucial cellular processes, such as ATP production through oxidative phosphorylation, cell division and cell cycle progression are all highly energy-demanding processes that require oxygen (Ortmann et al., 2014). Exposure to hypoxia activates a number of different responses at both the cellular and whole organism level. One of the crucial alterations mediated by hypoxia is a change in gene expression (Kenneth and Rocha, 2008).

Hypoxia activates a variety of transcription factors (Kenneth and Rocha, 2008), but the most important for survival and adaptation to this stress is a group of transcription factors known as the hypoxia-inducible factors (HIFs). These are heterodimeric transcription factors that comprise an oxygen-labile HIFα subunit and the constitutively expressed HIF1β subunit (also known as ARNT) (Moniz et al., 2014). Three different genes encode for the currently known isoforms of the HIFα subunit [HIF1α, HIF2α (also known as EPAS1) and HIF3α]. All of the three HIFα isoforms share some structural similarity, most notably they all contain a basic helix-loop-helix Per-Arnt-Sim (bHLH-PAS) domain, which is crucial for its interaction with its transcriptional partner HIF1β (To et al., 2006). In addition, they also contain an oxygen-dependent degradation domain (ODD), which renders these proteins sensitive to proteasomal degradation in the presence of oxygen. Although transcription and translation of the HIFα isoforms plays a role in the control of these transcription factors (Bernardi et al., 2006; Nayak et al., 2013; van Uden et al., 2008, 2011), the oxygen-dependent control of HIFα is achieved through protein degradation, which occurs very rapidly in the presence of oxygen (Fandrey et al., 2006).

During normoxia, when cells have access to oxygen, HIFα is hydroxylated on two key proline residues located within the ODD domain by a group of proline hydroxylase enzymes (PHDs). PHDs require molecular oxygen as a co-substrate to carry out hydroxylation, but they also have a requirement for α-ketoglutarate (α-KG) and Fe^{2+} as cofactors (Fandrey et al., 2006). As α-KG is a key component of the Krebs cycle, it is thought that, in addition to sensing oxygen levels, PHDs can also sense the metabolic state within the cell (Kaelin, 2012). More recent data have shown that the PHDs are also important for amino acid sensing (Durán et al., 2013). Currently there are three known isoforms in mammalian cells (PHD1, PHD2 and PHD3, also known as EGLN2, EGLN1 and EGLN3, respectively), all of which have the ability to hydroxylate HIFs. HIF1α is hydroxylated on P402 and P564, whereas HIF2α is hydroxylated on P405 and P531. Biochemical analysis of all three isoforms has shown that PHD2 has the highest affinity for HIFs, but, interestingly, the PHDs also possess preferential affinities for the proline that they target (Appelhoff et al., 2004; Berra et al., 2003). Genetic studies have shown that, out of the three isoforms, deletion of PHD2 is embryonic lethal (Takeda et al., 2006), whereas deletion of PHD1 and PHD3 are not. However, loss of PHD1 and/or PHD3 lead to developmental defects, most notably in the cardiovascular system (Fong and Takeda, 2008; Takeda et al., 2008).

During normoxia, hydroxylation of HIFα creates a binding site for the von Hippel-Lindau (VHL) tumor suppressor E3 ligase complex. Binding of VHL results in polyubiquitylation and proteasomal degradation of HIFα. During hypoxia, when oxygen levels are decreased, PHD activity is reduced, leading to stabilization of HIFα and dimerization with HIF1β, resulting in a transcriptionally active complex. HIFs have been shown to regulate a large number of genes involved in a variety of cellular processes, such as metabolism, apoptosis, autophagy, angiogenesis and cell proliferation (Moniz et al., 2014; Rocha, 2007). The response engaged during hypoxia promotes cell survival and turns off highly energy-consuming processes, such as cell proliferation and translation.

One of the most energy-consuming processes within the cell is the cell cycle and, hence, cell division. This process must be tightly regulated to ensure there is no hyperproliferation and/or mis-segregation of genetic information. Errors within the cell cycle can...
ultimately lead to disease states, such as cancer. Cell cycle control is achieved through multiple mechanisms, but amongst the most important regulators are cyclin-dependent kinases (CDKs) (Besson et al., 2008; Bloom and Cross, 2007; Obaya and Sedivy, 2002). CDKs are a family of serine/threonine kinases, which are activated when the cell chooses to enter the cell cycle. Their activation is dependent on multiple factors, but most important is their interaction with their regulatory cyclins (Bloom and Cross, 2007). In addition, CDK activity is also regulated through their interaction with inhibitory proteins, such as p21 (also known as CDKN1A) and p27 (also known as CDKN1B). These proteins interact directly with the CDKs and inhibit the interaction with their regulatory cyclin (Besson et al., 2008).

Several studies have shown that hypoxia also affects the cell cycle. Early work has demonstrated that, upon exposure to hypoxia, cells are reversibly arrested in G1 or S phase (Gardner et al., 2001; Ortmann et al., 2014). The mechanisms controlling this arrest have been shown to be both HIF dependent and independent (Ortmann et al., 2014). More recent studies have shown that HIF1α can inhibit DNA replication independently of its transcriptional activity (Hubbi et al., 2013).

In more recent years, the roles of the PHDs in processes other than hypoxia have become more apparent. Recently PHD1 has been shown to regulate the transcription factor FOXO3A (Zheng et al., 2014). PHD3 has also been shown to hydroxylate PKM2 and HCLK2 (Luo et al., 2011; Xie et al., 2012). Moreover, our recent work has shown that PHD1 can regulate mitotic progression through its ability to control the levels of the key centrosomal component Cep192 (Moser et al., 2013). However, the mechanism that determines whether PHD1 targets Cep192, HIFα, or both, is not known. In addition, there is no information on how these enzymes engage with the cell cycle.

Here, we show that PHD1 is regulated by the cell cycle at the post-translational level. We show that PHD1 is phosphorylated at S130 in a CDK-dependent manner. PHD1 phosphorylation reduces its interaction with HIF1α, but increases the association between PHD1 and Cep192. Functionally, this results in increased levels of HIF1α protein and increased transcriptional activity in response to hypoxia, and in reduced levels of Cep192 protein. These results indicate that the behaviour of PHD1 towards different substrates can be altered by specific post-translational modifications.

RESULTS

PHD1 is phosphorylated on S130

We performed mass spectrometry analysis to map PHD1 phosphorylation events by using extracts from U2OS cells expressing GFP-tagged PHD1. PHD1–GFP was immunoprecipitated from cells and then subjected to mass spectrometry analysis (Fig. 1A, Table 1). A good coverage of PHD1 peptides was obtained, and we found that S130 was phosphorylated in interphase cells (Fig. 1A). Mass spectrometry validation was achieved using antibodies that specifically recognize phosphorylated (phospho-) serine or threonine residues (Fig. 1B), confirming that PHD1 can be phosphorylated on serine residues but not on threonine residues (Fig. 1B). Sequence alignment of PHD1 from different organisms demonstrates that the S130 phosphorylation site in humans is highly conserved in higher mammals, mice and rats but is absent in organisms such as zebrafish, Xenopus and the fruit fly (Fig. 1C). An antibody against a synthetic phospho-peptide corresponding to the region around S130 was generated. Antibody specificity and validation was performed using U2OS cells stably expressing GFP, PHD1–GFP and two GFP-tagged PHD1 mutants cell lines where the S130 residue has been replaced with either an alanine (PHD1-S130A) or an aspartate (PHD1-S130D). All of these cells expressed PHD1 to similar levels (Fig. S1A). Immunoprecipitation of GFP from the GFP, PHD1–GFP, PHD1-S130A–GFP or PHD1-S130D–GFP cells, revealed that the phospho-specific antibody only detected a band in the extracts derived from wild-type PHD1, demonstrating its specificity (Fig. 1D). In addition, we knocked down PHD1 levels using several different small interfering RNA (siRNA) oligonucleotides directed against PHD1, and used the antibody to determine its specificity in cell extracts (Fig. 1E). A substantial loss of signal was detected specifically when PHD1 was depleted, further demonstrating the specificity of this antibody (Fig. 1E).

Although, endogenous PHD1 levels are difficult to detect in cell lines other than breast cancer lines (Moser et al., 2013; Zheng et al., 2014), we used an immunofluorescence approach in U2OS cells to determine whether the antibody against phospho-S130 PHD1 was able to detect endogenous protein (Fig. 1F). PHD1 phosphorylation was localised primarily within the nucleus, consistent with previous data showing that PHD1 is a nuclear protein (Metzen et al., 2003). The PHD1 phosphorylation signal was lost when cells were treated with a PHD1-specific siRNA. The same localisation pattern was also observed in PHD1–GFP cells (Fig. S1B). Collectively, these results show that PHD1 is subject to phosphorylation on S130 in cells.

PHD1 phosphorylation is regulated by interphase CDKs

Further analysis of the sequence surrounding the phosphorylation site revealed it falls into a consensus for CDK phosphorylation [S/T]pRx [R/K] (Endicott et al., 1999). To investigate whether CDKs can modify PHD1, we determined whether PHD1 could be detected by an antibody specifically directed towards phospho-CDK substrates. PHD1–GFP was immunoprecipitated from cells, and extracts were probed with both the antibody for the CDK substrates and with an anti-PHD1 antibody (Fig. 2A). This revealed that a small proportion of PHD1 was indeed recognised by the anti-CDK-substrate antibody, highlighting the potential for PHD1 to be a CDK substrate. In addition, PHD1 could interact with CDK2, CDK4 and CDK6, but not CDK1 in cells (Fig. 2B; Fig. S1C). We could also detect an interaction between PHD1 and CDK2 at the endogenous level (Fig. 2C).

Functionally, depletion of CDK2, CDK4 and CDK6, either individually or in combination, resulted in reduced levels of phospho-S130 PHD1 (Fig. 2D,E). CDK1 depletion led to increased levels of this phosphorylation on PHD1 (Fig. 2D), with a corresponding increase in cells arrested in the G2 and M phase of the cell cycle. However, CDK depletion did not alter PHD1 localisation in cells (Fig. S1D). By contrast, gain-of-function experiments revealed that when increased levels of CDK4 were present, there was a concomitant increase in the levels of S130 phosphorylation of PHD1 (Fig. 2F), indicating that indeed CDKs can change the PHD1 phosphorylation status.

To demonstrate that CDKs can phosphorylate PHD1, we also performed in vitro kinase assays with CDK2 and CDK1 (Fig. 2G, Fig. S1E). CDK2 was immunoprecipitated from cells, and kinase assays were performed using bacterially expressed recombinant PHD1, followed by western blot analysis using the anti-phospho-PHD1 antibody (Fig. 2G). This analysis showed that CDK2 phosphorylates PHD1 at S130. Interestingly, when a radioactive kinase assay was performed with recombinant CDK1–cyclin-B, a CDK for which we were unable to detect an interaction with PHD1, we could detect phosphorylation of PHD1 in vitro (Fig. S1E). In this case, mutation of S130 only slightly reduced the phosphorylation signal, suggesting that other sites on PHD1 are
being targeted by CDK1 in vitro. Taken together, these results suggest that no single CDK is exclusively responsible for the phosphorylation of PHD1, indicating a redundancy between them. This is to be expected, as genetic studies have shown that only CDK1 is essential, as knockout of CDK1 results in significant developmental defects in mice (Diril et al., 2012).
was regulated by growth factor stimulation (Fig. 3B). After starving the cells for 24 h, S130 phosphorylation was decreased when compared with the control, and this was coupled with an increase in the percentage of cells in the G1 phase of the cell cycle (Fig. 3B). Following release, there was a gradual increase in PHD1 phosphorylation, which coincided with an increase of cells moving into S and G2 phases of the cell cycle. Interestingly, this increase was delayed when compared with ERK1 and ERK2 (ERK1/2, also known as MAPK3 and MAPK1, respectively) phosphorylation, indicating that the kinase responsible is activated later than the ERK pathway.

A primary driver of cell proliferation and cell cycle progression in the context of cancer is the activation of oncogenes (Matsumura et al., 2003). We overexpressed the oncogene Myc and two mutant derivatives. One of the mutants contains a deletion in the Myc box II domain (ΔMBII), which prevents full transactivation of Myc (Cowling and Cole, 2008). The other mutant has a T58A mutation and acts as a Myc gain-of-function mutant (Welcker et al., 2004). T58 is a known GSK3 phosphorylation site and, without this site, Myc has increased stability, making it more active (Welcker et al., 2004). We also tested the effects of overexpression of E2F1 and Src. Overexpression of either wild-type Myc or either of the mutants, led to an increase in PHD1 phosphorylation, although the increase in PHD1 phosphorylation when overexpressing ΔMBII was not very pronounced (Fig. 3C, lanes 2–4). PHD1 phosphorylation also increased when E2F1 was overexpressed (Fig. 3C, lane 5), but not when we overexpressed Src (Fig. 3C, lane 6). This shows that S130 phosphorylation of PHD1 can be induced by increased levels of oncogenes, such as Myc and E2F1.

**PHD1 phosphorylation regulates HIFα levels and activity**

As PHD1 is a proline hydroxylase, we next determined whether phosphorylation of PHD1 at S130 could impact on its intrinsic enzymatic activity. Previous work performed in bacteria has shown that PHD1 is potentially phosphorylated on S132 (Li et al., 2008) and this can lead to a decrease in PHD1 activity in vitro. To analyse PHD1 activity we used an in vitro hydroxylation assay, using a peptide derived from the HIF1α ODD sequence, followed by mass spectrometry (Fig. 4A). This analysis revealed that all the mutants had similar activity to the wild-type enzyme in vitro. We also

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**Table 1. Fragmentation ions for phosphorylation of PHD1**

| b-H3PO4 | b-H3PO4+r2 | b | b+r2 | Sequence | y | y+2 | y+3 | y-H3PO4 | y-H3PO4+r2 |
|---------|------------|---|------|---------|---|-----|-----|--------|----------|
| –       | –          | – | –    | 1       | W | –   | –   | –      | –         |
| –       | –          | – | –    | 2       | A | 17  | 1876.8338 | 938.9205 | 626.2828 | 1778.8569 | 889.9321 |
| –       | –          | – | –    | 3       | E | 16  | 1805.7967 | 903.402  | 602.6037 | 1707.8198 | 854.4135 |
| –       | –          | – | –    | 4       | D | 15  | 1676.7541 | 838.8807 | 509.5895 | 1578.7772 | 789.8922 |
| –       | –          | – | –    | 5       | G | 14  | 1561.7271 | 781.3672 | 521.2472 | 1463.7502 | 732.3767 |
| –       | –          | – | –    | 6       | G | 13  | 1504.7057 | 752.8565 | 502.2401 | 1406.7288 | 703.8688 |
| –       | –          | – | –    | 7       | D | 12  | 1447.6812 | 724.3457 | 483.2329 | 1349.7073 | 675.3573 |
| –       | –          | – | –    | 8       | A | 11  | 1326.6572 | 666.8323 | 444.8906 | 1234.6046 | 617.8438 |
| –       | –          | – | –    | 9       | P | 10  | 1261.6201 | 631.3137 | 421.2116 | 1163.6432 | 582.3253 |
| 968.3745 | –         | – | –    | 10      | S(phospho) | 9 | 1164.5674 | 502.8783 | 388.8060 | 1066.5905 | 533.7989 |
| 1065.4272 | –        | – | –    | 11      | P | 8   | 997.569 | 499.2881 | 333.1945 | –         | –         |
| 1152.4592 | –        | – | –    | 12      | S | 7   | 900.5162 | 450.7816 | 300.8436 | –         | –         |
| 1280.5542 | –        | – | –    | 13      | K | 5   | 813.4842 | 407.2457 | 271.8329 | –         | –         |
| 1436.6553 | –        | – | –    | 14      | R | 6   | 685.3993 | 343.1983 | 229.1346 | –         | –         |
| 1533.7081 | –        | – | –    | 15      | P | 4   | 529.2881 | 265.1477 | –         | –         |
| 1719.7874 | –        | – | –    | 16      | W | 3   | 432.2354 | 216.6123 | –         | –         |
| 1790.8245 | –        | – | –    | 17      | A | 2   | 246.1561 | 123.5817 | –         | –         |
| –       | –          | 18 | R    | 1       | 175.119 | 88.0631 | –       | –         | –         |

**Summary of the theoretical fragment ions generated upon CID fragmentation of the triply charged peptide WAEDGGDAPPS(phospho)PSKRPRWAR derived from the tryptic digestion of PHD1. Fragment ions that were observed in our analysis are highlighted in bold (for y ions) and italics for b ions.**
Fig. 2. See next page for legend.
Fig. 2. PHD1 phosphorylation at S130 is regulated by CDKs. (A) 300 µg of U2OS GFP or PHD1–GFP cell extracts were subjected to immunoprecipitation (IP) using GFP-trap beads, and precipitated material was analysed by western blotting using the indicated antibodies. (B) 300 µg of U2OS GFP or PHD1–GFP cell extracts were subjected to immunoprecipitation using antibodies towards CDK4, CDK6 and CDK2, and precipitated material was analysed by western blotting for the presence of PHD1. (C) 500 µg of U2OS cell extracts were subjected to immunoprecipitation using an anti-PHD1 antibody crosslinked to Sepharose beads and processed as in A. (D) U2OS PHD1–GFP cells were transfected with control, CDK1 or CDK2 siRNA (siCDK) oligonucleotides alone or in combination for 48 h prior to lysis for western blotting or fixation for FACS analysis. Cell lysates were analysed for the levels of phosphorylated PHD1 at S130 and appropriate controls. The middle graph depicts the mean±s.d. of the quantification of the western blot analysis, representing a minimum of three independent experiments. The right panel depicts the corresponding cell cycle profile of cells treated as mentioned (mean±s.d. of a minimum of three independent experiments). (E) U2OS PHD1–GFP cells were transfected with control or the indicated CDKs siRNAs alone or in combination for a period of 48 h prior to being processed and analysed as in D. (F) U2OS PHD1–GFP cells were transfected with 1 µg of control or CDK4 expression constructs for 48 h prior to lysis and analysed by western blotting for the levels of phosphorylated PHD1 at S130 and appropriate controls. Ev, empty vector control. Graph depicts mean±s.d. of the quantification of the western blot analysis, from a minimum of three independent experiments. (G) CDK2 was immunoprecipitated from cells and used in a kinase assay with 2 µg of recombinant PHD1 and PHD1–S130A protein. Reactions were analysed by western blotting with the indicated antibodies. *P<0.05, **P<0.01 and ***P<0.001 compared to control conditions (Student’s t-test), p, phosphorylated form of the protein. See also Fig. S1.

performed this analysis using dot blot and an anti-HIF1α-hydroxylation antibody (Fig. S2A). We compared hydroxylation of the HIF peptide over time using recombinant purified GST–PHD1, GST–PHD1–S130A and GST–PHD1–S130D proteins. No significant difference in hydroxylation activity between the wild-type PHD1 and the two mutants could be detected (Fig. S2A). These data indicate that phosphorylation of PHD1 at S130 has little or no effect on the intrinsic enzymatic activity of PHD1.

To determine whether there is a functional role of PHD1 S130 phosphorylation in the cellular response to hypoxia, we started by analysing the levels of its targets, that is HIF1α and HIF2α. To this aim, we utilized the GFP, PHD1–GFP, PHD1–S130A–GFP and PHD1–S130D–GFP U2OS cells, and assessed their PHD activity by measuring both the levels of hydroxylated HIF1α and total levels of HIF1α and HIF2α when their degradation was blocked by a proteasomal inhibitor (Fig. 4B). Specificity of the anti-hydroxy-HIF1α antibody was confirmed by analysis of extracts from cells treated with the PHD inhibitors DFX and DMOG (Fig. S2B). In addition, endogenous PHD1 was depleted by siRNA and, as expected, when MG132 was added, we saw an accumulation of HIF1α in all cell types. However, there were decreased levels of HIF1α in the PHD1–GFP and PHD1–S130A–GFP cells, whereas HIF1α levels in PHD1–S130D–GFP cells were similar to the cells expressing GFP alone (Fig. 4B). Similarly, when we analysed HIF1α hydroxylation levels, less hydroxylation was detected in PHD1–S130D–GFP cells when compared with PHD1–S130A–GFP cells. This result suggests that phosphorylation of PHD1 on S130 does impact on the ability of PHD1 to hydroxylate HIF1α in cells. Interestingly, mutation of S130 alters the PHD1-mediated regulation of HIF2α, regardless of mutation to alanine or aspartate (Fig. 4B). As expected, none of the PHD1 mutations affected the total levels of HIF1β.

In cells exposed to hypoxia, PHD1 overexpression results in a reduction of HIF1α levels and its targets as expected (Fig. S2C,D). Hypoxia does not alter PHD1 phosphorylation at earlier times of exposure but does result in a significant reduction after 24 h (Fig. S2E). This is to be expected, as exposure to hypoxia for this period results in G1 arrest (Ortmann et al., 2014), a stage where PHD1 S130 phosphorylation is reduced. When all three cell lines were exposed to hypoxia, we detected an induction of HIF1α in all of them (Fig. 4C), although to lower levels than control cells (Fig. S2C,F). However, for the PHD1-overexpressing cells, the highest induction of HIF1α was observed in the PHD1–S130D–GFP cells (Fig. 4C). This highlights that the change in hydroxylation activity we observed when we treated cells with MG132 (Fig. 4B) is physiologically relevant. In addition, levels of HIF2α are higher in both of the PHD1 mutant cell lines, this being particularly evident in PHD1–S130D–GFP cells (Fig. 4C). Despite being less efficient at targeting HIF1α, PHD1–S130D–GFP is active, as the levels of HIF1α in these cells are lower than cells expressing GFP alone (Fig. S2F), indicating that PHD1 hydroxylase activity is still present when S130 is phosphorylated.

To understand whether the changes in HIF1α levels observed in the PHD1–S130D–GFP cells are altering HIF1α transcriptional activity, we assessed HIF activity by investigating the levels of several HIF target genes. Levels of HIF1α targets were always higher in PHD1–S130D–GFP cells when compared with cells expressing wild-type PHD1. However, additional differences were also observed for PHD1–S130A–GFP cells when compared with cells expressing wild-type PHD1, with higher levels of PHD2, PHD3 and CAIX (also known as CA9) observed in these cells (Fig. 4D). This could reflect the contribution of HIF2α to the regulation of certain targets (Elvidge et al., 2006). We also analysed mRNA levels for BNIP3, CAIX and Glut3 (also known as SLC2A3), at 24 h following exposure to hypoxia in these cells (Fig. S2G). These results show that levels of HIF1α targets are always higher in PHD1–S130D–GFP cells when compared with cells expressing wild-type PHD1. In this analysis, PHD1–S130A–GFP activity was comparable to wild-type PHD1, for all the genes analysed (Fig. S2G).

We have shown that PHD1 S130 phosphorylation is regulated by the cell cycle. As such, we next determined whether cells where PHD1 phosphorylation is high would have different levels of HIF1α. To this end, we synchronised cells with a double-thymidine block, and then released them into fresh medium containing the PHD inhibitor DFX, and then visualised HIF1α. We initially determined whether PHDs were still active under these conditions by investigating the levels of hydroxylated HIF1α in a time course of DFX treatment (Fig. S2H). We thus chose a 3-h DFX treatment because, at this time point, HIF is stabilised but still hydroxylated, and hence PHD activity changes could still be monitored. When we investigated HIF1α levels in the different stages of the cell cycle, we could observe that cells in G1 had lower levels of PHD1 S130 phosphorylation as well as lower HIF1α levels (Fig. 4E). By contrast, when cells were synchronised in either S or G2 and M phase, PHD1 S130 phosphorylation increased and so did HIF1α levels (Fig. 4E). Taken together, this analysis reveals that both the phosphorylation-mimicking mutation S130D and increased S130 phosphorylation results in reduced PHD1 activity towards HIF1α, leading to both increased HIF1α levels and increased activity of this transcription factor under hypoxia.

**PHD1 phosphorylation results in increased HIF1α half-life by reducing PHD1–HIF1α interaction**

Our analysis so far has revealed that PHD1 phosphorylation does not alter intrinsic enzymatic activity in vitro but does so in the context of cells, leading to increased levels of HIF1α levels and activity. To understand the mechanism behind these differences, we
started by measuring HIF1α half-life in GFP, PHD1–GFP, PHD1-S130A–GFP, and PHD1-S130D–GFP cells, using a cycloheximide chase approach (Fig. 5A; Fig. S3A). Cells were incubated under hypoxia for 4 h prior to treatment with cycloheximide for the indicated periods of time. We measured p53 levels as a positive control for the treatment, as it is known to have a high turnover rate
**Fig. 3. PHD1 phosphorylation at S130 is regulated by the cell cycle and responds to mitogenic signalling.** (A) U2OS PHD1–GFP cells were subject to a double-thymidine block release protocol prior to lysis or fixation after the indicated periods of time. The left panel depicts western blot analysis for the levels of phosphorylated PHD1 at S130, and appropriate controls. The right panel represents the cell cycle profile of matching samples analysed by flow cytometry. As, asynchronous. Graph depicts mean±d. of a minimum of three independent experiments. G1, S, and G2/M are the phases of the cell cycle that correspond to the indicated time points. (B) U2OS PHD1–GFP cells were serum starved for 24 h prior to addition of full serum medium and were harvested at the indicated times. Cells were lysed for western blot analysis or fixed for FACS analysis. Cell lysates were analysed for the levels of phosphorylated PHD1 at S130, and appropriate controls, where phosphorylated ERK1/2 was used as a marker of mitogenic signalling. The right panel depicts the mean±d. of the quantification of the western blot analysis, representing a minimum of three independent experiments. The bottom panel depicts the corresponding cell cycle profile of cells treated as mentioned above. The graph depicts the mean±d. of a minimum of three independent experiments. (C) U2OS PHD1–GFP cells were transfected with 1 µg of empty vector (Ev) control or the indicated oncogenes for 48 h prior to lysis for western blot analysis. Cell lysates were analysed for the levels of phosphorylated PHD1 at S130 and appropriate controls. The right graph depicts the mean±d. of the quantification of the western blot analysis, representing a minimum of three independent experiments. ***P<0.001 compared to control conditions (Student’s t-test).

**PHD1 S130 is important for the control of the cell cycle**

We previously identified Cep192 as a new PHD1 target (Moser et al., 2013). Cep192 is important for cell cycle progression, in particular for controlling the process of mitosis and spindle assembly (Joukov et al., 2014; Moser et al., 2013). We next determined the impact of PHD1 S130 on cell proliferation (Fig. 6A). In cells depleted of endogenous PHD1 and only expressing GFP, there was no evident cell proliferation. However, this was restored in cells expressing exogenous wild-type PHD1 and also in cells expressing PHD1 with the S130A mutation (Fig. 6A). Interestingly, PHD1-S130D was unable to restore proliferation in cells depleted of endogenous PHD1 (Fig. 6A). This suggests that phosphorylation of S130 in PHD1 alters PHD1 function, either towards Cep192 directly, or to an alternative, as yet unknown, substrate controlling cell cycle progression.

**Fig. 5.** (A) U2OS PHD1–GFP and PHD1-S130A–GFP cells were transfected with 1 µg of empty vector (Ev) control or the indicated oncogenes for 48 h prior to lysis for western blot analysis. The right graph depicts the mean±s.d. of the quantification of the western blot analysis, representing a minimum of three independent experiments. The **p**, phosphorylated form of the protein.

To determine the mechanism behind the loss of PHD1 targeting of HIF1α in PHD1-S130D–GFP cells, we next investigated whether this modification alters the ability of PHD1 to interact with HIF1α in cells. To this end, we treated cells with MG132, to stabilise HIF1α, and immunoprecipitated endogenous HIF1α from all the PHD1 cell lines. Normal IgG was used as a negative control for this approach (Fig. 5B). As can be seen in the left panel of Fig. 5B, there was a substantial immunoprecipitation of HIF1α, with higher levels of HIF1α recovered from the PHD1-S130D–GFP cells. However, the amount of interacting PHD1 was reduced in these cells, when compared with PHD1–GFP cells (Fig. 5B). Interestingly, a reduction in the interaction between HIF1α and PHD1 was also visible in PHD1-S130A–GFP cells. We also analysed this interaction using a transient transfection approach in HEK293 cells (Fig. 5B), again using MG132 treatment to stabilise HIF1α prior to immunoprecipitation. In this system, we could again detect a reduction in the level of PHD1 that interacts with HIF1α in the cells transfected with the PHD1-S130D–GFP construct (Fig. 5B). By contrast, there was no difference between PHD1–GFP- and PHD1-S130A–GFP-transfected cells in the levels of interaction between PHD1 and HIF1α. These results suggest that phosphorylation of PHD1 at S130 disrupts the interaction between HIF1α and PHD1.

We next determined whether the N-terminal region of PHD1 can bind HIF1α. Different PHD1 deletions were created according to the domain structure for PHD1 (Fig. 5C) and the subcellular localisation of these mutants was assessed by fluorescence microscopy. This revealed that all deletion constructs had a nuclear localisation, with the exception of the construct containing the hydroxylase domain only, which was diffusely distributed throughout the cell (Fig. 5C). This is to be expected, as this construct lacks the nuclear localisation signal. Furthermore, we could detect good expression levels of all of the constructs in cells when these were analysed by western blotting (Fig. S3D). Interaction assays with the deletion constructs revealed that both the N-terminal and C-terminal regions of PHD1 were able to interact with HIF1α in cells (Fig. 5D). We also analysed the effect of a phospho-mimicking mutation on the construct containing the hydroxylation domain but lacking the first 100 amino acids. Under these conditions, although the levels of immunoprecipitated HIF1α were similar for both PHD1 constructs, we observed a reduced interaction with the PHD1 construct containing the S130D mutation (Fig. 5E). These results suggest that the N-terminal region of PHD1, and in particular S130, is important for PHD1 binding to HIF1α.
Fig. 4. See next page for legend.
**RESEARCH ARTICLE**

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To determine whether S130 phosphorylation correlated with Cep192 levels, we analysed the synchronisation release samples described in Fig. 3A for the levels of Cep192 (Fig. 6D). There, it was possible to observe that Cep192 levels were inversely correlated with PHD1 phosphorylation at S130 and HIF1α levels (Fig. 6D).

Our results suggest that S130 phosphorylation of PHD1 results in reduced targeting of HIF1α and increased targeting of Cep192 by this enzyme. We next investigated whether alteration in the levels of Cep192 resulted in changes to the levels of HIF1α. To this end, we depleted either Cep192 or the ubiquitin ligase Skp2, which we have previously shown to regulate Cep192 levels in a manner dependent on PHD1 (Moser et al., 2013). Reducing the levels of Cep192 resulted in reduced levels of HIF1α. Conversely, in the absence of Skp2, more Cep192 was present, higher levels of HIF1α were observed (Fig. 6E). Skp2 itself is regulated by hypoxia, due to the fact that cells arrest in G1, and Skp2 levels are regulated by the cell cycle (Ortmann et al., 2014; Yamada et al., 2013). Similar results were also observed in cells overexpressing Cep192 (Fig. S4B). These results suggest that competition between the HIF and Cep192 substrates can occur.

Taken together, we conclude that S130 phosphorylation by interphase CDKs is an important determinant of which substrate PHD1 will target in cells (Fig. 6F).

**DISCUSSION**

In this report, we have identified and characterised the functional significance of a CDK-dependent phosphorylation site on PHD1. Our results show that phosphorylation of PHD1 on S130 is dynamic and regulated by CDK activity, cell cycle stage and oncogenic signals. Interestingly, although phosphorylation of PHD1 on S130 does not alter PHD1 hydroxylase activity in vitro, it regulates PHD1 activity in cells, determining target selection between HIF and Cep192. These results provide a mechanistic link between the cell cycle and the regulation of PHD1 activity in cells, allowing for different functions to be carried out at specific stages of the cell cycle.

Although PHD2 and PHD3 are transcriptional targets of HIF1α (Metzen et al., 2005; Pescador et al., 2005), PHD1 is not induced following HIF1α activation. However, the activity of all of these enzymes is regulated by availability of cofactors, such as molecular oxygen, Fe2+ and α-KG (Fandrey et al., 2006; Kaelin, 2012). Little information exists about how PHD protein levels are regulated. Some studies have demonstrated that PHD3 protein turnover is regulated by the Siah-2 ubiquitin ligase (Nakayama and Ronai, 2004), whereas the FKBP38 protein, a peptidyl-prolyl cis-trans isomerase, regulates the protein stability of PHD2 (Barth et al., 2009, 2007). More recently, PHD3 has been shown to be sumoylated, which although not altering PHD3 hydroxylase activity, was important for PHD3-mediated repression of HIF1α transcriptional activity (Nunez-O’Mara et al., 2015).

Despite these studies, there was no information on how these processes are regulated and how different signalling pathways and/or cellular processes integrate with PHD function. Here, we describe how PHD1 function is regulated by phosphorylation. S130 of PHD1 is well conserved in different mammalian species and, as such, we would predict that PHD1 could be regulated in a similar manner in these species. A closely related site is also present on PHD2, where S125 has been identified as a phosphorylation site in unbiased mass spectrometry screens (Olsen et al., 2010; Zhou et al., 2013). Although no functional characterisation has been done for PHD2, these mass spectrometry screens have suggested that phosphorylation of PHD2 S125 is also regulated by the cell cycle (Olsen et al., 2010; Zhou et al., 2013). However, further research on PHD2 regulation is needed before any conclusion can be made regarding the importance of this phosphorylation site in cells under physiological conditions.

Our data suggest that interphase CDKs (CDK2, CDK4 and CDK6) are involved in the regulation of PHD1 phosphorylation on S130, but not CDK1. Previous data have indicated that CDKs show a degree of functional redundancy and can compensate for each other, with only CDK1 being essential (Diril et al., 2012). Our data supports this view. In fact, we could detect CDK2, CDK4 and CDK6 binding to PHD1, but not CDK1, further indicating that S130 is likely not a CDK1 phosphorylation site in cells. However, CDK1 was still able to phosphorylate PHD1 in vitro. Phosphorylation of this site was induced by serum, and by particular oncogenes, such as Myc and E2F1, both of which are known to regulate the cell cycle (Matsumura et al., 2003; Obaya et al., 2002). This strengthens the notion that phosphorylation of PHD1 at S130 could be of relevance to cancer biology and future research should be directed to investigate this site in the context of this disease.

Although PHD2 is the main regulator of HIF1α, PHD1 is also involved (Appelhoff et al., 2004). Our results show that a phosphomimicking mutation of S130 (S130D), reduces PHD1 activity towards HIF1α, leading to increased HIF1α half-life and activity. Interestingly, this S130D mutation did not alter hydroxylase activity in vitro, when recombinant protein was analysed, but instead altered the interaction between PHD1 and HIF1α in cells. This could be due either to inability to bind a cofactor in cells or to a direct interference of the phosphorylation site with binding to HIF1α. Our analysis revealed that the N-terminal region of PHD1 is able to bind HIF1α without the help of the hydroxylation domain, suggesting that this...
latter point might be the case. The N-terminus of PHD2 has also been shown to be important for the regulation of HIF1α by PHD2, although in this case through an indirect mechanism involving a chaperone protein (Song et al., 2013). This highlights the fact that other domains of PHD enzymes can contribute to the hydroxylase activity of these enzymes in cells.

Fig. 5. PHD1 phosphorylation at Serine 130 alters the ability of PHD1 to target HIF1α. (A) U2OS PHD1–GFP, PHD1-S130A–GFP and PHD1-S130D–GFP cells were exposed to 1% O2 for 4 h prior to treatment with cycloheximide for the indicated periods of time. Whole-cell lysates were analysed by western blotting for the levels of HIF1α and appropriate controls. Western blots were quantified and the graph depicts mean±s.d. of a minimum of three independent experiments. (B) U2OS GFP, PHD1–GFP, PHD1-S130A–GFP and PHD1-S130D–GFP cells were treated with MG132 for 3 h prior to lysis. 300 µg of cell extracts were used to immunoprecipitate (IP) HIF1α, with normal mouse IgG used as a control. Precipitated material was analysed by western blotting for the indicated proteins. Western blots were quantified, and the graph depicts the mean±s.d. of a minimum of three independent experiments. (C) Schematic diagram of the PHD1 expression constructs used in this study. Highlighted are the nuclear localization signal (NLS), S130 and the hydroxylase domain (HD). (D) HEK293 cells were transfected with 1 µg of the indicated expression constructs for 48 h prior to treatment with MG132 and processed as in B. * non specific band. (E) HEK293 were transfected with 1 µg of the indicated expression constructs for 48 h prior to treatment with MG132 and processed as in B. Western blots were quantified, and the graph depicts mean±s.d. of a minimum of three independent experiments. *P<0.05, **P<0.01 and ***P<0.001 compared to control conditions (Student’s t-test). See also Fig. S3.
Recently, we have identified the centrosomal protein Cep192 as a target for PHD1 in cells (Moser et al., 2013). Cep192 requires a precise regulation of its expression level to allow for centrosome duplication and maturation (Joukov et al., 2014; Moser et al., 2013). As such, either too much or too little Cep192 results in a similar defect in centrosomes and causes cell cycle arrest. Given the finding...
Fig. 6. S130 of PHD1 is important for PHD1-mediated control of cell proliferation. (A) U2OS GFP, PHD1–GFP, PHD1–S130A–GFP and PHD1–S130D–GFP cells were transfected with siRNA oligonucleotides targeting the 3′UTR of endogenous PHD1 prior to proliferation being assessed. Total cell numbers were counted, and the graph depicts means±s.d. of a minimum of three independent experiments. Data were normalised to proliferation in GFP cells and expressed as a percentage. (B) U2OS GFP, PHD1GFP, PHD1–S130A–GFP and PHD1–S130D–GFP cells were transfected with siRNA oligonucleotides targeting the 3′UTR of endogenous PHD1 prior to fixation and immunostaining for Cep192 and pericentrin. Scale bars: 2 µm. Graph depicts box-and-whisker plots for Cep192 and Pericentrin intensity. Box-and-whisker plot, middle line shows the median value; the bottom and top of the box show the lower and upper quartiles (25–75%); whiskers extend to 10th and 90th percentiles, and all outliers are shown. n=22–38 cells per condition. (C) U2OS GFP, PHD1GFP, PHD1–S130A–GFP and PHD1–S130D–GFP cells were treated with MG132 for 3 h prior to lysis. 300 µg of cell extracts were used to immunoprecipitate (IP) Cep192, with normal mouse IgG used as a control. Precipitated material was analysed by western blotting for the indicated proteins. Western blots were quantified and the graph depicts means±s.d. of a minimum of three independent experiments. (D) Cell extracts from Fig. 3A were analysed by western blotting for the levels of Cep192 and HIF-1α. G1, S, and G2/M are the phases of the cell cycle that correspond to the indicated time points. (E) U2OS were transfected with the indicated siRNAs prior to treatment with 1% O2 for 24 h. Whole-cell lysates were analysed by western blotting using the depicted antibodies. (F) Schematic diagram for the proposed model with 1% O2 for 24 h. Whole-cell lysates were analysed by western blotting for the indicated proteins.}

Presented here concerning the regulation of PHD1 function by phosphorylation, we investigated how S130 phosphorylation of PHD1 impinged on cell cycle progression and Cep192 levels. In rescue experiments, where endogenous PHD1 was depleted, exogenous expression of either wild-type PHD1 or the unphosphorylatable S130A PHD1 mutant meant that cells were able to proliferate, and both proteins restored Cep192 levels. However, the exogenous expression of the phospho-mimic mutant S130D was unable to restore proliferation and resulted in reduced levels of Cep192 when compared with control cells. In addition, we observed increased interaction between PHD1 and Cep192 in the phospho-mimic mutant S130D cells. This suggests that PHD1 at phosphorylated S130 has increased activity towards Cep192, corroborating the notion that phosphorylation of this site can alter the target specificity of PHD1. This provides a mechanism by which PHD1 function could be changed throughout the cell cycle, directing it towards specific targets in response to signals, such as high interphase CDK activity, for example.

Taken together, our results suggest a new paradigm for the regulation of PHD1 function by post-translational modifications. We have focused on characterising the functional consequences of PHD1 phosphorylation on S130 due to its conservation across species, and because of the presence of a similar site on PHD2. Despite this conservation, the effects of phosphorylation of PHD1 at S130 will probably only alter HIF levels, as the Cep192 hydroxylation site is not conserved in mice (Moser et al., 2013). However, additional post-translational modifications might also occur in these enzymes that help regulate activity and control the targeting of PHDs to specific substrates in cells. Further investigation will reveal whether this is the case.

**MATERIALS AND METHODS**

**Cells**

U2OS osteosarcoma cancer cells and HEK293 human embryonic kidney cells were obtained from the European Collection of Cell Cultures and grown in Dulbecco’s modified Eagle medium (Lonza) supplemented with 10% fetal bovine serum (Gibco), 50 units/ml penicillin (Lonza) and 50 µg/ml streptomycin (Lonza) for no more than 30 passages at 37°C and 5% CO₂. Stable U2OS cell lines expressing GFP, PHD1–GFP, PHD1–S130A–GFP, and PHD1–S130D–GFP were maintained with 400 µg/ml G418. U2OS-HRE-luciferase cells were maintained in 0.5 µg/ml puromycin. Cells were routinely tested for contamination.

**Plasmids**

GFP-N1 was obtained from Clonetech. GFP–PHD1 was a kind gift from Eric Metzen (Essen University, Essen, Germany) and was used as a template to create the GFP-tagged PHD1–S130A and PHD1–S130D mutations by site directed mutagenesis. For the truncation mutants of PHD1, GFP–PHD1 and GFP–PHD1–S130D plasmids were used as templates. Primer sequences are available upon request.

**CMV-HIF1α and CMV-Sce expression constructs** were obtained from Origene. E2F1 and wild-type Myc and mutant expression constructs were a kind gift from Victoria Cowling (University of Dundee, Dundee, UK), HA–CDK1 (1888, Addgene), HA–CDK4 (1876, Addgene) and HA–CDK2 (1884, Addgene) were as previously described (van den Heuvel and Harlow, 1993).

**Hypoxia induction and chemical treatments**

Cells were incubated at 1% O₂ in an *in vivo* hypoxia workstation (Ruskin, UK). Cells were lysed for protein extracts and RNA extraction in the work station to avoid re-oxygenation. Whole-cell lysates were obtained using Triton lysis buffer (20 mM Tris-HCl pH 7.5, 150 mM NaCl, 1% Triton X-100, 250 mM NaVO₄, 10 mM NaF and phosphatase inhibitors 1 tablet, 10 ml (Roche Complete)). RNA was extracted using Peqlab Total RNA Kit (Peqlab) as per the manufacturer’s instructions.

MG132 was obtained from Merck/Millipore and used at the final concentration of 20 µM for 3 h. Desferroxamine mesylate (DFX) was obtained from Sigma and used at the final concentration of 200 µM.

**In vitro hydroxylation assay**

Hydroxylation assays were performed as in Moser et al. (2013).

**Kinase assays**

Kinases assays were performed as described in DeGregori et al. (1995). 2 µg of recombinant PHD1 protein was used per reaction.

**Antibodies**

Antibodies were against the following proteins: phospho-serine-CDK substrate (2324, Cell Signaling), 1:1000; phospho-threonine (9386, Cell Signaling), 1:1000; phospho-serine (05-1000, Millipore), 1:1000; HIF-1α (610958, BD Biosciences and sc-53546 Santa Cruz Biotechnology), 1:1000; HO-HIF1α (3434, Cell Signaling), 1:1000; HIF-2α (PA1-16510, Thermo Scientific), 1:1000; HIF-1β (#3718, Cell Signaling), 1:1000; β-actin (3700, Cell Signaling), 1:5000; CAIX (NB100-417, Novus Biologicals), 1:1000; HK2 (2867, Cell Signaling), 1:1000; BNP3 (ab10433, Abcam), 1:2000; GLUT3 (53520, Anaspec), 1:1000; PHD3 (A300-327A, Bethyl labs), 1:1000; PHD1 (Bethyl A300-326A; Novus NBP1-40773), 1:1000; PHD2 (Bethyl A300-322A), 1:1000; Cep192 (Bethyl A302-324A; Novus NBP-84634), 1:1000; H3K4me2 (AbCam), 1:2000; HDAC1 (1-10199, Millipore), 1:2000; CDK1 (9116, Cell Signaling), 1:1000; CDK2 (2546, Cell Signaling), 1:1000; CDK4 (sc-260, Santa Cruz Biotechnology; 2906, Cell Signaling), 1:1000; CDK6 (sc-203, Santa Cruz Biotechnology; 3136, Cell Signaling), 1:1000; phospho-ERK1/2, 1:1000; cyclin A (sc-53546 Santa Cruz Biotechnology), 1:1000; cyclin E (4129, Cell Signaling), 1:1000; E2F1 (3742, Cell Signaling), 1:1000; Src (2109, Cell Signaling), 1:1000; cMyC (gift from Victoria Cowling), 1:2000; p53 (2524, Cell Signaling), 1:2000; pericentrin (28144, Abcam), 1:100; phospho-S130-PHD1 [produced in rabbits by immunisation with phosphopeptide for S130 of PHD1 (CEDGGDAPSpHPSKR) and purified by Dundee Cell products], 1:200.

**Immunoprecipitation of PHD1 and sample preparation for LC-MS analysis**

PHD1–GFP cells were lysed as described previously (Moser et al., 2013) and PHD1 was immunoprecipitated using GFP-TRAP® magnetic beads (ChromoTek).

Immunoprecipitation eluates were separated on 1D SDS PAGE gels and stained (SimplyBlue; Invitrogen). The protein bands of interest were
excised, chopped into ∼1-mm² 1-mm pieces and destained at room temperature [2×30 min in 50:50, acetonitrile (ACN) and 100 mM triethylammonium bicarbonate buffer (TEAB) pH 8.5]. After 15 min dehydration in 100% ACN, proteins in the gel pieces were reduced by incubation in 25 mM tris(2-carboxyethyl)phosphine (TCEP) in 100 mM TEAB for 15 min at 37°C and alkylated by adding iodoacetamide to a final concentration of 50 mM and incubating in the dark at room temperature for 30 min.

After reduction and alkylaion, the gel pieces were washed with 50:50 acetonitrile and TEAB to remove excess iodoacetamide, dehydrated in acetonitrile then dried in a vacuum to remove residual organic solvent prior to digestion. For tryptic digestion, the dried gel pieces were rehydrated using sequencing grade modified trypsin (Promega) solution (15 µl, 1 ng µl⁻¹ in TEAB). Digestion was performed overnight at 37°C in 50 µl TEAB.

Digested peptides were extracted by adding 1% formic acid in acetonitrile (50 µl) to the gel pieces and incubating for 20 min at room temperature. The supernatant, now containing tryptic peptides, was transferred to a clean tube. The gel pieces were extracted further with two washes with 100 µl water: supernatant, now containing tryptic peptides, was transferred to a clean tube.

Author contributions
B.O. performed the majority of the experiments and analysed the data. D.B. performed the mass spectrometry analysis and analysed the data. S.C. performed the analysis of Cep192 and Pericentrum levels. S.C.M. helped with the mass spectrometry analysis. S.M. created stable cell lines. E.R.G., A.L.I. and J.R.S. helped writing the manuscript. S.R. conceived the project, performed experiments and wrote the manuscript.

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Supplementary information
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