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# Proximity proteome mapping reveals PD-L1-dependent pathways disrupted by anti-PD-L1 antibody specifically in EGFR-mutant lung cancer cells

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## Abstract

**Background** PD-L1, a transmembrane ligand for immune checkpoint receptor PD1, has been successfully targeted to activate an anti-tumor immune response in a variety of solid tumors, including non-small cell lung cancer (NSCLC). Despite the success of targeting PD-L1, only about 20% of patients achieve a durable response. The reasons for the heterogeneity in response are not understood, although some molecular subtypes (e.g., mutant EGF receptor tumors) are generally poor responders. Although PD-L1 is best characterized as a transmembrane PD1 ligand, the emerging view is that PD-L1 has functions independent of activating PD1 signaling. It is not known whether these cell-intrinsic functions of PD-L1 are shared among non-transformed and transformed cells, if they vary among cancer molecular subtypes, or if they are impacted by anti-PD-L1 therapy.

**Methods** Here we use quantitative microscopy techniques and APEX2 proximity mapping to describe the behavior of PD-L1 and to identify PD-L1's proximal proteome in human lung epithelial cells.

**Results** Our data reveal growth factor control of PD-L1 recycling as a mechanism for acute and reversible regulation of PD-L1 density on the plasma membrane. In addition, we describe novel PD-L1 biology restricted to mutant EGFR cells. Anti-PD-L1 antibody treatment of mutant EGFR cells perturbs cell intrinsic PD-L1 functions, leading to reduced cell migration, increased half-life of EGFR and increased extracellular vesicle biogenesis, whereas anti-PD-L1 antibody does not induce these changes in wild type EGFR cells.

**Conclusions** Growth factor acute regulation of PD-L1 trafficking, by contributing to the control of plasma membrane density, might contribute to the regulation of PD-L1's immune checkpoint activity, whereas the specific effects of anti-PD-L1 on mutant EGFR cells might contribute to the poor anti-PD-L1 response of mutant EGFR tumors.

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## Introduction

Therapies targeting the PD1/PD-L1 checkpoint axis are effective in many solid tumors, including non-small-cell lung carcinoma (NSCLC) [1, 2]. Despite the successes of these therapies, only about 20% of patients achieve durable response. The reasons for the heterogeneity in response are not understood. Cancer cell PD-L1 expression has proven to be an imperfect biomarker of response to immune checkpoint therapy [3]. Some molecular subtypes, most notably mutant EGFR NSCLC but ALK rearrangement tumors as well [4–6], respond poorly to anti-PD-L1 antibody therapy although the reason(s) for the resistance have not been described.

Although PD-L1 is best characterized as a transmembrane PD1 ligand, there is accumulating evidence that PD-L1 has functions independent of activating PD1 signaling. Recent studies support a role for PD-L1 in regulation of epithelial mesenchymal transition (EMT), resistance to chemotherapeutic agents as well as interferon-mediated cell toxicity [7–10]. It is not known whether these cell-intrinsic functions of PD-L1 are shared among non-transformed and transformed cells, if they vary among cancer molecular subtypes, or if they are impacted by anti-PD-L1 therapy.

In addition to being localized to the plasma membrane, PD-L1 will transit through a number of membrane compartments during its life time. CMTM6 has been described as a protein that contributes to the control of PD-L1 expression by regulating PD-L1 targeting to lysosomes [11, 12]. Additional proteins such as HIP1R and TRAPCC4 complex have been shown to target PD-L1 for lysosomal degradation or Rab11-mediated recycling to the plasma membrane respectively [13, 14]. ALIX, an ESCRT accessory protein, has been shown to sort PD-L1 into intraluminal vesicles of MVBs and control its abundance on extracellular vesicles [15]. Although these studies show PD-L1 degradation or trafficking can be modified through the expression or depletion of these proteins, it remains unclear whether steps of PD-L1 trafficking are regulated under physiologic conditions and how physiologic control of PD-L1 trafficking contributes to its function as an immune checkpoint or to its cell-intrinsic activities.

Here we use quantitative microscopy techniques and APEX2 proximity mapping to describe the behavior of PD-L1 in living human lung epithelial cells. APEX2 biotinylation mapping is a method to define the proteome within 20 nm of a protein linked to the APEX peroxidase [16, 17]. We leveraged the PD-L1 proximity map to provide a context for further investigations of potential cell-intrinsic roles of PD-L1. Our data reveal growth factor control of PD-L1 recycling as a mechanism for acute and reversible regulation of PD-L1 density on the plasma

membrane. In addition, we describe novel PD-L1 biology restricted to mutant EGFR cells. Anti-PD-L1 antibody treatment of mutant EGFR cells perturbs cell intrinsic PD-L1 functions, leading to reduced cell migration, increased half-life of EGFR and increased extracellular vesicle biogenesis, whereas anti-PD-L1 antibody does not induce these changes in wild type EGFR cells.

## Materials and methods

### Cell lines and culture

BEAS-2B (non-tumorigenic), A549 (KRAS mutant), and H23 (KRAS mutant) were purchased from ATCC. H1650, HCC827, HCC4006 and PC9 (all EGFR Ex19Del mutant) were a gift from Dr. Raffaella Sordella (Cold Spring Harbor Laboratory). H1975 (EGFR L858R/T790M double mutant) were a gift from Dr. Brandon Stiles (Weill Cornell Medicine). All primary human samples were obtained and used in accordance with an approved IRB protocol. Primary tumor cells were processed to obtain single cell suspensions as described previously [18] and plated onto glass-bottom dishes and analyzed 3–5 days later. All cells were cultured in DMEM media supplemented with 10% FBS and 1% penicillin–streptomycin at 37 °C and 5% CO<sub>2</sub>. Where indicated, cells were treated with serum-free DMEM media supplemented with 0.02 M HEPES and 0.03 M NAHCO<sub>3</sub>.

### Stable cell lines

Stable cell lines were generated via lentiviral infection using Lenti-X packaging system. Cells were either antibiotic selected via blasticidin (5 µg/mL) or puromycin (3 µg/mL) or sorted for expression of protein of interest via FACS. PD-L1 KO cells were generated via CRISPR/CAS9 with both targeting sequences (guide1: 5'-CGG GAAGCTGCGCAGAAGT-3'; guide2: 5' -TCTTTA TATTCATGACCTAC-3'). Live cells were stained with PD-L1 antibody and KO cells were sorted by FACS. Bulk population of KO cells was verified for PD-L1 KO via Western Blot. CMTM6 stable inducible shRNA knock-down cells were generated with one shRNA targeting sequence (5'-TTCAAAGAAGTAGAGGCCTCCA-3'), cells were selected with puromycin and mRNA knock-down was confirmed via qRT-PCR.

### Cell treatments

For PD-L1 PM to total measurements, cells were plated on glass bottom dishes in full growth media and assayed the next day. Cells were switched to either serum-free media, serum free media supplemented with growth factors or fresh full growth media for 4 h. EGF was used at 8 nM, IGF at 10 nM and insulin at 10 nM. For Akt inhibition experiments, cells were incubated in full growth media with 5 µM MK2206 or DMSO for 4 h. Growth

factors and MK2206 concentrations were chosen based on induction and inhibition of Akt phosphorylation via western blot. Interferon gamma treatment was performed at 40 ng/mL for 24 h, the lowest concentration and timepoint at which we detected maximum PD-L1 induction. CMTM6 shRNA knock-down was induced by doxycycline at 1 µg/mL for 48 h. For antibody treatment experiments, cells were treated with specified antibody for 7 h at a final concentration of 10 µg/mL, saturating concentration under 15 min of 37 °C staining, in serum free DMEM media. For protein turn-over experiments cycloheximide was used at 10 µg/mL. For stimulated EGFR degradation, cells were serum starved for 4 h and treated with EGF at 50 ng/mL for indicated timepoints respectively, while antibody was used at 10 µg/mL and maintained for the duration of the experiment. For EGFR signaling experiments, cells were incubated with antibody in serum free media for 7 h and stimulated with 0.5 ng/mL EGF for specified timepoints. For migration assays, cells were pre-incubated with Durvalumab antibody for 5 days before being plated for experiment.

### Microscopy

Cells were plated on glass-bottom dishes prior to all experiments. After corresponding experimental treatments, cells were washed and fixed in 3.7% formaldehyde for 5 min. Cells were stained with primary and fluorescent secondary antibodies for proteins of interest on the plasma membrane without permeabilization. Saponin (0.25 mg/mL) was used to permeabilize the cells for total protein stain. Cell nuclei were labeled with Hoechst stain. For quantifications, images were acquired on an inverted microscope with a 20 × air objective or a 40 × oil objective (Leica Biosystems DMI6000-B). Images were analyzed on MetaMorph image processing software and the background corrected average intensity under each cell's area was used to quantify protein amount. The values were normalized to the average of replicate coverslip dishes in control condition in each experiment to allow for comparison across experiments. Unpaired two tailed Student's t test or paired ratio t-test were performed on unnormalized raw data, and one sample t test was used on normalized data respectively as indicated. For comparison of three and more groups, one-way ANOVA was performed on raw data with post-hoc Tuckey test where indicated. For PD-L1 stain images, Z stack of the cells was acquired using Zeiss Confocal microscope (LSM880) with Airyscan on a 40 × oil objective.

### Exocytosis assay

Live cells were incubated with a saturating concentration of PD-L1 antibody (50 µg/mL) for indicated times, washed extensively with PBS, and then fixed. All cells

were permeabilized and stained with a secondary antibody to visualize all antibody bound to the cells. Separate set of dishes was used to quantify total PD-L1 stain in fixed cells in each experiment. Cells were imaged and quantified as described above. All values were normalized to the plateau, which is defined as the average of values at 60 and 120 min in each experiment.

### Colocalization analysis

For co-localization analysis cells were fixed and stained for total PD-L1 and the protein of interest with primary antibody followed by Cy3 and Alexa488-labeled secondary antibodies. Cells were imaged on Zeiss Confocal microscope (LSM880) with Airyscan on a 40 × oil objective. Images were analyzed in MetaMorph software. Intensity threshold was manually determined for each colocalization marker and kept constant for a binary mask of the marker. The binary mask was overlaid with PD-L1 staining resulting in PD-L1 stain only in pixels positive for the marker proteins. For each cell, integrated intensity of PD-L1 staining in the original images as well as the overlaid images were calculated. Ratio of overlaid intensity over original intensity was used as "fraction under mask". Unpaired t-test was performed on raw data.

### Quantitative RT-PCR

Cells were lysed after indicated treatments in each experiments and RNA was extracted using RNeasy kit, and cDNA was prepared from extracted RNA using the RNA to cDNA EcoDry Premix. Quantitative RT-PCR was performed using appropriate primer pairs indicated in Additional file 5: Table 4. 2<sup>(-Delta Delta C(T))</sup> method was used to quantify gene expression.

### Western blotting

Cells were lysed in laemmli buffer with protease/phosphatase inhibitors and the lysate was boiled at 95 °C for 7 min. Lysates were run on 8% or 10% SDS gels and transferred onto nitrocellulose membranes. The membrane was blocked with 5% milk or Licor blocking buffer for 1 h at room temperature, then incubated with primary antibody overnight at 4 °C followed by secondary antibody for 1 h at room temperature. Blots were visualized using Licor Odyssey Fc Imager for two color or Thermofisher MyECL Imager for single color assays and analyzed on ImageStudio or ImageJ software respectively. Actin or GAPDH expression was used as loading control where indicated. One sample t-test was performed on normalized data for comparison of two groups. One-way ANOVA and post-hoc paired ratio t-test was performed on raw data for three or more groups. For degradation assays, K rate constant for the non-linear regression and

slopes for linear regression were used to calculate statistical difference.

### Migration and proliferation

For scratch assays 35 K cells were plated on each side of the Ibidi 2-well scratch assay chambers. Cells were allowed to attach for 24 h and wound healing was assayed in complete or serum-free media for KO/rescue cells and Durvalumab treated cells respectively at specified timepoints. Cells were imaged in bright-field on Leica Microscope. 9 images of each sample were taken at each timepoint, and three distances were measured in each image using Image J. For migration assay, 30 k cells were plated on the top chamber of transwell-migration plates with 8 micron pore size in serum free media. Bottom chamber was filled with DMEM with 10% FBS and cells were allowed to migrate for 24 h. Cells were then fixed in 3.7% formaldehyde for 15 min, washed and then stained in Crystal Violet for 20 min. Unmigrated cells were scraped off with damp cotton swab and membrane was cut out and mounted on slides. Migrated cells were imaged on upright microscope (Zeiss Axioplan) with 10× objective and analyzed via ImageJ. For proliferation assay, 5 k cells were plated in triplicate 96 wells (day 0) and MTT was used to quantify viable cells on days 1–5. Paired t test was performed on raw transwell migration data, and slopes of nonlinear regression lines were used to calculate significance for scratch and MTT assays.

### Extracellular vesicle and whole cell extract preparation for mass spectrometry

Cells were adjusted to media supplemented with 10% exosome free FBS for 1 passage and plated at 60% confluency in exosome free media. After 48 h of Durvalumab treatment (10 µg/mL), cell media was collected for extracellular vesicle isolation and cells were lysed in 1× RIPA buffer with protease/phosphatase inhibitors. Extracellular vesicles were isolated by sequential ultracentrifugation (12000 g for 10 min to discard cell debris, 20,000 g for 20 min to discard microvesicles, 100,000 g for 70 min twice to isolate and wash the EVs) and resuspended in a final volume of 100 µL of RIPA buffer with protease/phosphatase inhibitors. Cell extract and EV lysates were quantified for protein concentration. A final amount of 100 µg of WCE, and equi-volume of EVs approximating 10 µg of protein in each sample was submitted for mass spectrometry analysis.

### PD-L1-APEX2 labeling

APEX2 proximity assay was performed as described previously [19]. Briefly, BEAS-2B cells stably expressing PD-L1 APEX2 were grown until confluency. Cells were preincubated with biotin phenol in full growth media

for 30 min and hydrogen peroxide was added for 1 min (except in negative control). H1650 PD-L1-APEX2 stably expressing cells were treated similarly with the exception of biotin phenol concentration (500 µM for BEAS-2B and 2.5 mM for H1650). Cells were washed three times with ice cold quenching solution (10 mM sodium ascorbate, 5 mM Trolox and 10 mM sodium azide solution in PBS) and cells were lysed in 1× RIPA lysis buffer with protease/phosphatase inhibitors. Protein concentration was measured, and biotinylated proteins were pulled down using streptavidin-agarose beads overnight at 4 °C. Beads were washed (twice with RIPA lysis buffer, once with 1 M KCl, once with 0.1 M Na<sub>2</sub>CO<sub>3</sub>, once with 2 M urea in 10 mM Tris-HCl, pH 8.0, and twice with RIPA lysis buffer) and pull-down proteins were submitted for mass spectrometry.

### Extracellular vesicle, whole cell extract mass spectrometry

The samples were digested in solution with trypsin overnight at 37°C following reduction with 5 mM DTT and 14 mM alkylation with iodoacetamide. The digests were vacuum centrifuged to near dryness and desalted by Sep-Pak column.

A Thermo Fisher Scientific EASY-nLC 1000 coupled on-line to a Fusion Lumos mass spectrometer (Thermo Fisher Scientific) was used. Buffer A (0.1% FA in water) and buffer B (0.1% FA in ACN) were used as mobile phases for gradient separation (1). A 75 µm × 15 cm chromatography column (ReproSil-Pur C18-AQ, 3 µm, Dr. Maisch GmbH, German) was packed in-house for peptides separation. Peptides were separated with a gradient of 3–28% buffer B over 110 min, 28%–80% B over 10 min at a flow rate of 300 nL/min. The Fusion Lumos mass spectrometer was operated in data dependent mode. Full MS scans were acquired in the Orbitrap mass analyzer over a range of 300–1500 m/z with resolution 120,000 at m/z 200. The top 15 most abundant precursors with charge states between 2 and 5 were selected with an isolation window of 1.4 Thomsons by the quadrupole and fragmented by higher-energy collisional dissociation with normalized collision energy of 35. MS/MS scans were acquired in the Orbitrap mass analyzer with resolution 15,000 at m/z 200. The automatic gain control target value was 1e6 for full scans and 5e4 for MS/MS scans respectively, and the maximum ion injection time is 60 ms for both.

The raw files were processed using the MaxQuant computational proteomics platform version 1.5.5.1 (Max Planck Institute, Munich, Germany) for protein identification. The fragmentation spectra were used to search the UniProt human protein database (downloaded on 01/16/2021). Oxidation of methionine and protein N-terminal acetylation were used as variable modifications for

database searching. The precursor and fragment mass tolerances were set to 7 and 20 ppm, respectively. Both peptide and protein identifications were filtered at 1% false discovery rate based on decoy search using a database with the protein sequences reversed.

#### PD-L1-APEX2 mass spectrometry

Peptides from streptavidin purified proteins were released via on-bead digestion using trypsin (Promega) and desalted on C18 STAGE Tips [PMID: 12585499]. Eluted peptides were dried and re-suspended in 5% formic acid (FA). Mass Spectrometric analysis was performed on a Thermo Orbitrap Fusion mass spectrometer equipped with an Easy nLC-1000 UHPLC. Peptides were separated with a 115 min gradient of 5–25% acetonitrile in 0.1% FA at 350 nl/min and introduced into the mass spectrometer by electrospray ionization as they eluted off a self-packed 40 cm, 100  $\mu$ m (ID) column packed with 1.8  $\mu$ m, 120  $\text{\AA}$  pore size, C18 resin (Sepax Technologies, Newark, DE). The column was heated to 60  $^{\circ}$ C. Peptides were detected using a data-dependent method. For each 3 s cycle, high resolution MS1 scans performed in the orbitrap and the most abundant ions per scan were isolated, fragmented by CID, and scanned in the ion trap. Ions selected for MS2 analysis were excluded from reanalysis for 60 s. Ions with +1 or unassigned charge were also excluded from analysis. MS/MS spectra were matched to peptide sequences using COMET (version 2019.01 rev. 5) (PMID: 23,148,064) and a composite database containing the 20,415 Uniprot reviewed canonical predicted human protein sequences (<http://uniprot.org>, downloaded 5/1/2019) and its reversed complement. Search parameters allowed for two missed cleavages, a mass tolerance of 25 ppm, a static modification of 57.02146 Da (carboxyamidomethylation) on cysteine, and a dynamic modification of 15.99491 Da (oxidation) on methionine. Peptide spectral matches (PSMs) were filtered to 1% FDR using the target-decoy strategy [20] and then to 1% protein FDR. Label-free quantification was performed using peptide intensities from the integrated areas under each corresponding extracted-ion-chromatogram peak. Intensities for all peptides mapping to each protein were summed for each sample.

#### Mass spectrometry data analysis

BEAS-2B PD-L1-APEX2 data were analyzed in two different ways. In one, proteins, in which protein intensity in the experimental sample was two-fold or greater than in the control sample, were identified to be in the proximity of PD-L1. Data from 5 independent experiments were analyzed individually and the final PD-L1 proximity map was determined as those proteins satisfying the criteria for PD-L1 proximity in each of the five experiments,

yielding 438 PD-L1 proximity proteins. This set of proteins was used for further analyses, including biological processes pathway analysis via geneontology.org (GO database released 11–16-2021) [21]. The data were also analyzed by performing a t-test on  $\text{Log}_{10}$  transformed spectral intensity values between labeled and negative control samples ( $-\text{H}_2\text{O}_2$ ) on the set of 1647 proteins common to all five experiments. Significant differences between the experimental and control were determined using Benjamini–Hochberg correction for multiple testing. Fold change was calculated as ratio of average spectral intensity in labeled samples over average in negative control samples (as shown in Fig. 2F). The two different methods yielded essentially the same group of proteins. For H1650 PD-L1-APEX2 data, proteins with intensities twofold or higher in experimental sample over control in all 3 replicate experiments were identified as PD-L1 proximal.

For extracellular vesicle (EV) mass spectrometry analysis, for each protein its intensity in EV was divided by the corresponding intensity in whole cell extract (WCE). Proteins that were not detected in either sample were excluded. The remaining proteins were rank ordered based on the EV/WCE ratio. This analysis was done for both conditions (+/– Durvalumab) and both biological replicate experiments (4 samples total for each cell line). Proteins falling in the top quartile in 3 out of 4 samples in each cell line were used for downstream analyses.

## Results

### PD-L1 is dynamically maintained on the plasma membrane of cells

PD-L1 was distributed between the plasma membrane (PM) (immunostaining of non-permeabilized cells) and intracellular compartments (immunostaining of permeabilized cells) of freshly isolated primary human NSCLC cells (Fig. 1A). Of the total cell PD-L1, approximately 30% was at the PM (Fig. 1B). The steady-state PD-L1 distribution between PM and interior of primary NSCLC cells was similar to its distribution in established NSCLC cell lines as well as its distribution in a non-transformed human lung epithelial cell line (BEAS-2B) (Fig. 1C). Because oncogenic KRAS and EGFR are the two most common mutations in NSCLC [22], we focused our studies on cell lines harboring either KRAS or EGFR mutations. PD-L1's distribution did not vary with the oncogenic driver, nor was it influenced by total expression of PD-L1 (Fig. 1B–D). In primary NSCLC cells and all the NSCLC cell lines examined, PM PD-L1 showed some enrichment in puncta, which was enhanced by bivalent secondary antibody staining compared to monovalent secondary Fab stain (Additional file 1: Fig. S1A). Intracellular PD-L1 was primarily in vesicles dispersed

throughout the cytoplasm, although in some cells there was a juxta-nuclear concentration typical of the Trans-Golgi Network/Golgi (Fig. 1A, E, Additional file 1: Fig. S1B). In all NSCLC cell lines examined, interferon gamma (IFN $\gamma$ ), a potent transcriptional regulator of PD-L1 [23, 24], increased the total amount of PD-L1 protein without affecting PD-L1 distribution between the interior and PM of cells, its localization to PM puncta, or its localization to cytoplasmic vesicles (Fig. 1E–F, Fig. S1B).

PM PD-L1 is poised to function as an immune checkpoint by binding in trans to PD1 on the surface of T cells [25, 26]. Intracellular PD-L1 could serve as a reservoir to rapidly modulate the amount of PD-L1 on the PM to fine-tune its function as an immune checkpoint on a shorter timescale than IFN $\gamma$  transcriptional regulation of PD-L1 expression. To interrogate the relationship between the PM and intracellular pools of PD-L1, we used a quantitative fluorescence microscopy assay in which trafficking of PD-L1 between intracellular compartments and the PM is monitored by incubating living cells in medium containing an antibody that binds to the extracellular domain of PD-L1 (Fig. 1G). At early time points of incubation, the amount of cell-associated anti-PD-L1 antibody predominantly reflects the amount of PD-L1 on the PM. An increase in cell-associated anti-PD-L1 antibody with increasing time of incubation reflects antibody binding to unbound PD-L1 newly recycled to the PM from intracellular compartments. Cell-associated anti-PD-L1 antibody will increase until all the intracellular PD-L1 that is in equilibrium with the PM has cycled to the PM (i.e., bound by anti-PD-L1 antibody). The rise to the plateau is the rate at which intracellular PD-L1 traffics to the PM. Cell-associated anti-PD-L1 antibody reached a plateau by 30 min, demonstrating that PM density of PD-L1 is dynamically maintained by constitutive PD-L1 cycling between intracellular compartments and the PM. The equilibration half-time for PD-L1 exchange between the

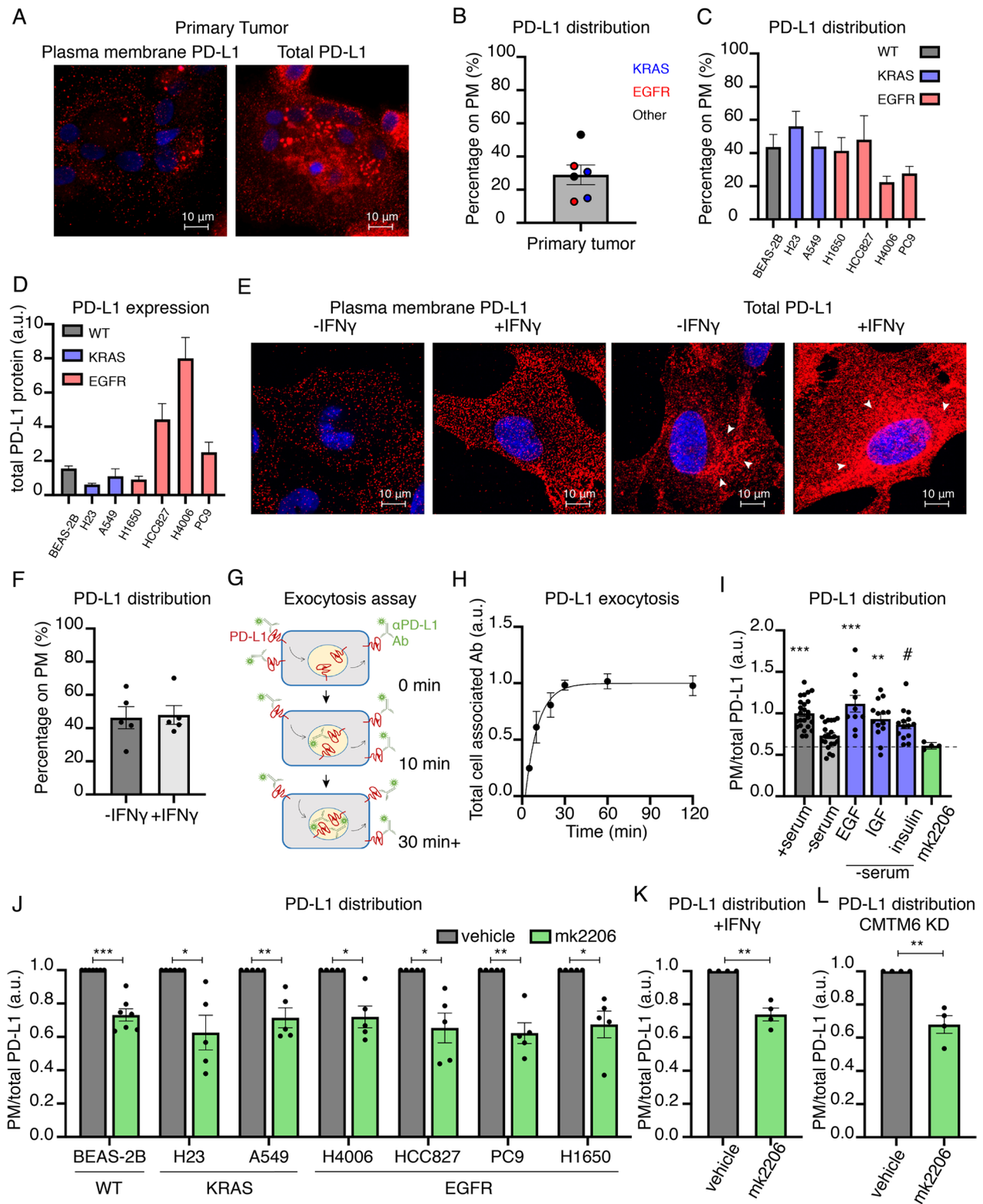
PM and intracellular compartments is approximately 10 min (Fig. 1H). These data suggest PD-L1's traffic to and from the PM as a mechanism to acutely and reversibly increase or decrease PM PD-L1 amounts, an effect that would impact PD-L1 immune checkpoint function.

We tested the hypothesis that PD-L1 trafficking is acutely regulated by determining the effect of growth factor stimulus on PM PD-L1. Removal of tonic growth factor signaling by incubation of cells in serum-free medium for 4 h resulted in a significant reduction of PM PD-L1 (Fig. 1I). This reduction of PM PD-L1 reflected a redistribution of PD-L1 from the PM to intracellular compartments without significant degradation of PD-L1 (Additional file 1: Fig. S1C). Addition of different growth factors (i.e., EGF, IGF or insulin) to serum-free medium restored PM PD-L1 without affecting the total amount of PD-L1 (Fig. 1I, Additional file 1: Fig. S1C). Thus, signaling downstream of various growth factors actively maintains PD-L1 on the PM by regulation of PD-L1 cycling to and from the PM, and identifying acute modulation of growth factor signaling as a mechanism that contributes to the control of PD-L1 PM. This acute and reversible effect of growth factors stimulation reflects a change in PD-L1 localization, not a change in total PD-L1 amounts.

Growth factors commonly regulate membrane protein trafficking downstream of activation of the serine/threonine AKT kinase [27, 28]. Consistent with a role for AKT in actively maintaining PD-L1 on the PM, inhibition of AKT reduced the amount of PD-L1 in the PM without affecting the total amount of PD-L1 expressed in cells (Fig. 1I, Additional file 1: Fig. S1C–E). Inhibition of AKT resulted in a near two-fold reduction of the PD-L1 recycling rate constant, accounting for reduced PM PD-L1 (Additional file 1: Fig. S1D). The effect of AKT inhibition on PD-L1 expression in the PM was independent of the driver oncogene (Fig. 1J). AKT-dependent regulation of PM PD-L1 was distinct from IFN $\gamma$  transcriptional regulation of PD-L1 expression; therefore, it is a complementary mode to regulate PM density of PD-L1 (Fig. 1K).

(See figure on next page.)

**Fig. 1** PD-L1 is dynamically maintained on the plasma membrane of cells. **A** Representative confocal images of primary human tumor cells immunostained for plasma membrane (PM) and total PD-L1. Red: PD-L1. Blue: Hoechst. **B** Quantification of percentage of PM PD-L1 in primary human tumors. Each data point is a patient sample color coded for known oncogenes. **C, D** Quantification of **C** percentage of PM PD-L1 and **D** total PD-L1 protein expression in human lung cell lines via immunofluorescence imaging. Cell lines in blue and red harbor mutant KRAS and mutant EGFR respectively. **E** Representative confocal images of BEAS-2B with or without IFN $\gamma$  treatment and immunostained for PM and total PD-L1. Red: PD-L1. Blue: Hoechst. Sum intensity of z-stacks is projected. Fluorescence intensity is equally scaled across panels. **F** Quantification of PM localization of PD-L1 in BEAS-2B cells treated with IFN $\gamma$ . Unpaired student's t-test. **G** Schematic of PD-L1 exocytosis assay. Live cells are incubated with unlabeled PD-L1 antibody. Cells are fixed, permeabilized and stained with fluorescently labeled secondary antibody depicted as a star. **H** PD-L1 steady state exocytosis in BEAS-2B cells. The line is a fit of the data to one-phase association exponential equation,  $N = 3$ . **I** PD-L1 PM to total ratio with growth factors treatment or Akt inhibition in BEAS-2B cells. One-way ANOVA and post-hoc Tukey test comparing each condition to -serum. **J** PD-L1 PM to total ratio in cell lines treated with vehicle (DMSO) or Akt inhibitor for 4 h. One sample t-test. **K** PD-L1 PM to total ratio in IFN $\gamma$  treated BEAS-2B cells incubated with vehicle (DMSO) or Akt inhibitor. One sample t-test. **L** PD-L1 PM to total ratio in CMTM6 knocked-down (KD) IFN $\gamma$ -treated BEAS-2B cells with vehicle (DMSO) or Akt inhibitor. Except panel B and H, each data point is an experiment. All error bars = SEM. \*  $\leq 0.05$ , \*\*  $\leq 0.01$ , \*\*\*  $\leq 0.001$ , # = 0.058



**Fig. 1** (See legend on previous page.)

CMTM6, a ubiquitously expressed protein, post-translationally controls PD-L1 PM amounts by determining whether internalized PD-L1 is recycled or targeted for degradation [11, 12]. In agreement with those studies, transient knockdown of CMTM6 resulted in a 30% reduction of total PD-L1 amounts. However, the distribution of the remaining PD-L1 between the plasma membrane and interior of cells was unchanged (Additional file 1: Fig. S1F). Regulation of PM PD-L1 amounts downstream of AKT was also unaffected by CMTM6 knock down. Thus, regulation of PD-L1 traffic downstream of AKT is a mechanism to acutely modulate PM PD-L1 independently of other known mechanisms (e.g., IFN $\gamma$  and CMTM6) (Fig. 1L).

### PD-L1 proximity mapping

Localization of PD-L1 to intracellular compartments might reflect cell-intrinsic function(s) of PD-L1 that are in addition to regulating the fraction of PD-L1 expressed on the surface of cells. A more complete knowledge of the proteins in proximity to PD-L1 will provide a framework for understanding potential cell intrinsic functions of PD-L1. To identify the PD-L1 proximal proteome we used APEX2 mapping [17]. We expressed a PD-L1 chimera in which the APEX2 peroxidase catalytic domain was fused to the cytosolic carboxyl terminus of PD-L1 (PD-L1-APEX2) (Fig. 2A). PD-L1-APEX2 expressed in BEAS-2B cells behaved similarly to endogenous PD-L1. Approximately 40% of PD-L1-APEX2 was in the PM, localized to puncta, whereas intracellular PD-L1-APEX2 was distributed between dispersed puncta and a concentration in the peri-nuclear region of cells (Fig. 2B, C). Furthermore, the amount of PD-L1-APEX2 in the PM was controlled by AKT (Fig. 2D). These data validate PD-L1-APEX2 as a reporter for PD-L1.

To catalog the PD-L1 proximal proteome, living cells were pre-incubated with biotin phenol, and the reaction was catalyzed by the addition of hydrogen peroxide to generate a biotin-phenoxy radical that covalently binds to electron rich amino acids on nearby proteins (Fig. 2E, Additional file 1: Fig. S2A). PD-L1 proximal proteins were identified by streptavidin beads enrichment and mass spectrometry analysis (Fig. 2E). In each of the five individual experiments, APEX2 biotinylated proteins were identified as streptavidin-absorbed proteins whose

intensities in the H<sub>2</sub>O<sub>2</sub>-treated samples were twofold or greater than in samples not treated with H<sub>2</sub>O<sub>2</sub> (no APEX reaction). The aggregate PD-L1 proximity proteome was determined as the overlapping set of 438 proteins from the five independent experiments (Fig. 2F, Additional file 2: Table S1). Of note, PD-L1 (CD274) was identified as an APEX2 biotinylated protein in all five experiments.

HLA class I proteins (HLA-A, -B and -C), which are involved in presentation of endogenous antigens, were identified as components of the PD-L1 proximity proteome (Fig. 2F, G, Additional file 2: Table S1). Immunofluorescence confirmed the proximity of PD-L1 to HLA-I, while using proximity of PD-L1 to transferrin receptor as a control transmembrane protein (Additional file 1: Fig. S2B). PD-L1's proximity to HLA-I will enhance the likelihood of binding PD1 when HLA-I presenting antigen is engaged by the T cell receptor, thereby potentially contributing to PD-L1's effectiveness as an immune checkpoint. Our data establish that the previously reported PD-L1 proximity to HLA-I proteins in lymphocytes [29] is conserved in non-immune cells.

Gene ontology analysis of the 438 protein PD-L1 proteome revealed multiple biological pathways enriched among the PD-L1 proximal proteome, including an enrichment of proteins that function in the traffic of membrane proteins to and from the PM as well as among intracellular compartments (Fig. 2G). These data agree with our functional data that PD-L1 traverses a number of intracellular compartments as it constitutively traffics between the PM and interior of cells.

### A role for PD-L1 in cell migration

Approximately 29% of the PD-L1 proximity map proteins are annotated to cytoskeleton organization, covering a broad range of both actin and microtubule biology (Fig. 2H, Additional file 2: Table S1). Because the actin and microtubule cytoskeletons have a role in the regulation of membrane protein traffic, this enrichment likely reflects, in part, the constitutive movement of PD-L1 among different cellular compartments. However, to explore the possibility that enrichment of cytoskeletal proteins in the PD-L1 proximity map also reflects a role for PD-L1 in cytoskeletal-dependent functions, we determined the effect of PD-L1 knockout (KO) on cell migration. PD-L1 KO was confirmed by western

(See figure on next page.)

**Fig. 2** PD-L1 proximity mapping. **A** Schematic of PD-L1-APEX2 labeling assay. **B** Representative epifluorescence image of PD-L1-APEX2 plasma membrane and total stain in BEAS-2B cells. **C** PM to total distribution of PD-L1-APEX2 in BEAS-2B cells. **D** PM to total ratio of endogenous PD-L1 and PD-L1-APEX2 construct in BEAS-2B cells treated with vehicle (DMSO) or Akt inhibitor. One-sample *t* test. **E** Schematic of experimental design for PD-L1-APEX2 proximity labeling. **F** Volcano plot of PD-L1-APEX2 labeled proteins. FC = fold change of (labeled/ctrl) spectral intensity. Dashed red lines represent fourfold change on the x-axis, and *p*-value corresponding to FDR = 0.1 after Benjamini-Hochberg correction. **G** Biological pathways and compartments identified in PD-L1-APEX2 proximity labeling. David gene ontology terms and identifiers are shown. **H** Additional Biological processes pathways enriched in PD-L1-APEX2 labeling in BEAS-2B. All error bars = SEM. \*  $\leq 0.05$ , \*\*  $\leq 0.01$ , \*\*\*  $\leq 0.001$

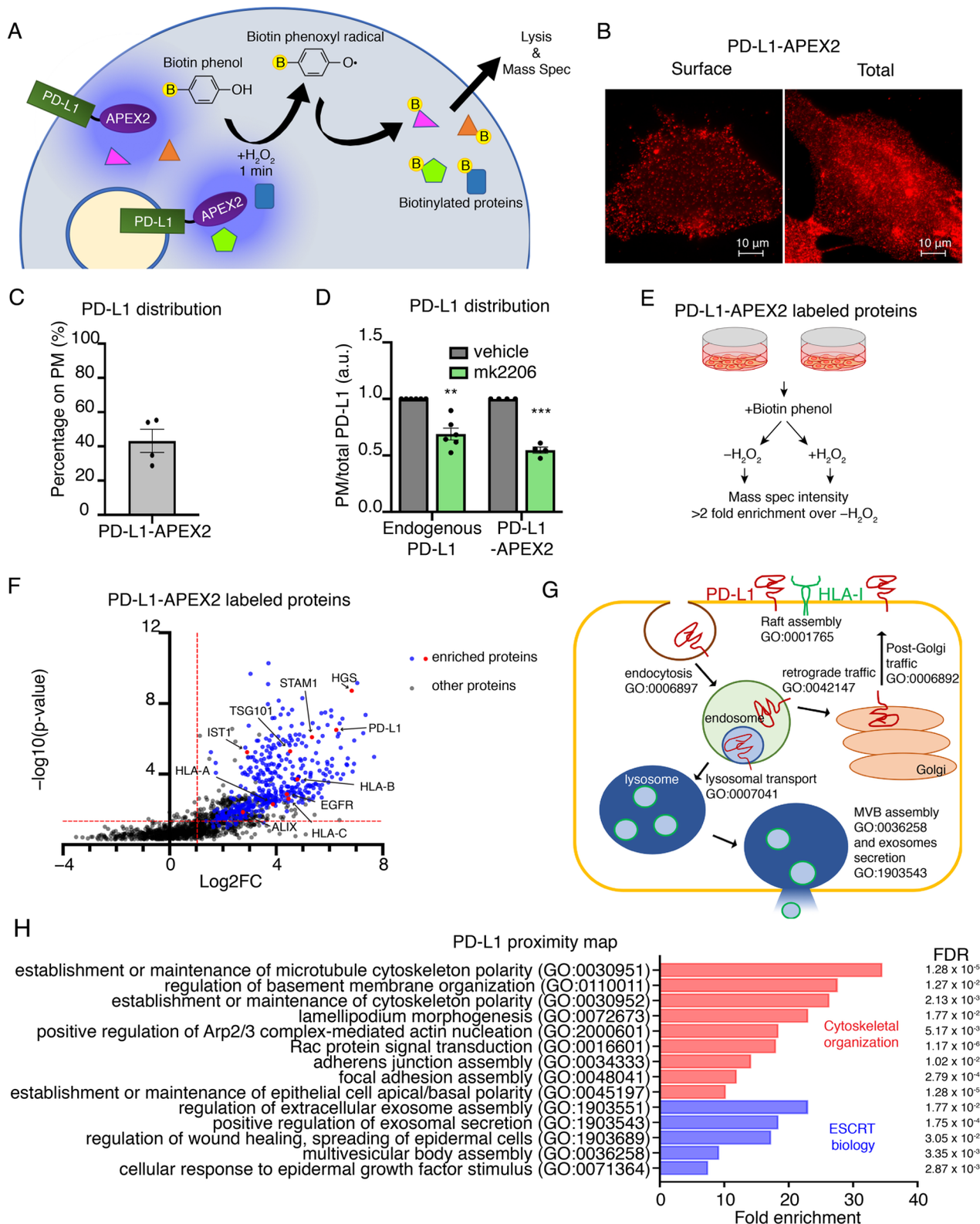


Fig. 2 (See legend on previous page.)

blotting (Additional file 1: Fig. S3A). PD-L1 KO inhibited the migration of BEAS-2B cells in a scratched monolayer assay (Fig. 3A, J, Additional file 1: Fig. S3B). This finding is in line with previous reports that PD-L1 has a function in the regulation of cell migration [30]. Durvalumab, a therapeutic anti-PD-L1 antibody [31, 32], did not affect migration of the BEAS-2B cells, suggesting that the anti-tumor activity of antibody binding PD-L1 does not involve disruption of PD-L1's control of migration (Fig. 3B, J, Additional file 1: Fig. S3B).

We next determined whether PD-L1 has a similar role in migration of cancer cells. PD-L1 KO and re-expression in H1650 cells were confirmed by western blotting (Additional file 1: Fig. S3A). PD-L1 KO blunted migration of H1650 cells (an EGFR mutant cell line), an effect that was rescued by re-expression of PD-L1 (Fig. 3C, J, Additional file 1: Fig. S3C). Unexpectedly, Durvalumab inhibited migration of H1650 to the same degree as PD-L1 KO (Fig. 3D, J, Additional file 1: Fig. S3D). PD-L1 KO or anti-PD-L1 antibody did not affect H1650 cell proliferation, confirming the effects were on cell migration into the wounded monolayer (Additional file 1: Fig. S3E-F). The different effects of anti-PD-L1 antibody on migration in non-transformed cells BEAS-2B and NSCLC cell line, H1650, suggest cell-context specific effects of  $\alpha$ PD-L1 binding to PD-L1.

To determine whether the effect of anti-PD-L1 antibody on migration was specific to cancer cells, we studied another human NSCLC cell line, A549 cells. These cells do not migrate into a scratched monolayer; therefore, we monitored migration by determining penetration of cells into a filter disk. H1650 migration into the filter disk was inhibited by PD-L1 KO and rescued by re-expression of PD-L1, similar to the effect on H1650 migration measured by the scratch assay (Fig. 3E, J). PD-L1 KO in A549 cells was confirmed by western blotting (Additional file 1: Fig. S3A). PD-L1 KO in A549 cells reduced penetration into the filter, confirming a role for PD-L1 in migration in a second cancer cell line (Fig. 3F, J).

Anti-PD-L1 antibody reduced H1650 cell migration into the filter disk, confirming the effect on migration of these cells into scratched monolayer (Fig. 3G, J). However, as was the case for non-transformed BEAS-2B cells, anti-PD-L1 antibody had no effect on migration of A549

cells (Fig. 3H, J). H1650 cells are EGFR mutant, whereas A549 are wild type for EGFR. Anti-PD-L1 antibody also reduced migration of another EGFR mutant NSCLC line, H1975, suggesting the effect of anti-PD-L1 antibody on migration is a feature of EGFR mutant cell lines, revealing the possibility of mutant EGFR cell context-dependent PD-L1 functions (Fig. 3I, J).

#### Anti-PD-L1 antibody treatment inhibits turnover of mutant but not wild type EGFR

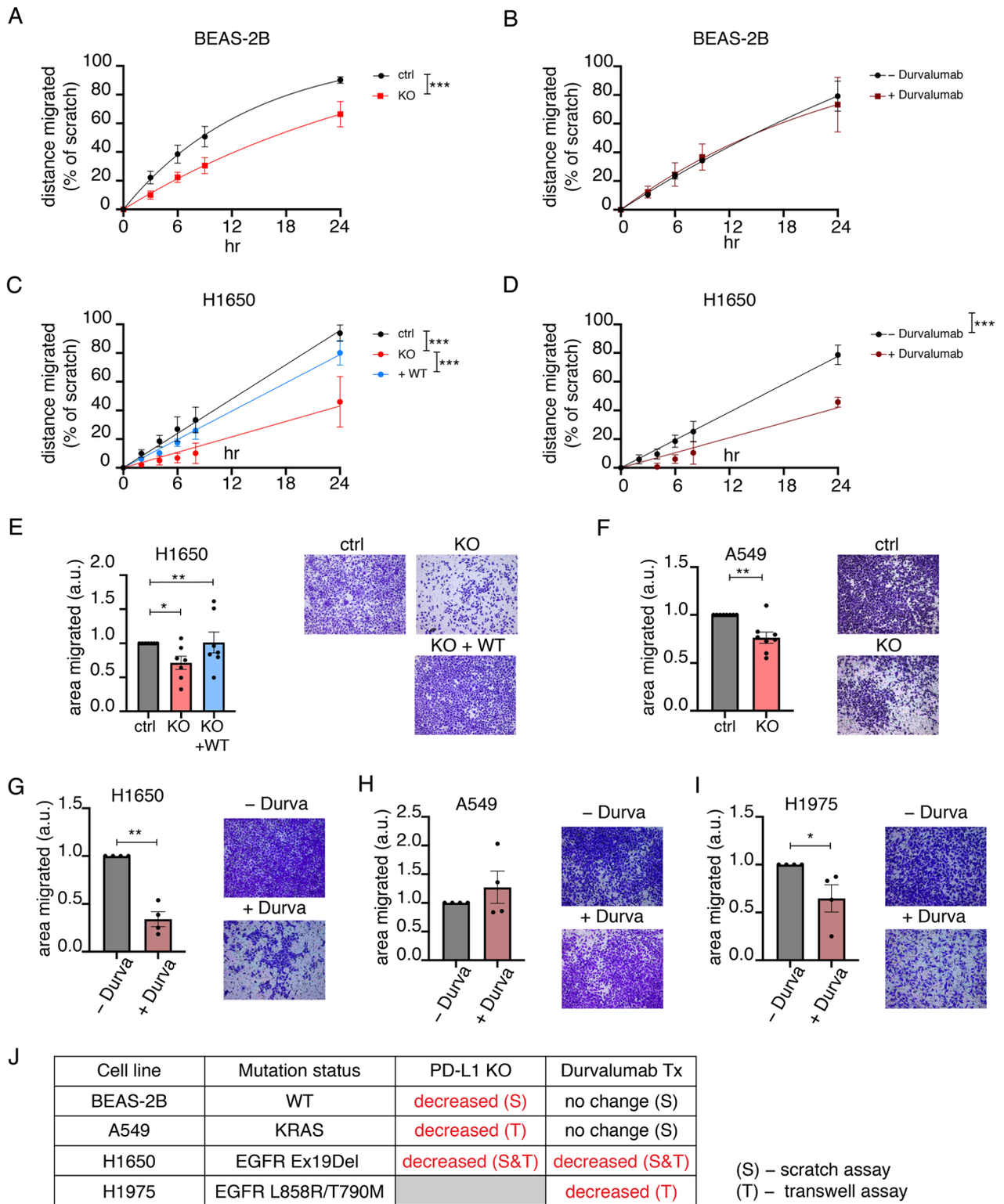
A potential cell context-specific link between PD-L1 and EGFR, as suggested by the anti-PD-L1 antibody effect on migration of mutant EGFR cell lines was also supported by EGFR being a component of the PD-L1 proximal proteome (Fig. 2F, Additional file 2: Table S1). In addition to the enrichment of EGFR, ESCRT complex proteins, which have a role in regulation of intracellular trafficking of ligand activated EGFR, were also enriched in the PD-L1 proximal proteome (Fig. 2F, H). We confirmed proximity to PD-L1 for EGFR and HGS (a subunit of ESCRT-0 complex) (Additional file 1: Fig. S2B). We also observed PD-L1 co-localized with constitutively active mutant-EGFR in H1650 cells. Interestingly, EGF stimulation reduced PD-L1 co-localization with wildtype but not mutant EGFR (Additional file 1: Fig. S2C).

The ESCRT complex is involved in targeting membrane proteins to the lysosomes for turnover, and one of its best studied client proteins is ligand-activated EGFR [33, 34]. Although proximity of PD-L1 to ESCRT proteins may reflect PD-L1 turnover, given the relative high enrichment of ESCRT0 and ESCRT1 proteins in the PD-L1 proximity map (Fig. 2F) we explored the possibility of a link between PD-L1 and ESCRT-dependent function(s) beyond the targeting of PD-L1 for degradation.

Knockdown (KD) of PD-L1 in BEAS-2B cells (two-fold decrease in PD-L1 protein expression with KD) delayed ligand stimulated turnover of EGFR (Fig. 4A, Additional file 1: Fig. S4A). In both wildtype and PD-L1 KD cells, within 30 min of EGF stimulation, EGFR moved from a diffuse distribution on the PM to concentration in endosomes, demonstrating PD-L1 KD did not affect EGF-stimulated EGFR internalization. However, 60 and 90 min after 50 ng/mL EGF stimulation, EGFR fluorescence was higher in the PD-L1 KD cells than control cells,

(See figure on next page.)

**Fig. 3** A role for PD-L1 in cell migration. **A, B** Quantification of scratch assay in **A** BEAS-2B PD-L1 KO cells, N = 6, and **B** BEAS-2B treated with Durvalumab, N = 4. For **A, B** rate constants are compared using extra sum-of-squares F test. Cells were in serum-free media for the duration of the assay. **C** Quantification of scratch assay in H1650 PD-L1 KO/rescue cells, N = 4. Cells were in serum complete media for the duration of the assay. **D** Quantification of scratch assay in H1650 cells treated with Durvalumab, N = 3. Cells were in serum-free media for the duration of the assay. For **C, D** difference in slope values is calculated using F-test. **E** Transwell migration in H1650 PD-L1 KO/rescue cells. One-way ANOVA and post-hoc Tukey test. **F** Transwell migration in A549 PD-L1 KO cells. **G–I** Transwell migration in (**G**) H1650, (**H**) A549 and (**I**) H1975 cells treated with Durvalumab. For **F–I** paired student's t-test was used on raw values. **J** Table summarizing cell lines' response to PD-L1 knock-out or Durvalumab treatment. All error bars = SEM. \*  $\leq 0.05$ , \*\*  $\leq 0.01$ , \*\*\*  $\leq 0.001$



**Fig. 3** (See legend on previous page.)

demonstrating EGFR turnover was delayed by PD-L1 KD at a post-internalization step, perhaps in transport from the endosomes to lysosomes, a step requiring ESCRTs. PD-L1 KO, in a western blot assay for EGFR expression, also delayed EGF-induced receptor turnover, confirming a role for PD-L1 in controlling the post-stimulation half-life of EGFR (Fig. 4B).  $\alpha$ PD-L1 antibody did not affect the half-life of activated EGFR in BEAS-2B cells (Fig. 4C). Thus, as was the case for migration of BEAS-2B cells, PD-L1 KO but not  $\alpha$ PD-L1 antibody has effects on EGFR behavior in BEAS-2B cells.

In H1650 cells, both PD-L1 KO and  $\alpha$ PD-L1 antibody increased mutant EGFR protein amounts by about 50%, and re-expression of PD-L1 fully rescued the effect of the knockout (Fig. 4D, Additional file 1: Fig. S4B). Durvalumab and lab reagent anti-PD-L1 antibody (clone# 130,021, Novus Biologicals), which bind different extracellular epitopes of PD-L1, were equally effective at increasing mutant EGFR protein amounts, whereas an antibody targeting the extracellular domain of the transferrin receptor, another PM protein, did not affect the amount of EGFR protein (Fig. 4E, F, Additional file 1: Fig. S4C,D). Analyses of a panel of NSCLC lines demonstrated that the effect of  $\alpha$ PD-L1 antibody on turnover of the ligand-stimulated EGFR protein amount was specific to cells harboring mutant EGFR (Fig. 4G).  $\alpha$ PD-L1 antibody caused a decrease turnover, evidenced by increase in the amount of EGFR in three different EGFR mutant lines, whereas the amounts of EGFR in two KRAS mutant lines and a non-transformed line were unaffected. The effect is not linked to a specific EGFR mutation because the EGFR mutant lines harbor different activating mutations, nor was the effect of  $\alpha$ PD-L1 antibody associated with differences in PD-L1 expression among the cell lines (Fig. 1D). Furthermore, anti-PD-L1 antibody treatment did not affect the distribution nor total amount of either PD-L1 or the transferrin receptor in EGFR mutant cells (Additional file 1: Fig. S4E, F).

To directly monitor the effect of anti-PD-L1 antibody on EGFR half-life, we compared EGFR turnover in cells treated with cycloheximide to inhibit translation. In these conditions the time dependent reduction of mutant EGFR reflects its turnover. anti-PD-L1 antibody

significantly increased mutant EGFR half-life without an effect on wildtype EGFR (Fig. 4H, I, Additional file 1: Fig. S4G). These data also demonstrate the increased mutant EGFR protein amount is not due to an increase in translation. In addition, anti-PD-L1 antibody did not alter mutant EGFR mRNA amounts, supporting the conclusion that anti-PD-L1 antibody modulates mutant EGFR protein levels by increasing its half-life (Additional file 1: Fig. S4H, I). The extension of mutant EGFR half-life by anti-PD-L1 treatment did not augment EGFR signal transduction measured by changes in gene transcription (Additional file 1: Fig. S4J). These data on EGFR turnover provide additional support for the hypothesis that EGFR mutant cells have a heightened sensitivity to perturbation of PD-L1 by anti-PD-L1 antibody binding, while PD-L1 KO, a more severe disruption of PD-L1, impacts both WT and mutant EGFR cell lines.

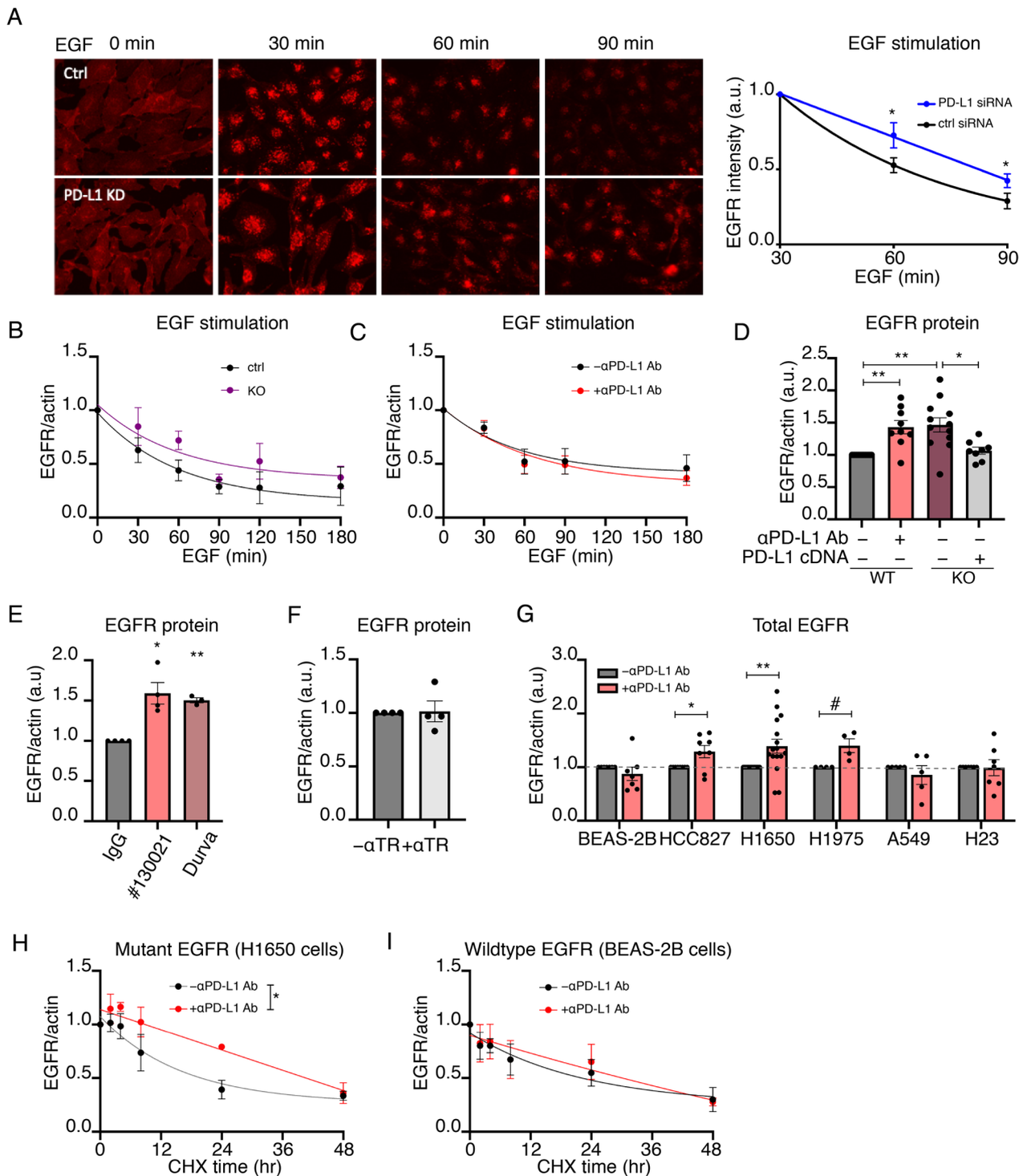
#### Anti-PD-L1 antibody treatment affects extracellular vesicles

ESCRTs, in addition to having a key role in targeting EGFR (and other proteins) for lysosomal degradation, are also involved in the formation and secretion of extracellular vesicles (EVs) [35]. To assay EV formation, we used mass spectrometry to identify proteins in EVs isolated from the growth media of BEAS-2B, A549 and H1650 cells treated with or without anti-PD-L1 antibody for 48 h. We also analyzed the whole cell extract collected from the same cells. To limit the analysis to most highly enriched EV proteins, we focused on proteins with EV to whole cell extract (WCE) ratio in the top quartile for each respective cell line. Although there was some variation, the lists largely overlapped among the three cell lines (Fig. 5A, Additional file 3: Table S2). The biological pathways enriched among the EV proteins of the 3 cell lines were similar and the biological pathways of the identified proteins were congruent with reported functions of EVs, including exosome formation, cell adhesion and migration as well as immune cell extravasation [36, 37] (analysis of H1650 EV is shown Fig. 5B).

Next, we analyzed if anti-PD-L1 antibody affects EV secretion or content. As a general measure of EV formation, we first examined a set of 6 canonical EV

(See figure on next page.)

**Fig. 4** Anti-PD-L1 antibody treatment inhibits turnover of mutant but not wild type EGFR. **A** Representative images and quantification of EGFR in BEAS-2B cells with EGF stimulation, N=6. Paired student's t-test at 60 and 90 min were calculated. **B, C** EGF-stimulated EGFR degradation in (B) BEAS-2B PD-L1 KO cells, N=4, and (C) BEAS-2B cells treated with  $\alpha$ PD-L1 antibody, N=4. Rate constants are compared using extra sum-of-squares F test. **D** EGFR protein in WT, PD-L1 KO and PD-L1-reexpressing H1650 cells. One way ANOVA and post-hoc Tukey test. **E** EGFR protein expression in H1650 cells treated with reagent grade  $\alpha$ PD-L1 antibody (clone # 130021, Novus Biologicals) and clinical Durvalumab as compared to isotype IgG control antibody. One-way ANOVA and post-hoc Tukey test. **F** EGFR protein in H1650 cells with anti-transferrin receptor antibody (aTfR) antibody treatment. One sample t-test. **G** EGFR protein expression in cell lines treated with  $\alpha$ PD-L1 antibody. One sample t-test. **H, I** Quantification of EGFR degradation in **H** H1650, N=3, and **I** BEAS-2B, N=3, in the presence of cycloheximide. Rate constants are compared using extra sum-of-squares F test. For **B-I** Western blot quantifications of EGFR protein normalized to actin expression are shown. All error bars = SEM. \*  $\leq 0.05$ , \*\*  $\leq 0.01$ , \*\*\*  $\leq 0.001$ , # = 0.051

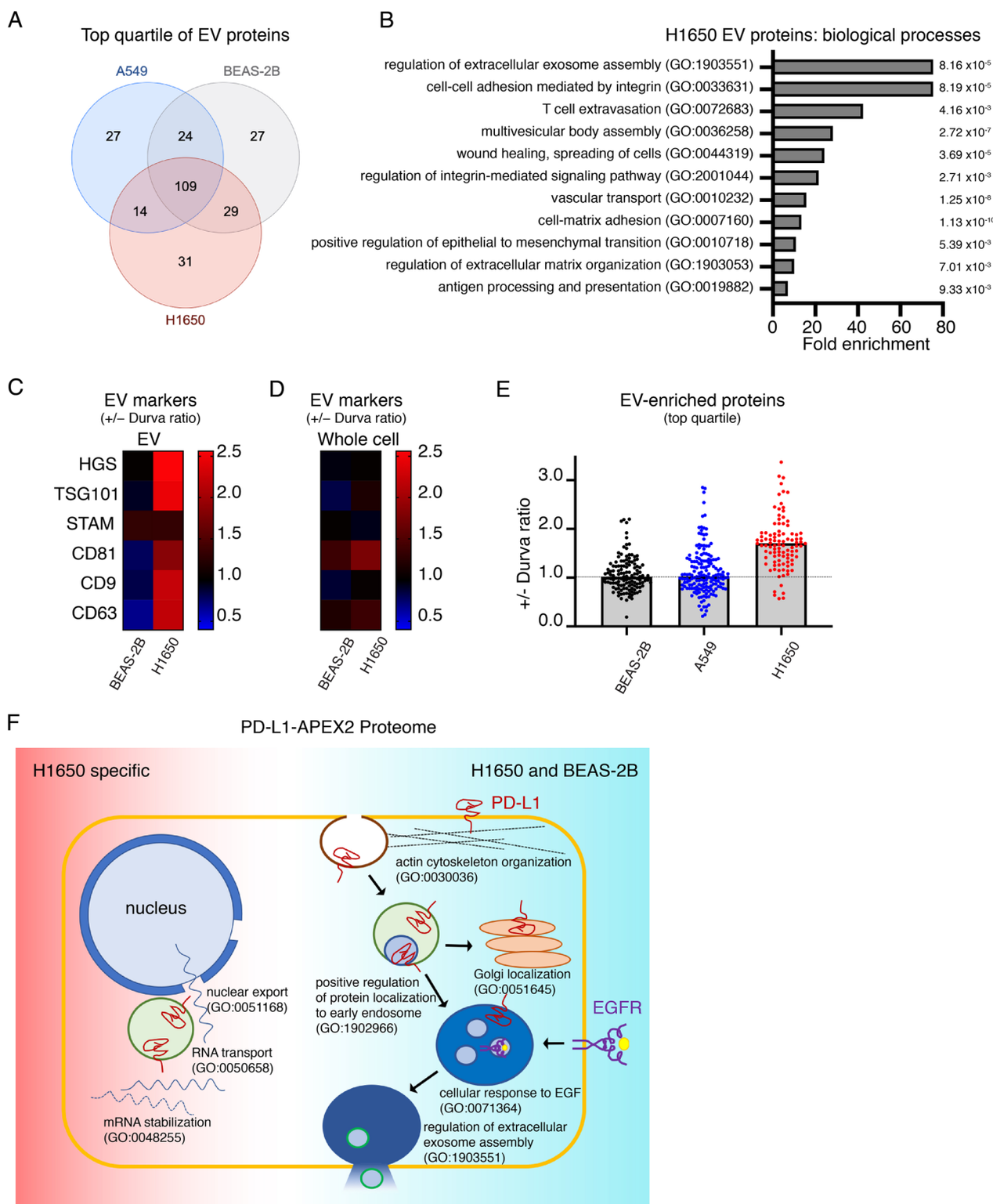


**Fig. 4** (See legend on previous page.)

proteins: HGS, TSG101, STAM, CD81, CD9 and CD63 [38] (Fig. 5C). Durvalumab did not affect the amounts of these proteins in EVs isolated from BEAS-2B cells, whereas Durvalumab increased these proteins, except for STAM, in EVs isolated from H1650 (Fig. 5C). Anti-PD-L1

antibody did not affect the amounts of these proteins in whole cell extracts (Fig. 5D).

Consistent with the effects of anti-PD-L1 antibody on the 6 canonical EV proteins, anti-PD-L1 antibody treatment caused a general increase of proteins detected in



**Fig. 5** Anti-PD-L1 antibody treatment affects extracellular vesicles. **A** Venn diagram of EV proteins with the highest intensities (top quartile) in each of the cell lines. **B** Biological processes representing proteins that are most enriched in the EVs of H1650 cells. Fold enrichment and FDR for the pathways are shown. **C** Heatmap of Durvalumab-induced fold change of extracellular vesicle marker proteins expression in **(C)** EV fraction and **D** whole cell lysate in BEAS-2B and H1650. **E** Durvalumab-induced fold change of extracellular vesicle-enriched proteins (top quartile for each cell line) in BEAS-2B, A549 and H1650. For **C–E** cells were treated with Durvalumab. Average +/- Durvalumab value for N=2 is shown. **F** Schematic of H1650 PD-L1-APEX2 proximity map shared with BEAS-2B and unique to H1650

EVs isolated from H1650 conditioned media and elicited no change in BEAS-2B or A549 (Fig. 5E). These data support  $\alpha$ PD-L1 antibody increasing EV biogenesis and or protein packaging selectively in EGFR mutant cells.

#### Anti-PD-L1 proximity proteome in EGFR mutant cells

Our functional data support the hypothesis that EGFR mutant cells are uniquely sensitive to anti-PD-L1 treatment. To determine whether this difference is revealed by a difference in PD-L1 protein in EGFR mutant cells, we repeated PD-L1-APEX2 proximity labeling in H1650 cells. There was considerable overlap in the PD-L1 proximity proteome between WT and EGFR mutant cells revealed at the level of biological pathways, including significant enrichments of proteins annotated to the control of membrane protein traffic pathways and to various aspects of cytoskeleton biology (Fig. 5F). At the individual protein level, there is greater than 60 percent correspondence in the PD-L1 proximal proteome between the WT and mutant EGFR cell lines, including proximity of PD-L1 to EGFR and ESCRT components (Additional file 4: Table S3). These data suggest that a difference in PD-L1 localization between wild-type and mutant EGFR does not account for the sensitivity of EGFR mutant cells to  $\alpha$ PD-L1 antibody treatment. Interestingly, PD-L1 proximity proteome in H1650 uniquely revealed proteins involved in nuclear export as well as RNA stability, which could be related to its role in regulation of RNA stability [39].

#### Discussion

PD-L1 is a transmembrane ligand for the PD1 immune checkpoint receptor [25, 26, 40, 41]. Tethering PD-L1 to the membrane provides spatial constraint on its immune checkpoint activity because the PD1-bearing cell must be close enough for PD-L1 on the surface of the expressing cell to bridge the distance between cells. Our proximity mapping reveals a second feature of spatial control of PD-L1 checkpoint activity. We identified HLA-I proteins are in close proximity to PD-L1 in non-transformed human lung epithelia cells and mutant EGFR NSCLC cell line (Additional files 2 and 4: Tables S1 & S3). This juxtaposition of HLA-I and PD-L1, formed in the absence of HLA-I binding to the TCR, ensures that PD-L1 is poised to engage PD1 once HLA-I has engaged the TCR, perhaps enhancing the efficiency of PD-L1 to attenuate TCR activation. This PD-L1 to HLA-I proximity is similar to PD1 to TCR proximity on the T cell [42–44]. Our findings extend the previous observation of PD-L1 adjacent to HLA-I in leukocytes by demonstrating this spatial control is not solely restricted to immune-to-immune cell interaction and that it exists in the absence of TCR-HLA-I engagement.

Because PD-L1 is a transmembrane protein, its PM density can be modulated by controlling membrane protein traffic. Our kinetic analyses demonstrate that the amount of PD-L1 is actively maintained by constitutive cycling between the interior of cells and the PM (Fig. 1). Proteins that regulate membrane protein traffic from and to the PM were enriched in the PD-L1 proteome, supporting our results that PD-L1 is constitutively transiting multiple membrane compartments rather than being statically localized to the PM and or a single intracellular site (Fig. 2, Additional file 2: Table S1). Others have previously probed the regulation of PD-L1 trafficking. Most notably, it has been shown that CMTM6 plays a key role in controlling the fate of internalized PD-L1 [11, 12]. CMTM6 KO results in increased degradation of PD-L1, reducing both PM PD-L1 and total cellular PD-L1. Our results established growth factor regulation of constitutive PD-L1 recycling as an additional mechanism for regulation of PD-L1 PM density (Fig. 1). Growth factor regulation, mediated by AKT activity, is independent of CMTM6 and IFN $\gamma$  regulation of total cellular PD-L1. Both CMTM6 and AKT-dependent control are acute modulators of PD-L1 density, acting more rapidly than IFN $\gamma$  transcriptional regulation, which could be advantageous in defending against normal tissue damage at a site of an active immune response, likely non-pathologic activity of PD-L1. AKT control of PD-L1 traffic was similar in non-transformed and transformed cells and it was unaffected by whether the driver oncogene was mutant EGFR or KRAS, indicating this is a fundamental feature of PD-L1 biology and not specific to cancer cells or specific driver mutation.

Regulation of PD-L1 recycling has an advantage compared to CMTM6 in the control of PD-L1 PM density because the changes are rapidly reversible since the location of PD-L1 not the total amount are modulated. Reduced growth factor signaling slows recycling of PD-L1, shifting PM PD-L1 to intracellular sites while maintaining constitutive cycling, albeit at a reduced rate (Additional file 1: Fig. S1D). Restoration of growth factor AKT activation redistributes intracellular PD-L1 to the PM by increasing recycling without requiring new synthesis of PD-L1, whereas reversal of CMTM6 depletion of PD-L1 requires new synthesis to overcome the effect of increased PD-L1 degradation.

Many studies have highlighted PD-L1 functions that are in addition to its role as an immune checkpoint ligand, including regulation of cell growth, EMT, and resistance to cellular stresses such as chemotherapy and IFN-mediated cytotoxicity [7, 8, 45]. Biological pathways enriched among the PD-L1 proximal proteome support the conclusions of many of those previous studies. Our studies of PD-L1 within different cell contexts (non-transformed,

EGFR mutant and KRAS mutant cells) advances the prior studies by revealing novel cell-context dependencies for PD-L1 role in cell migration as well as the formation and or release of exosomes.

PD-L1 has been shown to have a role in migration of cell lines with non-EGFR driver oncogenes [30]. Those studies relied on genetic manipulation of PD-L1. We confirm and extend those studies by showing genetic ablation of PD-L1 inhibits cell migration in non-transformed, EGFR mutant and KRAS mutant cell lines (Fig. 3). Strikingly, we found that anti-PD-L1 antibody treatment inhibited migration of EGFR mutant cells but had no effect on KRAS mutant (wild type EGFR) or non-transformed cells. We do not know why EGFR mutant cells are sensitive to anti-PD-L1 antibody in a way that other cells are not. In addition to possible changes in cell physiology arising from the constitutive EGFR signaling of mutant EGFR, wildtype and mutant EGFR have been described to have largely dissimilar downstream signaling [46, 47]. Our data suggest that the cellular context created by oncogenic EGFR signaling may make EGFR mutant cells more susceptible to anti-PD-L1 antibody-mediated signaling changes. In support of that view, a recent study showed PD-L1 induces phospholipase C- $\gamma$ 1 (PLC- $\gamma$ 1) activation only in the presence of activated EGFR [48]. However, additional experiments are required to uncover at a molecular level why anti-PD-L1 has specific effects specifically in mutant EGFR cells.

ESCRT proteins control the turnover of transmembrane proteins and extracellular vesicle formation (EV) by inducing the intraluminal budding of vesicles from the limiting membrane of endosomes to form multivesicular bodies (MVBs) [33, 49, 50]. The ESCRT pathway is key in the negative regulation of EGFR signaling by controlling EGFR degradation [51]. In agreement with previous studies identifying PD-L1 proximity with EGFR [52], and ESCRT accessory protein ALIX regulating PD-L1 sorting into intraluminal vesicles [15], we identified EGFR as well as several ESCRT complex proteins as components of the PD-L1 proximal proteome (Additional file 2: Table S1). Steady-state PD-L1 proximity to the ESCRT machinery might reflect normal turnover of PD-L1 by lysosomal degradation or EV secretion [11, 13, 15, 53]. However, our analyses strongly support the hypothesis that PD-L1 is functionally associated with the ESCRT pathway beyond being a client for MVB sorting. Anti-PD-L1 treatment had two effects on ESCRT-dependent functions in EGFR mutant cells: turnover of constitutively active EGFR was significantly reduced and biogenesis/secretion of EVs was significantly increased.

Notably, the effects of anti-PD-L1 treatment on ESCRT pathway functions, like the effect on cell migration, were only observed in EGFR mutant cells. Genetic ablation of

PD-L1 in wild type EGFR cells mimics the effects of anti-PD-L1 treatment on EGFR mutant cells, suggesting a more significant perturbation is required to disrupt cell-intrinsic PD-L1-dependent functions in wild type EGFR cells. We do not understand what it is about constitutive EGFR signaling that sensitizes cells to the effects of anti-PD-L1 therapy.

Our PD-L1-APEX2 proximity labeling in wild-type BEAS-2B and EGFR-mutant H1650 cells shows a largely overlapping list of proteins, including high enrichment for EGFR, ESCRT proteins as well as PLC- $\gamma$ 1. However, due to varying levels of biotin-phenol uptake and APEX2 labeling efficiency between the cell lines, we are unable to quantitatively compare the enrichment of these proteins in PD-L1 proximity. Future experiments involving exogenous expression of mutant EGFR in wild-type cells may be useful to describe quantitative changes in protein proximity and interaction with PD-L1 that may help explain how mutant EGFR signaling sensitizes cells to perturbations in PD-L1 biology.

The effect of antibody binding to PD-L1 has not been extensively studied in the context of PD-L1 reverse signaling. Our data suggest antibody binding in mutant EGFR cells results in a loss of function of PD-L1 as it phenocopies the effect of PD-L1 KO on EGFR expression, cell migration and EV secretion. Future studies are required to determine whether the effects of anti-PD-L1 antibody binding are due to a perturbation in PD-L1 transmembrane signaling or disruption in PD-L1 binding with partner proteins, which are responsible for signaling changes. Recent studies have identified PD-L1 signaling motifs and binding partners that result in broad transcriptional and signaling changes within the cells [8, 48, 54].

The effects of anti-PD-L1 antibody on EGFR mutant cells we discovered might be informative for understanding the increased resistance of EGFR mutant tumors to anti-PD-L1 therapy [5, 6, 55]. In particular, the anti-PD-L1 antibody-induced increase in EVs from mutant EGFR cells could have broad ranging effects on the tumor microenvironment and tumor immune landscape that could contribute to resistance to immunotherapy [56, 57]. PD-L1 was not detected by mass spectrometry in EVs from any of the cell lines, therefore our data do not speak to the possibility that PD-L1 on EVs serves to perturb immune activity or that it affects anti-PD-L1 therapy as reported [56].

PD-L1 expression on immune cells has an important immune-modulatory role [54, 58–60] that will also be affected by anti-PD-L1 therapy. For example, PD-L1 on dendritic cells (DCs) is necessary for efficient homing of dermal DCs to the lymph nodes through modulation of chemokine receptor signaling and actin remodeling [54]. Our finding that the effects of anti-PD-L1 are

cell-context dependent raise the possibility that anti-PD-L1 effects in immune cells extend beyond blocking PD1 engagement.

### Limitations

As a hypothesis generating study our work provides a foundation for future work, in particular studies designed to better understand the specific sensitivity of mutant EGFR cells to anti-PD-L1 antibody. A limitation of our work is that all studies were performed in cultured cells. However, we are technically constrained in this regard because there are no syngeneic mouse mutant EGFR cell lines that can be used for in vivo testing of hypotheses emerging from our in vitro data in an immune competent mouse.

### Conclusions

Here we have we focused on two aspects of cell-autonomous PD-L1 biology. We identify growth factor signaling as a mode to acutely and reversible control the density of PD-L1 in the plasma membrane, and by doing so potentially impact PD-L1's activity as an immune checkpoint ligand for PD1 receptor. This acute regulation of PD-L1 plasma membrane density is in addition to transcriptional control and previously described regulation of total PD-L1 levels by control of its degradation. We exploited the PD-L1 proximity proteome generated by APEX2 proximity mapping to further explore cell-autonomous PD-L1 biology. Most notably, our work reveals novel PD-L1 biology restricted to mutant EGFR cells. Anti-PD-L1 antibody treatment of mutant EGFR cells perturbs cell-intrinsic PD-L1 functions, leading to reduced cell migration, increased half-life of EGFR and increased extracellular vesicle biogenesis, whereas anti-PD-L1 antibody does not induce these changes in wild type EGFR cells. These effects might contribute to the relative resistance of EGFR mutant tumors to immune checkpoint blockade therapy. Our hypothesis generating study provides a foundation for future mechanistic work, in particular studies designed to better understand the specific sensitivity of mutant EGFR cells to anti-PD-L1 antibody.

### Abbreviations

NSCLC	Non-small-cell lung carcinoma
EMT	Epithelial mesenchymal transition
PM	Plasma membrane
KD	Knockdown
KO	Knockout
EGFR	Epidermal growth factor receptor
EV	Extracellular vesicles
WCE	Whole cell extract
MVB	Multivesicular bodies
IFN $\gamma$	Interferon gamma

### Supplementary Information

The online version contains supplementary material available at <https://doi.org/10.1186/s12964-023-01084-6>.

**Additional file 1: Fig. 1.** PD-L1 is dynamically maintained on the plasma membrane of cells. (A) Epifluorescent images of BEAS-2B cells treated with IFN $\gamma$  and stained for PM PD-L1 using bivalent secondary antibody or monovalent secondary Fab. (B) Representative confocal images of human lung cell lines with or without IFN $\gamma$  treatment and immunostained for PM and total PD-L1. Red: PD-L1. Blue: Hoechst. Sum intensity of z-stacks is projected. Fluorescence intensity is equally scaled across panels for each cell line. Insets in -IFN $\gamma$  PM are enhanced for better visualization. (C) Total PD-L1 protein expression in BEAS-2B cells with growth factor treatment or Akt inhibition. One-way ANOVA. (D) Exocytosis rate constants in BEAS-2B cells incubated with vehicle (DMSO) or Akt inhibitor. Paired student's t-test. (E) Immunoblots showing Akt T308 phosphorylation in cells treated with vehicle (DMSO) or Akt inhibitor. Corresponding quantification of phospho-T308/total Akt is shown in the graph. (F) CMTM6 and PD-L1 mRNA expression, PD-L1 protein expression and distribution in BEAS-2B expressing doxycycline inducible CMTM6 shRNA. One sample and unpaired student's t-tests. All error bars = SEM. \*  $\leq 0.05$ , \*\*  $\leq 0.01$ , \*\*\*  $\leq 0.001$ . **Additional file 1: Fig. 2.** PD-L1 proximity mapping. (A) Representative immunoblot of biotinylated proteins in labeled and control BEAS-2B cells. (B) PD-L1 co-localization with specified proteins quantified by fraction of PD-L1 intensity co-localized with each target protein. (C) PD-L1 co-localization with EGFR in BEAS-2B and H1650 under basal and EGF stimulated conditions. For (B-C), each data point is a collected from single plane images of individual cells under confocal microscopy. **Additional file 1: Fig. 3.** A role for PD-L1 in cell migration. (A) Immunoblots of PD-L1 in control and PD-L1 KO cell lines. Tubulin is used as loading control. (B) Timelapse images of scratch assay at 0, 8 and 24 h in BEAS-2B cells. (C-D) Timelapse images of scratch assay at 0, 9 and 24 h in (C) H1650 PD-L1 KO rescue cells and (D) H1650 cells treated with Durvalumab. For (B-D) Leading edge of the cells are marked in white. Cells were in (B, D) serum-free media and (C) serum-complete media for the duration of the assay. (E-F) Cell proliferation as measured by MTT assay in (E) H1650 cells treated with Durvalumab, N = 2, and (F) H1650 PD-L1 KO rescue cells, N = 3. For (E-F) difference in slope values is calculated using F-test. All error bars = SEM. \*  $\leq 0.05$ , \*\*  $\leq 0.01$ , \*\*\*  $\leq 0.001$ . **Additional file 1: Fig. 4.** Anti-PD-L1 antibody treatment inhibits turnover of mutant but not wild type EGFR. (A) PD-L1 protein expression in BEAS-2B cells transfected with control and PD-L1 siRNA. One sample t-test. (B-D) Immunoblots of EGFR expression in (B) H1650 PD-L1 KO/rescue cells and H1650 cells treated with (C) PD-L1 antibody and (D) TFR antibody. (E-F) Distribution of (E) PD-L1 and (F) TFR in H1650 cells treated with Durvalumab. Unpaired multiple t-tests. (G) Immunoblots of EGFR degradation in BEAS-2B and H1650 treated with cycloheximide in serum complete media. Actin is used as loading control. (H-I) mRNA expression of (H) EGFR and (I) PD-L1 in H1650 cells treated with  $\alpha$ PD-L1 antibodies. (J) EGF stimulated gene transcription in H1650 cells treated with  $\alpha$ PD-L1 antibody. qRT-PCR mRNA quantification of EGFR target genes is shown, N = 3. All error bars = SEM. \*  $\leq 0.05$ , \*\*  $\leq 0.01$ , \*\*\*  $\leq 0.001$

**Additional file 2.** Table S1. PD-L1-APEX2 BEAS-2B

**Additional file 3.** Table S2. EV proteins

**Additional file 4.** Table S3. PD-L1-APEX2 H1650

**Additional file 5.** Table S4. RT-PCR primer sets

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#### Author contributions

AL designed and performed the experiments, analyzed the data, and wrote the manuscript. EY designed experiments, generated the KO cells, and edited the manuscript. PC and ND performed and assisted with APEX2 mass spectrometry experiments and analysis. NKA conceptualized the project, contributed to experimental design and interpretations, and edited the manuscript. TEM conceptualized the project, contributed to experimental design, performed analysis and interpretation, and wrote the manuscript. All authors read and approved the final manuscript.

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#### Availability of data and materials

The processed mass spectrometry data are provided as supplemental tables. The datasets used and/or analyzed during the current study, including the raw mass spectrometry data, are available from the corresponding author on reasonable request.

#### Declarations

##### Ethics approval and consent to participate

Human lung tumor tissue samples were obtained from surgically resected material at New York-Presbyterian-Weill Cornell Medicine. The study was approved by the New York-Presbyterian-Weill Cornell Medicine institutional review board and patient consent was received for these studies.

##### Competing interests

The authors have no competing interests to disclose related to this study. NKA has equity in Angiocrine Bioscience. TEM receives research funding from Janssen and from Pfizer, Inc. The other authors have nothing to disclose.

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