Mutational analysis of Polycomb genes in solid tumours identifies PHC3 amplification as a possible cancer-driving genetic alteration

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Background: Polycomb group genes (PcGs) are epigenetic effectors implicated in most cancer hallmarks. The mutational status of all PcGs has never been systematically assessed in solid tumours.

Methods: We conducted a multi-step analysis on publically available databases and patient samples to identify somatic aberrations of PcGs.

Results: Data from more than 1000 cancer patients show for the first time that the PcG member PHC3 is amplified in three epithelial neoplasms (rate: 8–35%). This aberration predicts poorer prognosis in lung and uterine carcinomas (P<0.01). Gene amplification correlates with mRNA overexpression (P<0.01), suggesting a functional role of this aberration.

Conclusion: PHC3 amplification may emerge as a biomarker and potential therapeutic target in a relevant fraction of epithelial tumours.

Epigenetic aberrations have been widely reported in human neoplasms, and preclinical studies have established conclusive links between epigenetics and most cancer hallmarks (Baylin and Jones, 2011). Despite this, the only epigenetic drugs approved for clinical use are DNA-methyltransferase and histone-deacetylase inhibitors, whose employment is restricted to a few haematological malignancies (Piekarz and Bates, 2009). Even the clinical exploitation of epigenetic biomarkers as prognostic/predictive factors is currently limited to a few notable exceptions (Wick et al, 2009).

Polycomb group genes (PcGs) are epigenetic effectors organised into two main repressive complexes (PRC1, PRC2). The PRCs silence tumour-suppressor genes through histone modifications, thereby driving cancer cell proliferation, metastasis and drug resistance (Crea et al, 2012). Despite a plethora of preclinical studies demonstrating their crucial role in human neoplasms, PcGs still lack clinical value. Most molecular research has focused on the histone-methyltransferase EZH2 (PRC2 member), which acts as an oncogene in several neoplasms (Chase and Cross, 2011). To the contrary, the role of PRC1 in many solid tumours is still largely overlooked.

The recent identification of EZH2-specific inhibitors has been welcomed as a fundamental advancement in epigenetic therapy (Melnick, 2012). Those molecules are particularly effective on lymphoma cells harbouring EZH2-activating mutations, but not on cells with simple mRNA upregulation (McCabe et al, 2012). In addition, inhibition of the catalytic subunit of PRC1 induces cancer cell apoptosis (Wen et al, 2013), thereby paving the way for
the development of PRC1-targeting drugs. As PcG mutations have been described almost exclusively in haematological malignancies (Chase and Cross, 2011), there is currently no genetic evidence to predict their efficacy in solid tumours. Most reports on PcG status in solid neoplasms are restricted to gene expression analysis of a few popular targets (EZH2, BMI1), overlooking the evidence that human PcGs are encoded by at least 18 different genes (Christophersen and Helin, 2010), and not taking into account possible mutational events. For this reason, we explored publically available databases and patient data sets to investigate the mutational status and clinical significance of all PcGs in solid tumours.

METHODS

We conducted a two-step analysis on publically available databases. First, we queried the cBio database for somatic aberrations (gene amplification, insertion, deletion, point mutation) in each PcG (listed in Christophersen and Helin (2010), Supplementary Table 1). This database includes mutational data from exome sequencing, copy number calls from GISTC 2.0, and for some studies, mRNA levels expressed as Z score from RNA seq (Cerami et al, 2012). The cBio portal provides access to more than 5000 samples from over 20 cancer studies. Herein we report results only for those genes showing a mutational rate higher than 5%. We restricted our analysis to studies available for publication (bladder, breast, ovarian, glioblastoma, lung, prostate, and uterine carcinomas). To confirm our findings in an independent data set, we selected the most frequently mutated gene (PHC3), and investigated its clinical relevance in both cBio and Oncomine (Rhodes et al, 2007) databases (more than 100 additional data sets, more than 10000 patients screened), searching for correlations between gene expression/amplification and clinico/pathological characteristics in lung, uterus, and ovarian neoplasms. To further reduce the false discovery rate, we considered significant results with P-value<0.01 and fold-change >2.0. Amplification and overexpression of PHC3 in lung cancer was assessed in an additional data set comprised of 169 lung adenocarcinoma (AC) and 92 lung SCC samples, of which 35 AC and 13 SCC tumours had patient-matched normal tissue (GSE31800) (Starczynowski et al, 2011).

RESULTS

Our global mutational analysis is summarised in Figure 1A. We found no relevant deletions and only two missense mutations with a frequency higher than 5%. To the contrary, several loci were amplified in at least one investigated neoplasm. According to previous reports (Chase and Cross, 2011), EZH2 is not mutated in most common solid tumours, although we found an interesting rate of EZH2 amplification (5.6%) in ovarian serous carcinoma. This neoplasm shows relevant amplification of four different PcGs. Gene amplification was significantly correlated with higher mRNA levels (Supplementary Figure 1), thereby suggesting that these epigenetic effectors are crucial for its development.

The most striking result arising from our query was the high rate of PHC3 genetic amplifications. This aberration was found in lung (34.8%), ovarian (20.6%), and uterine ACs (7.7%). In order to explore the functional role of this genetic aberration, we computed the correlation between gene amplification and mRNA upregulation in those three neoplasms. Notably, gene amplification was significantly correlated with mRNA upregulation in each tumour type (Figure 1B–D).

Oncomine data confirmed that PHC3 is selectively upregulated in SCC, compared with other non-small-cell lung cancer subtypes (Figure 2A, \( P = 1.32 \times 10^{-11}\), fold-change = 2.137). Furthermore, PHC3 overexpression was found to be associated with shorter
3-year survival rate in lung cancer (Figure 2B, \( P = 0.005 \), fold-change = 2.267, Oncomine analysis). PHC3 mRNA was also upregulated in cervical, colorectal, gastric, and prostate ACs, compared with normal tissues (\( P < 0.01 \), fold-change > 2.0, Oncomine data not shown).

PHC3 is located at 3q62.2, a genomic region that has been described as amplified in solid tumours (Lavigne et al., 2004; Linxweiler et al., 2012). For this reason, we investigated amplification and expression level of eight genes flanking the PHC3 locus (Figure 2C). The rate of genomic amplification was very similar for all eight genes (34–35% in lung cancer, data not shown). However, only two genes (PHC3 and PRKCI) showed mRNA upregulation in more than 50% of tumours with genomic amplification, suggesting that the genetic aberration has a functional role in those loci. The remaining six genes showed an mRNA upregulation rate lower than 35%, even in the presence of an amplified gene.

We confirmed the high frequency of PHC3 amplification and overexpression in lung SCC using an additional publicly available data set of 169 AC and 92 SCC tumours. PHC3 was amplified in 71 out of 92 (77%) lung SCC samples and 36 out of 169 (21%) AC and overexpressed in 7 of 13 (54%) and 5 of 35 (14%) SCC and AC cases, respectively. Moreover, comparison of mRNA expression levels between AC and SCC revealed no difference in PHC3 mRNA levels in non-malignant tissue (Figure 2D, \( P = 0.76 \)), but significantly higher expression in SCC tumours relative to AC tumours (Figure 2E, \( P = 0.002 \)) and SCC tumours relative to matched non-malignant tissue (Figure 2F, \( P = 0.0024 \)).

A very interesting clinical correlation was identified in uterine carcinoma. In this neoplasm, PHC3 amplification also predicted shorter disease-free survival (Supplementary Figure 2). Notably, PHC3 amplification was detectable only in grade 3 neoplasms. Despite this, our analysis indicates that PhC3 amplification is an independent prognostic factor, even when contrasted with tumour grade (Table 1). To the contrary, PhC3 amplification was not significantly associated with prognosis in ovarian cancer (Supplementary Figure 3).

### DISCUSSION

Taken together, our analyses of data from three different data sets indicate that the PHC3 gene is amplified in three epithelial tumours, with percentages ranging from 8 to 35%. Gene

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**Table 1. Multi-parametric analysis of PHC3 amplification (uterine carcinoma)**

| Parameter | P-value | Odds ratio (95% CI) |
|-----------|---------|---------------------|
| PHC3 status | | |
| Non-amplified | | |
| Amplified | 0.0157 | 4.0 (1.3–12.3) |
| Grade | | |
| 1 | 0.99 | NC |
| 2 | 0.99 | NC |
| 3 | 0.99 | NC |

Abbreviations: CI = confidence interval; NC = not computable. Cox regression analysis was performed to determine whether PhC3 mutational status was an independent predictor of progression-free survival.
amplification is highly correlated with mRNA overexpression and, at least for lung and uterine cancer, associated with poorer prognosis. PHC3 is rarely deleted and almost never downregulated in cancer vs normal tissue. Oncomine analysis revealed that cancer-specific PHC3 upregulation might be a common phenomenon in several epithelial neoplasms. To the best of our knowledge, PHC3 genomic amplification and mRNA upregulation have never been described before. However, PHC3 has been found to be downregulated in human sarcomas, where it acts as a tumour suppressor (Deshpande et al., 2007).

PHC3 is a member of PRCl, which catalyses histone H2A ubiquitination. PRCl is composed of RING1B (catalytic subunit) and other proteins responsible for interactions with transcription factors, epigenetic regulators, and DNA sequences. PhC proteins bridge distant chromatin templates, thereby recruiting PRCl to selected loci (Lavigne et al., 2004). Currently, three PhC isoforms are known, with different tissue distribution and partially overlapping functions (Deshpande et al., 2007). PRCl has been implicated in tumorigenesis and progression of several cancer types. Tobacco smoke is able to activate PRCl genes, thereby increasing the tumorigenicity of lung cancer cells (Hussain et al., 2009). Amplification of the chromosomal region harbouring the PHC3 locus (3q26) has been described in preneoplastic and invasive lung SCC, and its rate increases with disease progression (Pelosi et al., 2007). Future molecular studies should dissect the role of PHC3 in lung tumorigenesis.

As genomic amplification can occur at a multigenic scale, we investigated if PHC3 copy number gain was associated with similar alteration in nearby loci. Our results indicate that the PHC3-proximity region is frequently amplified in solid tumours. However, when we contrasted mRNA expression with genomic amplification, we observed that only PHC3 and PRKCI mRNA levels were frequently upregulated in tumours with copy number gain (Figure 2D). The remaining six genes did not show a correlation between copy number gain and increased gene expression. Interestingly, PRKCI (protein kinase C iota) has been described as a potential oncogene in lung cancer (Regala et al., 2009; Linxweiler et al., 2012). Our data indicate that copy number gain may have a functional role in the PHC3 locus too, and that PHC3 upregulation is functionally relevant for cancer cells. In conclusion, we identified for the first time that PHC3 copy number gain as a novel and possibly clinically significant genetic alteration in three common epithelial neoplasms (lung SCC, uterine carcinoma, and ovarian serous carcinoma). Future molecular studies should dissect the role of PHC3 in each specific context. However, our data indicate that PHC3 may emerge as a novel oncogene, and possibly a prognostic marker and a target for epigenetic therapy, in epithelial neoplasms.

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REFERENCES

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