The Rare YAP1 Subtype of SCLC Revisited in a Biobank of 39 Circulating Tumor Cell Patient Derived Explant Models: A Brief Report

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ABSTRACT

Introduction: Recent consensus defines four SCLC subtypes on the basis of transcription factor expression: ASCL1, NEUROD1, POU2F3, and YAP1. The rare YAP1 subtype is associated with “neuroendocrine (NE)-low” cells among SCLC cell lines and patient samples. We evaluated YAP1 in 39 patients with phenotypically diverse circulating tumor cell–derived explant (CDX) models and revisited YAP1 in terms of prevalence, cell phenotype, and intertumor and intratumor heterogeneity.

Methods: YAP1 transcript and protein expression were assessed by RNA sequencing and immunohistochemistry or multiplexed immunofluorescence of NE and non-NE CDX subpopulations. Physically separated NE and non-NE CDX ex vivo culture lysates were Western blotted for YAP1, NE marker SYP, and AXL.

Results: RNA sequencing normalized for the four subtype transcription factors identified YAP1 expression in 14 of 39 CDX. A total of 10 CDX expressed YAP1 protein, and eight had strong YAP1 expression confined to rare non-NE cell clusters. This was confirmed in ex vivo CDX cultures in which adherent non-NE cells lacking SYP expression expressed YAP1. However, in two CDX, weaker cellular YAP1 expression was observed, widely dispersed in SYP-positive NE cells.

Conclusions: YAP1 was predominantly expressed in non-NE cell clusters in SCLC CDX, but two of 39 CDX expressed YAP1 in NE cells. CDX22P, with relatively high YAP1 expression, is an ASCL1 NE subtype with a low NE score and an outlier within this subtype in our CDX biobank. These descriptive data reveal subtly different YAP1 expression profiles, paving the way for functional studies to compare YAP1 signaling in non-NE and low NE cell contexts for potentially personalized therapeutic approaches.

Keywords: Small cell lung cancer (SCLC); YAP1; Neuroendocrine; Nonneuroendocrine; Intratumoral heterogeneity

Introduction

SCLC is an aggressive neuroendocrine (NE) cancer with poor prognosis, characterized by high circulating tumor cell (CTC) burden and early widespread
metastasis. Treatment for SCLC remained unchanged for decades until the recent incorporation of immune checkpoint inhibitors into first-line treatment in the United States, benefiting a poorly defined subset of patients. Effective targeted therapies for patients with SCLC remain urgently required. Emerging evidence suggests that characterization of SCLC molecular heterogeneity will identify specific therapeutic vulnerabilities within patient subtypes to enable biomarker-driven patient stratification to improve patient outcomes. Four SCLC subtypes have been proposed on the basis of expression of key transcriptional regulators in patient tumors, patient-derived preclinical models, and human cell lines: ASCL1 and NEUROD1, both master regulators of NE phenotypes; POU2F3 that identifies a tuft cell of origin; and YAP1, a regulator of Hippo signaling.

RNA sequencing (RNaseq) of 38 SCLC CTC–derived explants (CDXs) identified models representing three of the four consensus SCLC subtypes (ASCL1, NEUROD1, and POU2F3). We discovered a further subtype on the basis of the expression of the neurogenic transcription factor (TF) ATOH1, but our analysis did not reveal the YAP1 subtype. Four studies collectively reported the rarity of the YAP1 subtype, in which YAP1 was expressed in five of 126 tumors and seven of 51 SCLC cell lines with low or absent NE marker expression, a low NE score, and a variant, loosely adherent morphology.

The tumor-suppressive Hippo pathway controls cell proliferation, apoptosis, and organ size. When Hippo signaling is active, YAP1 is inactivated and sequestered in the cytoplasm for degradation. When Hippo signaling is inactive, YAP1 binds Transcriptional Enhanced Associated Domain family nuclear TFs to direct prosurvival gene expression, proliferation, and tissue growth. YAP1 nuclear activity correlates with chemoresistance, cancer stem cell renewal, and metastasis. YAP1 also responds to extracellular matrix stiffness and mechanical cues that promote cancer cell motility. Given these “hallmarks of cancer” YAP1 functions, we revisited YAP1 RNA and protein expression across 39 CDX models, evaluating the prevalence, cell phenotype, and tumor heterogeneity to provide a characterization of YAP1 at single-cell resolution.

Materials and Methods

CDX generation, RNaseq, and immunohistochemistry (IHC) were performed as described previously. CDX models were generated from patients’ CTCs enriched from blood samples pre-chemotherapy baseline or at posttreatment disease progression time points (designated P or PP). Transcriptomic analysis was performed with amendments to the previously described alignment (NF-core RNaseq pipeline with Spliced Transcripts Alignment to a Reference aligner) and annotation (mapped to eEnsembl version 99).

Three independent tumors per CDX were used for all analyses. Antibodies for IHC were the following: (1) YAP1 (1:100, ab52771; Abcam, Cambridge, United Kingdom); (2) REST (1:150, MA5-24606; ThermoFisher Scientific, Waltham, MA); and (3) SYP (pA0299; Leica Biosystems, Wetzlar, Germany). Digitally scanned slides were analyzed using HALO software (Akoya Biosciences, Menlo Park, CA); expression levels were reported as a percentage of positive cells within each tissue slice. Multiplex immunofluorescence was performed using PerkinElmer Opal 4-Color Automation IHC Kit (NEL800001KT) (PerkinElmer, Waltham, MA) and quantified using HALO.

For ex vivo analysis, CDX tumors were disaggregated, cultured, and lysates immunoblotted as previously described using the aforementioned antibody to YAP1 (1:1000), REST (1:500, LS-C668231; Lifespan Biosciences, Seattle, WA), SYP (1:20,000, ab32127; Abcam, Cambridge, United Kingdom), AXL (1:500, C89E7; Cell Signalling Technology, Danvers, MA), and tubulin (1:1000, 2144S, Cell Signalling Technology). CDX cultures were treated with a titration of cisplatin concentrations for 5 days, followed by cell number analysis using Cell-Titer Glo 3D viability assay (G9683; Promega, Madison, WI).

Results

YAP1 Is Expressed in SCLC CDX

Our biobank of 38 SCLC CDX recapitulates the ASCL1, NEUROD1, and POU2F3 subtypes. We have generated an additional model, CDX31P, belonging to the ASCL1 subgroup (Supplementary Data 1) and derived from a patient with extensive disease, postchemotherapy. Although unbiased clustering of CDX RNaseq data did not identify the YAP1 subtype, normalization of transcript levels (variance stabilizing transform) for each TF identified 14 CDX models with detectable YAP1 transcript, albeit at relatively low levels compared with ASCL1, NEUROD1, and POU2F3 (Fig. 1A) (seven of 14 CDX variance stabilizing transforms > 6.5). Of note, our analysis revealed a bisection of the ASCL1 subtype with detectable YAP1 expression in one subset; two of the four ATOH1 CDX models expressed YAP1, and neither of the NEUROD1 or POU2F3 models were YAP1 expressers. YAP1 protein was expressed in more than 0.2% cells in 10 of 39 CDX models (dotted line, Fig. 1B), comprising eight of 14 models identified by RNaseq and an additional two models (Fig. 1B). Overall, there was good...
concordance between RNAseq and IHC; analyses were performed on independent tumor replicates, likely explaining the lack of concordance for rare cells in some models. YAP1 expression was comparable by IHC in paired CDX models generated at prechemotherapy baseline and at posttreatment disease progression (Supplementary Data 2). Concordant with RNAseq, CDX22P had the highest YAP1 expression (9% cells) (Fig. 1B), and the YAP1 expressing cells were dispersed throughout the tumor, a distribution also noted in CDX04 (Fig. 1C). In the remaining models, YAP1 was observed within cell clusters.

**Figure 1.** YAP1 RNA and protein expression in CDX. (A) A heatmap illustrating relative RNA transcript expression of five SCLC TFs across the CDX biobank with YAP1-expressing models marked with an asterisk (*) and the VST values depicted for each. The average expression is illustrated for three independent CDX tumor replicates per model. (B) Immunohistochemical analysis of YAP1 protein expression in CDX (three independent tumors per CDX). The dashed line represents a cutoff of YAP1 expression in more than 0.2% cells. (C) Representative images of YAP1 expression by IHC (brown stain) in selected CDX models. Scale bars = 100 μm. CDX, circulating tumor cell-derived explant model; IHC, immunohistochemistry; TF, transcription factor; VST, variance stabilizing transform.

**YAP1 Expression in Non-NE Cell Clusters**

YAP1 RNA was expressed in seven of 51 SCLC cell lines, which also had low or absent NE marker expression. Therefore, we evaluated YAP1 protein alongside established SCLC NE and non-NE markers by IHC on serial tissue sections (Fig. 2). Where YAP1 expression was confined to cell clusters, it colocalized with REST, a known repressor of NE differentiation in SCLC, concordant with reduced expression of the SCLC diagnostic NE marker SYP, indicating non-NE cell expression of YAP1 (Fig. 2A). In CDX04 and CDX22P, YAP1 was expressed in non-NE cell clusters (Fig. 2A) and
throughout the tissue within NE cells (REST-low and SYP-positive) (Fig. 2B).

Multiplex immunofluorescence and quantification of YAP1 and REST colocalization was carried out in seven CDX models representing a range of YAP1 and REST expression profiles (Supplementary Data 3 and Fig. 3A). In CDX models in which YAP1 expression was less than 2% and expressed in cell clusters (CDX15P, CDX15PP, CDX17, CDX30P, and CDX42P), 57% of YAP1 cells were non-NE and colocalized with REST (Fig. 3B), despite the low abundance of REST-positive (RESTpos) cells (mean 15.5%). When YAP1 expression was greater than 3% and dispersed throughout the tumor (CDX04 and CDX22P), less YAP1-REST dual positivity was observed (mean 10%), indicating that 90% of YAP1-expressing cells were non-NE, whereas suspension cells were RESTneg and SYPpos, consistent with a NE phenotype (Fig. 3C). YAP1 was expressed in all non-NE subpopulations of CDX ex vivo cultures and by the NE cells from CDX04 and CDX22P that expressed relatively low levels of SYP (Fig. 4C). YAP1 cellular localization is dynamic, making a robust assessment of nuclear localization difficult in fixed tissue (Fig. 3C). However, the YAP1 downstream target AXL receptor tyrosine kinase was expressed in the YAP1pos cells only (Fig. 4C), and a YAP transcriptional target signature17 was enriched in CDX17 and CDX30P non-NE cells compared with NE cells (Fig. 4D), consistent with YAP1 nuclear activity. Concordant with a previous study,8 RNAseq of the physically separated non-NE and NE cells revealed reciprocal expression of YAP1 with three established NE markers (INSM1, SYP, CHGA) in non-NE CDX17 and CDX30P cells (Fig. 4E). In these CDX, YAP1, REST, VIM, CD44, and MYC were more highly expressed in the non-NE cell subpopulation, which also expressed the previously defined YAP1 SCLC gene signature (Supplementary Data 4).8 Overall, these data indicate that YAP1 expression is predominantly in non-NE cells but can also be present at lower levels in NE-low cells. Consistent with previous reports of increased chemoresistance in non-NE compared with NE cells in an SCLC genetically engineered mouse model15 and the association of YAP1 expression and chemoresistance,8,13 YAP1pos non-NE cells in CDX30P and CDX31P ex vivo cultures were fivefold to 7.5-fold more resistant than YAP1neg NE cells to cisplatin (Fig. 4F).

Figure 2. YAP1 protein expression in non-NE cell clusters. (A) Immunohistochemical analysis in serial CDX tissue sections showing clusters of YAP1 expressing cells colocalized with non-NE marker REST and with reduced expression of NE marker SYP. (B) Diffuse YAP1 expression in regions of CDX04 and CDX22P with low or absent REST and with low expression of SYP. Representative images are depicted from the staining of three biological replicates per CDX. Scale bars = 50 μm. CDX, circulating tumor cell-derived explant model; NE, neuroendocrine.
Discussion

Classification of SCLC as a recalcitrant cancer in 2012 by the U.S. government provoked a resurgence of SCLC research focused on developing and interrogating patient-derived preclinical models to more accurately reflect inter-tumoral and intra-tumoral heterogeneity.\(^4\) Identification of SCLC subtypes\(^4,5\) is beginning to reveal specific molecular vulnerabilities\(^6\) and identify novel targets and innovative biomarker-driven treatment strategies. YAP1 is a key mediator of the tumor-suppressive Hippo pathway,\(^5\) and although its expression defines one consensus SCLC subtype,\(^5,8\) there were only five studies on YAP1 in SCLC at time of writing. Although the YAP1 subtype was not represented within our published biobank of 38 CDX on the basis of unbiased hierarchical clustering of bulk RNAseq data,\(^4\) YAP1 transcript or protein was detectable in 16 CDX (Fig. 1).

Figure 3. YAP1 and REST multiplex immunofluorescence in CDX. (A) Multiplex immunofluorescence assay illustrating representative images of YAP1 (yellow) and REST (pink) expression with DAPI stained nuclei (blue). Scale bars = 10 μm. (B) Quantification of multiplex immunofluorescence showing the percentage of YAP1, REST, and dual YAP1 and REST-positive cells for each CDX. Bimodal YAP1 and REST colocalization is indicated by two means for % YAP1 cells coexpressing REST (57%, CDX15P, CDX15PP, CDX17, CDX30P, CDX42P; 10%, CDX04, CDX22P). Whole-tumor sections from three to four independent mice per CDX were analyzed and plotted individually. (C) YAP1 subcellular localization calculated for each immunofluorescence assay represented as % of cells with YAP1 expression in the nucleus and cytoplasm. The mean is plotted from three to four independent tumor replicates, and error bars represent SEM. (D) Intensity of YAP1 fluorescence in YAP1-positive and REST-negative (NE) cells versus YAP1-positive and REST-positive (non-NE cells) in CDX. Whole-tumor sections from three independent mice per CDX were analyzed. CDX, circulating tumor cell–derived explant; DAPI, 4',6-diamidino-2-phenylindole; NE, neuroendocrine.
However, expansion of the rare non-NE subpopulation by culture ex vivo resulted in detectable YAP1 protein (Fig. 4).

CDX04 and CDX22P had the lowest NE score and highest YAP1 expression, and in these models, NE and non-NE cell ex vivo cultures expressed YAP1 with absent or low SYP. CDX22P is a rare ASCL1 subtype with a low NE score (0.28) (Supplementary Data 3 and Simpson et al.4), more closely matching the “NE-low” phenotype described for YAP1-expressing SCLC cell lines and tumors,5,7–10 and the rest of the CDX also exhibited this correlation between YAP1 expression and lower NE score (Supplementary Data 6). Furthermore, CDX22P can be classified as a YAP1 subtype on the basis of the YAP1 50-gene signature8 (Supplemental Data 7) and provides the first example of YAP1 expression in both suspension NE-low and adherent non-NE cells (Figs. 3 and 4).

In summary, we found that high cellular YAP1 expression occurs predominantly in rare non-NE cell clusters within CDX, which explains why unbiased hierarchical clustering failed to detect a YAP1 subgroup.4 However, NE-low cells dispersed across CDX tumors can also express YAP1, albeit at lower cellular expression levels. These descriptive data from a biobank of patient-faithful CDX models reveal subtle differences within SCLC cellular phenotypes expressing YAP1. We speculate that YAP1 may fulfill different functional role(s) in NE-low versus non-NE phenotypes, warranting further study. SCLC subtyping has largely been performed using a single time point “snapshot” of intratumoral heterogeneity; yet recent evidence suggests SCLC subtypes are dynamic, and the non-NE YAP1 lineage can emerge during...
disease evolution. In future, SCLC subtyping may need to account for both temporal and spatial expressions of SCLC transcriptional drivers to fully appreciate intratumoral heterogeneity, especially as subtype-specific therapeutic vulnerabilities are discovered. The availability of large biobanks of patient-relevant models of SCLC, including longitudinal models, now allows the field to explore intertumoral and intratumoral heterogeneity in more detail in the search for improved and personalized treatment of this aggressive cancer.

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Supplementary Data
Note: To access the supplementary material accompanying this article, visit the online version of the Journal of Thoracic Oncology at www.jto.org and at https://doi.org/10.1016/j.jtho.2020.07.008.

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