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ABSTRACTS

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1869 Immunohistochemical Evaluation of Loss of Expression of BRCA-1 Associated Protein 1 (BAP1) on Tumors and Normal Tissues from Various Organs

Syeda Absar¹, Haiyan Liu¹, Jianhui Shi¹, Fan Lin¹ ¹Geisinger Medical Center, Danville, PA

Disclosures: Syeda Absar: None; Haiyan Liu: None; Jianhui Shi: None; Fan Lin: None

Background: Loss of expression of BRAC-1 associated protein 1 (BAP1) was reported in 69% of biphasic mesotheliomas and 15% of sarcomatoid mesotheliomas (Cigognetti M et al. Modern Pathology 2015; 28:1043-1057). However, available data on the BAP1 expression in carcinomas and other tumors were limited. Because differentiating a mesothelioma from an adenocarcinoma and other tumors is clinically significant, in this study we focused on the evaluation of BAP1 expression in a large series of carcinomas and tumors from various organs.

Design: Immunohistochemical analysis of BAP1 (sc-28383; Santa Cruz Biotechnology, Inc.) was performed on 1233 cases of tumor and normal tissues on tissue microarray (TMA) sections, including mesothelioma (N=18), lung neuroendocrine carcinoma (CA) (N=24), lung adenocarcinoma (ADC) (N=88), lung squamous cell CA (N=66), papillary thyroid CA (N=47), ENT squamous cell CA (N=28), breast fibroadenoma (N=20), pancreatic neuroendocrine tumor (N=33), pancreatic ADC (N=43), adrenal pheochromocytoma (N=14), endometrial CA FIGO II (N=59), ovary papillary serous CA (N=41), clear cell CA of uterus and ovary (N=22), melanoma (N=32), skin neuroendocrine CA (N=27), invasive urothelial CA (N=43), bladder small cell CA (N=24), prostatic ADC (N=38), germ cell tumors (N=60), colonic ADC (N=164), angiosarcoma (N=12), papillary renal cell carcinoma (RCC) (N=33), clear cell RCC, low grade (N=79), clear cell RCC, high grade (N=51), hepatocellular CA (N=47), liver metastatic neuroendocrine CA (N=18), glioblastoma multiforme (N=23), mantle cell lymphoma (N=13) and hairy cell leukemia (N=1). Samples from normal tissue included pancreas (N=13), rectum/appendix/colon (N=39) and ilium/duodenum/stomach (N=13). Nuclear expression for BAP1 was regarded as positive. Loss of expression of BAP1 was recorded when internal positive controls were present.

Results: Loss of expression of BAP1 was noted in 17% of mesothelioma cases (3 of 18). Intact BAP1 expression was present in all other tumors and normal tissues.

Conclusions: Our data demonstrated that BAP1 expression was not identified in carcinomas and other tumors from various organs, which further validated the previous report that loss of expression of BAP1 was highly specific for a malignant mesothelioma. However, the loss of expression of BAP1 was only seen in a minority of mesotheliomas. The low diagnostic sensitivity in this study is most likely due to the small number of cases, including some with sarcomatoid features.

1870 Comparison of Solid Tissue Sequencing and Liquid Biopsy: Identification of Clinically Relevant Gene Mutations and Rearrangements in Lung Adenocarcinomas

Douglas Allison¹, George Jour¹, Kyung Park¹, Deborah DeLair², Andre Moreira³, Matija Snuderl⁴, Paolo Cotzia⁵

¹NYU Langone Health, New York, NY, ²NYU Langone Medical Center, New York, NY, ³New York Langone Health, New York, NY, ⁴New York University, New York, NY, ⁵New York University Langone Medical Center, New York, NY

Disclosures: Douglas Allison: None; George Jour: None; Kyung Park: None; Deborah DeLair: None; Andre Moreira: None; Matija Snuderl: None; Paolo Cotzia: None

Background: Molecular screening for therapeutically targetable alterations is considered standard of care in the management of non-small cell lung cancer. However, most molecular assays utilize tumor tissue, which may not always be available. This has led to the development of "liquid biopsies": plasma-based Next Generation Sequencing (NGS) tests that use circulating tumor DNA as a substrate to identify relevant targets. In this study, we sought to determine the level of agreement between the two tests as they are used in clinical practice and to investigate the utility of concurrent plasma/tissue testing.

Design: We identified 47 cases of lung adenocarcinoma diagnosed over the past 2 years, who received concurrent testing (within 24 weeks) with both our institution's tissue (DNA and RNA based) NGS assay and a commercial plasma-based NGS assay. The results were reviewed to establish concordance in the identification of mutations or fusions deemed clinically relevant or for which a targeted therapy was available.

Results: Patients in our cohort represented both new diagnoses (31 cases, 66%) and disease progression on treatment (16 cases, 34%). The majority (83%) had stage 4 disease. Tissue NGS identified clinically relevant mutations in 39 cases (83%), including in 14 (88%) of the previously treated cases. By comparison, plasma NGS identified clinically relevant mutations in 20 cases (43%, p<0.001), including 6 treated cases (38%, p=0.01). Tissue NGS identified therapeutic targets in 55% of cases and 75% of previously treated cases; while plasma NGS identified targets in 28% and 25% respectively (p<0.001 and p=0.01 respectively). All clinically relevant mutations identified by plasma NGS were also detected by tissue NGS, while plasma NGS detected only 51% those identified by tissue NGS. Discrepant cases involved hotspot mutations and actionable fusions including those in *EGFR*, *KRAS*, and *ROS1* (Table 1).

Table 1. Discrepant Findings						
Gene Mutation/Fusion						
KRAS	G12C (x2), G12D, G12E, G13C, Q61K					
EGFR	Exon 19 deletion (x6), G719A, L858R, M766_A767insASV					
ALK	EML4-ALK fusion (x2)					
ROS1	SLC34A2-ROS1 fusion, ZCCHC8-ROS1 fusion					

Conclusions: Tissue NGS detects more clinically relevant alterations and therapeutic targets compared to plasma NGS, especially in the post-treatment setting, suggesting that tissue NGS should be the preferred method for molecular testing of lung adenocarcinoma. Additionally, all clinically relevant mutations identified by plasma NGS were also detected by tissue NGS, suggesting that tissue/plasma cotesting provides little additional benefit over tissue NGS alone. Plasma NGS can detect clinically relevant targets, and still plays an important role when tissue testing is impractical or not possible.

1871 D4D6 and SP384 Immunohistochemistry Clones for Screening of ROS1 Rearrangements in Non-Small Cell Lung Carcinoma: Comparison of Different Platforms with Focus on Specificity

Wajd Althakfi¹, Michèle Orain², Nathalie Bastien², Christian Couture³, Patrice Desmeules⁴, Philippe Joubert⁵

¹Quebec Heart and Lung Institute, King Saud University, Quebec City, QC, ²Institut Universitaire de Cardiologie et de Pneumologie de Quebec Research Center, Quebec, QC, ³IUCPQ, Quebec City, QC, ⁴IUCPQ - Quebec Heart and Lung Institute, Quebec, QC, ⁵Quebec Heart and Lung Institute, Quebec, QC

Disclosures: Wajd Althakfi: None; Michèle Orain: None; Nathalie Bastien: None; Christian Couture: *Advisory Board Member*, Pfizer; Patrice Desmeules: *Grant or Research Support*, Pfizer Canada; *Grant or Research Support*, Novartis; Philippe Joubert: None

Background: Detection of *ROS1* gene rearrangements is recommended in all advanced non-small cell lung cancer (NSCLC) patients for proper selection of candidates for targeted therapy. Immunohistochemistry (IHC) is useful as a screening tool for this rare alteration, but protocols and proper cut-offs are not standardized, resulting in modest specificity and high rate of required confirmatory testing. Until recently, D4D6 (Cell Signaling Technology) was the single antibody clone available but performance characteristics of the new SP384 clone (Ventana) are emerging. Here we present our experience with both clones using tissue microarrays (TMA) and routine samples with different IHC staining platforms.

Design: A retrospective cohort consisting of 540 lung adenocarcinomas was assessed using TMAs. Immunohistochemistry was performed in parallel with D4D6 clone on a Dako link48 autostainer using a laboratory developed IHC protocol, and with SP384 clone on a Ventana BenchMark Ultra using the manufacturer's instructions. A selection of 50 additional cases from a prospective screening cohort showing any level of immunostaining using on the D4D6 protocol was stained with SP384 for comparison. All tumors underwent ROS1 fluorescent in situ hybridization testing and next-generation sequencing was also performed on a subset of cases. Quantification of immunostaining was performed on digitalized slides using H-score.

Results: Performance was similar for detecting ROS1-rearranged cases (n=10) with D4D6 (median H-Score; range: 205; 0-300) and SP384 (230; 0-290) protocols. In the TMA set, any level of non-specific immunostaining was seen in 23,3% and 35,7% of tumors, and the sensitivity was 97,8% and 93,7% when using a H-score cut-off of 150, with D4D6 and SP384 protocols, respectively. Overall, H-score correlation was modest (R²=0,59) but mean H-score difference low (14) between protocols. However, differences greater or equal to 100 H-score was observed in 44/540 (9,8%) and 15/50 (30%) samples from both the retrospective and prospective cohorts and in both directions in regard to the clone used. Large differences could not be explained by sample type (biopsy or cytology specimens) or low number of tumor cells (2/15).

Conclusions: Using two different ROS1 IHC protocols with D4D6 and SP384 antibody clones on different automated platforms, we found that overall performance was similar with a slightly better specificity favoring the first protocol. However, we noted a subset of samples exhibiting striking differences

1872 CD5, CD117, BAP1, MTAP and TdT is a Useful Immunohistochemical Panel to Distinguish Thymoma from Thymic Carcinoma

Mounika Angirekula¹, Sarah Jenkins¹, Anja Roden¹

*Mayo Clinic, Rochester, MN

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Background: The distinction between thymic carcinoma and thymoma, specifically types B3, A and occasionally micronodular thymoma with lymphoid stroma (MNTLS) can be challenging as also has been shown in interobserver reproducibility studies. However, thymic

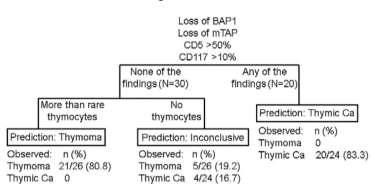
carcinomas have a worse prognosis with a mean 10-year survival of 29% vs 62% and 97% for types B3 and A thymomas, respectively. This study aims to identify immunohistochemical markers that might aid in the distinction between thymoma and thymic carcinoma.

Design: Thymic carcinomas, type A and B3 thymomas and MNTLS (1963-2019) were identified in the database of thymic epithelial tumors (TET) of one of the authors. Immunohistochemistry (IHC) was performed using antibodies to TdT, Glut-1, CD5, CD117, BAP1 and mTAP. Percent tumor cell staining was recorded (Glut-1, CD5, CD117); loss of expression (BAP1, mTAP) was considered if essentially all tumor cells were negative; TdT was recorded as thymocytes present or absent (which included cases with only rare thymocytes). Statistical analysis was performed.

Results: Sixty-six specimens included 55 completely and 8 incompletely resected and 3 biopsied thymomas (N=36; 18/36 type A, 12 B3, 6 MNTLS) and thymic carcinomas (N=30, including 18 squamous cell carcinomas). The results of immunostains are summarized in table 1. Complete IHC data were available in 50 patients. Using BAP1, mTAP, CD5, CD117 and TdT, 83.3% of thymic carcinomas and 80.8% of thymomas could be predicted (Figure 1). Glut-1 expression was not found to be useful in that distinction.

	Table 1. Results of immunostains							
		Thymic Carcinoma	Thymoma	p-value				
N		30	36					
BAP1	% cases with loss of expression	13.8	0	0.04				
MTAP	% cases with loss of expression	14.8	0	0.03				
Glut1	Median % positive tumor cells (range)	90 (0-100)	5 (0-90)	<0.001				
CD5	Median % positive tumor cells (range)	5 (0-100)	0 (0-30)	0.004				
CD117	Median % positive tumor cells (range)	90 (0-100)	0 (0-1)	<0.001				
TdT	% cases with more than rare thymocytes	O ^a	82.1	<0.001				
		^a a single case with rare thymocytes						

Figure 1 - 1872



6 thymic carcinomas (thymic Ca) and 10 thymomas excluded due to incomplete data for one or more IHC stains.

Conclusions: A panel of immunostains including BAP1, mTAP, CD5, CD117 and TdT can be useful in the distinction between thymomas and thymic carcinomas with only a minority of cases being inconclusive.

1873 Bronchiolar Adenoma: a Pneumocytoma-Like Component and Senescence Markers Expression Ana Aula Olivar¹ Marta Sesé² Jorge Luis Rodríguez³ Luisa Sofía Silva Alcoser⁴ Mario Giner Pichel⁴ Santiago Ramon V Caial

Ana Aula Olivar¹, Marta Sesé², Jorge Luis Rodríguez³, Luisa Sofía Silva Alcoser⁴, Mario Giner Pichel⁴, Santiago Ramon Y Cajal⁴, Irene Sansano⁵

¹Vall d'Hebron University Hospital (HUVH), Barcelona, Spain, ²Molecular Pathology Group. Vall d'Hebron Institute Research. Barcelona. Spain, Barcelona, Spain, ³Hospital Universitario San Ignacio, Pontificia Universidad Javeriana, Bogotá, D.C., Colombia, ⁴Vall d'Hebron University Hospital, Barcelona, Spain, ⁵Hospital Universitari Vall d'Hebron, Barcelona, Spain

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Background: Bronchiolar adenoma (BA) is a recently described benign lung tumor defined as a nodular proliferation of bronchiolar-type epithelium with a continuous layer of basal cells. They can reach several cm in size, harbor oncogenic mutations and be misdiagnosis as adenocarcinoma (ADK). As these tumors show a very good clinical prognosis we hypothesized that could be senescent. There are not yet

feasible markers of cellular senescence on paraffin but the expression of p16 protein and a low ki67 are surrogate markers of that cellular stage. Sclerosing pneumocytoma (SP) is a benign tumor with a combination of patterns and a characteristic dual population of surface cells and round cells. The first express TTF1 and CK and the last only TTF1. No driver mutations have been described in this lesion.

Design: We reviewed the histological characteristics and immunohistochemical findings of 4 BA from 3 patients and compared it to 1 SP and 1 ADK (from the patient with 2 BA). The antibodies used were: TTF1 (Ventana, clone 8G7G3), Ki67 (Ventana, 30-9), p16 (CINtec, E6H4) and p40 (Ventana, BC28). We also performed next-generation sequencing (NGS) (Thermofisher, panel Oncomine solid tumor) for all the BA and the ADK.

Results: All 4 BA displays high expression of p16 protein in over 80% cells and a proliferative index below 3%. One of the BA contained round cells CK- and TTF1+, similar to round cells in SP. SP, SP-like cells and ADK did not show a significant p16 expression. See table 1.

Case	1	2	3a	3b	3c	4
Age	51	51	75		*	46
Sex	\$	\$	<i>d</i>	*	*	3
Smoking history	No	No	Yes	*	*	Unk
Size (cm)	1,7	0,8	2,9	0,6	0,5	Unk
Morphology	Distal	Proximal (Papillary)	ADK	Proximal (flat)	Distal	SP
Cilia	Present	Present		Present	Present	
Round cells (TTF1+, CK-)	No	Yes	No	No	No	Yes
Frozen section diagnosis	-	ADK	-	-	-	-
TTF1	+	-	+	-	+	+
P16	++	+++	+ focal	++	+++	-
Ki67	<1%	<1%		<1%	<1%	<1%
Putative driver mutation	KRAS c.35G>A p.Gly12Asp VAF 6.3 %	FGFR3 c.1958A>C p.Asn653Thr VAF 48.9 %	KRAS c.35G>A p.Gly12Asp VAF 26.8 %	-	-	-

Conclusions: We have identified a SP-like component that has not been described in classic BA.

BA may represent another kind of benign tumor with senescent features and oncogenic alterations, similarly to nevus or pylocitic astrocytomas. These findings can be relevant in the differential diagnosis with adenocarcinomas as well as in the understanding of the malignant transformation in lung tumors. Further research in a larger number of cases is needed.

1874 Histologic Patterns of Lung Adenocarcinoma Reported on Biopsy Accurately Predict Composition of Resected Tumor

Marina Baine¹, William Travis¹, Darren Buonocore¹, Jason Chang¹, Gross Daniel¹, Natasha Rekhtman¹, Prasad Adusumilli¹, Jennifer Sauter¹

¹Memorial Sloan Kettering Cancer Center, New York, NY

Disclosures: Marina Baine: None; William Travis: None; Darren Buonocore: None; Jason Chang: None; Gross Daniel: None; Natasha Rekhtman: None; Prasad Adusumilli: None; Jennifer Sauter: None

Background: Solid (S) and micropapillary (MP) histologic patterns are known poor prognostic features in patients with lung adenocarcinoma, suggesting that more extensive surgery, such as segmentectomy or lobectomy, may be indicated for early stage adenocarcinoma when these patterns are found on biopsy. Thus, accurate recognition of these high-grade patterns on biopsy may become critical for surgical management. Currently, little data are available on accuracy of recognition of adenocarcinoma histologic patterns on biopsy and how they relate to choice of surgery and patient outcomes.

Design: Pathology reports of paired biopsies and resections of non-mucinous lung adenocarcinomas (n=100) between January-July 2019 were retrospectively reviewed. Digital slides were reviewed from cases where a pattern reported on biopsy was not documented on resection.

Results: In most cases (n=85; 85%), patterns identified on biopsy were also seen on subsequent resection. In remaining cases, papillary (n=5;5%), S (n=5;5%), MP (n=4;4%) and acinar (n=1;1%) patterns were reported on biopsy but not on resection. In 40 (40%) and 75 (75%) of discrepant cases, S and MP patterns, respectively, were seen on re-review of resection slides but comprised ≤5% of total tumor. Remaining discordant S (n=2) and MP (n=1) patterns were due to biopsy artifact including acinar collapse/telescoping and acinar-lining tumor cells shedding into the lumen, respectively. Only 5 (5%) cases in this study could have ultimately resulted in potentially unnecessary more extensive surgery. Conversely, in 83 (83%) cases, patterns seen on resection were not all initially reported on biopsy, with MP being the most common pattern with this discrepancy. Most discrepancies were due to sampling, as the patterns missing on biopsy were often minor components (mean of 25%) of total tumor.

Conclusions: Histologic patterns of adenocarcinoma are accurately recognized on biopsy in the majority of cases. Often when S or MP are reported on biopsy but not on resection, these patterns comprise a small component of total tumor. Biopsies with interpretive discrepancies from subsequent resection specimens are primarily due to biopsy artifacts. Thus, while documenting the presence of S or MP on biopsy is important, our findings suggest considering artifact in this setting. Although most missed patterns on biopsy can be attributed to sampling, among the remaining minority, MP is the most commonly underrecognized pattern.

1875 The Distinction between Spread through Air Spaces (STAS) and Artifacts in Selected Images is Highly Reproducible

Marina Baine¹, Mary Beasley², Darren Buonocore¹, Jason Chang¹, Teh-Ying Chou³, Mari Mino-Kenudson⁴, Natasha Rekhtman¹, Jennifer Sauter¹, Arne Warth⁵, William Travis¹

¹Memorial Sloan Kettering Cancer Center, New York, NY, ²Mt. Sinai Medical Center, New York, NY, ³Taipei Veterans General Hospital, Taipei, Taiwan, ⁴Massachusetts General Hospital, Boston, MA, ⁵Institute for Pathology, University Hospital, Wetzlar, Hessen. Germany

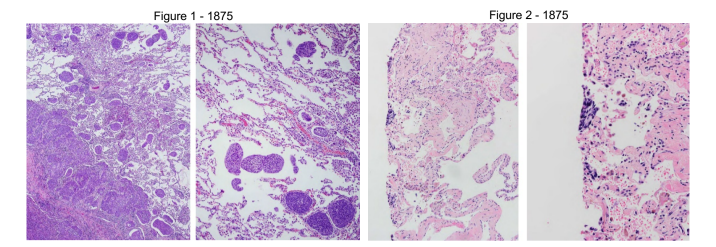
Disclosures: Marina Baine: None; Mary Beasley: None; Darren Buonocore: None; Jason Chang: None; Teh-Ying Chou: None; Mari Mino-Kenudson: None; Natasha Rekhtman: None; Jennifer Sauter: None; Arne Warth: None; William Travis: None

Background: Spread through air spaces (STAS) is a pattern of spread beyond the edge of the main tumor associated with increased recurrence and poor survival in all major histologic types of lung cancer studied. The definition of STAS includes criteria for excluding artifacts which have not been tested with reproducibility studies.

Design: 30 sets of images of artifacts and STAS were distributed as a PowerPoint file in PDF format to 10 observers from 5 institutions and 3 countries to evaluate for reproducibility. A defined set of criteria were distributed. The criteria for STAS were: 1) spread of tumor cells beyond the edge of the main tumor into the air spaces surrounding the tumor; 2) three patterns include: micropapillary, solid nest and single cell; 3) STAS tumor cells should be at least one airspace beyond the tumor edge and 4) there should be two or more air space clusters. The criteria favoring artifacts included: 1) Mechanically induced tumor floaters are randomly situated often at the edge of the tissue section or out of the plane of section, 2) Jagged edges of tumor cell clusters suggests tumor fragmentation or edges of a knife cut during specimen processing; 3) Isolated tumor clusters at a distance from the tumor rather than spreading in a continuous manner from the tumor edge; 4) Linear strips of tumor cells that are lifted off alveolar walls.

Results: Of the 30 cases 16 were artifacts and 14 were STAS. In 24/30 (80%) there was unanimous agreement: 13/16 (81%) and 11/14 (79%) were unanimously interpreted to be artifacts and STAS, respectively. Kappa values averaged 0.857 (range 0.614-1.00) which is almost perfect (Table). In the 6 cases with disagreements the number of discordant opinions was in 4 (n=1), 2 (n=3) and 1 (n=2) cases. In communicating with the participants, it seemed that these disagreements were mostly due to study design issues. The greatest disagreement involved a case of alveolar macrophages where tumor cells were not present in the photograph for comparison.

	P2	P3	P4	P5	P6	P7	P8	P9	P10
P1	0.933	0.933	0.933	0.796	0.933	0.933	0.933	0.737	1.000
P2	0.000	1.000	0.867	0.733	1.000	0.867	0.867	0.867	0.933
P3			0.867	0.733	1.000	0.867	0.867	0.800	0.933
P4				0.862	0.867	0.864	0.867	0.675	0.933
P5					0.733	0.862	0.773	0.614	0.796
P6						0.867	0.867	0.800	0.933
P7							0.867	0.675	0.933
P8								0.800	0.933
P9									0.737



Conclusions: In summary this study of selected images suggests that the distinction between STAS and artifacts is achievable and reproducible. Further work on reproducibility is needed using glass slides and frozen sections. These data suggest that these originally proposed criteria to make the distinction between artifacts and STAS are readily applicable. This likely has helped the many independent investigators who have performed studies demonstrating the clinical significance of STAS.

1876 Novel Biological Subsets of Small Cell Lung Carcinoma Defined by ASCL1 and NeuroD1: Immunohistochemical and Histopathological Characterization

Marina Baine¹, Wei-Chu Lai¹, Jacklynn Egger¹, Hira Rizvi¹, Amanda Beras¹, William Travis¹, Jennifer Sauter¹, Jason Chang¹, Darren Buonocore¹, Achim Jungbluth¹, Charles Rudin¹, Natasha Rekhtman¹, John Poirier²

**Memorial Sloan Kettering Cancer Center, New York, NY, **NYU Langone School of Medicine, New York, NY

Disclosures: Marina Baine: None; Wei-Chu Lai: *Advisory Board Member*, G1 Therapeutics; Jacklynn Egger: None; Hira Rizvi: None; Amanda Beras: None; William Travis: None; Jennifer Sauter: None; Jason Chang: None; Darren Buonocore: None; Achim Jungbluth: None; Charles Rudin: None; Charles Rudin: None; Non

Background: Recent preclinical investigations have proposed the existence of biological subsets in small cell lung carcinoma (SCLC) defined primarily by expression of ASCL1 and NeuroD1 transcription factors, which are associated with distinct downstream transcriptional programs and therapeutic liabilities. In model systems expression of these markers is inversely correlated. Currently, there are only limited data on protein expression of these markers and associated pathologic characteristics in clinical samples of SCLC.

Design: A total of 99 SCLC were analyzed for expression of ASCL1 and NeuroD1 by immunohistochemistry either in tissue microarrays (TMA; n=39) or full samples (n=60). Subsets defined by these markers were compared for expression of conventional neuroendocrine markers (CNM; chromogranin, synaptophysin, CD56, and INSM1) and TTF1.

Results: Of 99 SCLC, ASCL1 was positive in 83%, and NeuroD1 in 48%. Combined expression was as follows: ASCL1+/NeuroD1+42.4%, ASCL1+/NeuroD1-38.4%, ASCL1-/NeuroD1+7.1%, and ASCL1-/NeuroD1-12.1%. For a subset of cases (n=39) relative expression level of these markers was analyzed by quantitative H scores (range 0-300), revealing that in most cases one of the markers was strongly dominant over the other (≥100 differential in H-scores in 80% of cases). Using quantitative scores, 72% of cases were ASCL1-dominant, 15% NeuroD1-dominant, and 13% double-negative. ASCL1- vs NeuroD1-dominant cases showed no differences in expression of CNMs, TTF1 or histopathologic characteristics. Consistent with published gene expression observations, double-negative cases were strongly associated with expression of fewer CNMs (P=0.0001), overall lower level of expression of CNMs as defined by combined H score (P=0.007), and lack of TTF1 expression (P=0.003) compared to other cases. Notably, all ASCL1/NeuroD1 double-negative cases did express at least one CNM and had classic SCLC morphology.

Conclusions: Expression of ASCL1 and NeuroD1 is not mutually exclusive in a significant number of SCLC (42%), yet in most cases there is clear dominance of one marker over the other. While we did not identify overt differences in tested parameters between subsets with dominant ASCL1 vs NeuroD1 expression, we did find that the double-negative subset is highly distinct, and represents pan-NE marker low SCLC subset. Validation of clinical significance of these novel subsets of SCLC is under investigation.

1877 Collagen V Expression Pattern as a Proposed Surrogate Marker for the Diagnosis of Malignant Mesothelioma

Marcelo Balancin¹, Leticia Clemente², Lucas Reis², Mario Marques-Piubelli³, Alexandre Ab´Saber⁴, Vitória Contini⁵, Ana Paula Pereira Velosa⁵, Walcy Teodoro⁵, Paola Da Souza⁶, Vera Luiza Capelozzi⁷

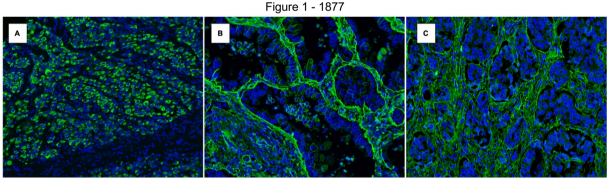
¹University of São Paulo, São Paulo, SP, Brazil, ²Hospital das Clínicas da Faculdade de Medicina da USP, São Paulo, SP, Brazil, ³The University of Texas MD Anderson Cancer Center, Houston, TX, ⁴University of São Paulo, Cotia, SP, Brazil, ⁵Faculty of Medicine - University of São Paulo, São Paulo, SP, Brazil, ⁶Faculty of Medicine - University of São Paulo, Maringá, PR, Brazil, ⁷Faculty of Medicine University of São Paulo, São Paulo, SP, Brazil

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Background: The diagnosis of Malignant Mesothelioma (MM) is a challenge in surgical pathology. The varied phenotype in both epithelioid and sarcomatoid types, immunohistochemical markers not always are reliable and ample differentials compose a challenging panorama. Type V Collagen (Col (V)) expression has been studied in breast and lung cancer as matrix remodeling elements associated with prognosis. While characterizing extracellular matrix in MM, we have found a not previously described pattern of Col (V) deposition. We have compared with other tumor types, revealing its potential diagnostic utility.

Design: A retrospective multi-center retrieve was performed for a ten-year period. Ninety cases of MM, 50 cases of non-small cell lung adenocarcinoma (NSCLA) and 50 cases of breast carcinoma (BC) were retrieved from the pathology archives. All slides were reviewed and reclassified by expert pathologists. A tissue microarray (TMA) was built from three sections of each case, resulting in 570 cores. Immunofluorescence (IF) was performed with a Col (V) in-house clone (described by Atayde et al, PLOS 2018). Images were captured through a microscope camera and the percentage of the positive area was accessed by threshold properties in the ImageJ software. Patterns of deposition were defined as fibrillar (a linear pattern in fine bundles of intercellular deposition), surrounding (a membrane pattern, surrounding and isolating each cell) and mixed. For the MM TMA, a double Col (V) D2-40 stain was also performed to enhance mesothelial cell specificity. Statistical analysis was performed using SPSS 25 by ANOVA followed by Spearman's test. A p-value of less than 0.05 was considered significant.

Results: The analysis was possible in 419 spots (74% of total):126 NSCLA (84%), 102 BC (68%), 229 MM (70%). The fibrillar type was expressed in 117 (93%) of NSCLA, 100 (98%) of BC and 9 (0.05%) of all spots of MM. The surrounding pattern was expressed in 182 (95%) of MM spots (p-value <0.05). The mixed pattern was expressed in 9 (7.7%) and 2 (1.2%) of NSCLA and BC spots, and when present, fibrillar dominated. Average Col (V) expression was: 3.58% (NSCLA), 12.1% (BC), and 10.65% (MM). Lung expressed significantly less than the other groups (p<0.05).



Collagen V Immunofluorescence Pattern: A. Surrounding (membrane pattern, surrounding and isolating each cell); B. Mixed; C. Fibrillar (linear pattern in fine bundles of intercellular deposition) (x400, IF, Green: Col (V), Blue DAPI).

Figure 2 - 1877

Collagen V and D2-40 Double Staining in MM (x400, IF, Green: Col (V), Red D2-40, Blue, DAPI)

Conclusions: IF expression of Col (V) in MM reveals a unique pattern, surrounding and isolating neoplastic cells, not previously described. Although IF is not employed in routine pathology practice, this study sets grounds for translating these finding into immunohistochemistry.

1878 Utility of Transbronchial Cryobiopsy in Diagnosis of Interstitial Lung Disease from a Histopathologic Perspective: A Single Institutional Experience

Janet Basinger¹, Rosana Eisenberg¹, Joyce Johnson², Mitra Mehrad¹

¹Vanderbilt University Medical Center, Nashville, TN, ²Vanderbilt Medical Center, Nashville, TN

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Background: Given the significant morbidity and mortality associated with surgical lung biopsy (SLB), transbronchial cryobiopsy (TBC), a minimally invasive technique, has recently gained increasing popularity for evaluation of interstitial lung disease (ILD). However, the diagnostic accuracy of TBC remains controversial. Our goal was to assess the diagnostic utility of TBC from a histologic standpoint at a large academic center.

Design: Pathology files were retrospectively searched for consecutive patients who underwent TBC for indeterminate ILD from 2012-2016. Slides were reviewed by pathologists with thoracic expertise (RE, JJ). The TBC diagnoses were compared to subsequent SLB or explant if available. Rates of definite histologic diagnosis (defined as diagnostic or strongly suggestive), pattern recognition with a broad differential diagnosis, and non-diagnostic biopsies were calculated.

Results: Ninety-seven patients were identified. The majority were male (60%) and Caucasian (92%). Age range was 25-84 years (median: 62). Definite pathologic diagnoses were rendered in 35/97 (36.1%) while 50/97 (51.5%) were given a broad differential diagnosis. The remaining 12/97 (12.4%) were interpreted as non-diagnostic with no significant pathologic change. Among the definite diagnoses, usual interstitial pneumonia (UIP) was the most common (16/35; 45.7%), followed by hypersensitivity pneumonitis (4/35; 11.4%) and non-specific interstitial pneumonia (4/35; 11.4%), constrictive bronchiolitis in the setting of burn-pit exposure in veterans (4/35; 11.4%), sarcoidosis (2/35; 5.8%), infection (2/35; 5.8%), Langerhans cell histiocytosis (1/35; 2.8%), pneumoconiosis (1/35; 2.8%), and smoking-related interstitial fibrosis (1/35; 2.8%). Three cases had subsequent SLB, one of which had concordant diagnoses of UIP on TBC and SLB. The 2nd case was non-diagnostic on TBC but had a bronchiolocentric pattern of disease on SLB and was given a wide differential diagnosis. The 3rd case was interpreted as a bronchiolocentric process on TBC, but had a UIP diagnosis with no histologically identifiable underlying etiology on SLB. One patient underwent transplantation and the diagnoses on TBC and explant were UIP.

Conclusions: In our series, UIP is the most common disease confidently identified on TBC. Our study suggests that TBC has histopathologic diagnostic yield in approximately 1/3 of patients and may potentially be helpful in arriving at a correct diagnosis in an additional ~50% of patients through multidisciplinary discussion.

1879 ZEB1 Expression in Peritoneal Malignant Mesothelioma

Amelia Baxter-Stoltzfus¹, Kartik Viswanathan², Michael Kluger³, Hanina Hibshoosh⁴, Alain Borczuk⁵

¹New York-Presbyterian Weill Cornell, New York, NY, ²New York-Presbyterian Hospital/Weill Cornell Medical Center, New York, NY, ³New York-Presbyterian/Columbia University Medical Center, New York, NY, ⁴Columbia University Medical Center, New York, NY, ⁵Weill Cornell Medicine, New York, NY

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Background: Malignant mesotheliomas are morphologically characterized by epithelioid, biphasic, or sarcomatoid histology. Sarcomatoid histology is associated with worse prognosis, suggesting that this epithelioid-to-mesenchmyal transition (EMT) contributes to more aggressive tumor behavior. Nuclear expression of ZEB1, a transcription factor which promotes EMT, has been reported in malignant mesothelioma. In addition, ZEB1 has been reported in the spindle and giant cell component of sarcomatoid carcinoma. Our goal was to evaluate ZEB1 immunohistochemistry in peritoneal mesotheliomas as a marker of sarcomatoid histology and potentially as a prognostic biomarker.

Design: One hundred seventy-three peritoneal mesotheliomas were investigated, of which 41 were biphasic and 132 were epithelioid. Immunohistochemical analysis using ZEB1 antibody (Sigma, HPA027524, 1:500) was performed on tissue microarrays and scored semiquantitatively for intensity (0, 1+, 2+). Percent of cells staining was also recorded. A ZEB1 H-score in the top tertile of scores (>10) was designated ZEB1-high for survival analysis. Statistics were performed as Chi square and Kaplan-Meier with log rank statistic for intensity and for H score.

Results: Twenty-seven of the biphasic tumors (66%) were positive for ZEB1 (staining intensity 1+ or 2+), as opposed to 41 of the epithelioid tumors (31%; p < 0.001). As expected, median survival for biphasic tumors (214 days) was significantly lower than for epithelioid tumors (1664 days, p < 0.001). Among epithelioid tumors, median survival of patients with ZEB1-positive tumors was significantly shorter

than that of patients with ZEB1-negative tumors (781 days and 1798 days, respectively; p = 0.008). Among epithelioid tumors, median survival of patients with ZEB1-high tumors was significantly shorter than that of patients with ZEB1-low tumors (476 days and 1798 days, respectively; p = 0.001).

Conclusions: As expected, biphasic morphology was associated with ZEB1 staining and poorer survival. ZEB1-positive epithelioid tumors had shorter median survival. Furthermore, strong and diffuse positive staining in epithelioid mesotheliomas was associated with worse prognosis. Use of this immunohistochemical stain may help to risk-stratify patients with epithelioid mesotheliomas, even in the absence of sarcomatoid morphology.

1880 Primary Mediastinal Germ Cell Tumors with High Prevalence of Somatic Malignant Transformation: An Experience from a Single Tertiary Care Cancer Center

Arshi Beg¹, Ayushi Sahay¹, Rajiv Kumar²

¹Tata Memorial Hospital, Mumbai, Maharashtra, India, ²Mumbai, India

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Background: Primary mediastinal germ tumors (PMGCT) constitutes, a mere 3-4 % of all germ cell tumors(GCT). Although, they account for approx. 16% of mediastinal tumors in adults and 19-25% in children as per western literature, there is hardly any large series on PMGCT reported from Indian subcontinent.

Design: We have retrospective analysed clinico-pathological features of 101 MGCT cases, diagnosed over a period 10years(2010-2019) from a single tertiary care oncology centre. Metastatic tumors to mediastinum were excluded.

Results: The study group (n=101) comprised predominantly of males(95)(M:F-15:1), with age range between 3 months to 57 years (mean: 25 years). The tumors were predominantly located in the anterior mediastinum(n=99) and cough/dyspnea were common symptoms. Non-seminomatous germ cell tumors (NSGCT) (composed of admixture of embryonal, yolk sac tumor(YST) &teratoma) was the most common histological subtype(n=26), followed by seminoma(n=25), mature teratoma(n=21), YST(n=13), immature teratoma(IT)(n=6) and GCT;NOS(n=6). Mixed seminomatous and NSGCT was seen only in 4 cases. Interestingly, 11 cases revealed secondary somatic malignancy (6 carcinomas and 5 sarcoma), including one case with adenoid cystic carcinoma. Somatic malignancies were seen predominantly in third decade. All 6 female patients had exclusively teratomas(5 mature and one IT). Serum AFP levels were elevated in all YST and LDH levels in 75 cases. Chemotherapy(CT) and radiotherapy(RT) details were available for 85 patients, out of which 64 received CT and 10 received RT. Fifty eight patients underwent surgical excision mostly post neo-adjuvant therapy. Histological examination revealed residual viable (RV) GCT in 25 cases; 9 cases showed teratoma only and 3 cases complete response. Follow-up was available of 67 patients ranging from 1week to 115 months (mean 27months). Six patients died/ had progressive disease and one of them as a consequence of transformation due to angiosarcoma.

Conclusions: Interestingly, in this series, PMGCT was seen predominantly in young adult males and somatic malignancies was noted in as high as in 10% of cases. Patient with somatic malignancy have aggressive clinical course, hence, extensive sampling and careful histopathological evaluation is recommended for definitive characterization.

1881 Correlation of VEGF-D Levels and Radiological Findings in Cystic Lung Diseases Clinically Suspicious of Lymphangioleiomyomatosis (LAM): An Institutional Study of Lung Wedge Biopsies

Swati Bhardwaj¹, Vijayalakshmi Ananthanarayanan², Swati Mehrotra², Stefan Pambuccian², Daniel Dilling³

¹Vardhman Mahavir Medical College and Safdarjung Hospital, Gurgaon, Haryana, India, ²Loyola University Medical Center, Maywood, IL, ³Loyola University Chicago Stritch School of Medicine, Maywood, IL

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Background: Lymphangioleiomyomatosis (LAM) is a rare cystic lung disease in young females that leads to progressive deterioration in lung function. Surgical biopsy is considered a gold standard for diagnosis. It is, thus, a grave diagnosis of immense clinical significance, as most patients with the diagnosis go on to require oxygen therapy and eventually, lung transplantation. Currently VEGF-D levels >800pg/ml is considered as diagnostic of LAM. In the current study, we evaluated cases clinically suspicious for LAM and correlated with surgical biopsy as well as VEGF-D levels.

Design: Pathology database was queried to identify cases with clinical suspicion of LAM over the past 10 years. Clinical and imaging findings of these cases were retrieved from electronic patient records. Histological diagnosis and immunostains were reviewed and compared with clinico-radiological findings and VEGF-D levels. We excluded cases in which the diagnosis of LAM was made on a transbronchial biopsy, or a lobectomy or pneumonectomy specimen.

Results: Using the above criteria, we identified 19 cases, of which 11 were diagnosed as LAM on histology. All cases were females, ranging from 25-62 years, with an average age of 44 years (Table 1). 1. The VEGF-D levels ranged from 85 to 794 pg/dl. No case was above the present diagnostic cut-off of 800 pg/ml. All 7 cases with VEGF-D levels >400pg/ml were diagnosed as LAM on histology. 7 out of 9 cases with VEGF-D levels <400 pg/ml did not have LAM. (PPV=100%, NPV=77.7%) 2. All positive cases had a typical radiological findings of multiple, diffuse, bilateral small cysts, usually <1cm. Radiological findings in cases not diagnosed as LAM were variable, including predilection for lower lobes, coexisting nodular lesion, varying size cysts, etc. 3. Cases where the radiology was characteristic but histologically not proven to be LAM, were noted to have VEGF-D levels of <300pg/ml

S. No.	Age	Sex	VEGF-D (pg/ml)	Radiological Findings	Histopathology
1	33	F	NA	Multiple <2cm cysts with subpleural nodular opacities	Pulmonary LCH
2	40	F	NA	Multiple, bilateral, randomly distributed <1cm cysts	LAM
3	50	F	364	Multiple, bilateral, randomly distributed <1cm cysts	LAM
4	40	F	538	Multiple, bilateral, randomly distributed <1cm cysts	LAM
5	39	F	265	Multiple, bilateral, thin walled cysts primarily at bases	Minimal change pattern of injury with small airway disease
6	64	F	351	Mulitple, bilateral, thin walled cysts <5mm to 3cm, along with bilateral nodular lesions	DLBCL arising in low grade lymphoma
7	58	F	438	Multiple, bilateral, randomly distributed <1cm cysts	LAM
8	48	F	299	Solitary, and cavitary nodules in both lungs	Placental transmogrification
9	25	F	794	Multiple, bilateral, randomly distributed cysts	LAM
10	51	F	290	Scattered cysts throughout both lungs	Emphysematous changes, non-necrotizing granulomas, meningothelial nodules
11	46	F	691	Multiple, bilateral, randomly distributed cysts	LAM
12	62	F	311	Multiple, bilateral, randomly distributed <1cm cysts	LAM
13	40	F	725	Multiple, bilateral, randomly distributed cysts	LAM
14	31	F	495	Multiple, bilateral, randomly distributed cysts	LAM
15	72	F	85	Multiple thin walled cysts in mid and lower lobes	Emphysematous changes, lymphocytic bronchitis
16	27	F	285	Numerous, bilateral randomly distributed cysts	Cystically dilated airspaces with peribronchial inflammation
17	56	F	351	Pneumothorax, scattered blebs and bullae throughout the lungs	Emphysematous changes, subpleural blebs
18	38	F	501	Multiple, bilateral, randomly distributed cysts	LAM
19	56	F	NA	Multiple, bilateral, randomly distributed cysts	LAM

Conclusions: It is important to correlate radiological findings and VEGF-D levels when considering the need for a surgical lung biopsy, as those cases with non-characteristic radiological findings and VEGF-D <400pg/ml were unlikely to be diagnosed as LAM on histology. Considering that all patients with biopsy proven LAM had a VEGF-D levels of greater than 400 pg/ml, we speculate that this level might be used as new cut-off for triaging patients with cystic lung disease for a biopsy, in an appropriate clinical setting.

1882 Intratumour Heterogeneity and Tumour Evolution in Pulmonary Neuroendocrine Cancer Lorelle Brownlee¹, Robert Bentham², Nicholas McGranahan³, Charles Swanton⁴, David Moore², Mariam Jamal-Hanjani², TRACERx Consortium⁵

¹University College London Hospital, Tonbridge, Kent, United Kingdom, ²University College London Cancer Institute, London, United Kingdom, ³The Francis Crick Institute, UCL Cancer Institute, London, United Kingdom, ⁴The Francis Crick Institute, London, United Kingdom, ⁵Multiple UK Centres, United Kingdom

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Background: Primary pulmonary neuroendocrine tumours comprise up to 25% of all lung cancer diagnoses. Pulmonary neuroendocrine tumours lie on a spectrum of malignant behaviour, ranging from low grade classical carcinoid tumours to high grade aggressive, small cell lung carcinoma. The evolutionary relationships between neuroendocrine tumours of different grades, and the evolutionary trajectories within tumour subtypes, remain poorly understood.

Design: The TRACERx study (TRAcking non-small cell lung Cancer Evolution through therapy (Rx)) is a UK multi-centre prospective cohort study which explores the genomic and phenotypic evolution of non-small cell lung cancer from early to late stage disease. In this substudy, tissue and blood samples were taken from 20 patients with pulmonary neuroendocrine tumours. Multi-region fresh tissue sampling was performed from primary tumour resections and subjected to deep whole-exome sequencing. Histological assessment of each tumour region was performed, including description of architecture, morphology and tumour infiltrating lymphocytes (TILs). Reference germline DNA for was obtained from whole blood. The data were analysed using the established TRACERx bioinformatics pipeline.

Results: Evolutionary phylogenetic trees illustrate genomic divergence for each tumour and demonstrate genetic intratumour heterogeneity. Differences were found between tumour grades across a number of parameters, including whole genome instability index, tumour mutational burden, clonal and sub clonal mutational burden, and loss of heterozygosity of human leucocyte antigen (HLA). Predicted driver events were also identified, both early (clonal) and late (subclonal) in tumour evolution. These included specific mutations in *ATM*, *TP53* and *MAP3K1*.

Conclusions: Intratumor heterogeneity is a feature of the different members of the primary pulmonary neuroendocrine tumour family, with high grade tumours demonstrating greater whole genome instability and subclonal divergence, compared with low grade tumours. Across the spectrum, both clonal and subclonal driver events are present indicating potentially different processes involved in early and late tumour evolution.

1883 Does Histologic Assessment Accurately Distinguish Separate Lung Adenocarcinomas from Intrapulmonary Metastasis? A Study of Paired Resected Lung Nodules in 28 Patients Using a Routine Next-Generation Sequencing Panel for Driver Mutations

Frido Karl Bruehl¹, Erika Doxtader², Carol Farver³, Daniel Farkas⁴, Sanjay Mukhopadhyay⁴

¹Cleveland Clinic Foundation, Cleveland, OH, ²Cleveland Clinic, Pepper Pike, OH, ³University of Michigan School of Medicine, Ann Arbor, MI, ⁴Cleveland Clinic, Cleveland, OH

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Background: Various molecular approaches have been reported for distinguishing synchronous primary adenocarcinomas from intrapulmonary metastasis. However, some of these approaches use testing that is not available at many institutions. Our aim was to investigate the accuracy of consensus histologic assessment (CHA) compared to a routine next generation sequencing (NGS) panel for driver mutations.

Design: We searched our database for cases of two or more resected lung adenocarcinomas in different lobes in order to avoid inadvertent sampling of the same nodule. Each pair of tumors was reviewed by three pathologists and categorized as related or unrelated based on CHA alone. Most tumors underwent routine NGS testing for common driver mutations. Three cases only underwent PCR for aberrations in *EGFR*, and 15 cases underwent limited NGS testing. Tumors with non-identical drivers were classified as unrelated, while those with identical low-frequency drivers were classified as related. Pairs with identical high-frequency *KRAS* mutations (34G>T or 35G>T) and with unknown driver gene status in a corresponding mutated tumor were considered indeterminate. CHA and the original pathology report were analyzed using molecular analysis as the gold standard.

Results: Of 28 pairs of lung adenocarcinomas (Table 1), 14 were classified as related by CHA, and 14 were classified as unrelated. Of the 14 cases classified as related by CHA, five were classified as related by molecular analysis, four as unrelated and five as indeterminate. Of 14 cases classified as unrelated by CHA, 10 were classified as unrelated by molecular analysis, none as related, and four as indeterminate. Of the 12 cases staged as related in the original pathology report, five were classified as related by molecular analysis, two were unrelated, and five were indeterminate. Of 16 cases classified as unrelated in the original pathology report, 12 were classified as unrelated by molecular analysis, none as related, and 4 were indeterminate. The CHA and the staging in the original pathology report were inaccurate in 4/28 and 2/28 cases, respectively. There was no discordance between cases deemed unrelated by CHA or original staging and molecular analysis.

Table 1: Characteristics of cases with two lung adenocarcinomas arising in separate lobes.

Case	Consensus Histologic Assessment	Original Pathologic Staging	Molecular Assessment	Mutation Tumor 1	Mutation Tumor 2
1	Related	Related	Unrelated	Negative	KRAS c.35G>T
2	Related	Related	Indeterminate	KRAS c.34G>T 66 AF	KRAS c.34G>T
3	Unrelated	Unrelated	Unrelated	EGFR c.2240T>C	KRAS c.34G>T
4	Unrelated	Unrelated	Indeterminate	KRAS c.34G>T	KRAS c.34G>T
5	Related	Related	Unrelated	BRAF c.1406G>T	KRAS c.35G>T
6	Unrelated	Unrelated	Unrelated	Negative	KRAS c.35G>T
7	Related	Related	Related	MET c.3029C>T	MET c.3029C>T
8	Related	Related	Related	EGFR c.2573T>G	EGFR c.2573T>G
9	Unrelated	Unrelated	Unrelated	KRAS c.34G>T	Negative
10	Unrelated	Related	Indeterminate	KRAS c.35G>T	KRAS c.35G>T
11	Related	Related	Related	ERBB2 c.2264T>C	ERBB2 c.2264T>C
12	Related	Related	Related	EGFR c.2240_2257del	EGFR c.2240_2257del
13	Related	Related	Related	KRAS c.57G>T	KRAS c.57G>T
14	Related	Related	Indeterminate	Negative	Negative
15	Related	Related	Indeterminate	Negative	Negative
16	Related	Related	Indeterminate	Negative (BRAF not tested)	BRAF N581S
17	Related	Unrelated	Unrelated	KRAS c.183A>T	KRAS c.34G>T
18	Unrelated	Unrelated	Unrelated	EGFR exon 19 p.Glu746_Ala750del	EGFR - c.2303_2304insTGTGGCCAG
19	Unrelated	Unrelated	Unrelated	KRAS c.34G>T	Negative
20	Related	Related	Indeterminate	ERBB2 p.A775_G776insYVMA	Negative (ERBB2 not tested)
21	Unrelated	Unrelated	Unrelated	KRAS c.35G>T	Negative
22	Unrelated	Unrelated	Unrelated	KRAS c183A>C	MET c.3028+2T>C
23	Unrelated	Unrelated	Unrelated	KRAS c.35G>T	KRAS c.34G>T
24	Related	Unrelated	Unrelated	KRAS c.38G>A	KRAS c.34G>A
25	Unrelated	Unrelated	Unrelated	KRAS c.37G>T	KRAS c.34G>T
26	Unrelated	Unrelated	Unrelated	Negative	KRAS c.34G>T
27	Unrelated	Unrelated	Unrelated	KRAS c.35G>T	KRAS c.34G>T
28	Unrelated	Unrelated	Unrelated	Negative	KRAS c.34G>T
	1				

Conclusions: NGS testing for routine driver mutations can aid in the differentiation between synchronous primaries and intra-pulmonary metastasis. A subset of cases judged as related by morphology is shown to be inaccurate by molecular analysis, causing significant downstaging.

1884 When is Inflammation just Inflammation? Pulmonary Nodules with an Initial Core Needle Biopsy Showing Non-Specific Inflammation: Radiological-Pathological Features and Correlation with Final Diagnosis

Aurelia Busca¹, Carolina Souza², Ashish Gupta², Susan John², Aleksandra Paliga³, Marcio Gomes⁴
¹EORLA- University of Ottawa, Ottawa, ON, ²The Ottawa Hospital - University of Ottawa, Ottawa, ON, ³The Ottawa Hospital, General Campus Site, Ottawa, ON, ⁴The Ottawa Hospital/University of Ottawa, ON

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Background: Management of patients with lung nodules whose initial core needle biopsy (CNB) shows non-specific inflammation is challenging and often result in rebiopsy, resection or prolonged follow-up (FU). The findings may represent the periphery of undersampled malignancy, low grade lymphoproliferative disorder or true inflammatory lesion (e.g. nodular lymphoid hyperplasia). The study purpose was to identify radiological and pathological features associated with benign outcome.

Design: We retrospectively identified patients with focal pulmonary lesions on chest CT and CNB diagnosis of non-specific inflammation. Cases with features suggestive of a specific benign diagnosis (e.g. necrotizing granuloma) were excluded. The reference standard for final diagnosis was resection, radiological resolution or at least 6-month FU. Chest CTs/PET-CTs were reviewed by 2 chest radiologists and biopsy slides by senior resident, thoracic pathologist and hematopathologist. Descriptive radiologic and pathologic data were collected, including lesion measurements on CT and biopsy slides to estimate representativeness of samples.

Results: 45 patients were identified. On FU (mean 31 months) 6 cases (13.3%) had a final diagnosis of malignancy (2 adenocarcinomas, 1 squamous cell carcinoma, 1 MALT lymphoma, 1 DLBCL, 1 NSCLC). Of these, 3 were diagnosed on repeat CNB, 1 on wedge resection after repeat benign CNB and 2 on FU FNA. Other 11 cases (24.4%) with repeat biopsy and 2 cases (4.4%) with FU resection were benign (1 wedge resection and 1 lobectomy, both nodular lymphoid hyperplasia).

All nodules were solid (86.7%) or part-solid on CT. Most benign and all malignant cases had spiculated (24.4%) or irregular margins (55.1%). Initial biopsies in malignant cases had either inflammatory (5/6) or lymphoma-like pattern (1/6). All cases with scar-like pattern (7/45, 15.5%) were benign. Shorter solid lesional component on CNB correlated with final benign diagnosis (p=0.01; solid component of all malignant cases measuring ≤10 mm). Cases with final malignant diagnosis also had CNB with shorter lesional tissue and higher SUVmax, which approached statistical significance despite low number.

Table 1. Clinical, radiological and pathological characteristics of study population according to final diagnosis.

	Benign (N=39)	Malignant (N=6)	p value
Gender	Male- 20	Male- 3 (50%)	1**
	(51.3%)	Female-3 (50%)	
	Female-19		
	(48.7%)		
Age (mean ± SD)	63.74 ±11.75	72 ± 6.63	0.08*
Number of lesions on CT (mean ± SD)	2.59 ±1.73	2 ± 1.55	0.53*
Size index lesion on CT (mm± SD)	29.31±16.17	32.33 ± 18.41	0.86*
SUVmax (mean ± SD)	4.66 ± 2.87	10.42 ± 6.71	0.06*
·	(N=29)	(N=5)	
Consistency:	34 (87.2%)	5 (83.3%)	1**
- Solid	5 (12.8%)	1 (16.7%)	
- Part solid			
Margins:	4 (10.3%)	0	0.88**
- Smooth	3 (7.7%)	0	
- Lobulated	10 (25.6%)	1 (16.7%)	
- Spiculated	22 (56.4%)	5 (83.3%)	
- Irregular			
Radiologic impression:	10 (25.6%)	0	0.38**
- Probably benign	13 (33.3%)	2 (33.3%)	
- Indeterminate	16 (41.0%)	4 (66.7%)	
- Probably malignant			
Total lesional length on CNB (mm± SD)	15.38 ±8.74	8 ± 5.54	0.05*
Solid component length on CNB (mm± SD)	14.87 ± 8.88	6.08 ± 3.44	0.01*
Histologic pattern:	7 (17.9%)	0	0.62**
- Scar-like	28 (71.8%)	5 (83.3%)	
- Inflammatory	4 (10.3%)	1 (16.7%)	
- Lymphoma-like			

^{*} Wilcoxon non-parametric test; ** Fisher's exact test

Conclusions: Combined radiological-pathological approach may improve the diagnostic accuracy of patients with focal lung lesions and non-specific inflammation on initial CNB. Good representation of the solid component of the lesions is associated with final benign diagnosis and should be considered when sampling and making management decisions.

1885 Histopathologic and Radiologic Features of Interstitial Lung Disease in Patients with Short Telomeres

Matthew Cecchini¹, Tara Tarmey¹, Allison Ferreira¹, Thomas Hartman¹, Eunhee Yi¹, Anja Roden¹ *Mayo Clinic, Rochester, MN

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Background: Many forms of interstitial lung disease (ILD) are diseases of aging secondary to repetitive injury or exposures that causes an abnormal cellular senescence. A subset of ILD occurs in patients with short telomeres who can also have premature greying of hair, liver fibrosis and bone marrow failure. Cells with critically shortened telomeres can enter into a cellular senescence mediated in part by upregulation of the CDK inhibitor p16. Despite our increasing understanding of the underlying pathogenetic mechanisms, histopathologic and radiologic features of ILDs in patients with confirmed short telomeres have not been well characterized.

Design: Cases diagnosed as positive for short telomeres (2009-2019) were identified by testing of peripheral blood granulocytes and/or lymphocytes with 10th percentile of telomeres or less. Lung explants or wedge biopsies from the patients with short telomeres were reviewed independently by 2 pathologists using defined morphologic parameters. Discordant cases were reviewed by a third pathologist. High resolution CT scans were reviewed by 2 radiologists independently. Immunohistochemistry for p16 (clone E6H4) was performed on a section of each case showing the most areas of active fibrosis with abundant interfaces between fibrotic and architecturally preserved lung parenchyma. All foci showing p16 positivity in both epithelial cells and associated fibroblasts were counted. Cases without short telomeres were used as controls.

Results: The morphologic and radiologic features of cases with short telomeres (n=15) as compared to those without short telomeres (n=5) are outlined in table 1. Among 8 cases with short telomeres that were tested for mutation, 5 (62.5%) cases had a germline mutation in a gene related to telomere maintenance (NAF1, TERT, RTEL1, TERC). The cases in the short telomere group often demonstrated features atypical for usual interstitial pneumonia (UIP) on both histopathologic and radiologic examination. Nine of 15 (60%) cases with short telomeres were classified as non-UIP, while 3 of 5 (60%) without short telomeres were diagnosed as UIP. The average number of p16-positive foci was higher in the cases with short telomeres though not statistically significant (p=0.4) (Table 1).

Telomere Pathologic Diagnosis p16 +ve Radiologic Fleischner Classification for UIP NSIP status UIP CHP Unclassified PPFE foci (average) typical probable indeterminate nonconsistent Short 5 6 2 1 8.4 5 5 5 0 3 0 0 0 4.4 0 Non-short 3

Table 1. Histopathologic and radiologic features of ILD in patients with short and non-short telomeres

Conclusions: The majority of ILDs in patients with short telomeres showed morphologic and radiologic features atypical for UIP and were often diagnosed as CHP or unclassifiable ILD. Both groups with and without short telomeres demonstrated p16-positive foci.

1886 Histologic Challenges in Distinguishing Separate Primary Lung Carcinomas from Intrapulmonary Metastases Using Broad Next-Generation Sequencing as a Gold Standard for Determining Tumor Clonal Relationships

Jason Chang¹, Deepu Alex², Matthew Bott¹, Kay See Tan¹, Andrew Golden¹, Jennifer Sauter¹, Darren Buonocore¹, Chad Vanderbilt³, Sounak Gupta⁴, Patrice Desmeules⁵, Francis Bodd¹, David Jones¹, Maria Arcila¹, William Travis¹, Marc Ladanyi¹, Natasha Rekhtman¹

¹Memorial Sloan Kettering Cancer Center, New York, NY, ²BC Cancer, Vancouver, BC, ³Memorial Sloan Kettering Cancer Center, Denver, CO, ⁴Mayo Clinic, Rochester, MN, ⁵IUCPQ - Quebec Heart and Lung Institute, Quebec, QC

Disclosures: Jason Chang: None; Deepu Alex: None; Matthew Bott: None; Kay See Tan: None; Andrew Golden: None; Jennifer Sauter: None; Darren Buonocore: None; Chad Vanderbilt: *Consultant*, Docdoc Ltd. (Singapore); *Consultant*, Paige Al; *Consultant*, OncoKB; Sounak Gupta: None; Patrice Desmeules: *Grant or Research Support*, Pfizer Canada; *Grant or Research Support*, Novartis; Francis Bodd: None; David Jones: *Advisory Board Member*, Diffusion Pharmaceuticals; *Consultant*, Merck; *Consultant*, AstraZeneca; Maria Arcila: *Speaker*, Biocartis; *Speaker*, Invivoscribe; William Travis: None; Marc Ladanyi: None; Natasha Rekhtman: None

Background: We have recently reported on the utility of comprehensive next-generation sequencing (NGS) for distinguishing separate primary lung carcinomas (SPLC) from intrapulmonary metastases (IPM) in clinical practice. Here, we report on detailed review of histologic challenges in determining the relationships between multifocal adenocarcinomas using NGS as a gold standard.

Design: A total of 70 surgically-resected adenocarcinoma pairs underwent molecular profiling using 341-468 gene hybridization-capture based NGS assay. Comparative genomic profiles were used to stratify tumor pairs into clonally-unrelated (SPLC) and clonally-related

(IPM). Relationship of tumor pairs predicted by prospective histopathologic assessment was compared with NGS-based classification. Histopathologic features contributing to challenges in distinguishing tumor relationships were assessed retrospectively.

Results: Of 70 adenocarcinoma pairs, NGS classified 47 as SPLCs and 23 as IPMs. Prospective histologic prediction was discordant with NGS in 16 cases (23%), with significantly higher histologic misclassification rate for NGS-confirmed IPMs than SPLCs (43% vs 13%, P=0.0003). The discordance rate was significantly higher when histologic prediction was regarded as potentially equivocal by a pathologist and confirmation by NGS was suggested (p=0.02). Retrospective review highlighted several specific factors contributing to misinterpretation of NGS-defined IPMs as morphologically unrelated tumors, including 1) morphologic progression leading to higher proportion of solid and micropapillary patterns in secondary tumors (n=6), and 2) presence of significant amounts of non-predominant lepidic pattern in both primary and secondary tumors (n=4). NGS-defined SPLCs that were initially misinterpreted as morphologically related tumors (n=6) showed closely overlapping architectural or cytologic features.

Conclusions: Comprehensive histopathologic assessment is adequate for distinguishing SPLCs from IPMs in most cases, but has notable limitations in the recognition of a subset of cases, particularly IPMs. Our results support the adoption of molecular testing to supplement histologic assessment for robust discrimination of clonal relationships of multifocal adenocarcinomas in clinical practice, and we propose an algorithm incorporating specific histopathologic scenarios where molecular profiling may be most helpful.

1887 Germline Mutated Malignant Mesotheliomas: Are These Morphologically Different from Sporadic Malignant Mesotheliomas?

Heather Chen¹, Meghna Gadiraju², Jane Churpek³, Hedy Kindler¹, Yin Hung⁴, Mari Mino-Kenudson⁴, Richard Attanoos⁵, Jefree Schulte¹, Jeffrey Mueller⁶, Thomas Krausz⁷, Aliya Husain¹

¹The University of Chicago, Chicago, IL, ²The University of Chicago Medicine, Chicago, IL, ³The University of Wisconsin School of Medicine and Public Health, Madison, WI, ⁴Massachusetts General Hospital, Boston, MA, ⁵Cardiff University, Cardiff, Wales, United Kingdom, ⁶University of Chicago Medical Center, Chicago, IL, ⁷University of Chicago Hospital, Chicago, IL

Disclosures: Heather Chen: None; Meghna Gadiraju: None; Jane Churpek: None; Hedy Kindler: *Consultant*, Inventiva; *Consultant*, AstraZeneca; *Advisory Board Member*, Boehringer Ingelheim; *Consultant*, Kyowa; *Grant or Research Support*, Aduro, AstraZeneca, Bayer, BMS, Deciphera, GSK, Lilly, Merck, Polaris, Verastem, Blueprint; Yin Hung: None; Mari Mino-Kenudson: None; Richard Attanoos: None; Jeffree Schulte: None; Jeffrey Mueller: None; Thomas Krausz: None; Aliya Husain: None

Background: Malignant mesothelioma (MM) is rare but lethal; some affected individuals develop MM in the setting of predisposing inherited mutations. Our group found a 12% prevalence of pathogenic or likely pathogenic variants in prospectively tested MM patients. These patients showed longer survival after platinum therapy, suggesting distinct biology. Germline mutated *BAP1* MM, the most prevalent mutated gene in MM, appears to be more indolent than wild-type. This study aims to compare the morphology of epithelioid (E-) MMs with germline mutations to those without.

Design: Eligible patients with pathologically confirmed MM underwent panel-based hereditary cancer susceptibility germline genetic testing. There were two control groups with no germline mutation matched by MM histologic subtype, age at diagnosis, and sex; 22 patients had retained BAP1 and 9 had loss of BAP1 on IHC. In E-MM, nuclear grade (3-tiers comprised of mitotic count and nuclear grade), presence of necrosis (yes/no), and patterns were compared using GraphPad Prism via Fisher's Exact Tests.

Results: Of 265 MMs tested, 25 had germline mutations; an additional 5 cases were from other institutions (n=30). Pathology was unavailable in 7. Of the 23 left (Table 1), there were 22 E-MM and 1 biphasic MM with 10 gene variants. *BAP1* was most common (n=9). Among all MMs with a nuclear grade of 1 (n=79), 15 (19%) had a germline mutation. The E-MMs had a lower nuclear grade than those without (p=0.01). The mitotic count was a significant contributor to lower grade (p=0.03). By site, there was a trend in lower grade for peritoneal MMs (n=13) between germline mutated and non-mutated cases (p=0.06); that difference was not observed for pleural MMs (n=9, p=0.24). When focusing on *BAP1*, lower nuclear grade was also present when comparing BAP1 mutated MMs to sporadic BAP1 IHC loss MMs (p=0.05) and BAP1 IHC retained MMs (p=0.05). There was a trend of lower mitotic count in the *BAP1* germline mutated cases compared to the BAP1 IHC loss and the retained cases (p=0.16, p=0.13). There was a trend of less necrosis in *BAP1* germline mutated MMs vs MMs with retained BAP1 (p=0.11). Solid and trabecular patterns were most frequently observed in both study cases and controls.

Table 1. Mutations of Patients Included in Study

Mutation	Number of Patients
BAP1	9
ATM	1
BRCA1	1
BRCA2	4
CDKN2A	1
CHEK2	2

MMR (MLH1, MLH2, MSH6, PMS2)	2
SDHA	1
VHL	1
WT1	1

Conclusions: Germline mutated MMs (BAP1 and others) have a lower nuclear grade than those without and are mostly E-MM, thus correlating with the better prognosis and slower clinical course reported in the literature.

1888 High Expression of PKM2 Synergizes with PD-L1 to Predict Worse Survival in the Patients with Lung Adenocarcinoma

Honglei Chen¹, Sufang Tian², Long Long³, Mengxi Chen¹, Yuhan Yang¹

¹Wuhan University, Wuhan, Hubei, China, ²Wuhan University Zhongnan Hospital, Los Angeles, CA, ³Wuhan University Zhongnan Hospital, Wuhan, China

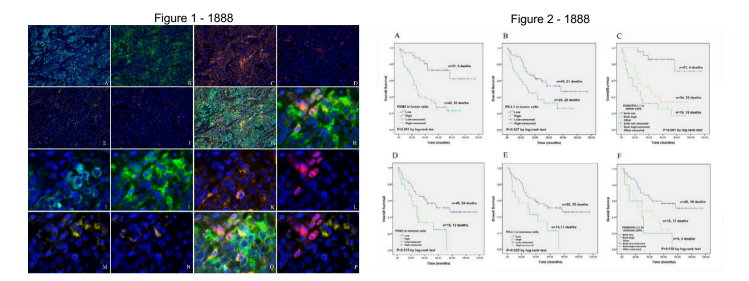
Disclosures: Honglei Chen: None; Sufang Tian: None; Long Long: None; Mengxi Chen: None; Yuhan Yang: None

Background: Lung cancer has been the most prevalent and the most deadly malignant cancer globally, which is still incurable disease in essence. Immunotherapy targeting PD-1/PD-L1 represents a breakthrough in the treatment of lung cancer. Nevertheless, the response rate of anti-PD-1/PD-L1 immunotherapy remains unsatisfactory, due to tumor resistance and complexity of immune microenvironment. To enable more patients to benefit from immunotherapy, a thorough understanding of the regulatory mechanisms of PD-L1 expression will be pivotal for novel combinational immunotherapies. Pyruvate kinase M2 (PKM2) is a critical player of glycolysis, conducing to tumor progression and immune response. However, the correlation and clinical significance of PKM2 and PD-L1 expression in human lung adenocarcinoma (LUAD) remain not entirely explored.

Design: Expression of PKM2 and PD-L1 were detected by immunohistochemistry in 74 cases of LUAD and the corresponding noncancerous tissues. Simultaneously, multiplex immunofluorescence was used to detect PKM2, PD-L1, CD3, CD68, CD163 and pan-CK by using the Opal 7-Color IHC Kit, combined with multi-spectral imaging system and InForm software. We measured expression patterns and co-localization of these targets, evaluating their correlation with clinicopathological features and overall survival. Validation of findings was conducted using mRNA expression data from The Cancer Genome Atlas (TCGA) of 515 lung adenocarcinoma cases.

Results: Co-expression of PKM2 and CK, CD3, CD68 were found. PD-L1 protein expression was detected in the tumor cells and immune cells including T cells and tumor associated macrophages (CD68+CD163+). High expression of PKM2 in tumor cells (TCs) was significantly related to lymph node metastasis and TNM stage. Moreover, PKM2 expression in TCs was positively correlated with PD-L1 expression in TCs. High expression of PKM2, PD-L1, as well as both PKM2 and PD-L1 in TCs and immune cells predicted high mortality rate and worse survival, respectively. Additionally, multivariate Cox regression models indicated that high expression of PKM2 in TCs was an independent prognostic factor. Based on TCGA genomic data, high PKM2 mRNA expression was significantly associated with poorer survival.

	PKM2 in	tumor cells		PD-L1 in	tumor cells		PKM2 and P	D-L1	
Parameters	High	Low	P	High	Low	P	Both high	Others	P
Gender			0.253			0.197			0.974
Male	22	20		15	27		8	34	
Female	21	11		7	25		6	26	
Age			0.235			0.567			0.208
<=64	31	26		16	41		9	48	
>64	12	5		6	11		5	12	
Depth of invasion (T)		0.249			0.218			0.027
T1,T2	37	30		18	49		10	57	
T3,T4	6	1		4	3		4	3	
Lymph node metasta	asis (N)		0.035			0.297			0.075
N0	23	24		12	35		6	41	
N1,N2	20	7		10	17		8	19	
TNM stage			0.017			0.151			0.029
	20	23		10	33		4	39	
III	23	8		12	19		10	21	
Smoking history			0.628			0.686			0.603
Yes	17	14		10	21		5	26	
No	26	17		12	31		9	34	
Histological type		0.383			0.096			0.339	
Invasive	40	26		22	44		14	52	
Variant	3	5		0	8		0	8	



Conclusions: High expression of PKM2 synergizes with PD-L1 to predict worse survival in patients with LUAD. Combinational therapy targeting PKM2 and PD-L1 may be a hotspot for future immunotherapy of LUAD.

1889 PD1 and PD-L1 Distribution Patterns in Non-Small Cell Lung Cancer: Validation of Two Novel Immunohistochemistry Markers

Richard Chiu¹, Khaled Algashaamy², Hadi Yaziji³

¹Larkin Community Hospital, Hialeah, FL, ²University of Miami/Jackson Memorial Hospital, Miami, FL, ³Vitro Molecular Laboratories, LLC, Miami, FL

Disclosures: Khaled Algashaamy: None; Hadi Yaziji: None

Background: Immunotherapeutic agents have revolutionized the standard of care in patients with advanced non-small cell lung cancer (NSCLC) and have emerged as novel treatment strategies as they have demonstrated promising treatment advantage over traditional chemotherapeutic agents. Immunohistochemical (IHC) detection of PD-L1 expression in NSCLC cases is considered to be a clinical decision-making tool to support the use of checkpoint inhibitors in NSCLC patients. FDA-approved assays and other antibodies are commercially available. This validation study is designed to compare a commercially available assay (28-8) with another clone (CAL10), and to also determine the distribution patterns of expression of PD-L1 and PD1 in the tumor and tumor-associated environment.

Design: 134 consecutively diagnosed NSCLCs with available whole tissue sections were evaluated for the study, which includes testing with the FDA-approved PD-L1 assay 28-8 (clone 28-8; Agilent/Dako), PD-L1 clone CAL10 (Biocare Medical) and PD1 clone (NAT105). Whole sections were chosen over tissue microarrays to adequately assess intra-tumor heterogeneity. Deparaffinized tissue sections were pretreated using either online (for the two non-FDA approved tests) or offline (for 28-8), followed by antibody incubation, polymer detection and DAB visualization. For the 2 PD-L1 assays, the 2019-revised FDA TPS scoring criterion of ≥1% was employed. For the PD1 assay, immunoe cell score was employed.

Results: 50% of tumors (n = 67) showed TPS positive expression of ≥1% using 28-8 as a gold standard assay for PD-L1. Agreement between 28-8 and CAL10 was 100%, with a consistently higher TPS score observed with CAL10 compared to 28-8. Immune cell score of NAT105 anti-PD1 antibody was positive in 8% of tumors (n = 12). When comparing the distribution of PD-L1 and PD1 expression on the same tumor, Out of the 12 PD1 positive cases, 4 showed positive PD-L1 expression, and 8 showed a mutually exclusive expression with only PD1 positive expression.

Conclusions: The validation study shows that PD1 and PD-L1 expression on the same tumors in NSCLC is uncommon, a phenomenon that can be exploited to expand targeted therapy treatment strategies in this setting, and could possibly be expanded into other oncology settings.

It also demonstrates the slightly higher sensitivity of detection achieved by clone CAL10 compared to 28-8 using the tumor proportion score, which warrants an expanded evaluation.

1890 Conida in Post-Transplant Lung Biopsies and Associated Infections

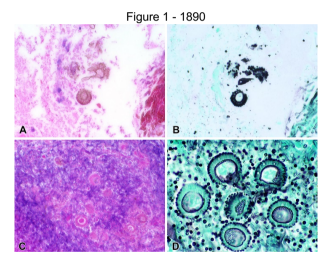
Haley Davis¹, David Howell¹, Elizabeth Pavlisko¹, John Carney¹, Matthew Pipeling², Carolyn Glass¹ Duke University Medical Center, Durham, NC, ²Duke University Hospital, Durham, NC

Disclosures: Haley Davis: None; David Howell: None; Elizabeth Pavlisko: None; John Carney: None; Matthew Pipeling: None; Carolyn Glass: None

Background: Infectious complications significantly contribute to morbidity and mortality in lung transplant patients with a one-year cumulative incidence of 8.6%. Opportunistic fungal infections with *Aspergillus* and *Candida* species are most frequent. Colonization with conidia, asexual non-motile spores of fungi also known as "fruiting heads," has been predominantly associated with *Aspergillus* species. Recent studies have implicated small conidia species with higher rates of bronchiolitis obliterans syndrome (BOS). To date, only 2 reports have described conidia in the setting of lung transplant patients. We review here our case series from a high-volume lung transplant center.

Design: Post-lung transplant biopsies with fungal organisms exhibiting conidia were collected over a 15 year period at a single institution. A GMS or PAS special stain was performed on each case. Clinical data to include age, indication for lung transplant, onset of fungal infection, corresponding culture data, treatment outcome and the presence of BOS was reviewed.

Results: In the last 15 years 1,532 lung transplants were performed. Fungal infection with conidia was present in 10 lung transplant cases (mean age 52; range 23-77 years). The mean time to infection from date of transplant was 205 days (range 48 to 757 days). Indications for transplant included cystic fibrosis, chronic obstructive pulmonary disease, and idiopathic pulmonary fibrosis. Nine cases had positive fungal cultures with the following species: Aspergillus fumigatus (5, figure 1C and 1D), Aspergillus ochraceus (1), Aspergillus niger (1), Aspergillus flavus (1), and Scopulariopsis (1, figure 1A and 1B). Fungal infection resolved with treatment in 9 of the cases. However, 4 patients died after resolution of infection from rejection, bacterial pneumonia, and BOS. One patient died with ongoing fungal pneumonia, and rejection. BOS was present in 6 patients, 4 of whom grew small conida species, one of whom grew Scopulariopsis, and the last wasn't cultured.



Conclusions: The presence of conidia with fungal organisms has been predominantly associated with *Aspergillus* species in lung transplant patients. We report for the first time, an association with invasive *Scopulariopsis*, a less common but highly resistant species with extremely high mortality. In our cohort, the majority of conidia seen were from small conidia species, which may confer a higher risk for BOS. All patients with BOS and a speciated culture had the presence of a small conidia species.

1891 EZH2 Expression in SMARCA4-mutant Primary Lung Malignancies

Richard Davis¹, Kristen Deak¹, John Carney², Carolyn Glass²

¹Duke University, Durham, NC, ²Duke University Medical Center, Durham, NC

Disclosures: Richard Davis: None; Kristen Deak: None; John Carney: None; Carolyn Glass: None

Background: SMARCA4-mutant lung carcinomas and thoracic sarcomas have recently been identified as distinct entities involving the SWI/SNF chromatin modifying complex. To date, over 20% of human malignancies have been associated with defects in the SWI/SNF complex. SMARCA4 is the critical ATPase subunit that drives the complex, leading to modification of chromatin and transcriptional regulation. SMARCA4 mutations in particular, have shown to have prognostic and potential therapeutic implications in lung and mediastinal tumors. Recent studies show inhibition of Enhancer of Zeste Homolog 2 (EZH2), an H3K27 histone methyltransferase that regulates the

SWI/SNF complex, sensitizes SMARCA4-mutant lung tumors to chemotherapy (Fillmore, Nature 2015). However, the expression of EZH2 in SMARCA4-mutant lung tumors is unknown.

Design: We reviewed a cohort of 282 thoracic specimens sampled between 2014-19 that previously underwent next generation sequencing (NGS) of over 320 cancer-associated genes as part of routine clinical care. SMARCA4 frameshift, nonsense and splice site mutations were defined as loss of function (LOF). EZH2 immunohistochemistry (IHC) was performed and expression was quantitatively assessed on tumor cells based on the extent of staining as follows: loss (0), weak (1+), moderate (2+) and strong (3+). A cohort of 5 lung carcinomas without SMARCA4-mutations were also stained as controls.

Results: We identified 13 patients (mean age 62, range 42 to 81 years, 64% male) with SMARCA4-mutant lung cancers (incidence 4.6%). LOF mutations were present in 8 (62%) cases and included 3 nonsense, 3 frame shift and 2 splice site variants. Two additional cases had SMARCA4 duplications. All but two cases (92%) were lung adenocarcinomas, with 1 poorly differentiated non-small cell carcinoma and 1 sarcomatoid carcinoma. All cases with LOF mutations exhibited a solid growth pattern and 40% presented as distant metastatic disease. Tumor tissue was available for 8 cases; EZH2 was expressed in 100% of cases with confirmed SMARCA4-mutations (mean 2.5 intensity). Of 5 SMARCA4 non-mutated tumors, only 1 tumor demonstrated EZH2 expression (mean 0.2 intensity, p<0.01)

Conclusions: EZH2 is overexpressed in SMARCA4-mutant lung cancers and may serve as a potential prognostic and predictive factor for those treated with epigenetic inhibitor therapy. This is the first study to assess EZH2 expression in this subset of tumors that are known to be resistant to conventional chemotherapy.

1892 RNA Targeted Analysis for the Detection of Druggable Fusions in Lung Adenocarcinoma: One-Year Experience

Thomas Denize¹, Audrey Mansuet-Lupo², Laure Gibault³, Diane Damotte⁴, Karen Leroy⁵, Christos Chouaid⁶, Elizabeth Fabre⁷, Francoise Le Pimpec-Barthes⁷, Marie Wislez⁵, Hélène Blons⁸, Simon Garinet⁷

¹APHP.5 Hôpital Européen Georges Pompidou, boston, MA, ²Hôpital Cochin, Paris, France, ³European Hospital Georges Pompidou, Paris, France, ⁴University Hospital Cochin, AP-HP, Paris, France, ⁵APHP.5 Hôpital Cochin, Paris, Ile de France, France, ⁶Centre Hospitalier Intercommunal de Créteil, Créteil, Ile de France, France, ⁷APHP.5 Hôpital Européen Georges Pompidou, Paris, Ile de France, France, ⁸Hôpital Européen Georges Pompidou, Paris, France

Disclosures: Thomas Denize: None; Audrey Mansuet-Lupo: None; Laure Gibault: None; Diane Damotte: None; Karen Leroy: None; Christos Chouaid: None; Elizabeth Fabre: None; Francoise Le Pimpec-Barthes: None; Marie Wislez: None; Hélène Blons: *Consultant*, AstraZenece, MSD; Simon Garinet: None

Background: Gene fusions are major targetable oncogenic events in non-small cell lung cancer (NSCLC). Main fusions involve *ALK* (4-5%), *RET* (1-2%), *ROS1* (1-2%) and *NTRK1* (<1%). Fluorescent *in situ* hybridization (FISH) is the gold standard for fusion detection but remains challenging. Optimized targeted next generation sequencing methods allow multiplex fusion detection from formalin fixed paraffin embedded (FFPE) RNA samples. Here, we present a one-year experience of routine fusion analysis in *EGFR/KRAS* wild-type metastatic NSCLC using ArcherDX® technology.

Design: 63 NSCLC samples were analyzed with ArcherDX technology. Inclusion criteria were metastatic spread, wild-type status for *KRAS/EGFR/NRAS/BRAF/ERBB2* determined by Next Generation Sequencing (NGS) and negative ALK staining. RNA was extracted using the Maxwell® 16 FFPE LEV RNA Purification Kit (Promega, Charbonnières-les-Bains, France). NGS was performed using the FusionPlex® lung kit (Archer DX Inc., Boulder, CO, USA) targeting *ALK, RET, ROS1, NGR1, NTRK1/2/3, FGFR1/2/3, BRAF, KRAS, EGFR* fusions and *MET* exon 14 skipping on a MiSeq platform (Illumina Inc., San Diego, CA, USA). This technology uses a specific 3'primer for the targeted gene and a universal 5'primer, enabling the detection of any partner.

Results: Oncogenic rearrangements were found in 16/58 samples (28%) with contributive analyses. The most frequent event was a *KIF5B-RET* translocation (6 patients). We also found 4 *ROS1* cases (*CD74* and *EZR*), 2 *NRG1* (*SLC3A2* and *FLYWH1*), 2 *EGFR* (*VOPP1*), 1 *ALK* (*EML4*) and 1 *NTRK1* (*EGFR*). Moreover, 4 patients had *MET* exon14 skipping confirmed by the identification of a splice mutation. All but *EGFR-VOPP1* fusions were targetable (18/20, 88%). RET FISH was negative or equivocal in 3/6 cases, *ROS1* FISH was negative in 1/4 positive case. ALK (5A4) and NTRK1 IHC were negative in both positive cases. Patients with fusions were younger than usually described and the sex ratio was close to 1. The proportion of smockers was higher in the group without oncogenic events (p = 0.045). Patients without an oncogenic event were more likely to express PD-L1 (p < 0.05). Five patients were treated, with partial response.

Conclusions: Our study showed that 1/3 of patients with a *KRAS/EGFR/NRAS/BRAF/ERBB2/ALK-IHC* wild type tumor had a targetable fusion oncogenic event, while <10% are expected in an unselected NSCLC population. Sequential analysis based on mutational status is efficient to test for fusions regardless of patient's clinical features or tobacco exposure.

1893 Vaping-Induced Pulmonary Injury: Histopathologic and Clinical Characteristics at a Single Referral Institution

Megan Desai¹, Rosana Eisenberg², Margaret Compton², Fabien Maldonado², Mitra Mehrad²

¹Nashville, TN, ²Vanderbilt University Medical Center, Nashville, TN

Disclosures: Megan Desai: None; Rosana Eisenberg: None; Margaret Compton: None; Fabien Maldonado: None; Mitra Mehrad: None

Background: Electronic (e-) cigarettes are battery-powered devices that heat a liquid, commonly containing nicotine, tetrahydrocannabinol (THC) and cannabidiol (CBD) as well as varying solvents into an aerosolized form to be inhaled. Use of these e-cigarettes, also known as 'vaping', is marketed as safer than smoking of conventional cigarettes; however, there are now over 500 reported cases of severe vaping-induced pulmonary injury (VAPI), including multiple deaths. Few studies have been recently published, predominantly focusing on the clinical aspect of VAPI, however, pathologic features are understudied.

Design: Slides from clinically confirmed cases of VAPI at a tertiary care hospital, who underwent lung biopsy and/or bronchoalveolar lavage (BAL) were identified and reviewed. Patients were excluded if an infectious or rheumatologic etiology was confirmed. Clinicoradiologic findings were obtained by chart review.

Results: Four patients (3 male; 1 female; age range 20-31 years) were identified over a course of one month (August 2019) (Table 1). Three patients underwent transbronchial biopsies with BAL and one only had a wedge biopsy. All patients had history of vaping within one month of onset of symptoms. The main histologic pattern seen was organizing acute lung injury with mild alveolar septal mononuclear infiltrate. Two cases had sparse neutrophilic infiltrate within the alveolar septa and spaces and 3 showed foamy and pigmented macrophages. BALs were notable for lipid-laden macrophages identified by Oil-Red-O stain. GMS stain for fungal organisms was negative in all biopsies.

Age/Sex/Race	21y/Male/Caucasian	28y/Male/Caucasian	20y/Male/Caucasian	31y/Female/Caucasian
Clinical History	-THC (daily use, 5-6 bands/week x 6-7	-nicotine and THC (x9 months)	-THC (1 cartridge/day x several	- History of vaping (unknown
	years)	-8 days prior to presentation	months)	duration)
	-14 days prior to presentation developed	developed SOB, cough, fever,	-14 days prior to presentation	- Cough/SOB
	SOB, cough, fever, malaise	malaise, nausea	developed SOB, cough, fever,	
	-last use 2 weeks prior to presentation	-last use 1 month prior to	nausea	
		presentation	-last use 21 days prior to	
			presentation	
Laboratory Findings	WBC 8.2 (81% PMNs)	WBC 20.7 (94% PMNs)	WBC 44.8 (95% PMNs)	None available
	AST 50, ALT 56, AlkPhos 53	AST 41, ALT 102, AlkPhos 144	AST 161, ALT 254, AlkPhos 199	
			Legionella serology inconclusive	
Radiology (CT)	-diffuse GGO, most pronounced in lower	-diffuse GGO	- diffuse GGO, basilar	-Bilateral persistent GGO
	lobes	-mediastinal and hilar LAD	dependent	
	-mediastinal and hilar LAD		-mediastinal LAD	
Pathology	TBBX: Organizing pneumonia, reactive	TBBX: Organizing pneumonia,	TBBX: Organizing pneumonia,	Wedge biopsy: Organizing
ratifology	pneumocytes, alveolar septal	fibrin exudate, reactive	fibrin exudate, reactive	pneumonia, fibrin exudate,
	mononuclear infiltrate, pigmented	pneumocytes, alveolar septal	pneumocytes, alveolar septal	reactive pneumocytes,
	macrophages	mononuclear and neutrophilic	mononuclear and neutrophilic	interstitial chronic inflammation.
	macropriages	· ·		
		infiltrate, foamy and pigmented	infiltrate, foamy and pigmented	foamy and pigmented
		macrophages	macrophages	macrophages
SOB: shortness of breath;	GGO: ground glass opacities; LAD: lymphadenopa	athy; TBBX: transbronchial biopsy		
Laboratory reference rang	es: WBC 3.9-10.7 x10^3/uL, AST 5-40 U/L, ALT 0-	55 U/L, AlkPhos 40-150 U/L		

Conclusions: The clinicoradiologic presentation of these four cases are similar to those recently reported in the literature. Our small series suggest that VAPI is characterized by organizing acute lung injury and should be considered in the differential diagnosis of diffuse alveolar damage of unknown etiology, particularly in younger patients. While foamy macrophages were found in 3 cases, the characteristic features of lipoid pneumonia were not evident on histologic examination. As such, the significance of lipid-laden macrophages on BAL requires further investigation.

1894 Next Generation Sequencing of Type 1 Congenital Pulmonary Airway Malformations Identifies Oncogenic KRAS Mutations in Mucinous and Non-Mucinous Epithelium

Walter Devine¹, Anatoly Urisman², Kirk Jones²

¹University of California San Francisco, Berkeley, CA, ²University of California San Francisco, San Francisco, CA

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Background: Congenital Pulmonary Airway Malformations (CPAMs) are rare developmental malformations of airway branching in the lung, typically diagnosed antenatally. They are classified into 4 subtypes (Type 1, Type 2, Type 3, and Type 4) based on their histologic appearance. Their origin, natural history, and genetics remain poorly understood as does their malignant potential. A subset of Type 1 CPAMs are thought to possess malignant potential and eventually go on to develop into invasive mucinous adenocarcinoma. These lesions are characterized by areas of mucinous epithelium that can often represent a very small fraction of the total Type 1 CPAM. It remains to be determined if Type 1 CPAM lesions with mucinous epithelium are molecularly distinct from Type 1 CPAMs that lack mucinous epithelium.

Design: A total of 10 Type 1 CPAMs and 10 Type 2 CPAMs with available slides and tissue blocks were identified. Each Type 1 CPAM was assessed for the presence or absence of mucinous epithelium. Four cases of Type 1 CPAM had microscopic evidence of mucinous epithelium and these areas were separately micro-dissected from areas without mucinous epithelium. Genomic DNA was extracted and used for molecular testing. Capture-based next generation sequencing was performed at the UCSF Clinical Cancer Genomics Laboratory, using an assay (UCSF500 panel) that targets the coding regions of 480 cancer-related genes, select introns from approximately 40 genes, and the TERT promoter with a total sequencing footprint of 2.8 Mb

Results: Among the four Type 1 CPAMs that contained mucinous epithelium, a pathogenic hotspot mutation in KRAS codon12 was detected in all four cases. This pathogenic mutation was detected in both the mucinous and non-mucinous epithelium at roughly similar mutant allele frequency. None of the remaining 6 cases of Type 1 CPAMs that lacked mucinous epithelium or the 10 Type 2 CPAMs contained a KRAS mutation.

Conclusions: 1.) Type 1 CPAM lesions represent two molecular subtypes: lesions with mucinous epithelium characterized by recurrent oncogenic KRAS mutations and lesions that lack mucinous epithelium that have no recurrent KRAS mutations. 2.) For Type 1 CPAM lesions with mucinous epithelium, oncogenic KRAS mutations are present in both mucinous and non-mucinous epithelium, and 3.) Recurrent genetic alterations in Type 1 CPAMs without mucinous epithelium or KRAS mutations remain to be discovered.

1895 Pathologic Findings in Lung Allograft Surveillance Transbronchial Biopsies with Donor-Specific Antibodies: A Preliminary Report

Muhammad Elahi¹, Aileen Grace Arriola¹, Steven Geier¹, Amandeep Aneja²

¹Temple University Hospital, Philadelphia, PA, ²Temple University Hospital, Bryn Mawr, PA

Disclosures: Muhammad Elahi: None; Aileen Grace Arriola: None; Steven Geier: None; Amandeep Aneja: None

Background: Difficulty with enumerating specific histopathologic criteria for lung transplant antibody mediated rejection (AMR) is well recognized. The 2016 Banff study confirmed capillary inflammation, acute lung injury (ALI), and endothelialitis as morphologic features in lung allograft biopsies that correlate with the presence of donor-specific antibodies (DSAs). Alveolar septal widening (ASW) is another AMR feature recently described in a multicenter pilot study. We performed a retrospective study to review the histologic features in surveillance transbronchial biopsies and determine the statistical significance with the presence of DSAs.

Design: Surveillance biopsies of patients with available serologic data between June 2018-19 were retrieved from departmental archives. A total of 30 biopsies were available, of which 15 were with DSAs and 15 with no antibodies (NABs). Two transplant pathologists blinded to all serologic and clinical data reviewed the biopsies for various histologic parameters. The readings were pooled and correlated with DSA status using Fisher's exact test.

Results: Biopsies with DSA had a statistically significant difference versus NABs with regards to ALI (p=0.0421), presence of capillary neutrophilic inflammation (p= < 0.00001), and the presence of ASW (p=0.0007). Similar to earlier reports, we found a positive correlation between the capillary neutrophils, ALI, and DSAs. Frequent ASW was noted in our patients with DSAs and was an easily recognized feature at low-power magnification. Endothelialitis was not appreciated in any of our cases, likely due to a small sample size.

Complement 4d (C4d) was available in 16.6 % of specimens and showed only focal staining (<50%). C4d, acute cellular rejection, and airway inflammation did not reveal any significant statistical association with DSA status.

Conclusions: ALI, capillary neutrophilic inflammation, and ASW were morphological features found to be statistically significant in this small sample size study. These are not pathognomonic, and currently, the triple test (clinical allograft dysfunction, DSAs, pathologic findings) is the best approach for the diagnosis of AMR. Further research investigating ASW, exploring its likelihood to be included among conventional histopathologic patterns prompting clinical and serologic evaluation is needed.

1896 Histologic Features of Desmoplastic Mesothelioma (DMM)

Katsura Emoto¹, Marina Baine², William Travis², Kelly Butnor³, David Klimstra², Kay See Tan², Prasad Adusumilli², Jennifer Sauter²

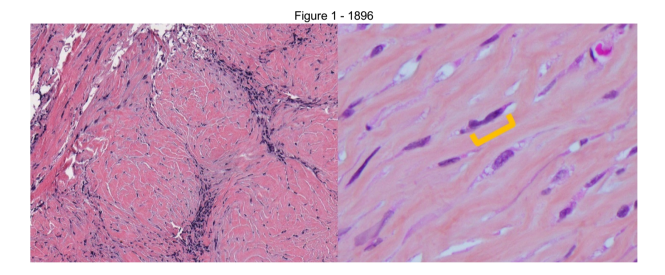
¹Keio University Hospital, Shinjuku-ku, Tokyo, Japan, ²Memorial Sloan Kettering Cancer Center, New York, NY, ³The University of Vermont Medical Center, Burlington, VT

Disclosures: Katsura Emoto: None; Marina Baine: None; William Travis: None; Kelly Butnor: None; David Klimstra: None; Kay See Tan: None: Prasad Adusumilli: None: Jennifer Sauter: None

Background: The diagnosis of DMM is often challenging because tumor cells may be bland and desmoplasia can mimic chronic fibrous pleuritis (CFP). The aim of this study is to identify histologic features useful for distinguishing DMM from CFP.

Design: 54 patients with malignant mesothelioma (MM) with at least focal desmoplastic component and 13 patients with CFP were identified from surgical databases and pathology consultation files at the study institutions. Two thoracic pathologists re-reviewed slides for the following: desmoplastic component (%), invasion, bland necrosis, proliferation nodules (defined as desmoplastic round nodules (<1mm), left figure), storiform pattern, myofibroblastic proliferation, granulation tissue, maximum nuclear length (defined in relation to lymphocyte size, right figure), nuclear atypia (grades 1-3), mitotic score of 1(0-1/2mm²), 2(2-4/2 mm²) or 3(≥5/2 mm²) and infiltrating inflammatory cells (%). Overall survival (OS) was determined by Kaplan-Meier method and compared between groups using log-rank test.

Results: Invasion, bland necrosis, proliferation nodules, nuclear atypia (grade ≥2), mitotic score (≥2), nodular growth and storiform pattern showed 100% specificity (all P<0.001) but variable sensitivity (75%, 67%, 69%, 81%, 88%, 89% and 96%, respectively) for MM. Nuclear length was significantly greater in MM than in CFP (median 4.0 vs 3.0x size of lymphocyte, respectively; P<0.0001). Myofibroblastic proliferation and granulation tissue were more frequently observed in CFP (69% and 77%, respectively) than in MM (both 19%) (P=0.001 and P<0.001, respectively). Percentage of infiltrating inflammatory cells was not significantly different (P=0.3) between MM and CFP (median 10% and 15%, respectively). There were no significant differences in gender (P=0.4), age (P=0.6), stage (P=1) and OS (P=0.13) between patients with sarcomatoid MM (n=26) and DMM (n=28, defined as ≥50% desmoplastic component by WHO).



Conclusions: In the distinction from CFP, several features appear specific for MM, among which nuclear atypia, mitotic score, nodular growth and storiform pattern are most sensitive. Myofibroblastic proliferation and granulation tissue, while not exclusive to CFP, may be a reliable indicator of benignity in absence of features specific for MM. Based on this limited cohort, there appear to be no significant survival differences in patients whose tumors meet WHO criteria for DMM compared to those with MM showing <50% desmoplastic component.

1897 Diagnostic Value of L1 Cell Adhesion Molecule (L1CAM) Immunostaining in Well-Differentiated Papillary Mesothelioma of the Peritoneum and Diffuse Malignant Peritoneal Mesothelioma

Mona Farahi¹, Alain Borczuk², Kartik Viswanathan³, Michael Kluger⁴, Hanina Hibshoosh⁵

¹New York-Presbyterian/Weill Cornell, New York, NY, ²Weill Cornell Medicine, New York, NY, ³New York-Presbyterian

Hospital/Weill Cornell Medical Center, New York, NY, ⁴New York-Presbyterian/Columbia University Medical Center, New York, NY, ⁵Columbia University Medical Center, New York, NY

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Background: Well-differentiated papillary mesothelioma (WDPM) is a rare indolent mesothelial tumor that is usually found incidentally in the peritoneal cavity. There is controversy about the prognosis of the disease with few reported cases of invasive and malignant behaving WDPM. Thus, some consider it a tumor of uncertain malignant potential. In contrast, diffuse malignant peritoneal mesothelioma (DMPM) is an aggressive tumor associated with asbestos exposure with a poor prognosis. The distinction between these two entities relies heavily on histopathological findings. Prior studies identified TRAF7 and CDC42 (mutually exclusive) somatic mutations in WDPM. L1 cell adhesion molecule (L1CAM) an NF-kB activated gene was overexpressed in WDPM. The goal of the present study is to assess the diagnostic utility of L1CAM in a larger cohort of peritoneal mesotheliomas.

Design: Immunohistochemistry utilizing mouse monoclonal anti-L1CAM antibody (clone UJ127.11, Sigma, L4543, 1:1000 dilution) was performed on formalin-fixed, paraffin-embedded tissue sections in 197 total cases consisting of: 41 cases of biphasic/sarcomatoid DMPM, 16 cases of WDPM, 8 cases of benign multicystic mesothelioma (BCM), and 132 cases of epithelioid DMPM. The staining was scored as either positive or negative but since 15% was the minimum percentage to show complete and intense membranous staining among the WDPM, this was chosen as the true positive cutoff.

Results: Immunohistochemistry for L1CAM was positive in 88% (14/16) of WDPM, 19% (25/132) of epithelioid DMPM, 20% (8/41) of biphasic/sarcomatoid DMPM and 0% (0/8) of BCM. Of the 25 epithelioid DMPM, the predominant patterns were: 4 papillary, 2 micropapillary, 8 solid, 1 solid and papillary, 1 deciduoid and solid, 6 tubulopapillary and 3 with no pattern information. Of the 25 L1CAM positive epithelioid DMPM patients, 19 were dead from disease with a median survival of 1060 days. L1CAM showed 88% sensitivity for WDPM, but a specificity of 81%, PPV of 36% and NPV of 98% when including only epithelioid DMPM for comparison (P-value <0.0001).

Table 1. Result Summary							
L1CAM Status by IHC	WDPM (%)	Epithelioid DMPM (%)	Biphasic/Sarcomatoid DMPM (%)	BCM (%)			
Positive (>15% membranous staining)	14 (88%)	25 (19%)	8 (20%)	0 (0%)			
Negative (≤15% membranous staining)	2 (12%)	107 (81%)	33 (80%)	8 (100%)			
TOTAL	16 (100%)	132 (100%)	41 (100%)	8 (100%)			

Conclusions: L1CAM is more frequently positive in WDPM than in epithelioid DMPM, biphasic/sarcomatoid mesothelioma or BCM. However, the immunoreactivity in a significant proportion of the epithelioid DMPM even with a cutoff of 15% reduces the value of this test diagnostically with only 36% PPV and specificity of only 81%.

1898 Dual-Staining Adenosquamous Lung Carcinoma with p16 Overexpression- A Distinct Variant?

Taylor Forns¹, Alexis Musick², Elizabeth Pavlisko¹, Elin Hughes³, Thomas Sporn⁴, Kristen Deak³, Carolyn Glass¹

¹Duke University Medical Center, Durham, NC, ²Duke University School Medicine, Durham, NC, ³Duke University, Durham, NC, ⁴Duke University Medical Center, New Bern, NC

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Background: Adenosquamous lung carcinoma (ASC) is a rare yet aggressive malignancy thought to potentially arise from a bipotential undifferentiated precursor cell. A topic of controversy until coined a distinct entity in the 2015 5th edition of the WHO Classification of Lung Cancer, ASC is defined as a tumor with two separate malignant populations (>10% glandular and squamous components). The coexistence of glandular and squamous differentiation *within the same tumor cells* has also previously been reported (*Pelosi, Journal of Thoracic Oncology 2015*) and termed "ASC immunophenotype" due to its amphicrine phenotypic nature. The single case report showed dual TTF-1 and p40 immunohistochemical (IHC) expression, along with specific electron microscopy and molecular findings. We report here for the first time a case series with dual differentiation and further characterize the findings with molecular analysis.

Design: From 58 primary lung ASCs diagnosed between 2017 and 2019 at our institution, we identified 7 patients (mean age 72.1 years) with dual glandular and squamous differentiation within the same population of malignant tumor cells. IHC stains for CK5/6, CK7, TTF-1, p40, and p16 were performed. The lon AmpliSeq[™] Cancer Hotspot Panel v2 (CHP2) and VariantPlex Myeloid Kit (ArcherDX) next generation sequencing (NGS) assays targeting 125 oncogenes were performed on two cases. Fluorescence in-situ hybridization (FISH) was also performed on the two sequenced cases.

Results: ASCs with dual differentiation occurred predominantly in women (85.7%, age range 60 to 85 years) with variable smoking history. All cases were poorly differentiated with a solid growth pattern and 43% (3/7) presented with distant metastasis. The following IHC stains were positive in the tumor population: CK5/6 (4/4, 100%), CK7 (3/3, 100%), TTF-1 (7/7, 100%), p40 (7/7, 100%), and p16 (7/7, 100%).

NGS revealed CDKN2A and TP53 mutations in both cases; one CDKN2A variant is known to disrupt p16INK4A- and p14ARF-dependent regulation of CDK4/6 and p53. Loss of function ATRX, BCORL1 and NOTCH1 mutations were also identified. FISH revealed CDKN2A mutations and relative loss of 9p21 in one case, and chromosome 9 monosomy in the other.

Conclusions: We report for the first time a case series of these rare lung ASCs with dual differentiation and show an association with CDKNA2 mutations and corresponding p16 overexpression.

1899 Finding "Ground Truth" for the Diagnosis of UIP with Artificial Intelligence - Standardization of Diagnoses by 16 Expert Pathologists from 9 Countries

Junya Fukuoka¹, Yoshiaki Zaizen², Mutsumi Ozasa³, Mano Soshi⁴, Rosane Duarte Achcar⁵, Amna Almutrafi⁶, Jaroslaw Augustyniak², Sabina Berezowska⁶, Luka Brcic⁶, Alberto Cavazza¹⁰, Alexandre Fabro¹¹, Kaori Ishida¹², Andre Moreira¹³, Alberto Marchevsky¹⁴, Anja Roden¹⁵, Frank Schneider¹⁶, Maxwell Smith¹७, Angela Takano¹⁶, Tomonori Tanaka¹⁶, Yasuhiro Kondoh²⁰, Andrey Bychkov²¹

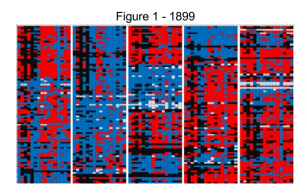
¹Nagasaki University, Nagasaki, Japan, ²Nagasaki University, Nagasaki City, Japan, ³Nagasaki Harbor Medical Center, Nagasaki, Japan, ⁴BonBon Ink, Kyoto, Japan, ⁵National Jewish Health, Denver, CO, ⁶King Abdulaziz Medical City, Riyadh, Saudi Arabia, ⁷Helios Clinics Schwerin, Schwerin, MV, Germany, ⁸Institute of Pathology, University of Bern, Bern, Switzerland, ⁹Medical University of Graz, Graz, Austria, ¹⁰Azienda USL Reggio Emilia, Reggio Emilia, RE, Italy, ¹¹Ribeirão Preto Medical School - University of São Paulo, Ribeirão Preto, SP, Brazil, ¹²Kansai Medical University, Hirakata, Osaka, Japan, ¹³New York Langone Health, New York, NY, ¹⁴Cedars-Sinai Medical Center, West Hollywood, CA, ¹⁵Mayo Clinic, Rochester, MN, ¹⁶Emory University, Atlanta, GA, ¹⁷Mayo Clinic Arizona, Scottsdale, AZ, ¹⁸Singapore General Hospital, Singapore, Singapore, ¹⁹Kobe University Hospital, Kobe, Hyogo, Japan, ²⁰Tosei General Hospital, Seto, Aichi, Japan, ²¹Kameda Medical Center, Kamogawa, Japan

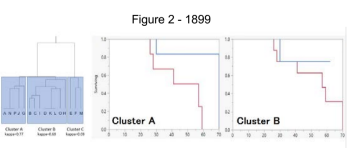
Disclosures: Junya Fukuoka: *Advisory Board Member*, N Lab, Co Ltd; Yoshiaki Zaizen: None; Mutsumi Ozasa: None; Mano Soshi: None; Rosane Duarte Achcar: None; Amna Almutrafi: None; Jaroslaw Augustyniak: None; Sabina Berezowska: None; Luka Brcic: None; Alberto Cavazza: None; Alexandre Fabro: None; Kaori Ishida: None; Andre Moreira: None; Alberto Marchevsky: None; Anja Roden: None; Frank Schneider: None; Maxwell Smith: None; Angela Takano: None; Tomonori Tanaka: None; Yasuhiro Kondoh: None; Andrey Bychkov: None

Background: There is considerable interobserver variability in the pathological diagnosis of UIP and other ILD. The variability in finding "ground truth" presents a difficult problem during the development of deep learning platforms designed to diagnose ILD using AI. We describe the use of a smartphone application that allows for the collection of multiple opinions from sixteen pulmonary pathologists from nine countries as a novel method for finding "ground truth" for UIP diagnoses.

Design: Whole slide images from 14 consecutive interstitial lung disease cases were scanned by Aperio CS2 at 20x objective and sliced into multiple individual 7x7mm images, resulting in 355 JPEG image patches. These individual images were randomized and shown over the internet to 16 expert pathologists from 9 countries using the novel smartphone application, BonBon system. Expert lung pathologists were asked to classify each image into one of 8 categories: UIP/IPF; CTD/UIP; CHP/UIP; UIP/other cause; Non-UIP; "not sure"; "normal"; and "exclude". Images classified as "exclude" by any of the experts were deleted from analysis. All individual diagnostic classes selected for individual images were grouped by case, and the predominant class for each case, was used as "diagnosis". All diagnoses were analyzed using clustering analysis and Kaplan Meier statistics using JMP. Interobserver agreement was calculated by Fleiss kappa coefficient using R.

Results: The diagnosis of each of the 14 cases by each of the 16 pathologists showed that interobserver agreement using the 7 categories was poor (k=0.19). None of the cases were classified as CHP-UIP or UIP/others by a majority of pathologists. In order to simplify the diagnoses into more clinically relevant classes, the categories UIP/IPF, CHP-UIP, and UIP/others were grouped into UIP and Non-UIP and CTD-UIP into Non-UIP as shown in Figure 1. Clustering analysis stratified the diagnoses of each case into 3 clusters as shown in Figure 2A (k= 0.77, 0.63, and 0.03, respectively), validating cluster A and B were meaningful. Log rank test showed significant survival difference between UIP and Non-UIP groups only for cases in cluster A (P=0.017) (Fig 2B) but not for cases in cluster B (Fig 2 C). Eventually, 269 H&E images with high agreement of UIP and non-UIP group were selected as "ground truth" for Al training.





Conclusions: Classification of image patches from WSI over APP offers a useful method to standardize UIP diagnosis and to develop diagnostic AI.

1900 Validation of the IdyllaTM EGFR Assay for Rapid Assessment of EGFR Mutation Status in Nonsmall Cell Lung Cancer

Ramya Gadde¹, Donald Green², Kelley Godwin², Konstantin Dragnev², Gregory Tsongalis², Laura Tafe²

¹Dartmouth Hitchcock Medical Center, Lebanon, NH, ²Dartmouth-Hitchcock Medical Center, Lebanon, NH

Disclosures: Ramya Gadde: None; Donald Green: None; Kelley Godwin: None; Konstantin Dragnev: None; Gregory Tsongalis: *Advisory Board Member*, Bioartis: Laura Tafe: None

Background: In advanced non-small cell lung cancer (NSCLC), presence of a sensitizing *EGFR* mutation confers a more favorable prognosis and strongly predicts response to EGFR tyrosine kinase inhibitors (TKIs). Having *EGFR* results available within a day of diagnosis allows for fast management decisions and quicker access to EGFR TKIs in patients with *EGFR*-mutated NSCLC. The Idylla™ system (Biocartis, Mechelen, Belgium) is a fully integrated, cartridge-based platform that provides automated sample processing (deparaffinization, tissue digestion and DNA extraction) and real-time PCR-based multiplex mutation detection within a single-use cartridge. The Idylla™ *EGFR* Mutation Assay was validated against next-generation sequencing (NGS) results for use as a rapid screen for NSCLC to detect actionable *EGFR* mutations.

Design: Twenty-seven archived formalin-fixed paraffin-embedded (FFPE) NSCLC tissue samples (13 excisions, 8 FNA cell blocks, 6 small biopsies) and HD300 FFPE cell line were included. NGS analysis was previously performed using the lon AmpliSeq 50-gene Cancer Hotspot Panel v2 (Thermo Fisher Scientific). For each case, 1-4 4 μm unstained FFPE tissue sections were macrodissected and introduced into the cartridge. All cases met Idylla™ minimum tumor content requirement of 10%. Precision, accuracy, limit of detection (LOD) and reproducibilty were assessed.

Results: Precision, reproducibilty and LOD were assessed using HD300 in triplicate by 2 operators on 2 instruments. The 27 FFPE and HD300 were used for accuracy. Idylla™ results were in complete agreement with those obtained by NGS for *EGFR* mutations targeted by Idylla™ except for one sample. These include L858R (1), G719A (3), G719C (2), L861Q (1), S768I (2), L861Q (1), T790M (3), exon 19 deletions (16) and exon 20 insertion (1). The discrepant sample had an exon 19 deletion identified by NGS but was wild-type by Idylla™. Upon review of NGS data, an *EGFR* synonymous SNV was present downstream of the deletion which might impair the probe binding, leading to false negative results. LOD of 5% VAF and tumor content of 10% were confirmed. No *EGFR* mutations were detected by Idylla™ in the samples determined by NGS as having wild-type *EGFR*.

Conclusions: The fully automated Idylla™ system offers rapid (turnaround time of approximately 2.5 hours) and reliable testing of clinically actionable mutations in *EGFR* directly from FFPE tissue sections. In our center, it will complement NGS testing by providing rapid *EGFR* results within 1-2 days of diagnosis.

1901 Exploring Genomic Heterogeneity of Synchronous Lung Adenocarcinomas by Whole-Exome Sequencing

Andréanne Gagné¹, Venkata Manem², Michèle Orain², Maxime Lamontagne³, Yohan Bossé⁴, Philippe Joubert⁵

¹Laval University, Quebec City, QC, ²Institut Universitaire de Cardiologie et de Pneumologie de Quebec Research Center, Quebec, QC, ³Institut Universitaire de Cardiologie et de Pneumologie de Quebec Research Center, Québec City, QC, ⁴IUCPQ, Québec, QC, ⁵Quebec Heart and Lung Institute, Quebec, QC

Disclosures: Andréanne Gagné: None; Venkata Manem: None; Michèle Orain: None; Maxime Lamontagne: None; Yohan Bossé: None; Philippe Joubert: None

Background: Identification of patients with synchronous lung adenocarcinomas has crucial implications for their staging and clinical management. These tumors are thought to represent independent primary neoplasms, but their oncogenesis is still poorly characterized. A few studies based on the analysis of limited panel of genes suggest that synchronous tumors have a high level of genetic heterogeneity. We sought to investigate the genetic mutations in a well-defined cohort of synchronous adenocarcinomas using whole-exome sequencing (WES).

Design: A retrospective cohort of 34 Caucasian patients that underwent surgical resections for synchronous lung adenocarcinomas between 2013 and 2016 in our center was selected. All slides were reviewed by a thoracic pathologist to confirm the diagnosis based on a comprehensive histologic assessment. Clinical characteristics and outcome were extracted from the electronic medical files. For each patient, three formalin-fixed paraffin embedded blocks were selected: one from two synchronous tumors and one from normal lung tissue. DNA was extracted and samples were submitted to WES on an Illumina HiSeq platform (Agilent SureSelect XT). Data processing was made according to GATK best practices and somatic variants (single nucleotide variants and insertion-deletions) were called using Strelka and MuTect.

Results: The cohort comprised 44.1% men with a mean age of 65.0 years, 91.2% had a stage I or II disease and 94.1% were smokers or ex-smokers. Mean follow-up was of 38.4 months with an 82.4% overall and 67.7% progression-free survival rates. WES showed an average of 317 called somatic variants per tumor. Variants were identified in known driver genes at the following prevalences: KRAS 35.3%, EGFR 8.8%, BRAF 8.8%, ROS1 5.9%, ALK 1.5%, MET 4.4%, RET 1.5%. Synchronous tumors from the same patients showed a high level of heterogeneity, as pairs shared 0 to 12 (0% to 3.3%) variants. All shared variants were likely passenger mutations, but one pair of tumors shared three variants including one in a driver gene, a KRAS p.G12C mutation.

Conclusions: We showed a high level of genomic heterogeneity between two synchronous adenocarcinomas from the same patient using WES, supporting independent primary tumors. One pair of tumors had an identical KRAS mutation with a high level of genomic heterogeneity, emphasizing the fact that synchronous tumors can share a mutation in a frequent driver gene randomly.

1902 Characterization of Patients with Synchronous Lung Adenocarcinomas in a Caucasian Population According to the IASLC Criteria

Andréanne Gagné¹, Raphaëlle Brière², Michèle Orain³, Philippe Joubert⁴

¹Laval University, Quebec City, QC, ²Institut Universitaire de Cardiologie et de Pneumologie de Quebec Research Center, Québec City, QC, ³Institut Universitaire de Cardiologie et de Pneumologie de Quebec Research Center, Quebec, QC, ⁴Quebec Heart and Lung Institute, Quebec, QC

Disclosures: Andréanne Gagné: None; Raphaëlle Brière: None; Michèle Orain: None; Philippe Joubert: None

Background: Identification of patients with synchronous lung adenocarcinomas (MSLA) is important for staging and clinical management purposes. To improve their recognition, the International Association for the Study of Lung Cancer (IASLC) recently proposed a classification based on clinicopathologic criteria. It divides MSLA in three categories: second primary, multiple ground glass opacities (GGO) and pneumonic type. However, the prevalence of these three subgroups remains poorly described and their clinical characteristics have been mostly described in Asian cohorts. We aimed to establish the prevalence, clinicopathologic characteristics and prognosis of patients with MSLA in a Caucasian population according to the IASLC criteria.

Design: We selected a retrospective cohort of 1012 consecutive patients, including 432 surgical patients, with a diagnosis of lung adenocarcinoma in our center between 2011 and 2014. The cohort was divided according to the IASLC classification: a group of sporadic tumors, comprising patients with one tumor and those with intrapulmonary metastasis (IPM), and a MSLA group further divided in second primary, multiple GGO and pneumonic type. Prevalence of each group was calculated in the whole cohort. Chi-square and t-tests were used to evaluate the associations between clinicopathological characteristics and the IASLC groups in surgical patients. Overall survival of surgical patients was compared using a Cox proportional model.

Results: 143 patients (14.1%) with MSLA were identified, including 52 (5.1%) second primary, 84 (8.3%) multiple GGO and 7 (0.7%) pneumonic type. Age (p=0.74), gender (p=0.38) and smoking status (p=0.77) were not associated with MSLA. MSLA patients had significantly more metachronous lung tumors (p=0.006), atypical adenomatous hyperplasia foci (p<0.001) and tumors with lepidic and acinary predominant patterns (p=0.03). There were no significant differences between the MSLA groups. Compared with patients with a single tumor, the multiple GGO group tended to have the best prognosis (HR=0.75, p=0.41) and the second primary (HR=1.97, p=0.02) and pneumonic type (HR=2.33, p=0.41) had worse survival when using a multivariate model.

Conclusions: To our knowledge, this is the first report to assess MSLA prevalence using the IALSC criteria to define patients with multiple adenocarcinomas. Even though there were no differences in the clinicopathological variables between the MSLA groups, their survival disparity supports the IASLC classification.

1903 Most Peripheral Large Cell NeuroEndocrine Carcinomas Have Carcinoid Features

Stéphane Garcia¹, Kevin Caselles², Jerome Winkler², Josephine Chenesseau³, Laurent Greillier², Donatienne Bourlard⁴
¹AP-HM, Marseille, France, ²AP-HM, Marseille, Bouches du Rhone, France, ³AP-HM, Marseille, Sud, France, ⁴AP-HM, Marseille, Bouches du Rhones, France

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Background: Unreported in the 2015 WHO classification, a subset tumor classified as Large Cell NeuroEndocrine Carcinoma but with carcinoid clinical and pathologic features has been described. We have reviewed resected LCNEC from our archives to identify this new entity.

Design: Histologic slides of pulmonary lobectomies for high grade NECs from the 2007-2018 period were reviewed. High grade was characterized by a mitotic count >10 per 2 mm². Necrosis was subtyped as massive or punctuate. Carcinoid morphology was identified

according to the WHO morphological criteria. Tumors with diffuse architecture, nuclear polymorphism and marked nucleoli were classified as "classic" LCNEC. Proliferation was also assessed with immunodetection of ki67.

Results: Ten cases were available, 4 were classic LCNEC, 6 had carcinoid-like features. Clinical and pathological data are listed in Table 1. There were no differences between the 2 groups for clinical data. Carcinoid-like subgroup had lower mitotic count (mean 28 vs 70), ki67 index (mean 54% vs 84%) and better survival (61 vs 18 months) than classic LCNEC respectively. No reliable statistical analysis was allowed due to the small size of our cohort.

	Classic LCNEC (n = 4)	Carcinoid-like LCNEC (n = 6)
Age mean (range)	57 (50 – 76)	71 (60 – 72)
Male/ Female	2/2	4/1
Tumor Size mean mm (range)	20,5 (15 – 99)	43 (28 – 80)
Architecture : Diffuse/ Organoïd	4/0	0/5
Nécrosis : Massive / Punctuate	3/1	0/5
Ki67 % (range)	84,4 (60 – 100)	54 (10 – 90)
Mitotic count (range)	70 (65 – 80)	28 (15 – 40)
Survival mean (months)	18	61

Conclusions: LCNECs with carcinoid features form a homogeneous subtype with precise morphological criterias. It is rare (0.3% of resected lung tumors in our center), nevertheless its real incidence is probably underestimated. Indeed, for sampling reasons, we chose to report a series including only surgical specimens but we have already identified this tumor-type on biopsies from on resectable mediastinal lesions.

Proliferation markers are lower in carcinoid-like LCNECs than in classic LCNECs. Nevertheless, results differ depending on the method of assessment. There is no overlap between the mitotic count values in both subgroups whereas ki 67 index overlaps with no upper limit value in carcinoid-like group.

These results rise two questions: 1/ the choice of the method of evaluation of proliferation between mitotic count or ki67, 2/ the place of carcinoid-like LCNEC within pulmonary NETs classification: an intermediate group between atypical carcinoids and classic LCNECs or a high grade well differentiated NET equivalent to that described in the digestive tract.

However, the interests of individualizing the carcinoid-like LCNEC subgroup are: diagnostic (must be distinguished from atypical carcinoid), prognostic (better than classic LCNEC) and probably therapeutic (therapeutic response)

1904 Detection of EGFR Exons 18-21 Hotspot Mutations Using a Fully-Automated, Cartridge-Based Platform with Ultra-Rapid Turnaround Time: A Comparison Study with Conventional Next Generation Sequencing

Jonathon Gralewski¹, Laura Toth¹, Andrew Judd¹, Emily Volpicelli¹, Mohammad Vasef¹ *University of New Mexico, Albuquerque, NM*

Disclosures: Jonathon Gralewski: None; Laura Toth: None; Andrew Judd: None; Emily Volpicelli: None; Mohammad Vasef: None

Background: Lung cancer is a leading cause of cancer death worldwide. However, the introduction of targeted therapies in recent years has led to improved overall prognosis and survival. Current guidelines for lung adenocarcinoma require epidermal growth factor receptor (*EGFR*) exons 18-21 mutational testing. Furthermore, it outlines molecular methods, such as next generation sequencing (NGS) with an acceptable turnaround time (TAT) up to two weeks. This prolonged TAT delays treatment decisions and increases healthcare costs. Recently, a fully-automated, cartridge-based platform has been introduced with an ultra-rapid (i.e. <3 hours) sample-to-result TAT that does not require traditional sample preparation, such as DNA extraction, library preparation, and PCR amplification.

Design: Twenty-one archived formalin-fixed paraffin embedded lung adenocarcinoma including 5 cytology and 16 surgical samples with previously characterized exon 18-21 *EGFR* mutations were selected for this study. A single unstained section of the selected paraffin blocks was macrodissected, placed between filter papers, and subsequently placed into the cartridge and run for 51 EGFR variant mutational analysis as per the manufacturer instructions. Results were generated in <3 hours without the need for complex bioinformatics analysis and interpretation.

Results: Of 23 previously characterized *EGFR* mutations in 21 analyzed samples, 22 *EGFR* variants were successfully detected by fully integrated ultra-rapid platform (96% concordance). The automated platform failed to identify an *EGFR* G719C variant with allelic frequency of 6% that had been detected using conventional NGS platform. An additional *EGFR* L858R variant was only seen by the cartridge based PCR platform. This variant was re-interrogated on the NGS-based platform and deemed likely a false positive result.

EGFR mutations	NGS platform (# of cases)	Cartridge platform (# of cases)	Concordance (%)
Exon 18 G719A/S/C	3	2	67
Exon 19 deletion	8	8	100
Exon 20 T790M	1	1	100
Exon 20 S768I	1	1	100
Exon 20 mutation	1	1	100
Exon 21 L858R	8	9*	100
Exon 21 L861Q	1	1	100
Total	23	22	96

Conclusions: The cartridge-based fully integrated platform with ultra-rapid TAT demonstrated high concordance with the conventional NGS-based platform. The fully-automated and integrated platform has minimized the need for significant molecular expertise and laboratory infrastructure. However, this platform does have its limitations, such as only detecting the most common *EGFR* mutations. Overall, this fully automated platform provides an ultra-rapid, and reliable cost-effective method in detecting the most common *EGFR* variants, as outlined in the current guidelines with a detection sensitivity comparable with the conventional NGS platform.

1905 Inflammation on Bronchoalveolar Lavage Cytology is Associated with Decreased Chronic Lung Allograft Dysfunction-Free Survival

Nancy Greenland¹, Fred Deiter¹, Jonathan Hoover¹, Steven Hays¹, Jeffrey Golden¹, Jonathan Singer¹, G. Zoltan Laszik¹, John Greenland¹

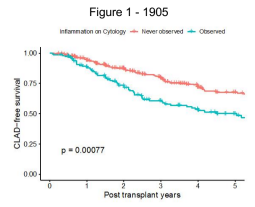
¹University of California San Francisco, San Francisco, CA

Disclosures: Nancy Greenland: None; Fred Deiter: None; Jonathan Hoover: None; Steven Hays: None; Jeffrey Golden: None; Jonathan Singer: None; G. Zoltan Laszik: None; John Greenland: None

Background: Lung transplant recipients undergo bronchoalveolar lavage (BAL) and biopsies to detect rejection and infection that may be antecedents of chronic lung allograft dysfunction (CLAD), the major limitation to long term survival. BAL cytology is routinely performed but recently some centers have advocated abandoning this practice because of the low diagnostic yield. We hypothesized that inflammation observed on BAL cytology would predict CLAD risk.

Design: We grouped diagnostic findings on BAL cytology between 2012 and 2019. Bronchoscopy indication, infection treatment, BAL and biopsy results, and CLAD-free survival were abstracted from medical records. Cytology associations with clinical characteristics were compared using generalized-estimating equation-adjusted logistic regression. The association between BAL inflammation and CLAD or death were determined using time-dependent Cox proportional hazards models adjusted for age, gender, diagnosis, lung allocation score, and transplant type.

Results: We evaluated 3,578 cytology reports from 491 subjects. Inflammation was the most common finding (6.4%), followed by fungi (5.2% of which 0.8% were likely pathogenic). There were 5 cases of malignancy and 2 cases of CMV. Inflammation on BAL cytology was more common in procedures performed for symptoms (11%) versus surveillance (3%, P<0.001), associated with antimicrobial initiation (a proxy for clinically significant infection, 9% vs 5%, P<0.001), associated with acute cellular rejection (P=0.01), and linked to increased BAL neutrophil and lymphocyte concentrations (P<0.001). Inflammation on BAL cytology was present for 32% of subjects on at least one sample, was more frequent around the time of CLAD onset, and was associated a 2.6-fold hazard ratio (CI 1.2-5.3) for CLAD or death (Figure 1). However, this association was not significant after adjusting for BAL cell counts and acute cellular rejection (P=0.22).



Conclusions: The presence of inflammation on BAL cytology specimens is clinically significant, suggesting acute rejection or infection and increased risk of CLAD or death. However, other indicators of allograft inflammation can substitute for some of the information provided by BAL cytology.

1906 Primary Pulmonary Myxoid Sarcoma and Myxoid Variant of Angiomatoid Fibrous Histiocytoma, Two Entities or One Entity with Two Faces

Hongxing Gui¹, John Brooks², Paul Zhang³

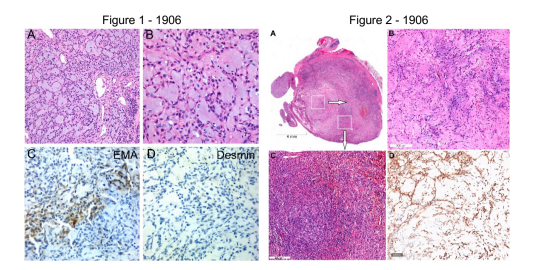
¹Philadelphia, PA, ²University of Pennsylvania Perelman School of Medicine, Swarthmore, PA, ³Hospital of the University of Pennsylvania, Media, PA

Disclosures: Hongxing Gui: None; John Brooks: None; Paul Zhang: None

Background: Primary pulmonary myxoid sarcoma (PPMS) is an exceedingly rare low-grade lung neoplasm characterized by reticular/lace-like growth of spindle to epithelioid cells embedded in an abundant myxoid matrix. It overlaps with myxoid variant of angiomatoid fibrous histiocytoma (AFH) morphologically. In terms of molecular genetics, they both have *EWSR1* gene rearrangements, with *EWSR1-CREB1* fusion in PPMS and either *EWSR1-ATF1* or *EWSR1-CREB1* fusion in AFH. It is unclear whether they are distinctive entities or the same entity with a spectrum of histomorphologies.

Design: We evaluated two cases of low-grade myxoid spindle cell tumor of the lung by histomorphology, immunohistochemistry and FISH studies.

Results: Case 1 was a 49-year-old man with a 3.8 cm right lower lobe mass extending into bronchus. Sections of the mass revealed proliferation of spindle cells in cords, strands and reticular patterns within an abundant myxoid stroma (Figure 1). The tumor cells were positive for EMA and negative for Desmin, AE1/3, Cam5.2, CK7, CK20, TTF-1, C-KIT, Vimentin, CD1a, p63, and S100. Case 2 was a 44-year-old man with left main bronchial mass. The periphery of the mass was partially encapsulated with a lymphoid cuff. The central portion contained juxtaposed myxoid and nodular components. The former was composed of spindle cells in myxoid matrix and the latter consisted of histiocytic and spindle cells in whorled and storiform patterns (Figure 2). The tumor cells from both areas were diffusely positive for Desmin and negative for EMA, PanCK, AE1/3, S100, SMA, CD34 and ALK1. FISH analysis demonstrated positive *EWSR1* gene rearrangements in both cases, showing *EWSR1-ATF1* fusion gene in case 1 and *EWSR1-CREB1* fusion in case 2.



Conclusions: We reported for the first time a case of PPMS with a novel *EWSR1-ATF1* translocation, which is usually common in AFH. The second case represented a hybrid of jaxtaposed PPMS and AFH components. These findings provide new evidence supporting that PPMS and myxoid AFH may represent a continuum with overlapping histologic, immunohistochemical and genetic features.

1907 PAX1 is a Sensitive and Specific Marker for Thymic Epithelial Neoplasms

Sarika Gupta¹, Sagar Vishal², Anthony Snow³, Andrew Bellizzi¹

¹University of Iowa Hospitals and Clinics, Iowa City, IA, ²Coralville, IA, ³North Liberty, IA

Disclosures: Sarika Gupta: None; Sagar Vishal: None; Anthony Snow: None; Andrew Bellizzi: None

Background: The differential diagnosis for epithelial tumors in the mediastinum includes carcinomas (CA) of the lung and thymus and thymoma. CD5 and KIT are often used to distinguish thymic (+) from lung (-) CA, but these markers are rarely positive in thymoma. Prior studies demonstrated frequent polyclonal PAX8-positivity in thymoma (≥90%) and thymic CA (67%). We recently switched to monoclonal PAX8, which has been shown to be negative in thymic tumors. Given prior experience, we hypothesized that polyclonal PAX8-positivity in the thymus represents cross-reactivity with another PAX-family transcription factor. PAX1 is normally expressed during thymic development, and a high-quality monoclonal antibody recently became commercially available.

Design: PAX1 immunohistochemistry (IHC) (clone 5A2) was performed on tissue microarrays of 79 thymomas, 10 thymic CAs, and 396 other CAs with an emphasis on differential considerations and PAX8-positive tumors: 175 squamous cell CAs (25 lung); 41 urothelial CAs; 36 lung adenoCAs; 28 renal cell, 25 serous, 22 endometrioid, 22 and 22 papillary and follicular thyroid, 10 breast, 7 colon, 5 esophagus, 2 poorly differentiated neuroendocrine, and 1 prostate CA. Intensity (0-3+) and extent (0-100%) of expression was evaluated, and an H-score was calculated. Fisher's exact and Mann Whitney tests were used with p<0.05 considered significant.

Results: PAX1 was expressed by 91% of thymic tumors, including 94% of thymomas (mean/median H-score 198/210) and 70% of thymic CAs (mean/median H-score 144/145). The differences in frequency (p=0.08) and H-score (p=0.19) were not significant. PAX1-positivity was noted in only 3.5% of non-thymic CAs at a mean (median) H-score of 15 (5) (both p<0.0001 compared to thymic tumors); these included 6 thyroid CAs (14%), 4 SCCs (2%) [including 2 lung (8%)], 3 urothelial CAs (7%), and 1 (10%) breast CA. Two of the PAX1-negative thymic CAs were KIT-positive; all 3 were CD5-negative. Detailed PAX1 expression data by thymoma type are presented in the Table; PAX1 was very frequently, strongly expressed across types.

Thymoma Type	% Positive	Mean (Median) H-score (if positive)
A (n=3)	100%	221 (200)
AB (n=15)	100%	199 (215)
B1 (n=19)	89%	190 (220)
B2 (n=22)	95%	191 (190)
B3 (n=10)	90%	217 (245)

Conclusions: PAX1 IHC using a novel monoclonal antibody is sensitive and specific for thymic epithelial neoplasms. Occasional weak positivity in thyroid tumors may represent low-level cross-reactivity with PAX8. This study exemplifies "next-generation IHC," which seeks to apply knowledge from developmental biology and molecular genetics to "intelligently design" novel IHC markers.

1908 Prevalence and Prognostic Significance of Tumor Spread through air spaces(STAS) in Resected Lung Cancers; Special Reference on Grading of STAS

Yeon Bi Han¹, Hyun Jung Kwon², Jeonghyo Lee³, Hyojin Kim⁴, Jin-Haeng Chung⁵
¹Seoul National University Bundang Hospital, Seongnam, Gyeonggi, Korea, Republic of South Korea, ²SNUBH, Seongnam-Si, Korea, Republic of South Korea, ³Seoul National University Bundang Hospital, Seongnam, Gyeonggi-do, Korea, Republic of South Korea, ⁴Seongnam Gyeonggi-do, Korea, Republic of South Korea, ⁵Seongnam, Korea, Republic of South Korea

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Background: Tumor spread through the air space (STAS) is an invasive pattern of lung cancer recently described. But there are some debates on its definition, quantification and clinical impact. In this study, we investigated the association between STAS grade and clinicopathological characteristics, as well prognostic impact in resected lung cancers.

Design: STAS has been prospectively described from 2008 and it was graded according to the distance from the edge of tumor margin as 2-tier system, I or II from 2011. Correlations between STAS grade and clinicopathologic characteristics and prognostic significance were analyzed in 2000 surgically resected lung cancers.

Results: Histologic subtypes of the 2000 cases were as follows; Adenocarcinoma (ADC)/Squamous cell carcinoma (SqCC)/Neuroendocrine tumors (NETs)/Others, 1544(77.2%)/325(16.3%)/57(2.9%)/74(3.7%). Among 2000 cases, STAS was observed in 829 cases (41.4%), and Grade (Gr) I/II, 471(23.5%)/358(17.9%). STAS was more frequently found in patients with NETs (p<0.001), pleural invasion (p<0.001), lymphovascular invasion (p<0.001), necrosis (p<0.001), radical resection (p<0.001) and higher pathologic stage (p<0.001). In STAS presence group, Gr II tumors more frequently showed these features than Gr I tumors. In ADCs, micropapillary (MP)-predominant ADCs showed highest prevalence of STAS, which was predominantly Gr II. On survival analysis, patients with higher grade STAS had worse Recurrence free survival (RFS) and Overall survival (OS) rates in total cohort. In stage IA non-mucinous ADC, multivariate analysis revealed that STAS grade stratified the prognosis for RFS in both of limited resection group and radical resection group (p=0.007, p<0.001, respectively). Moreover, the prognosis of patients with stage IA/STAS-Gr II was similar to that of patients with stage IB. (p=0.853)

Conclusions: STAS was more frequently found in NETs and MP-predominant ADCs. It was associated with well-known aggressive features, and STAS-Gr II tumors more frequently showed these features than Gr I tumors in ADCs. In stage IA non-mucinous ADC, multivariate analysis revealed that STAS grade was independent prognostic factor for RFS regardless of the extent of surgery. Moreover, comparable RFS rates were observed in patients with stage IA/STAS-Gr II and those with stage IB.

1909 Correlation of PDL1 Expression with Genomic Alterations in Non-Small Cell Lung Cancer: A 7-Month Retrospective Analysis

Susan Harley¹, Sophia Yohe¹, Robyn Kincaid¹, Matthew Schomaker¹, Bradley Johnson¹, Anup Tilak², Faqian Li¹, Andrew Nelson³, Stephen Michel¹, Pawel Mroz¹

¹University of Minnesota, Minneapolis, MN, ²Iowa City, IA, ³University of Minnesota, Saint Paul, MN

Disclosures: Susan Harley: None; Sophia Yohe: None; Robyn Kincaid: None; Matthew Schomaker: None; Bradley Johnson: None; Faqian Li: None; Andrew Nelson: None; Stephen Michel: None; Pawel Mroz: None

Background: PD1/PDL1 inhibitors have significantly improved overall survival in non-small cell lung cancer (NSCLC). PDL1 is now a standard biomarker along with molecular testing for therapy selection. Checkpoint therapy is given front-line for patients with high PDL1 and no actionable mutations or fusions, or after progression with any PDL1 expression. Studies comparing PDL1 status in the context of actionable mutations or fusions in NSCLC are sparse.

Design: All patients undergoing molecular testing for a next-generation sequencing (NGS) lung cancer panel at our institution in a 7 month period were reviewed. This included an NGS DNA panel for actionable/driver genes: EGFR, ERBB2, BRAF, K/NRAS, PIK3CA, and IDH1/2; and a NGS RNA fusion panel for ALK, ROS1, RET, NTRK1/3, NRG1, and MET e14 skipping. PDL1 expression by IHC was scored for tumor proportion score (TPS). Statistical comparison was performed using Fisher's exact test.

Results: 184 patients with NSCLC were included (138 adenocarcinoma, 29 squamous/adenosquamous, 14 poorly differentiated, and 3 NSCLC-NOS); 140 of those had concurrent PDL1 IHC. Most were tested at disease presentation; 5 had recurrent/progressive disease. 108 (59%) patients had a genomic alteration, most commonly RAS [22%], EGFR [12%], BRAF [4%], ALK [4%], and MET [3%]. Cases with RAS-driven tumors were compared to cases both with and without clinically relevant genomic alterations (Table 1). In both cases the prevalence of high PDL1 expression was greater in RAS-driven tumors: 50% vs 24% with actionable genomic alterations (p=0.02), and 50% vs 28% genomically negative tumors (p=0.02). The rate of any PDL1 positivity was not significantly different between these categories (p=0.78 and 0.77, respectively). The genomically actionable group also had a higher rate of PDL1 positivity than the genomically negative cases (83% vs. 60%; p=0.02), though the prevalence of high PDL1 positives was not different (24% vs. 28%; p=0.65).

Comparison of PDL1 Expression in RAS-driven NSCLC with NCCN Guideline Actionable					
	and Negative NSCLC				
	RAS-positive tumors (n=38)	Genomically actionable tumors*(n=42)	Genomically negative tumors (n=60)		
	% (n)				
		% (n)	% (n)		
PDL1 negative	21% (8)	16% (7)	40% (24)		
(TPS<1%)					
PDL1 positive (TPS>=1%)	79% (30)	83% (35)	60% (36)		
PDL1 high (TPS>=50%)	50% (19)	24% (10)	28% (17)		

TPS=tumor proportion score. Note that PDL1 high cases are also counted in the PDL1 positive category.

Conclusions: Our data indicate that at least half of RAS-driven NSCLC have high PDL1 expression, an important consideration for front-line therapy selection. The majority of genomically-actionable tumors have at least low-positive expression of PDL1, highlighting the potential importance of immune checkpoint therapy in the resistant-progression setting. Less than one third of genomically-negative cases have high PDL1 expression, and a significant percent are completely negative; additional innovations are needed to better tailor therapy for this patient subset.

1910 ROS1 RNA In Situ Hybridization (RNA-ISH) for Detecting ROS1 Rearrangements in Lung Adenocarcinoma

Timothy Helland¹, Azfar Neyaz², Vikram Deshpande¹, Yin Hung¹

¹Massachusetts General Hospital, Boston, MA, ²Massachusetts General Hospital, Malden, MA

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Background: Identification of ROS1 rearrangements in advanced lung cancer carries therapeutic implications, given the available ROS1-targeted therapy, but can be technically challenging. Immunohistochemistry (IHC) for ROS1 show expression in ROS1-rearranged tumors, but staining may be weak in some cases and seen in reactive pneumocytes. Fluorescence in situ hybridization (FISH) using break-apart probes may be difficult to interpret in cases with subtle intrachromosomal rearrangements. Next-generation sequencing (NGS) can be useful but requires sufficient tissue and a turnaround time of at least 1-2 weeks. Given that these current methods may have drawbacks, this study explores the utility of RNA in situ hybridization (RNA-ISH) in detecting ROS1 rearrangements in lung adenocarcinomas.

Design: This cohort included 7 lung adenocarcinomas with ROS1 rearrangements confirmed by both IHC and RNA-based anchored-multiplex NGS (with the following fusion partners: CD74 [n=2], EZR [n=2], MSN [n=1], SDC4 [n=1], and SLC34A2 [n=1]). The controls included 8 tumors confirmed to lack ROS1 rearrangements. RNA-ISH assay was performed using custom-designed ROS1 probes with branched DNA technology on an automated platform. The average number of ISH signal dots was quantified in 25 cells in each tumor using a standard bright-field microscope.

Results: Using the ROS1 RNA-ISH, all seven (100%) genetically-confirmed ROS1-rearranged lung adenocarcinomas showed positivity; whereas none of the controls (0%) was positive. The fraction of cells in ROS1-rearranged lung adenocarcinoma showing positivity ranged 60-90% (median 80%). The average number of ISH signal dots was far higher in confirmed ROS1-rearranged lung adenocarcinomas than controls (average: 29.6 per cell vs. 0.058 per cell). Background lung showed minimal ISH signal (average: 0.71 per cell). Also, the positive ISH signals were easily observed and could be seen using 10x objective in all cases or with a 4x objective in 6 of 7 cases.

Conclusions: The automated ROS1 RNA-ISH assay appears sensitive and specific in identifying ROS1 rearrangements in lung adenocarcinoma, with ease of use for minimally-trained eyes. ROS1 RNA-ISH may be a quick orthogonal tool in confirming ROS1 rearrangements in clinically problematic cases. Nonetheless, systematic comparison on its performance with that of ROS1 IHC and FISH may be warranted.

^{*}Tumors with one of the following: ALK fusion, NRG1 fusion, MET exon 14 skipping, EGFR mutation, or mutation in ERBB2, BRAF, RET, PIK3CA, or IDH1/2

1911 Clinicopathologic Features of Mismatch Repair-Deficient Diffuse Malignant Pleural Mesotheliomas Yin Hung¹, Corinne Gustafson², Assunta De Rienzo², Marina Kem¹, Mari Mino-Kenudson¹, Lucian Chirieac³, Raphael Bueno² 1 Massachusetts General Hospital, Boston, MA, 2 Brigham and Women's Hospital, Boston, MA, 3 Brigham and Women's Hospital, Harvard Medical School, Boston, MA

Disclosures: Yin Hung: None; Corinne Gustafson: None; Assunta De Rienzo: None; Marina Kem: None; Mari Mino-Kenudson: None; Lucian Chirieac: None; Raphael Bueno: *Grant or Research Support*, Roche, Genentech, Siemens, Merck, Gristone, Epizyme, Verastem; *Speaker*, Johnson and Johnson, IASLC, AATS, The Alliance for Clinical Trials; *Stock Ownership*, Navigation Sciences

Background: Diffuse malignant mesothelioma is a rare and highly aggressive tumor arising from the serosal lining, most commonly the pleura. Mismatch repair (MMR) deficiency has been described in diverse tumor types but is generally considered to be rare in mesothelioma. Given that MMR-deficient tumors are eligible for treatment with immune checkpoint blockade, this study aimed to identify mismatch repair-deficient malignant mesotheliomas and characterize their clinicopathologic features.

Design: Immunohistochemistry (IHC) for MMR proteins (MLH1, PMS2, MSH2, and MSH6) was performed in 2 institutional cohorts of malignant mesothelioma using formalin-fixed paraffin-embedded sections from tissue microarrays: cohort #1: 898 cases (each triplicate 0.7 mm-cores) and cohort #2: 65 cases (each duplicate 2 mm-cores). Candidates identified by IHC on tissue microarrays were re-tested on whole-tissue sections to confirm expression loss. Next-generation DNA sequencing was performed in one MMR-deficient tumor.

Results: Of 963 patients with malignant mesotheliomas (81% men; age 28-84; 64% epithelioid), mismatch repair deficiency was identified in 7 (0.7%) patients (4 men, 3 women; age 58-73 [median 71] years). The 7 MMR-deficient diffuse malignant pleural mesotheliomas included 3 epithelioid-, 3 biphasic-, and 1 sarcomatoid-type histology. PD-L1 IHC showed membranous staining in <1% of tumor cells in 5 tumors, 1-10% in 1 tumor, and >50% in 1 tumor. The MMR IHC patterns included isolated MSH6 loss in 1 tumor, loss of both MSH2 and MSH6 in 3 tumors, and loss of both MLH1 and PMS2 in 3 tumors (including 1 tumor with sub-clonal near-total loss of MLH1/PMS2, aside from focal intact expression). Sequencing in another tumor with MLH1/PMS2 loss identified mutation in *BAP1* and deletions of *NF2* and *SETD2*, but no mutations in MMR proteins. Of the 5 patients with lymph node sampled, 2 harbored metastatic disease. Of the 5 patients with available follow-up, all died at 3-55 (median 11) months after diagnosis.

Conclusions: MMR-deficient diffuse malignant pleural mesothelioma is rare (accounting for <1% of cases) and shows no apparent gender predilection, diverse histologic patterns, and a dismal prognosis. MMR loss identified included isolated MSH6, both MSH2/MSH6, and both MLH1/PMS2. The presence of sub-clonal near-total MLH1/PMS2 loss in one tumor and the lack of MMR mutations detected in another tumor may suggest a role of non-germline mechanisms leading to MMR inactivation in mesotheliomas.

1912 Toward a Comprehensive Prognostic Model for Malignant Peritoneal Mesothelioma: An International Collaboration

Aliya Husain¹, Jefree Schulte¹, Richard Attanoos², Luka Brcic³, Kelly Butnor⁴, Lucian Chirieac⁵, Andrew Churg⁶, Kenzo Hiroshima⁷, Yin Hung⁸, Hedy Kindler¹, Thomas Krausz⁹, Alberto Marchevsky¹⁰, Kazuki Nabeshima¹¹, Ann Walts¹², David Chapel¹³
¹The University of Chicago, Chicago, IL, ²Cardiff University, Cardiff, Wales, United Kingdom, ³Medical University of Graz, Graz, Austria, ⁴The University of Vermont Medical Center, Burlington, VT, ⁵Brigham and Women's Hospital, Harvard Medical School, Boston, MA, ⁶University of British Columbia/Vancouver General Hospital, Vancouver, BC, ⁷Chiba University Graduate School of Medicine, Chiba, Chiba, Japan, ⁸Massachusetts General Hospital, Boston, MA, ⁹University of Chicago Hospital, Chicago, IL, ¹⁰Cedars-Sinai Medical Center, West Hollywood, CA, ¹¹Fukuoka University School of Medicine and Hospital, Jonan-ku, Fukuoka, Japan, ¹²Cedars-Sinai Medical Center, Los Angeles, CA, ¹³Brigham and Women's Hospital, Boston, MA

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Background: Malignant peritoneal mesothelioma (MPeM) accounts for 10-15% of mesothelioma diagnoses, but comprehensive prognostic models for MPeM are lacking, with current literature largely limited to single-institution studies examining one to few prognostic variables. We report the results of an international collaboration to develop a clinicopathologic prognostic model for MPeM.

Design: Clinical data and H&E-stained slides from 207 MPeMs were collected from nine academic medical centers. Central histomorphologic review was performed. Univariate (log-rank) and multivariate (Cox proportional hazards regression) analyses of overall and progression-free survival (OS, PFS) were performed.

Results: The 207 MPeMs included 117 men and 90 women. Median age at diagnosis was 62 years (range 16-87). 134 patients died over follow-up of 1-166 months, with median OS of 25 months and 21% alive 96 months after diagnosis. Epithelioid, biphasic, and sarcomatoid histotypes were seen in 172, 26, and 8 cases, respectively.

On univariate analysis of all MPeMs, shorter OS was significantly associated with year of diagnosis, asbestos exposure, poorer performance status, lymph node metastasis, higher peritoneal disease burden, absence of cytoreduction and hyperthermic intraperitoneal chemotherapy, biphasic or sarcomatoid histotype, and tumor necrosis. On univariate analysis of epithelioid MPeM only, shorter OS was additionally associated with nuclear pleomorphism, higher mitotic rate, higher composite nuclear grade, and non-tubulopapillary architecture. On multivariate analysis, sarcomatoid and biphasic histotypes predicted shorter OS when adjusted for sex, asbestos exposure, and year of diagnosis. On multivariate analysis of epithelioid MPeM only, nuclear grade and non-tubulopapillary growth were independently predictive of shorter OS when adjusted for sex, year of diagnosis, and tumor necrosis.

Among patients with epithelioid MPeM, shorter PFS after cytoreduction was associated with lymph node metastasis, tumor necrosis, nuclear pleomorphism, higher mitotic rate, higher composite nuclear grade, and solid growth. On multivariate analysis, nuclear grade and solid growth were independently predictive of shorter PFS when adjusted for year of diagnosis, sex, and necrosis.

	Univariate Analyses of Overall Survival			
	All	Malignant Mesothelioma		Epithelioid Only
	(n=207)			(n=172)
	P value	Median OS (months)	P value	Median OS (months)
Year of Diagnosis	<0.0001	Wiedlan CC (Mentile)	<0.0001	modan oo (monalo)
After 2000	10.0001	36	10.0001	37
Before 2000		8		13
Sex	0.064	<u> </u>	0.11	19
Female	0.004	32	0.11	37
		15		
Male Asbestos	0.018	15	0.22	19
	0.016	4.4	0.22	40
Exposed		14 36		18
Not exposed	0.0000	36	0.0007	36
ECOG Performance Status	0.0006		0.0027	75
0 or 1		75		75
2 or 3	0.40	18.5		21
Synchronous pleural mesothelioma	0.46		0.5	
No		44		44
Yes		39		39
Lymph node metastasis	0.043		0.033	
No		64		76
Yes		39		39
Cytoreduction	<0.0001		<0.0001	
No		9		13
Yes		51		51
Peritoneal Carcinomatosis Index Group	0.02		0.023	
0-13		112		112
14-26		36		36
27-39		34		34
HIPEC	<0.0001		<0.0001	<u> </u>
No	10.0001	11	10.0001	14
Yes		62		62
Histotype	<0.0001	02		02
EMM	<u> </u>	30		
BMM		6		
SMM				
	0.000	4	0.040	
Necrosis	0.003	0.4	0.043	
No		34		36
Yes		9		15
Nuclear Pleomorphism Score			<0.0001	
1				44
2				34
3				12
Mitotic Rate (Score)			0.05	
0-1 per 10 hpf (Score 1)				36
2-4 per 10 hpf (Score 2)				32
>4 per 10 hpf (Score 3)				16
Composite Nuclear Grade Score			<0.0001	
2-3				40
4-5				28
6				8
Solid Architecture			0.8	
No No			0.0	30

Yes				29
Tubulopapillary Architecture			0.009	
No				19
Yes				62
		Multivariate Analyses o	f Overall Su	ırvival
		All Malignant Mesothelioma		EMM Only
	P value	Hazard Ratio	P value	Hazard Ratio
Diagnosis before 2000	<0.0001	2.8 (1.7-4.5)	0.0009	3.2 (1.7-5.9)
Sex	0.23		0.76	
Prior asbestos exposure	0.08	1.6 (0.95-2.5)		
Histotype	<0.0001	BMM vs EMM: 3.1 (1.7-5.9)		
		SMM vs EMM: 8.9 (3.2-25)		
		SMM vs BMM: 2.9 (1.1-7.9)		
Necrosis			0.38	
Composite Nuclear Grade Score			0.0001	3 vs 1: 5.8 (2.5-13.3)
				3 vs 2: 5.0 (2.1-11.7)
				2 vs 1: 1.2 (0.64-2.1)
Non-tubulopapillary Architecture			0.01	2.9 (1.5-5.5)

HIPEC: Hyperthermic intraperitoneal chemotherapy

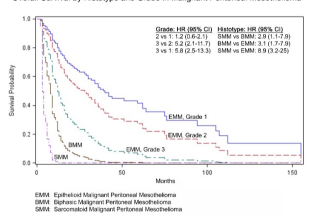
EMM: Epithelioid malignant mesothelioma

BMM: Biphasic malignant mesothelioma

SMM: Sarcomatoid malignant mesothelioma

Figure 1 - 1912

Overall Survival by Histotype and Grade in Malignant Peritoneal Mesothelioma



Conclusions: Prognostic histopathologic features as previously published for pleural mesothelioma overlap considerably with MPeM. Routine application of a standardized grading system to MPeM would provide useful clinical information.

1913 Pathologic Findings in Vaping-Induced Lung Injury: A Study of Four Patients

Peter Illei¹, Ankit Rajgariah²

¹Johns Hopkins University School of Medicine, Baltimore, MD, ²The Johns Hopkins University School of Medicine, Baltimore, MD

Disclosures: Peter Illei: None; Ankit Rajgariah: None

Background: There is emerging evidence that vaping can result in significant lung injury, the severity of which can be variable with few cases resulting in the patient's death also been reported. There is limited data on the pathologic findings in vaping-induced lung injury. Here, we report histologic and cytologic findings of four patients who developed lung disease following vaping.

Design: Review of pathologic material and clinical information of four patients with a history of vaping (6 specimens, 3 male and 1 female patient, age 16-37 years old).

Results: We have identified 4 patients with a history of vaping who presented to the hospital with severe respiratory dysfunction. Three of the patients are teenagers, while one is a younger adult and all of them required admission to the ICU. The three younger patients had bronchoscopy with bronchoalveolar lavage (BAL) with transbronchial biopsy in two. BAL showed lipid laden macrophages in all three with acute inflammation in one. Transbronchial biopsy showed intraalveolar fibrin with acute inflammation and organization in one of the biopsies. The adult patient is a known chronic alcohol and drug abuser who presented to the ED with dry cough and had subsequent wedge resection showing organizing pneumonia and chronic interstitial changes with the organizing pneumonia attributed to vaping.

Table 1. Summary of demographic information, clinical information, and imaging and pathologic findings of the fo	our patients.
--	---------------

Patient	1	2	3	4
Sex	M	M	M	F
Age	16	17	17	36
ICU Stay	Yes	Yes	Yes	Yes
Inhalant	THC	THC	Nicotene	Black market vaping product
CT scan	GGO, consolidations R > L	Bilateral consolidations	Bilateral infiltrates	Bilateral confluent GGO
Pathology	BAL: lipid laden macrophages	BAL: Lipid laden macrophages TBBX: Intraalveolar fibrin and reactive pneumocytes	BAL: Lipid laden macrophages, Acute inflammation TBBX: AFOP	Organizing pneumonia in a background interstitial lymphohistiocytic infiltrates

THC: Tetrahydrocannabinol; GGO: Ground Glass Opacity; AFOP: Acute Fibrinous and Organizing Pneumonia and acute hypxemic

Conclusions: Patients presenting with severe respiratory dysfunction following vaping have similar pathologic findings including lipid laden macrophages in BAL and intraalveolar fibrin with or without organization.

1914 VEGFR Family and Vasohibin in Lymphangioleiomyomatosis (LAM) Cells Associated with its Early Onset and Progression

Chihiro Inoue¹, Yasuhiro Miki², Ryoko Saito³, Kazuma Kobayashi⁴, Yoshinori Okada⁵, Hironobu Sasano⁶
¹Tohoku University Hospital, Sendai, Miyagi, Japan, ²Disaster Ob/Gyn, Int Res Inst of Disaster Sci, Tohoku Univ, Sendai, Miyagi, Japan, ³Tohoku University Graduate School of Medicine, Sendai-Shi Aoba-Ku, Miyagi-ken, Japan, ⁴Department of Thoracic Surgery, IDAC, Tohoku University, Sendai, Miyagi, Japan, ⁵Institute of Development, Aging and Cancer, Tohoku University, Sendai, Miyagi Prefecture, Japan, ⁶Tohoku University, Sendai-shi, Miyagi-ken, Japan

Disclosures: Chihiro Inoue: None; Yasuhiro Miki: None; Ryoko Saito: None; Kazuma Kobayashi: None; Yoshinori Okada: None; Hironobu Sasano: None

Background: Lymphangioleiyomomatosis (LAM) is a rare low-grade neoplasm associated with widespread interstitial infiltration of spindle cells and subsequent cystic changes of the lesions. LAM is also uniformly distributed in the lungs. Target therapies such as mTOR inhibitor have been to manage LAM but there are no curative therapies at this juncture. LAM cells are known to produce VEGF-C and VEGF-D with their receptor, VEGFR3, which promote proliferation of LAM cells also present in tumor cells. However, roles of other angiogenic factors have remained unclear. Therefore, in this study, we examined the expression of angiogenic factors such as VEGFR family and vasohibin (VASH) and examined the correlations between these angiogenic factors and histological and clinical findings.

Design: 36 LAM cases were obtained from 32 patients who underwent lung transplantation in Tohoku University Hospital from 2006 to 2018. We performed hierarchical clustering analysis to classify the cases based on the results of VEGFR1, VEGFR2, VEGFR3, VASH-1, and VASH-2 immunoreactivity in LAM cells. We also immunolocalized VASH-1/2, CD31, and D2-40 in microvessels of the lesions.

Results: One of these clusters harbored higher VEGFR1/3 and lower VEGFR2 and VASH-1/2, and the patients in this cluster clinically manifested symptom much older and higher P/F ratio was detected at the time of lung transplantation than those in the cluster with higher expression of all of the factors above. The cluster with higher VEGFR1/2/3 expression and lower VASH-1/-2 expression also demonstrated

significantly higher VASH-1/CD31 and VASH-2/CD31 ratios in microvessels than the cluster with lower expression of all these factors above. However, there were no significant differences of lymphatic vessel strength or other histological characteristics detected in our present clustering analysis.

Conclusions: Angiogenic factors such as VEGFR2 and VASH-1/-2 influenced on an early onset and progression of the clinical symptoms of LAM. In addition, LAM cells expressing VEGFR1/2/3 promoted angiogenesis. Therefore, not only VEGF-C/D-VEGFR3 axis but also other angiogenic factors may also enhance LAM progression.

1915 The Roles of Whole Block Imaging with Micro-Computed Tomography in Lung Adenocarcinoma

Takashi Inoue¹, Alexei Teplov², Natasha Rekhtman², William Travis², Yukako Yagi²

¹Memorial Sloan Kettering Cancer Center, Shunan, Yamaguchi, Japan, ²Memorial Sloan Kettering Cancer Center, New York, NY

Disclosures: Takashi Inoue: None; Alexei Teplov: None; Natasha Rekhtman: None; William Travis: None; Yukako Yaqi: None

Background: Micro-computed tomography (Micro-CT), is a non-invasive method which allows 3-dimensional morphometric analysis of tissues in formalin fixed paraffin-embedded (FFPE) tissue blocks without any sectioning or loss of sample. Lung adenocarcinoma has a major five tissue patterns according to the 2015 WHO classification (lepidic, acinar, papillary, solid and micropapillary). Each tissue pattern has different prognostic indicators and outcomes. In practice, pathologists have to diagnose a predominant tissue pattern and measure each percentage of tissue pattern. The aim of this study is to analyze the structure of lung adenocarcinoma tissue patterns using Micro-CT images from FFPE tissue blocks. The FFPE blocks were then sectioned, stained with hematoxylin-eosin (H&E) and scanned to create whole slide images for comparison.

Design: FFPE tissue blocks from five lung adenocarcinoma cases were scanned using a custom-built Micro-CT scanner (Nikon Metrology) and digitally re-constructed for visualization and analysis using a digital image system. All H&E slides were scanned with 0.5um/pixel by NanoZoomer S60 (Hamamatsu Photonics, Japan) Whole Slide Imaging Scanner. We then investigated features of each tissue pattern and correlate between Micro-CT images and whole slide images as well as histology 3D.

Results: Micro-CT of FFPE blocks highlighted the structure of lung adenocarcinoma (Figure.1) and normal lung tissue (Figure.2) in 3D images. We can detect lung adenocarcinoma in a Micro-CT image, and detect the tissue pattern, such as lepidic, acinar, papillary, solid, and micropapillary pattern as well as spread through air spaces (STAS) (Figure 1). We could detect the co-relation of normal lung tissue (bronchus, alveolar wall, pulmonary vessels, and visceral pleura) on H&E slides and micro-CT images (Figure 2).

Figure 1 - 1915

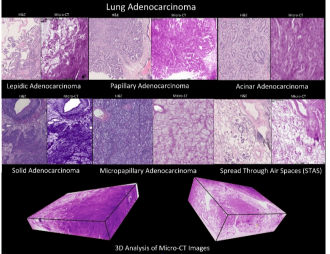
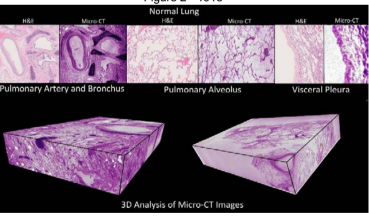


Figure 2 - 1915



Conclusions: Whole block imaging by Micro-CT allows for the identification of lung adenocarcinoma tissue patterns in FFPE blocks in a non-invasive and non-destructive manner. Correlation between micro-CT images of FFPE blocks and H&E histology images suggests that there is potential for (1) detecting pathologic features without sectioning and staining of the tissue and (2) to measure the volume of each adenocarcinoma tissue pattern including the invasive component in a block accurately.

1916 High-Level Expression of miR-19a is Associated with Advanced Stage and Higher Histologic Grade in Patients with Lung Adenocarcinoma

Seungyun Jee¹, Jongmin Sim², Yeseul Kim³, Seongsik Bang¹, Seong-Eon Park¹, Kiseok Jang¹

¹Hanyang University Hospital, Seoul, Korea, Republic of South Korea, ²Gangman-Gu, Seoul, Seoul, Korea, Republic of South Korea, ³Dong-Jak Gu, Seoul, Korea, Republic of South Korea

Disclosures: Seungyun Jee: None; Jongmin Sim: None; Yeseul Kim: None; Seongsik Bang: None; Seong-Eon Park: None; Kiseok Jang: None

Background: In our previous study, loss of mitochondrial tumor suppressor 1, or microtubule-associated scaffold protein 1 (MTUS-1) expression in lung adenocarcinoma was significantly associated with worse clinical outcome (unpublished data), implying the role of MTUS-1 as a tumor suppressor. This study investigated the clinicopathological significance of candidate microRNAs that were expected to inhibit MTUS-1 gene expression in lung adenocarcinoma.

Design: Computational prediction tools (Diana Tools, Target San, and Pictar) were used to select microRNAs expected to inhibit MTUS-1 expression. Quantitative RT-PCT (qRT-PCR) was used to detect candidate microRNA expression, and immunohistochemical staining was used to assess MTUS-1 expression in lung adenocarcinoma tissue samples. The chi-square test was used to analyze the association between microRNA expression and MTUS-1 expression, and with clinicopathological features. Kaplan-Meier survival curve was used to evaluate the prognostic value of microRNA expression levels.

Results: Three microRNA candidates were selected by the computational prediction tools: miR-19a, miR-19b, and miR-183. High levels of miR-19a was significantly associated with lower MTUS-1 expression (p = 0.008), whereas miR-19b and miR-183 levels showed no associations (p = 0.60 and p = 0.38, respectively; p = 132). High miR-19a expression was also significantly associated with worse histologic grade (p = 0.007), lymph node metastasis (p = 0.044), and higher AJCC stage (p = 0.008). Low miR-19b expression was significantly associated with perineural invasion (p = 0.048), and high miR-19b expression significantly associated with lymph node metastasis (p = 0.044). High miR-183 expression significantly associated with pleural invasion (p = 0.001). Patients with high miR-19a expression showed a trend of shorter disease-free survival than patients with low miR-19a expression, although the difference was not statistically significant (p = 0.094). Patients with high miR-19b or miR-183 did not show significantly worse disease-free survival (p = 0.86 and p = 0.10, respectively).

Conclusions: These results show that high-level expression of miR-19a is associated with adverse clinicopathological features, such as histologic grade, lymph node metastasis, and stage, possibly through downregulation of MTUS-1, an emerging tumor suppressor gene.

1917 Micropapillary or Solid Subtype of Lung Adenocarcinoma is a Single Significant Predictive Factor for Recurrence and Death Regardless of Proportion

Ji Yun Jeong¹, Jae Hui Kim¹, Tae In Park², Man Hoon Han³, Shin Yup Lee⁴, Sunha Choi⁴

¹Department of Pathology, Kyungpook National University Chilgok Hospital, School of Medicine, Kyungpook National University, Daegu, Korea, Republic of South Korea, ²Department of Pathology, Kyungpook National University Hospital, School of Medicine, Kyungpook National University, Daegu, Korea, Republic of South Korea, ³Kyungpook National University Hospital, School of Medicine, Kyungpook National University, Daegu, Korea, Republic of South Korea, ⁴Lung Cancer Center, Kyungpook National University, Daegu, Korea, Republic of South Korea, School of Medicine, Kyungpook National University, Daegu, Korea, Republic of South Korea

Disclosures: Ji Yun Jeong: None; Jae Hui Kim: None; Man Hoon Han: None

Background: The International Association for the Study of Lung Cancer, the American Thoracic Society, and the European Respiratory Society (IASLC/ATS/ERS) proposed a new histologic classification of lung adenocarcinoma in 2011. Although Micropapillary or solid (MP/S) subtypes are known to be a poor prognostic factor, differences by proportion of MP/S subtypes and underlying biologic mechanism are not fully elucidated yet. The aim of this study was to investigate the clinical characteristics and outcomes of lung adenocarcinoma with MP/S subtypes in surgically treated patients through detailed analysis by proportion of MP/S subtypes.

Design: We retrospectively reviewed medical records and pathologic reports of 410 patients with lung adenocarcinoma who underwent curative resection. Clinical features and recurrence free survival were compared between groups with MP/S subtypes (MP/S+, n=203) and without MP/S subtypes (MP/S-, n=207).

Results: MP/S subtypes were present in 203 patients (49.5%), and 141 (34.4%) had \geq 5% of S/MP proportion. MP/S subtypes were more frequently observed in patients with male gender (57.6% vs. 42.0%, P=0.002), tumor > 3cm (62.3% vs. 37.4%, P<0.001), and positive lymph node metastasis (85.9 vs. 40.9%, P<0.001) compared to female gender, tumor ≤ 3cm, and negative lymph node metastasis, respectively. Recurrence and death were more frequent in MP/S+ group compared with MP/S- group (24.1% vs. 1.9%, P<0.001 and 5.4% vs. 1.4%, P=0.02). The tumor with MP/S proportion of < 5% (MP/S<5%) were more frequent in patients with tumor > 3 cm (34.8% vs. 14.3%, P<0.001) and positive lymph node metastasis (60.9% vs. 18.7%, P<0.001) compared to tumor ≤ 3cm and negative lymph node

metastasis, respectively, and more frequently associated with recurrence and death (22.6% vs. 1.9%, *P*<0.001 and 6.5% vs. 1.4%, *P*=0.05) compared with MP/S- group. Survival analysis indicated that MP/S+ and MP/S <5% were associated with shorter recurrence free survival compare with MP/S- (HR=9.1, 95% CI=3.1-26.8, *P*<0.001; and HR=10.1, 95% CI=2.9-35.5, *P*<0.001, respectively). MP/S+ and MP/S<5% were more powerful predictor of recurrence than T or N stage in multivariate analyses.

Conclusions: Even very small proportion of MP/S subtype component was a significant predictive factor for recurrence in surgically resected lung adenocarcinoma. Further investigation on the underlying biological mechanism of poor prognostic effect of MP/S subtype is warranted.

1918 Significance of Major Pathologic Response (MPR) as Pathological Assessment of Therapeutic Response for Neoadjuvant Therapy in Non-Small Cell Lung Carcinoma

Ryutaro Kawano¹, Junko Watanabe², Hidehito Horinouchi², Yuko Nakayama³, Shun-ichi Watanabe⁴, Yuichiro Ohe⁵, Noriko Motoi¹ National Cancer Center Hospital, Chuo, Tokyo, Japan, ²Department of Thoracic Oncology, National Cancer Center Hospital, Chuo, Tokyo, Japan, ³Department of Radiation Oncology, National Cancer Center Hospital, Chuo, Tokyo, Japan, ⁴National Cancer Center Hospital, ⁵National Cancer Center Hospital, Tokyo, Japan

Disclosures: Ryutaro Kawano: None; Hidehito Horinouchi: None; Yuichiro Ohe: None; Noriko Motoi: None

Background: Recent progress of therapeutic option of lung cancer expands neoadjuvant setting for advanced-stage patients. Pathological response to neoadjuvant therapy is highlighting as a surrogate endpoint for survival or recurrence. Major pathologic response (MPR) has been proposed as a surrogate marker that was defined as a 10% residual viable tumor following neoadjuvant therapy (Hellmann M.D. et al. Lancet Oncol. 2014). Here, we examined the pathological significance of MPR in resected lung cancer with neoadjuvant chemo-radio-therapy.

Design: Seventy-seven primary lung cancer patients who received neoadjuvant therapy prior to surgery were recruited from 2003 until 2016. Clinicopathological factors, including age, gender, clinical stage, treatment regimen, disease-free and overall survival, and histological classification, were collected from medical chart. Two pathologists reviewed the archived HE stained slides of all the cases to assess residual viable tumor cell proportion and MPR, according to the Hellmann's criteria. pCR definition was a complete loss of viable tumor cells. In discrepancy cases, consensus score was reached with discussion under reviewing slides. Cohen's kappa values of the judgment of MPR and pCR were calculated using SPSS (version26).

Results: Patients' characteristics of 77 cases were as following; median age was 60 years old (range 33-77); 67 male and ten female; 16/16/41/4 of clinical Stage I/II/II/VI; 21 chemotherapy, 52 chemoradiation and 4 radiotherapy; 52 adenocarcinomas, 18 squamous cell carcinomas and seven other types of histology. MPR was observed in 42 (55%), and pCR in 8(10%). The concordance rate of MPR and pCR assessment among two pathologists was high (96% and 96%). Inter-observer agreement was high in MPR (kappa 0.928, p<0.001) and pCR (kappa 0.825, p<0.001). The discrepancy of MPR/pCR was due to the different judgment of tumor bed area and atypical cells whether they are benign or malignant. Pathological findings of discrepancy cases had temporal and spatially heterogeneity of fibrosis, active inflammation with reactive stromal and epithelial cell changes. Survival analysis will be updated at the time of presentation.

Conclusions: Our results revealed high reproducibility of MPR, and higher incidence compared to pCR, indicating MPR as a useful method for pathological therapeutic response.

1919 p16-positive Senescent Foci are Specific for Usual Interstitial Pneumonia and Predictive of Reduced Overall Survival in Interstitial Lung Disease

Jonathan Keow¹, Matthew Cecchini², Christopher Howlett³, Mariamma Joseph⁴, Marco Mura¹

¹London Health Sciences Centre, London, ON, ²Mayo Clinic, Rochester, MN, ³London Health Sciences Centre, Western University, London, AB, ⁴London Health Sciences Centre, Western University, London, ON

Disclosures: Jonathan Keow: None; Matthew Cecchini: None; Christopher Howlett: None; Mariamma Joseph: None; Marco Mura: None

Background: Interstitial lung disease (ILD) encompasses a spectrum of conditions with distinct clinical and pathologic features. A subset of ILD has been linked to abnormal cellular senescence which can induce a pro-fibrotic senescence associated secretory phenotype that results in progressive pulmonary fibrosis. The senescence is mediated in part by activation of the CDK inhibitor p16 which can arrest the cell cycle and can be used to mark senescent cells. We aim to demonstrate that a distinct subset of ILD associated with a senescent phenotype can be identified by expression of p16.

Design: 70 cases of ILD diagnosed with surgical lung biopsy were identified between 2003 and 2018 at a large tertiary-care level hospital (Figure 1). Additional p16 staining (clone E6H4, Roche) was performed on representative sections with the most active fibrosis. P16 positive senescent foci were defined as a loose collection of p16-positive fibroblasts with an overlying p16-positive epithelium and scored

as p16-low (0-2 foci/slide) or p16-high (≥3 foci/slide). The diagnosis was verified with the original pathology report and outcome data by time of biopsy to time of death or lung transplant.

Results: The presence of any p16-positive senescent foci was highly specific (92%) for the diagnosis of usual interstitial pneumonia (UIP). In the UIP group there was variable expression of p16 with a range of senescent foci between 0 and 23 per slide with 30 (67%) cases expressing some level of p16 and 20 (44%) cases expressing high levels of p16. Comparing cases with high levels of p16 to cases with low to absent p16, there was a reduced survival (HR 2.26; 95% CI, 1.17 to 4.24; p = 0.016) in the p16 high group that was an independent predictor of lung transplant-free survival (Figure 2). In a sub-group analysis of only cases with a diagnosis of UIP, p16 status trended towards significance (HR 2.06; CI, 0.99 to 4.24)(Table 1).

Table 1. Predictors of lung transplant-free survival after surgical lung biopsy. Univariate analysis.

Cox proportional	Hazard	Confidence	p-value
hazard analysis	ratio	interval	
p16 expression	2.26	1.17-4.24	0.0160
Age (years)	1.00	0.98-1.03	n.s.
Gender (female)	0.75	0.55-1.02	n.s.
Pack-years	1.02	0.99-1.04	n.s.
FVC (% pred.)	0.98	0.96-1.01	n.s.
TLC (% pred.)	0.97	0.93-1.01	n.s.
DLCO (% pred.)	0.93	0.90-0.97	0.0003

Figure 1 - 1919

Variable Mean ± SD Diagnosis (UIP/NSIP/other) Age (vears) 38/33 (54%) Male/Female (% male) BMI (kg/m²) 30±6 Pack-years 23±19 FVC (% pred) 73+17 67±17 TLC (% pred) DLCO (% pred) 45±16 Alive/transplanted/died 20/11/39

Abbreviations: BMI = body mass index, FVC = forced vital capacity. TLC = total lung capacity. DLCO = diffusion lung capacity for carbon monoxide.

Figure 2 - 1919

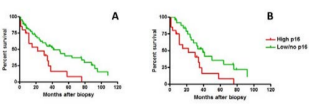


figure 2: p16 status predicts a distinct clinical outcome. Kaplan-Meer survival curves demonstrate that the presence of or more p16 positive foci per slide (High p16) correlate to decreased overall survival in all ILD cases (A) and in cases of UIP lone (B) when compared to cases with 0-2 p16 positive foci per slide (Lowino p16).

Conclusions: Increased levels of p16 positive foci are highly specific for the diagnosis of UIP and identify a subset of cases with a significantly worse transplant-free survival. These cases may represent an important subset of ILD that may be most effectively treated with emerging classes of drugs that target senescent cells.

1920 PD-L1 Expression and T-Cell Infiltrate Density and Distribution are Predictive Biomarkers of Response to Immune Checkpoint Inhibitors in Non-Small Cell Lung Cancer

Jong Kim¹, Corey Chang¹, Shimon Farber¹, Jianbo Song¹, David Engman¹, Jean Lopategui²

¹Cedars-Sinai Medical Center, Los Angeles, CA, ²Cedars-Sinai Medical Center, West Hollywood, CA

Disclosures: Jong Kim: None; Corey Chang: None; Shimon Farber: None; Jianbo Song: None; David Engman: None; David Engman: None; Jean Lopatequi: None

Background: Pembrolizumab is the first tissue agnostic FDA-approved therapy in microsatellite instability-high or mismatch repair deficient (dMMR) solid tumors. For NSCLC, FDA-approved pembrolizumab is indicated as a single agent for first-line treatment of advanced NSCLC that expresses PD-L1 (Tumor Proportion Score ≥ 1%). At our institution, we observed some patients with NSCLC expressing PD-L1 who didn't respond to the immune checkpoint inhibitors (ICI). It is not well-studied if additional biomarkers (i.e. MMR proteins, CD3/4/8 T-cells, regulatory T-cells, NK-cells, tumor-associated macrophages) are useful in predicting response to ICI in NSCLC. Here we evaluate potential biomarkers that may help better predict response to ICI.

Design: 16 cases of NSCLC patients who received ICI were identified from departmental surgical archive and response was retrieved from the electronic medical records, including 7 cases of responders and 9 cases of non-responders. Responders were defined as achieving a progression free response for more than 4 months. Immunohistochemistry (IHC) studies for PD-L1, CD3, CD4, CD8, CD21, CD34, CD56,

CD163, FOXP3, and MMR proteins (MMRp MLH1/MSH2, PMS2/PMS6) were performed. For CD3/4/8 IHC, the lymphocytic density (%) as well as the location (peripheral vs infiltrating) of positive T-cells were evaluated.

Results: All responders displayed a high PD-L1 expression (avg. of 68.8%, range 5% to 100%). The CD3/4/8 positive T-cells heavily infiltrated the tumor (avg. of 46.6%, range 30% to 80%). For non-responders, only 4 out of 9 cases displayed a lesser PD-L1 expression (avg. of 35.25%, range 1% to 70%). The CD3/4/8 positive T-cells were much less dense and infiltrative (<5%), and tended to concentrate at the periphery. The IHC results for CD56, FOXP3, CD21 were negative. CD163 and CD34 had a similar pattern in both responders and non-responders. MMRp showed intact MLH1, MSH2, PMS2, and PMS6 except for one non-responder.

Conclusions: In addition to high PD-L1 expression, the percentage and pattern of CD3/4/8 positive T-cells appear to be more predictive of ICI response in NSCLC than PD-L1 alone and may easily be implemented in routine testing. The tumor microenvironment may also play an important role in facilitating immunotherapy response. However, NK-cells (CD56), tumor-associated macrophages (CD163), regulatory T-cells (FOXP3), and MMRp were not predictive of ICI response in our pilot study. Additional studies are needed to further investigate our preliminary findings.

1921 The Usefulness of NF2 Fluorescence In Situ Hybridization in the Diagnosis of Mesothelioma in Tissue and Cytological Specimens

Yoshiaki Kinoshita¹, Makoto Hamasaki², Shinji Matsumoto³, Masayo Yoshimura¹, Ayuko Sato⁴, Tohru Tsujimura⁴, Akinori Iwasaki³, Kazuki Nabeshima⁵

¹Fukuoka University School of Medicine and Hospital, Fukuoka, Japan, ²Fukuoka University Hospital, Fukuoka, Japan, ³Fukuoka University Hospital and School of Medicine, Fukuoka, Japan, ⁴Hyogo College of Medicine, Nishinomiya, Hyogo, Japan, ⁵Fukuoka University School of Medicine and Hospital, Jonan-ku, Fukuoka, Japan

Disclosures: Yoshiaki Kinoshita: None; Makoto Hamasaki: None; Shinji Matsumoto: None; Masayo Yoshimura: None; Ayuko Sato: None; Tohru Tsujimura: None; Akinori Iwasaki: None; Kazuki Nabeshima: None

Background: *Neurofibromatosis type 2 (NF2)* is a tumor suppressor gene located on chromosome 22q12.2. Malignant pleural mesothelioma (MPM) is characterized by frequent genetic alterations in several genes including *NF2, cyclin-dependent kinase inhibitor 2A (CDKN2A)/p16* (found in the 9p21 locus), and *BRCA1 associated protein (BAP1)*. Recent studies using fluorescence *in situ* hybridization for *NF2* (*NF2* FISH) have identified hemizygous loss (chromosome 22 monosomy or hemizygous deletion) as the most common molecular alteration of *NF2*. However, the effectiveness of *NF2* FISH for differentiating MPM from reactive mesothelial hyperplasia (RMH) or reactive mesothelial cells (RMC) is unknown. In this study, we investigated whether *NF2* FISH, either alone or in a combination with other diagnostic assays [9p21 FISH, methylthioadenosine phosphorylase (MTAP) immunohistochemistry (IHC), and BAP1 IHC], is effective for distinguishing MPM from RMH or RMC in tissue specimens and cell blocks obtained from pleural effusions.

Design: We examined tissues from 53 MPM and 27 RMH cases and cell blocks from 22 MPM and 20 RMC cases. We used FISH to examine the deletion status of *NF*2 and 9p21 and IHC to determine the expression of MTAP and BAP1.

Results: Histologically, hemizygous *NF2* loss as detected by FISH showed 49.1 % sensitivity and 100% specificity in differentiating MPM from RMH. A combination of *NF2* FISH, 9p21 FISH, and BAP1 IHC yielded greater sensitivity (98.1%) than that detected for either assay alone (75.4% for 9p21 FISH, 67.9% for MTAP IHC, or 52.8% for BAP1 IHC). In cytological analysis, FISH detection of hemizygous *NF2* loss showed 50.0 % sensitivity and 100% specificity in differentiating MPM from RMC. A combination of *NF2* FISH, 9p21 FISH, and BAP1 IHC yielded greater sensitivity (100%) than that detected for either assay alone (59.1% for 9p21 FISH, 54.5% for MTAP IHC, or 83.3% for BAP1 IHC). The FISH determination of *NF2* status showed identical results between tissue samples and cell blocks in each case.

Conclusions: NF2 FISH alone or in combination with other diagnostic assays has application in differentiation of MPM from RMH or RMC.

1922 Pure High-Grade Fetal and Pure Enteric Adenocarcinomas of the Lungs Harbor Similar Genetic Alterations

Satsuki Kishikawa¹, Takuo Hayashi², Kazuya Takamochi¹, Yuka Yanai³, Ayako Ura⁴, Monami Kishi¹, Kieko Hara⁵, Tsuyoshi Saito¹, Kenji Suzuki¹, Takashi Yao⁶

¹Juntendo University, School of Medicine, Tokyo, Japan, ²Juntendo University Graduate School of Medicine, Tokyo, Japan, ³Juntendo University, Bunkyo-ku, Tokyo, Japan, ⁵Juntendo University, School of Medicine, Bunkyo-Ku, Tokyo, Japan, ⁶Juntendo University, Tokyo, Japan

Disclosures: Satsuki Kishikawa: None; Takuo Hayashi: None; Kazuya Takamochi: None; Yuka Yanai: None; Ayako Ura: None; Monami Kishi: None; Kieko Hara: None; Tsuyoshi Saito: None; Kenji Suzuki: None; Takashi Yao: None

Background: High-grade fetal adenocarcinoma (HFA) and enteric adenocarcinoma (EA) are both rare histopathological subtypes of lung adenocarcinoma. HFA and EA are occasionally combined with conventional-type lung adenocarcinoma, but pure types of both malignancies exist. The fact that the lungs and colon develop from the primitive striatum led us to speculate that these subtypes might share a common pathogenesis.

Design: Among the 2253 cases of primary lung adenocarcinomas reported in our hospital, we identified 4 and 5 pure (p) HFAs (0.18%) and EAs (0.22%), respectively. All pHFA tumors were high-grade adenocarcinomas with fetal lung morphology, necrosis, and immunopositivity for at least one of the following markers in addition to lacking morule formation: α-fetoprotein, SALL-4, or glypican-3. All pEA cases involved tumor cells that resembled colonic adenocarcinoma with no history of colorectal cancer. We evaluated the clinicopathological and molecular characteristics of these pHFA and pEA tumors. Next-generation sequencing was performed using the lon-Torrent Personal Genome Machine Platform and the lon Ampliseg Cancer Hotspot Panel v2.

Results: Both pHFA and pEA were associated with several characteristic clinicopathological features such as smoking exposure, high incidence of lymphovascular invasion, and frequent expression of CDX2 and HNF4α. Furthermore, nuclear accumulation of β-catenin was observed in 2 cases of pHFA (50%) and 3 cases of pEA (60%), which indicate activation of Wnt signaling. The most frequently mutated gene was *TP53* (3 pEAs), and other mutated genes that lead to the activation of Wnt signaling were *CTNNB1* (1 pHFA) and *APC* (1 pEA). An analysis of copy number variations revealed that *SMAD4* deletion was the most frequently detected mutation, regardless of Wnt activation (2 pHFAs; 2 pEAs). Additionally, amplifications of *FGFR3* were detected in 3 cases (1 EA; 2 pHFAs), which tend to be mutually exclusive with Wnt activation. Common mitogenic mutations in lung adenocarcinoma, such as *EGFR* and *KRAS*, were not detected.

Conclusions: pHFA and pEA have similar clinicopathological features and oncogenic alterations which included frequent association with the activation of Wnt signaling, amplifications of *FGFR3*, and *SMAD4* deletion. Recognition of these genetic subsets may help distinguish between lung adenocarcinoma with fetal and enteric morphology.

1923 Clinicopathological Characteristics of GNAS-Mutated Invasive Mucinous Adenocarcinoma of the Lung

Satsuki Kishikawa¹, Takuo Hayashi², Kazuya Takamochi¹, Taisei Kurihara¹, Kei Sano¹, Kieko Hara³, Yoshiyuki Suehara¹, Kenji Suzuki¹, Tsuyoshi Saito¹, Takashi Yao⁴

¹Juntendo University, School of Medicine, Tokyo, Japan, ²Juntendo University Graduate School of Medicine, Tokyo, Japan, ³Juntendo University, School of Medicine, Bunkyo-Ku, Tokyo, Japan, ⁴Juntendo University, Tokyo, Japan

Disclosures: Satsuki Kishikawa: None; Takuo Hayashi: None; Kazuya Takamochi: None; Taisei Kurihara: None; Kei Sano: None; Kieko Hara: None; Yoshiyuki Suehara: None; Kenji Suzuki: None; Tsuyoshi Saito: None; Takashi Yao: None

Background: *GNAS* hotspot mutations have been described in indolent and slow-growing mucinous epithelial neoplasms in several organs, such as the pancreas and appendix. Large genomic databases show that a subset of mucinous and non-mucinous lung adenocarcinomas harbor *GNAS* mutations. However, the clinicopathological impact of *GNAS* mutations on invasive mucinous adenocarcinoma of the lungs (IMA) is not fully determined.

Design: We evaluated the clinicopathological and molecular characteristics of IMAs with *GNAS* mutations in comparison with *GNAS* wild-type cases. We examined *EGFR*, *KRAS*, *GNAS*, and *TP53* mutations by PCR-direct sequencing in 80 IMAs. Subsequently, a NanoString-based screen for 90 tyrosine kinase fusions was performed for all IMAs with wild type *EGFR* and *KRAS*. Next-generation sequencing using the lon-Torrent Personal Genome Machine Platform and Ion Ampliseq Cancer Hotspot Panel v2 or RNA sequence were performed to confirm *GNAS* mutations and tyrosine kinase fusions. Mucin core proteins (MUC1, MUC2, MUC4, MUC5AC, and MUC6) and differentiation transcription factors, particularly differentiating on the basis of the cellular lineage (TTF-1, CDX-2, and HNF4α) were detected by immunohistochemical staining.

Results: Three of 80 IMAs (3.8%) harbored *GNAS* mutations (2 R201H and 1 R201C). Other mitogenic alterations including *KRAS* mutations (53 cases, 66.3%), *TP53* mutations (11 cases, 13.8%), *CD74–NRG1* fusions (2cases, 2.5%), and *MET* exon 14 skipping (1 case, 1.25%) were detected. Among 53 cases of *KRAS* mutations, G12V was the most frequent (42%), followed by G12D (34%) and G12C (13%). Neither *EGFR* mutations nor rearranged *ALK*, *ROS1*, *RET*, and *NTRK* were detected. All cases of *GNAS* mutations were found in women who were never or light smokers with wild-type *TP53*. Furthermore, *GNAS* R201H mutations co-occurred with *KRAS* G12D mutations in two cases. In comparison with *GNAS* wild-type cases, *GNAS*-mutated cases were significantly associated with the female sex (p<0.05) and immunopositivity for MUC4 (p<0.05). However, no significant differences were observed in other clinicopathological and immunohistochemical features, and progression-free and overall survival among both groups.

Conclusions: *GNAS*-mutated IMAs are rare, but frequently co-occur with *KRAS* G12D mutations and immunopositivity for MUC4, and are predominantly found in never- or light-smoking women; however, the prognostic impact of *GNAS* mutations in IMAs is unclear.

1924 Using Two MicroRNA Panels to Discriminate Subtypes of Lung Cancer in Sputum Specimens

Hui Kong¹, Jian Zhou², Shaohua Lu³

¹Fudan University Zhongshan Hospital, Shanghai, China, ²Department of Pulmonary Medicine, Fudan University Zhongshan Hospital, Shanghai, China, ³Department of Pathology, Fudan University Zhongshan Hospital, Shanghai, China

Disclosures: Hui Kong: None; Jian Zhou: None; Shaohua Lu: None

Background: Lung cancer is classified into small cell lung cancer (SCLC) and non–small cell lung cancer (NSCLC), which mainly contains adenocarcinoma (AC) and squamous cell carcinoma (SQ). Lung cancer subtyping plays an important role in choosing therapeutic schemes. Sputum is a kind of noninvasively accessible biologic fluids containing exfoliated airway epithelial cells. Although cytopathological examination of sputum is now available, its positive-rate of malignant cells is very low. It is hard to classify subtype via classic cytology on sputum specimens. We had reported that microRNA panels could accurately discriminate between three subtypes of lung cancer in bronchial brushing specimens. The diagnostic value of microRNAs in sputum specimens is need to be explored.

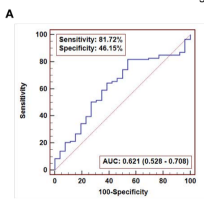
Design: In this study, 114 sputum specimens (49 AC, 39 SQ, and 26 SCLC) were investigated. Reverse-transcriptase quantitative polymerase chain reaction (RT-qPCR) was performed to evaluate expression of 7 candidate microRNAs discovered via microarrays previously. Two logistic regression models constructed before was validated in the cohort of 114 sputum specimens. The area under the receiver operating characteristic curve (AUC) was used to assess the diagnostic accuracy of microRNA panels. The diagnostic performance was compared between microRNA panels and cytology.

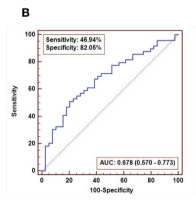
Results: Panel A, consisting of miR-29a and miR-375, was built to discriminate SCLC from NSCLC. In the cohort of 114 sputum specimens, the AUC value was 0.621 with sensitivity of 81.72% and specificity of 46.15%. Similarly, panel B, consisting of miR-34a and miR-205, was used to discriminate SQ from AC. In the cohort of 114 sputum specimens, the AUC value was 0.678 with sensitivity of 46.94% and specificity of 82.05%. Compared with cytology, microRNA panels or the combination of microRNA panels and cytology were of higher sensitivity and specificity in diagnosis of AC, SQ and SCLC.

Figure 1 - 1924

			,	- '-		
	Diagnosed SCLC			Diagnosed		
	Sensitivity	95% CI	P Value	Specificity	95% CI	P Value
Cytology	23.1% (6/26)	6.6% - 39.6%	0.092	19.3% (17/88)	10.4% - 26.2%	0.000
Panel A	50.0% (13/26)	30.4% - 69.6%		72.7% (64/88)	59.3% - 78.3%	
Combination	61.5% (16/26)	42.2% - 80.6%	0.250	75.0% (66/88)	61.7% - 80.3%	0.500
	Diagnosed SQ			Diagnosed AC		
	Sensitivity	95% CI	P Value	Specificity	95% CI	P Value
Cytology	12.8% (5/39)	2.2% - 23.4%	0.999	24.5% (12/49)	12.3% - 36.7%	0.000
Panel B	15.4% (6/39)	3.9% - 26.9%		95.9% (47/49)	90.3% - 100%	
Combination	25.6% (10/39)	11.7% - 39.5%	0.125	98.0% (48/49)	94.0% - 100%	0.999

Figure 2 - 1924





Panel A =-3.6303+2.5935*(miR-29a)-0.9431*(miR-375)

Panel B =2.8974+1.0175*(miR-34a)-1.1748*(miR-205)

Conclusions: In sputum specimens, those two microRNA panels for lung cancer subtype discrimination could achieve high sensitivity and specificity. Moreover, the combination of microRNA panels and cytology could improve the diagnostic accuracy in further. These findings could be helpful in therapy of lung cancer.

1925 Quantification of Diagnostic Tissue Obtained by Endoscopic Ultrasound-Guided Fine Needle Biopsy (EBUS/EUS) in Patients with Lymphoma

Vimal Krishnan¹, Gabriel Dayan², Moishe Liberman¹, Philippe Stephenson³

¹Centre Hospitalier de l'Université de Montréal, Montreal, QC, ²Université de Montreal, Montreal, QC, ³Toronto, AB

Disclosures: Vimal Krishnan: None; Gabriel Dayan: None

Background: Endoscopic ultrasound-guided fine needle biopsy (EBUS/EUS) as a diagnostic procedure for the diagnosis of lymphoproliferative disorders (LPDs) remains controversial. A recent meta-analysis evaluating 14 studies reported that the overall specificity of EBUS/EUS for LPDs was 99.3%, with an overall sensitivity of 66.2%. This rather high rate of false negatives remains unexplained. We sought to characterize and quantify the diagnostic tissue obtained by EBUS/EUS for LPD.

Design: We conducted a retrospective review of all patients for 2005 to 2019 that underwent EBUS/EUS (19G or 22G needle) for suspected mediastinal lymphadenopathy. Only patients with a final clinical diagnosis of lymphoma were included. For quantification, patients with a histology specimen (core needle biopsy or cell block) were selected (n=25). Quantification was performed using *Calopix* software (*TRIBVN healthcare*), on a single slide that contained the most lymphoid tissue by light microscopy evaluation. Diagnostic tissue was defined as any lymphoid tissue fragment with a minimal surface area of 0.01 mm². Acellular fragments and fragments with marked cautery artefact were excluded.

Results: Out of 459 patients that underwent EBUS/EUS for suspected mediastinal lymphadenopathy, 75 had a final diagnosis of LPD. Median age was 62 years old. Diagnostic categories were Hodgkin's lymphoma (n=26), low grade non-Hodgkin's lymphoma (n=33), high grade non-Hodgkin's lymphoma (n=11), T cell lymphoma (n=2) and unspecified B cell lymphoma (n=3). Eight patients had a false negative EBUS/EUS (Table). On image analysis, median total diagnostic area was 1.36 mm² and the median size of the largest fragment was 0.29 mm² (n=25). The largest fragment size was obtained for a case of diffuse large B-cell lymphoma (total area: 48.40 mm², Figure 1). There was no significant correlation between LPD diagnostic category and total fragment size (p=0.153) or largest fragment size (p=0.139). Preliminary data on surgical specimens obtained by mediastinoscopy show a total diagnostic area ranging from 50.86 mm² to 95.30 mm², with largest fragment size varying from 14.61 to 22.53 mm² (n=3).

Patient	EBUS/EUS	Final diagnosis on surgical biopsy
1	Negative	Diffuse large B cell lymphoma, NOS
2	Negative	Follicular lymphoma
3	Negative	Hodgkin's lymphoma
4	Negative	Hodgkin's lymphoma
5	Negative	Marginal zone lymphoma
6	Negative	Small lymphocytic lymphoma
7	Negative	Hodgkin's lymphoma
8	Negative	Diffuse large B cell lymphoma, NOS

Figure 1 - 1925

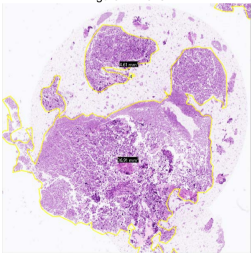


Figure 1. Quantification of diagnostic tissue for a patient with diffuse large B cell lymphoma.

Conclusions: We report here a series of lymphoma cases diagnosed by EBUS/EUS, ranging across several diagnostic categories, based primarily on small size fragments. Quantification of endoscopic diagnostic tissue provides insight on tissue yield, on its relationship with lymphoma subtype, and supports EBUS/EUS as an acceptable procedure for lymphoma diagnosis.

1926 Trichrome and Sulfated Alcian Blue Stains: Utility in Distinguishing Light Chain Deposition Disease (LCDD) from Amyloidosis in the Lung

Shajo Kunnath Velayudhan¹, Brandon Larsen², Govind Bhagat³, Dominick Santoriello¹, Shana Coley⁴, Thomas Colby², Anjali Sagi³

¹New York-Presbyterian/Columbia University Medical Center, New York, NY, ²Mayo Clinic, Scottsdale, AZ, ³Columbia University Medical Center, New York, NY, ⁴Columbia University, New York, NY

Disclosures: Shajo Kunnath Velayudhan: None; Brandon Larsen: None; Govind Bhagat: None; Dominick Santoriello: None; Shana Coley: None; Anjali Saqi: None

Background: Light chain deposition disease (LCDD) is characterized by amorphous "glassy" deposits of Immunoglobulin light chains in organs such as kidneys, heart, and liver. Since these amyloid-like deposits lack the secondary structure consisting of beta-pleated sheets, they don't stain salmon-pink with Congo Red. Additionally, a negative Congo Red may be attributed to suboptimal staining, especially when the deposits primarily involve organs where LCDD is rarely encountered. Moreover, Congo Red stains are compromised when the available sections are 5 micron thick. We noticed in a case of primary LCDD of the lung that the light chain deposits stained bright red with Masson trichrome and pink with sulfated alcian blue (SAB) stains. In the current study, we tested whether these two stains can distinguish between amyloidosis and LCDD.

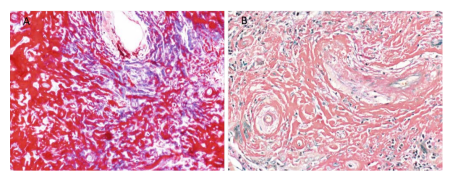
Design: We reviewed lung cases with eosinophilic deposits and immune infiltrates with negative Congo Red stain. We assessed trichrome and SAB staining on 9 cases of LCDD and 1 control case of amyloidosis involving the lung.

Results: 9 cases were identified (Table 1). The age of the patients ranged from 44 to 75 years and 56% were females. Imaging studies of 7 cases showed nodular deposits and the remaining cases showed cystic lesions and/or nodular lesions. Underlying hematologic abnormalities were detected in 7 LCDD patients which included MALT lymphomas (n = 6) and plasma cell neoplasm (n = 1). Systemic involvement was absent in 7 cases with available information. Hematoxylin and eosin stained slides of all cases showed "glassy" eosinophilic amyloid-like deposits predominantly around airways and vasculature. Congo Red staining was negative in all LCDD cases, while it showed salmon-pink staining of amyloid deposits in the control case. For all LCDD cases tested, trichrome stained the deposits as bright red and the SAB stained the deposits as pink (Figure 1). In contrast, amyloid deposits in the control case stained greyish blue with trichrome and bright green with SAB stains, as expected.

Patient No.	Age	Sex	Congo Red	Trichrome	SAB	Underlying disorder	SPEP	Radiology (diffuse, nodular, cystic)	lgH gene rearrangement	Light chain restriction (ISH)	Light chain restriction (pronase)
1	75	F	Neg	Red	Pink	MALT lymphoma	Lambda	Nodular	NP	NP	NP
2	61	F	Neg	Red	Pink	MALT lymphoma	Neg	Nodular	Polyclonal	Карра	Neg
3	54	М	Neg	Red	Pink	MALT lymphoma	Neg	Nodular	Clonal	Карра	Карра
4	67	М	Neg	Red	Pink	Unknown	NP	Nodular	NP	NP	NP
5	44	F	Neg	Red	Pink	Plasma cell neoplasm	Карра	Cystic	NP	Карра	NP
6	55	М	Neg	Red	Pink	MALT lymphoma	Neg	Cystic/Nodular	Clonal	Lambda	Lambda
7	56	F	Neg	Red+Blue	pink	MALT lymphoma	Unknown	Nodular	NP	kappa	kappa
8	58	М	Neg	Red	pink	Unknown	Unknown	Nodular	NP	kappa	Карра
9	55	F	Neg	Red	NP	MALT lymphoma	Unknown	Nodular	NP	kappa	kappa

Figure 1 - 1926

Figure 1. Trichrome (A) and SAB (B) staining of light chain deposits.



Conclusions: When Congo Red fails to stain "glassy" amorphous eosinophilic material salmon-pink, trichrome and SAB stains might indicate the non-amyloid, light chain nature of LCDD deposits when red and pink staining is seen, respectively. These screening studies can inform downstream testing such as immunofluorescence staining for light chains, transmission electron microscopy, and typing of light chains by mass spectrometry.

1927 Genetic Alterations of Pulmonary Pleomorphic Carcinoma and Their Metastatic Lesions Using Whole Exome Sequencing

Hyun Jung Kwon¹, Jin-Haeng Chung², Yeon Bi Han³, Hyojin Kim⁴, Sejoon Lee⁵, Jeonghyo Lee⁶
¹SNUBH, Seongnam-Si, Korea, Republic of South Korea, ²Seongnam, Korea, Republic of South Korea, ³Seoul National University Bundang Hospital, Seoul, Korea, Republic of South Korea, ⁴Seongnam Gyeonggi-do, Korea, Republic of South Korea, ⁵SNUBH, Seongnam-si, Gyeonggi-do, Korea, Republic of South Korea, ⁶Seoul National University Bundang Hospital, Seongnam, Gyeonggi-do, Korea, Republic of South Korea

Disclosures: Hyun Jung Kwon: None; Jin-Haeng Chung: None; Sejoon Lee: None; Jeonghyo Lee: None

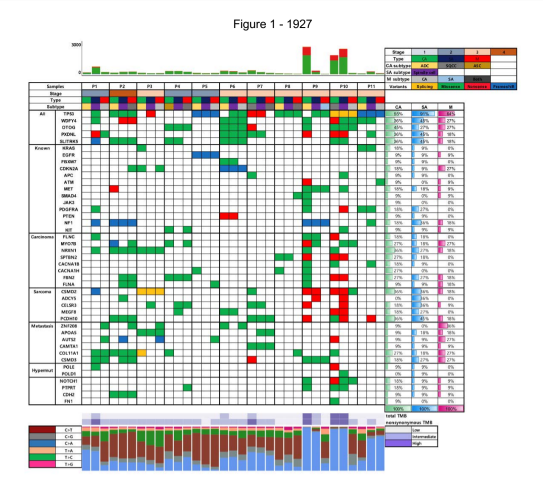
Background: Pulmonary pleomorphic carcinoma (PPC) is known for its aggressiveness and poor prognosis than other subtypes of non-small cell carcinoma. To better understand the molecular characteristics of PPC, we analyzed genetic alterations of PPCs and their metastatic lesions by whole exome sequencing.

Design: We included 11 PPC patients who underwent surgical resection for both primary lung cancer and metastatic lesions at Seoul National University Bundang Hospital. Carcinomatous and sarcomatous components of each primary PPC along with the metastatic lesions were microdissected. Somatic mutation profiles were generated by whole exome sequencing.

Results: The majority of PPC patients were male (10 of 11, 90.9%) and smokers (10 of 11, 90.9%). Carcinoma components of PPC consisted of adenocarcinoma (8 of 11, 72.7%), squamous cell carcinoma (2 of 11, 18.2%) or adenosquamous carcinoma (1 of 11, 9.1%). *TP53* (70%) was the most frequently recurrent genetic alteration, followed by *WDFY2* (36%), *OTOG*, *PXDNL*, and *SLITRK5* (33%). *KRAS* mutation was found in 2 cases (18.1%) and *EGFR* mutation in one (9.1%).

In addition, mutations discovered to be richer in either carcinomatous, sarcomatous or metastasis included various genes known to be associated with tumor progression and poor prognosis (FLNC, NRXN1, CSMD2, ZNF208, AUTS2). Gene alterations associated with somatic hypermutation (POLE, POLD1) and epithelial-mesenchymal transition (NOTCH1, CDH2, FN1) were also found. Four cases (36.4%) had high tumor mutation burden (TMB).

Of the 43 altered genes investigated in depth in this study, roughly half (n=18) were shared by carcinomatous, sarcomatous and metastasis in at least one case. Significant overlap of the mutation profiles between metastasis and either carcinomatous or sarcomatous component was not found.



Conclusions: This is the first study to analyze the molecular profile of PPC in both metastatic lesions and primary carcinomatous and sarcomatous components. Sporadic genetic alterations appeared to occur among carcinomatous, sarcomatous components and metastatic lesions of PPC. Various genes associated with tumor progression were altered in both primary and metastatic lesions.

1928 Histologic Characterization of Small Cell Transformation from EGFR-Mutant Non-Small Cell Lung Carcinoma after EGFR-Targeted Therapy

Christin Lepus¹, Julia Rotow², Pasi Janne², Lynette Sholl¹

¹Brigham and Women's Hospital, Boston, MA, ²Dana Farber Cancer Institute, Boston, MA

Disclosures: Christin Lepus: None; Julia Rotow: Consultant, AstraZeneca; Pasi Janne: Consultant, AstraZeneca; Consultant, Boehringer Ingelheim; Consultant, Roche/Genentech; Consultant, LOXO Oncology; Consultant, Eli Lilly; Lynette Sholl: Consultant, LOXO Oncology

Background: Histologic transformation of *EGFR*-mutant non-small cell lung carcinoma (NSCLC) to small cell lung carcinoma (SCLC) is a mechanism of acquired resistance to EGFR-tyrosine kinase inhibitors (TKIs) that occurs in approximately 5% of *EGFR*-mutant NSCLCs. However, the natural history/histologic progression of SCLC transformation from NSCLC is poorly understood. A retrospective analysis was conducted to characterize the morphologic variation during transformation of *EGFR*-mutant NSCLC to SCLC.

Design: We identified 12 patients with *EGFR*-mutant NSCLC that transformed to SCLC during treatment with EGFR-targeted tyrosine kinase inhibitor therapy. Histologic evaluation of longitudinal specimens (n=1-5 per patient) from both lung and distant metastases was performed to characterize the morphologic spectrum of lesions obtained from the initial diagnosis of NSCLC to SCLC transformation.

Results: All 12 patients had lung adenocarcinoma harboring *EGFR* mutations at initial diagnosis (exon 19 deletion, 91.6%; L858R, 8.4%) and had been treated with at least one EGFR TKI prior to development of SCLC. Most patients (10 of 12; 83.3%) were on second-line osimertinib at the time of transformation. When transformation was first documented, four patients (33.3%) were reported to have combined adenocarcinoma and SCLC or poorly differentiated carcinoma with mixed adenocarcinoma and SCLC morphologic features; the remaining 8 patients (66.6%) were reported to have SCLC. Retrospective review of this latter group demonstrated focal NSCLC-like morphology (mildly increased cytoplasm, variably prominent nucleoli, and rare gland formation) in 3 patients (37.5%). Conversely, 4 of

6 patients who had post-treatment, pre-transformation biopsies reported as adenocarcinoma showed classic adenocarcinoma architecture with superimposed small cell carcinoma-like cytomorphology, including increased nuclear-to-cytoplasmic ratio, nuclear hyperchromasia, and finely granular chromatin. Overall, a hybrid/transitional phenotype was captured in 7 of 12 patients (58.3%). Among 7 patients who had tumor genotyping upon SCLC transformation, all maintained the sensitizing *EGFR* mutation and showed acquired *RB* loss.

Conclusions: A spectrum of intermediate/transitional histologic phenotypes underly *EGFR*-mutant NSCLC transformation to SCLC. Recognition of these transitional morphologies may facilitate earlier detection of SCLC-driven resistance to EGFR-targeted therapy and inform timely selection of alternate treatment regimens.

1929 Predicting Specific Causes of Death in Lung Cancer Patients: Random Forest versus Multinomial Models

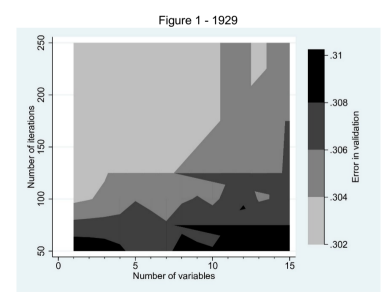
Yuan Li¹, Haijun (Steve) Zhou², Yong Lin³, Fei Deng⁴, John Heim⁵, Lanjing Zhang⁵
¹Fudan University Shanghai Cancer Center, Shanghai, China, ²Houston Methodist Hospital, Houston, TX, ³Department of Biostatistics, Rutgers School of Public Health, New Brunswick, NJ, ⁴School of Electrical and Electronic Engineering, Shanghai Institute of Technology,, Shanghai, China, ⁵Princeton Medical Center, Plainsboro, NJ

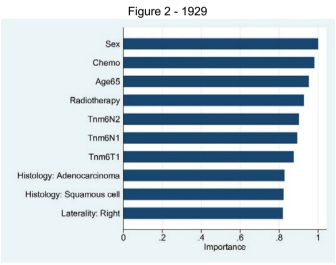
Disclosures: Yuan Li: None; Haijun (Steve) Zhou: None; Yong Lin: None; Lanjing Zhang: None

Background: Random forest model is a recently developed machine-learning algorithm, and superior to other machine learning and regression models for its classification function and better accuracy. But it is rarely used for predicting causes of death in cancer patients. On the other hand, specific causes of death in lung cancer patients are poorly classified or predicted, largely due to its categorical nature (versus binary death/survival).

Design: We therefore tuned and employed a random forest algorithm (Stata, version 15) to classify and predict specific causes of death in lung cancer patients, using the surveillance, epidemiology and end results-18 and several clinicopathological factors. The lung cancer diagnosed during 2004 were included for the completeness in their follow-up and death causes. The patients were randomly divided into training and validation sets (1:1 match). We also compared the accuracies of the final random forest and multinomial regression models.

Results: We identified and randomly selected 40,000 lung cancers for the analyses, including 20,000 cases for either set. The causes of death were, in descending ranking order, were lung cancer (72.45 %), other causes or alive (14.38%), non-lung cancer (6.87%), cardiovascular disease (5.35%), infection (0.95%), and. We found more 250 iterations and the 10 variables produced the best prediction, whose best accuracy was 69.8% (error-rate 30.2%, Figure 1). The final random forest model with 300 iteration and 10 variables reached an accuracy higher than that of multinomial regression model (69.72% vs 64.58%). The top-5 most important factors in the random-forest model were sex, chemotherapy status, age, radiotherapy status and nodal status (Figure 2).





Conclusions: We optimized a random forest model of machine learning to predict the specific cause of death in lung cancer patients using a population database. The model also appears more accurate than multinomial regression model.

1930 Clinicopathological Features of Thymic Epithelial Tumors with both Type B3 and C Characteristics Mei Liu¹. Yang Li². Zenghui Jiang³

¹First Medical Center of PLA General Hospital, Beijing, China, ²Chinese PLA General Hospital, Beijing, China, ³Qiqihaer Medical College, Qiqihaer, Heilongjiang, China

Disclosures: Mei Liu: None; Yang Li: None; Zenghui Jiang: None

Background: Most type B3 thymomas are easily differentiated from type C thymoma/thymic carcinoma. For the thymic epithelial tumors with both type B3 and C features (Borderlands Between Thymomas and thymic carcinomas), the fourth edition of WHO does not provide a detailed introduction. Only individual reports can be seen to describe this aspect.

Design: 807 cases of thymic epithelial tumors were reviewed in the Department of Pathology from June 2005 to June 2019. Thymic epithelial tumors with both type B3 and C characteristics were selected based on any of the following characteristics in addition to type B3 morphological characteristics: 1. The nests of tumor cells invasive with pushing borders within desmoplastic stroma. 2. Tumor cells express CD5 or CD117. 3. The expression of CD1a and TDT is absent in interstitial lymphocytes. Tumor staging was performed according to the Masaoka-Koga criteria.

Results: This group contains 56 cases of thymoma with type B3 components, 8 cases of thymic epithelial tumors with B3 and C characteristics and 84 cases of thymic carcinoma. The mean age of onset of patients with type B3 components was significantly lower than that of thymic epithelial tumors with both type B3 and C characteristics (P = 0.001). There was no significant difference in the proportion of each stage in the three kinds of thymic epithelial tumors (P=0.169). A total of 120 patients completed follow-up, and the follow-up period ranged from 1 month to 124 months. Survival analysis showed that the disease-specific survival of patients with thymomas containing B3 components was significantly higher than that of thymic carcinoma (P=0.004). Disease-specific survival of thymic epithelial tumors with both B3 and C characteristics were not significantly different from those of thymomas containing B3 components and thymic carcinomas with its survival curve closer to thymoma with B3 components. In the multivariate survival analysis, Masaoka-Koga staging was an independent prognostic factor.

Conclusions: The bland morphology and low proliferation index are the main indicators for *thymic epithelial tumors with B3 and C characteristics differentiated from thymic carcinoma. The survival of thymic epithelial tumors with both B3 and C characteristics is closer to that of B3 thymoma, suggesting that the malignancy of this tumor is closer to that of B3 thymoma, which can provide a basis for clinical selection of treatment.*

1931 Next Generation Sequencing in the Diagnostic Reevaluation of Non-Small Cell Lung Carcinomas: A Systematic Review

Ying-Chun Lo¹, Lynette Sholl¹, Fei Dong¹
¹Brigham and Women's Hospital, Boston, MA

Disclosures: Ying-Chun Lo: None; Lynette Sholl: Consultant, LOXO Oncology; Fei Dong: None

Background: Non-small cell carcinomas may exhibit diverse histological features and varying degrees of differentiation. The heterogeneity of histology findings can complicate the distinction of primary lung carcinoma from metastasis. Panel next generation sequencing is now widely available for treatment selection in lung cancers. In this study, we review a cohort of clinically suspected primary lung cancers that have undergone sequencing to evaluate molecular evidence of tumor origin.

Design: We systematically reviewed a cohort of 907 consecutive cancers that have undergone clinical next generation sequencing for the indication of non-small cell lung cancer. Molecular findings were reviewed and compared with clinical history and surgical pathology reports.

Results: 9 of 907 cases (1%) were found to have convincing sequencing evidence to support a diagnosis other than primary lung cancer. 4 cases had ultraviolet radiation-associated mutation signatures, indicating metastases from cutaneous malignancies, including malignant melanoma and basal cell carcinoma. 3 cases bore gene rearrangements classical for other entities (*SSX1-SS18* rearrangement for synovial sarcoma, *NCOA4-RET* for thyroid carcinoma, and *EWSR1-SSX1* for synovial sarcoma). 2 cases were found to have variants combinations extremely uncommon for primary lung cancer but were molecularly and clinically consistent with hepatocellular carcinoma and urothelial carcinoma. Diagnostic reclassification was supported by immunohistochemistry when applicable. Factors that may be associated with revised pathological diagnosis included biopsies (6 of 9), tissue from extrapulmonary sites (5 of 9), and consults from external institutions (5 of 9).

Conclusions: Next generation sequencing testing of non-small cell lung cancers may not only guide clinical management by finding actionable targets but also by revealing surprising molecular findings that revised pathological and clinical diagnosis.

1932 Comparison of 3 Immunohistochemistry Assays for Detection of ROS1 Rearrangements in a Series of 158 Consecutive Surgically-Resected Adenocarcinomas of the Lung

Francesca Locatelli¹, Giulia Courthod², Genny Jocollé², Giovanni Donati³, Giulio Rossi⁴

¹Pathology Unit "Degli Infermi" Hospital, Azienda USL della Romagna, Cesena, FC, Italy, ²Oncology Unit, Regional Hospital, Aosta, AO, Italy, ³Unit of Thoracic Surgery, Regional Hospital "Parini", Azienda USL della Valle d'Aosta, Aosta, Aosta, AO, Italy, ⁴Pathology Unit, Rimini, RI, Italy

Disclosures: Francesca Locatelli: None; Genny Jocollé: None

Background: ROS1 rearrangements characterize about 1-2% of lung adenocarcinomas and represent a molecular target for specific targeted therapy. Immunohistochemistry is used as a rapid and cost-effective screening test to detect ROS1 rearrangements, although positive cases require confirmation by FISH or other extractive molecular methods.

Design: The diagnostic accuracy of 3 primary antibodies anti-ROS1, namely clone EPMGHR2 (Abcam), clone SP384 (Ventana) and clone D4D6 (Cell Signaling Technology), in a series of 158 consecutive and surgically-resected adenocarcinomas of the lung was evaluated. All investigations were performed in an automated immunostainer (ULTRA, Ventana). All positive cases and 57 randomly-selected negative cases were re-tested by FISH technique using the ROS1 dual color break-apart probe (ZytoVision). Positive staining was quantified by H-score, then recording the percentage (0-100) and the intensity (0-3) of immunoreactivity in tumor cells (cytoplasm and/or membrane localization). Cases were considered positive with EPMGHR2 and D4D6 using a H-score > 150, while SP384 positivity was quoted when at least >30% of tumor cells with 2+/3+ of intensity were recorded.

Results: Only 2 adenocarcinomas (1.2%) showed ROS1 positivity at IHC and FISH. All 3 clones have a complete sensitivity, while specificity ranged from 96.3% (EPMGHR2) to 97.5% (clone SP384) and 99.3% (D4D6). Of note, 6, 4 and 1 false positive cases were detected with EPMGHR2, SP384 and D4D6, respectively. Most interesting, 49 cases (31%) showed borderline immunoreactivity with EPMGHR2 and 27 cases (17%) with SP384, whereas no indeterminate results were noted with D4D6.

Conclusions: The IHC screening to detect ROS1 rearrangements in lung cancer is effective, but pathologists should consider that the clone of anti-ROS1 primary antibody may affect the specificity, with clone D4D6 showing the best performance.

1933 Post-Transplant Lymphoproliferative Disorder Following Lung Transplant in the Current Era: Five-Year Experience in a High Volume Lung Transplant Center

Catherine Luedke¹, Carolyn Glass¹, Elizabeth Pavlisko¹, Jadee Neff¹ Duke University Medical Center, Durham, NC

Disclosures: Catherine Luedke: None; Carolyn Glass: None; Elizabeth Pavlisko: None; Jadee Neff: None

Background: While post-transplant lymphoproliferative disorder (PTLD) is a known complication of solid organ transplant, often linked to immunosuppression and Epstein-Barr virus (EBV), its characteristics remain highly variable resulting in decreased understanding of pathogenesis and outcomes.

Design: We reviewed lung transplants performed between 2014 and 2019 and identified patients who subsequently developed PTLD. Their clinicopathologic features were retrospectively analyzed.

Results: Of 325 lung transplant patients, 7 (2.2%) developed PTLD. 4 were male, and the median age at lung transplant was 30 years (range: 20-75 years). Indications included cystic fibrosis (5), chronic obstructive pulmonary disease (1), and usual interstitial pneumonia (1). At the time of transplant, 2 were EBV seronegative. All 7 received basiliximab induction and started a maintenance regimen of tacrolimus, prednisone, and mycophenolate (MMF). 1 switched from MMF to azathioprine while another discontinued MMF without replacement due to EBV viremia. The median interval from transplant to PTLD diagnosis was 6 months (range: 3-38 months). Of 5 early-onset PTLDs, 4 involved the allograft and 1 the colon; of 2 late-onset PTLDs, 1 involved the allograft and 1 the bladder. 4 had evidence of wide-spread disease by imaging while 3 were localized to the biopsy site. The PTLD subtypes were plasmacytic hyperplasia (3), polymorphic (1), and monomorphic with a diffuse large B-cell lymphoma (DLBCL) phenotype (3). 6 of the PTLDs were positive for EBV. The EBV-negative case did not have EBV viremia, was late-onset, and was classified as monomorphic DLBCL type. Treatment involved reduction of immunosuppression and rituximab, plus chemotherapy in 3 cases of monomorphic or polymorphic subtype. Median follow-up after PTLD diagnosis was 21 months (range: 1-29 months). Both elderly patients died prior to or during therapy, and 1 patient died of infection 21 months after diagnosis with no evidence of PTLD. 3 patients are alive with no evidence of PTLD, and 1 patient is alive with excellent response to therapy.

Conclusions: Within our cohort, the rate of PTLD is low, consistent with a reported trend of decreasing PTLD rates in recent years, possibly as a result of improved immunosuppression regimens. Response to therapy appears extremely promising, but the overall mortality rate remains high. Current histologic parameters do not predict disease extent or clinical outcome, highlighting the importance of continued biomarker development for PTLD.

1934 Utility of Flow Cytometry in the Diagnosis of Pulmonary Hematolymphoid Neoplasms: A Single Institution Experience

Malary Mani¹, May Fu², Weina Chen¹, Jesse Jaso¹, Mingyi Chen¹, Franklin Fuda²

¹University of Texas Southwestern Medical Center, Dallas, TX, ²University of Texas Southwestern, Dallas, TX

Disclosures: Malary Mani: None; May Fu: None; Weina Chen: None; Jesse Jaso: None; Mingyi Chen: None; Franklin Fuda: None

Background: Primary pulmonary hematolymphoid neoplasms (pHLNs) are rare, and the incidence is increasing with modern diagnostic advances and treatments. Flow cytometry (FC) is a proven powerful tool in the diagnosis of hematolymphoid disease, however its role in pHLN is not well represented. In the current study, we aim to assess the utility of FC in diagnosis of patients with pHLNs.

Design: We retrospectively reviewed the FC analyses of pulmonary specimens from our institutional database between 2016-2019. The specimens are comprised of pleural fluid, bronchoalveolar lavages, bronchoscopy or CT guided FNAs, and VATS or surgical biopsies of lung mass. pHLNs were detected by 10-color panel FC by identification of clonal or immunophenotypical aberrant hematolymphoid populations. Primary neoplasms are classified if no extrapulmonary lesions were detected by clinical radiological work up at the time of diagnosis or within three months of diagnosis. Lung involvement is also a frequent site in involvement of lymphoproliferative diseases and are referred to as secondary involvement of the lung.

Results: We retrieved 471 pulmonary specimens submitted for FC. We identified 111 cases that were positive for pHLNs. Median age was 49 years with M:F ratio=1.54/111 specimens aided in the diagnosis of a hematolymphoid disease. Distribution are as follows: 17 pleural fluids, 33 tissue biopsies, 4 bronchoalveolar lavages. Cases were classified as either primary (9/54 (17%)) or secondary (45/54 (84%)), and categorized as myeloid neoplasms, B-cell neoplasm, T-cell neoplasm, primary effusion lymphoma (PEL), post-transplant lymphoproliferative disorder (PTLD); and plasma cell neoplasms (Table 1). Of the pHLNs, diffuse large B-cell lymphoma (DLBCL) was the most common diagnosis (12/54 (22%)). Immunosuppression were notable in 14 patients. The remaining 57/111 cases represent followups of known lymphoproliferative diseases where FC analysis of lung specimen played a role in assessment of disease status. All FC positive cases were confirmed by tissue examination with positive predictive value (PPV) of 100%.

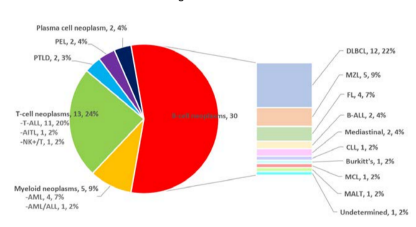


Figure 1 - 1934

Conclusions: FC demonstrates a clear utility for immunophenotyping in the diagnosis of pHLN, with a perfect PPV. Appreciating the spectrum of pHLN is helpful in disease detection. In addition, FC provides the ability to monitor residual disease and response to therapy; especially in immunosuppressed patients where expedited treatment can impact prognosis.

1935 Role of piR-796 in Early Stage Non-Small Cell Lung Carcinoma

Daniel Martinez¹, Marta Casadevall², Jose Ramirez³, Joan Castellano⁴, Laureano Molins⁵, Ramon Marrades⁶, Nuria Viñolas⁷, Jorge Moises⁸, Mariano Monzo⁹, Alfons Navarro⁴

¹Barcelona, Spain, ²University of Barcelona, Unit of Anatomy, Barcelona, Spain, ³Hospital Clinic, Barcelona, Spain, ⁴University of Barcelona, Barcelona, Spain, ⁵General Thoracic Surgery Department, Hospital Clinic, University of Barcelona, Barcelona, Spain, ⁶Respiratory Medicine, Hospital Clinic, Barcelona, Spain, ⁷Medical Oncology, Hospital Clinic, University of Barcelona, Barcelona, Spain, ⁸Hospital Clinic de Barcelona, Barcelona, Spain, ⁹University of Barcelona, Unit of Human Anatomy, Barcelona, Spain

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Background: PIWI-interacting RNAs (piRNAs) are small non-coding RNAs (24-35nt) that play an essential role in maintaining genome integrity trough regulation of transposable elements. Their expression was tough to be limited to germinal cells and early embryogenesis but recently an implication of piRNAs in cancer biology has been reported.

Design: The aim of this study was to explore the expression of piR-796 piRNA, which was identified by small RNAseq, in resected non-small cell lung cancer (NSCLC) patients, and to analyze the correlation with the clinic-pathological features.

We have analyzed 191 resected NSCLC samples from patients who underwent surgery in Hospital Clínic between 2007 and 2015. PiR-796 piRNA was quantified using custom TaqMan non-coding RNA assays in tumor and normal tissue. In vitro studies using siRNAs to inhibit piRNA expression were performed in two lung adenocarcinoma cell lines: HCC44 and A549.

Results: PiR-796 was overexpressed in tumor tissue compared to normal tissue (p<0.001). It also appeared to have higher expression in squamous cell carcinoma compared to adenocarcinoma histological subtype. Expression of piR-796 had prognosis impact in the group of early stage (I-II) adenocarcinoma patients. Higher piR-796 levels were associated with shorter disease-free survival (p=0.045). In vitro analysis showed that the silencing of piR-796 was associated with decreased cell migration in A549 cell line (p<0.0001) and increased apoptosis in HCC44 cell line (p=0.03).

Conclusions: In conclusion, piR-796 may have an important role in carcinogenesis in NSCLC, promoting cell migration and regulating apoptosis and may be a potential new prognostic biomarker for NSCLC.

1936 Pulmonary Nodular Light Chain Deposition Disease: Clinicopathologic and Ultrastructural Characterization of 10 Cases

Lucas Massoth¹, Martin Selig¹, Mari Mino-Kenudson¹, Lawrence Zukerberg², G. Pétur Nielsen³, Yin Hung¹ Massachusetts General Hospital, Boston, MA, ²Auburndale, MA, ³Harvard Medical School, Boston, MA

Disclosures: Lucas Massoth: None; Martin Selig: None; Mari Mino-Kenudson: None; Lawrence Zukerberg: None; G. Pétur Nielsen: None; Yin Hung: None

Background: Light chain deposition disease (LCDD) is characterized by accumulation of immunoglobulin light chains in viscera/soft tissue. LCDD rarely involves the lungs in a nodular or diffuse pattern. We present herein a large series of pulmonary nodular LCDD.

Design: In an institutional review from >2000 patients in 2000-2019, we identified 12 specimens from 10 patients with pulmonary nodular LCDD. We reviewed clinicopathologic features and performed electron microscopy in all cases.

Results: Of 10 patients (6 women, 4 men; age 42-74 [median 68] years), 7 were smokers, 4 had a history of autoimmune disease, but none had a history or evidence of a systemic lymphoproliferative/plasma cell disorder. Clinical presentations were most often incidental. Of 9 patients with radiology available, 3 showed a solitary nodule each; 6 showed multiple nodules (including 3 with associated cystic changes). Each nodule ranged 0.6-2.1 (median 1.2) cm. By light microscopy on hematoxylin-eosin staining, each nodule appeared as amorphous eosinophilic deposits; all cases showed multinucleated giant cells engulfing the deposits and prominent plasma cell infiltrates. While the morphology was reminiscent of amyloidoma, staining for congo red was negative for amyloid in all cases. PAS was positive in all 5 cases tested, and trichrome was distinctly bright red in 5 of 6 cases tested. Of 8 cases with plasma cell cytoplasmic light chain expression tested by immunohistochemistry and/or in-situ hybridization, 6 were kappa-predominant, 1 lambda-predominant and 1 polytypic. Immunofluorescence studies in 2 cases confirmed kappa and lambda light chain restriction in 1 case each. Ultrastructurally, all cases showed extracellular electron-dense granular deposits, variably admixed with entrapped collagen fibers, often appearing perivascular, and lacking non-branching fibril formation characteristic of amyloid. Of 6 patients with >2 years of follow-up (median 4.3 years), 5 were alive (the single death was unrelated to LCDD).

Conclusions: Pulmonary nodular LCDD appears clinically indolent, radiologically solitary or multifocal, and mimics amyloidoma by light microscopy. Yet, special stains (negative congo red, bright red trichrome) and ultrastructural features (granular deposits with variable entrapped collagen) aid the diagnosis of LCDD and its distinction from amyloidoma. The high rate of autoimmune disease in this cohort may suggest a role of chronic immunologic stimulation in the pathogenesis of pulmonary nodular LCDD.

1937 TFF-1 (trefoil factor-1) as a Marker for Non-TRU (terminal respiratory unit) Type Lung Adenocarcinoma with Gastrointestinal Features

Daisuke Matsubara¹, Taichiro Yoshimoto², Yusuke Amano³, Atsushi Kihara², Tomoki Shibano², Toshiro Niki⁴
¹Shimotsukeshi, Tochigi, Japan, ²Jichi Medical University, Shimotsukeshi, Tochigi, Japan, ³Jichi Medical University, Shimotsukeshi City, Tochigi ken, Japan, ⁴Jichi Medical University, Shimotsuke, Japan

Disclosures: Daisuke Matsubara: None; Yusuke Amano: None; Atsushi Kihara: None

Background: Molecular targeted therapies against *EGFR* and *ALK* have improved the quality of life of lung adenocarcinoma patients. However, targetable driver mutations are mainly found in TTF-1/NKX2-1-positive terminal respiratory unit (TRU) types and rarely in non-TRU types.

Design: To elucidate the molecular characteristics of the major subtypes of non-TRU-type adenocarcinomas, we analyzed 19 lung adenocarcinoma cell lines (11 TRU types and 8 non-TRU types). A characteristic of non-TRU-type cell lines was the strong expression of *TFF-1* (*trefoil factor-1*), a gastric mucosal protective factor. By immunohistochemistry, we examined TFF-1, TTF-1/NKX2-1, HNF4-alpha, and MUC5AC expressions using 238 primary lung adenocarcinomas resected at Jichi Medical University Hospital. We also examined the correlation between TFF-1 expression and clinico-pathological factors and genetic abnormalities.

Results: An immunohistochemical analysis of revealed that TFF-1 was positive in 31 cases (13%). TFF-1 expression was frequently detected in invasive mucinous (14/15(93%)), enteric (2/2(100%)), and colloid (1/1(100%)) adenocarcinomas, less frequent in acinar (5/24(21%)), papillary (7/120(6%)), and solid adenocarcinomas (2/43(5%)), and negative in micropapillary (0/1(0%)), lepidic (0/23(0%)), and microinvasive adenocarcinomas or adenocarcinoma *in situ* (0/9(0%)). TFF-1 expression correlated with the expression of HNF4alpha and MUC5AC (p<0.0001, p<0.0001, respectively) and inversely correlated with that of TTF-1/NKX2-1 (p<0.0001). These results indicate that TFF-1 is characteristically expressed in non-TRU-type adenocarcinomas with gastrointestinal features. TFF-1-positive cases harbored KRAS mutations at a high frequency, but no *EGFR* or *ALK* mutations. TFF-1 expression correlated with a poor prognosis in advanced stages. Moreover, the knockdown of TFF-1 inhibited cell proliferation and induced apoptosis in a TFF-1-positive and KRAS-mutated lung adenocarcinoma cell line.

Conclusions: These results indicate that TFF-1 is not only a biomarker, but also a potential molecular target for non-TRU-type lung adenocarcinomas.

1938 Clinical Impact of Rapid Biomarker Testing in Non-Small Cell Lung Cancer in a Community Setting Priscilla Matthews¹, Andrea Beharry¹, Massey Nematollahi¹, Anna Bendzsak¹, Kashif Irshad¹, Parneet Cheema¹, Brandon Sheffield¹

¹William Osler Health System, Brampton, ON

Disclosures: Priscilla Matthews: None; Andrea Beharry: None; Massey Nematollahi: None; Anna Bendzsak: None; Kashif Irshad: None; Parneet Cheema: None; Brandon Sheffield: *Grant or Research Support*, Pfizer; *Speaker*, Merck; *Consultant*, Roche; *Primary Investigator*, Boehringer Ingelheim; *Grant or Research Support*, Astra Zeneca

Background: Accurate and timely biomarker results are essential for the modern-day treatment of non-small cell lung cancer (NSCLC). Nonetheless, many institutions around the world, particularly those without in-house testing capabilities, are faced with prolonged turnaround time. This study investigates the clinical impact of implementing a rapid biomarker testing strategy.

Design: Rapid in-house biomarker testing was implemented utilizing immunohistochemical assays for PD-L1, ALK, ROS, and BRAF V600E, as well as qPCR for EGFR using the Biocartis Idylla technology. After a 6-month period, we performed a retrospective chart review of NSCLC patients presenting pre- and post-implementation of rapid biomarker testing at our facility.

Results: 198 patients were included in the study (85 underwent rapid biomarker testing and 113 underwent traditional testing). The median (IQR) turnaround time for biomarker reports decreased from 61.5 (31-100) to 4 (2-7) days (p < 0.001), with turnaround time defined as the total number of days between the diagnosis and biomarker result appearing on the medical record. The mean time to initiation of systemic therapy for advanced-stage patients decreased from 52.5 days to 31.9 days (p < 0.01) The proportion of patients who had a complete biomarker report at the time of their first consult with a medical oncologist increased from 26.8% to 58.0%. Similarly, the proportion of patients who had a complete biomarker report at the time of initiation of systemic therapy increased from 56.7% to 86.4%.

Conclusions: Timely biomarker results are essential for the delivery of targeted therapy and immunotherapy to NSCLC patients. Many centres experience delays in biomarker results for a number of reasons. In this study, we demonstrate that a small panel of rapidly-reported biomarkers can significantly reduce delays in the initiation of systemic therapy, ultimately leading to superior patient outcomes.

1939 Primary Thoracic Sarcomas have a Worse Clinical Outcome Compared to Soft Tissue Sarcomas

Mitra Mehrad¹, Justin Cates¹, Jennifer Gordetsky¹

¹Vanderbilt University Medical Center, Nashville, TN

Disclosures: Mitra Mehrad: None; Justin Cates: None; Jennifer Gordetsky: None

Background: Primary sarcomas of the thorax (heart, lung, pleura, thymus, mediastinum) are rare and little information is available on predictors of their prognosis. The National Cancer Institute's SEER (Surveillance, Epidemiology, and End Results) database contains

abundant information on the natural history of soft tissue sarcomas. Therefore, we investigated the clinical outcomes of primary thoracic sarcomas (TS) utilizing the SEER database.

Design: The SEER database was queried for TS entered between 1985 and 2013. TS was compared to sarcomas arising in soft tissues of the extremities and trunk. Staging was performed based on the AJCC 8th edition. Multivariable Cox regression was used to identify prognostic factors. Kaplan-Meier curves were plotted to assess cancer-specific survival (CSS). Patients <19 years of age and those without confirmed surgical resection were excluded.

Results: A total of 1308 TS and 20160 soft tissue sarcomas were available for analysis. Most TS arose in the lung (785; 60.0%) followed by mediastinum (224; 17.1%), heart (165; 12.6%), pleura (132; 10.1%) and thymus (2; 0.2%). The 5 most common TS were malignant solitary fibrous tumor (SFT) (147; 11.2%), leiomyosarcoma (LMS) (139; 10.6%), synovial sarcoma (128; 9.7%), undifferentiated pleomorphic sarcoma (UPS) (126; 9.6%), and angiosarcoma (122; 9.3%), whereas in the soft tissue were UPS (5477; 27.1%), LMS (2357; 11.6%), myxoid liposarcoma (LPS) (1544; 7.6%), well-differentiated LPS (1413; 7.0%), and myxofibrosarcoma (1331; 6.6%). Comparing the two groups by multivariate analysis, the TS were larger and more frequent in males (p<0.0001). They were also of higher histologic grade and stage and were less likely to receive adjuvant therapy (p<0.001). Among the TS, tumors of heart origin (p<0.0001), greater than 5 cm, stage III-IV, and histologic grade 3 had worse CSS (p<0.01). the more common TS tumors including malignant SFT, LMS, synovial sarcoma, and angiosarcoma still had worse CSS compared to their soft tissue counterparts even when adjusting for stage, size and treatment.

Conclusions: Primary thoracic sarcomas show worse clinical outcome compared to soft tissue sarcomas. A thoracic specific sarcoma staging system may be more predictive of CSS.

1940 Cytokeratin 7 is Superior to Lineage-Specific Markers SATB2, CDX2, and Villin in Distinguishing Primary Pulmonary Mucinous Adenocarcinoma from Metastatic Colorectal Carcinoma

Cherise Meyerson¹, Gregory Fishbein²

¹UCLA Medical Center, Reseda, CA, ²UCLA Health, Los Angeles, CA

Disclosures: Cherise Meyerson: None; Gregory Fishbein: None

Background: Primary pulmonary mucinous adenocarcinoma (PMA) may be difficult or impossible to distinguish from colorectal adenocarcinoma (CRC) due to considerable morphologic and immunohistochemical (IHC) overlap. Since the lung is a common site of metastasis, the need to distinguish PMA from CRC in the lung is a routine challenge. Commonly used lineage-specific IHC markers like CDX2, TTF-1, and napsin A are helpful to distinguish non-mucinous lesions but are either insensitive or nonspecific when applied to mucinous lesions. SATB2 is a relatively new IHC marker that distinguishes CRC from upper gastrointestinal and pancreaticobiliary tumors. Its ability to distinguish CRC from PMA is not yet completely elucidated.

Design: Three tissue microarrays of lung resections containing primary pulmonary mucinous adenocarcinomas (PMAs), metastatic colorectal carcinomas (CRCs), and primary pulmonary non-mucinous adenocarcinomas (PNMAs) were stained with CK7, CK20, SATB2, CDX2, villin, TTF-1, and napsin A. Two pathologists evaluated the number of positive neoplastic cells semiquantitatively, regardless of intensity: 0 (no staining), 1+ (<10%), 2+ (10-50%), 3+ (>50%).

Results: Thirty-one PMAs, 32 CRCs, and 34 PNMAs were assessed (Table 1). Thirty (97%) of PMAs and 34 (100%) of PNMAs were positive (3+) for CK7, while all CRCs were negative for CK7. Twenty-seven (84%) of CRCs and 2 (6%) of PMAs were positive (3+) for SATB2, and 29 (91%) of CRCs and 2 (6%) of PMAs were positive (3+) for CDX2. Both PMAs and CRCs had high rates of villin positivity, with 23 (74%) and 32 (100%) positive (3+), respectively. Only 4 (13%) and 7 (23%) of PMAs were positive (3+) for TTF-1 and napsin A, respectively. No CRCs were positive for TTF-1 or napsin A. In deciding PMA vs. CRC, CK7 was 97% sensitive and 100% specific for PMA. SATB2 was superior to CDX2 and Villin but was only 96% sensitive and 94% specific for CRC.

		CK7 n (%)	CK20 n (%)	SATB2 n (%)	CDX2 n (%)	Villin n (%)	TTF-1 n (%)	Napsin A n (%)
PMA	31	30 (97)	2 (6)	2 (6)	2 (6)	23 (74)	4 (13)	7 (23)
CRC	32	0 (0)	21 (66)	27 (84)	29 (91)	32 (100)	0 (0)	0 (0)
PNMA	34	34 (100)	0 (0)	0 (0)	0 (0)	7 (21)	34 (100)	33 (97)

Table 1. Number and percentage of cases with positive staining (3+) for CK7, CK20, SATB2, CDX2, villin, TTF-1, and napsin A in primary pulmonary mucinous adenocarcinoma (PMA), metastatic colorectal carcinoma (CRC), and primary pulmonary non-mucinous adenocarcinoma (PNMA)

Conclusions: Although the lineage-specific markers SATB2 and CDX2 were fairly specific for CRCs, a few PMAs were also positive. In contrast, all CRCs were negative for CK7 while almost all of the PMAs were positive. Lineage-specific markers TTF-1 and napsin A had a low sensitivity for PMAs. Villin showed a low specificity with the majority of CRCs and PMAs staining positively. Our results suggest that a CK7-positive tumor in the lung, whether mucinous or non-mucinous, is unlikely to be of colorectal origin. Lineage-specific markers such as SATB2 are of questionable value when evaluating mucinous lesions in the lung.

1941 Morphology and Ki-67 Proliferative Index Drive Prognosis in Combined Large Cell Neuroendocrine Carcinomas of the Lung

Massimo Milione¹, Patrick Maisonneuve², Federica Grillo³, Alessandro Mangogna⁴, Giovanni Centonze⁵, Giovanna Garzone⁶, Laura Cattaneo⁷, Ketevani Kankava⁸, Adele Busico⁶, Paola Spaggiari⁹, Alessandro Del Gobbo¹⁰, Luisa Bercich¹¹, Luigi Rolli¹², Elisa Roca¹¹, Natalie Prinzi⁶, Giancarlo Pruneri⁵, Alfredo Berruti¹¹, Ugo Pastorino¹², Carlo Capella¹³

¹Fondazione IRCCS Istituto Nazionale Tumori Milano, Milano, Italy, ²IEO, Milan, Italy, ³University of Genova, Genova, Italy, ⁴University of Trieste, Trieste, Friuli Venezia Giulia, Italy, ⁵Fondazione IRCCS Istituto Nazionale Tumori Milano, Milan, Italy, ⁶IRCCS Foundation, Istituto Nazionale dei Tumori, Milano, Italy, ⁸Teaching, Scientific and Diagnostic Pathology Laboratory, Tbilisi State Medical University, Tbilisi, Georgia, ⁹Istituti Clinic Humanitas, Rozzano, Milano, Italy, ¹⁰Fondazione IRCCS ca Granda Ospedale Maggiore Policlinico, Milan, Italy, ¹¹University of Brescia at ASST-Spedali Civili, Brescia, Italy, ¹²IRCCS Foundation National Cancer Institute, Milan, Italy, ¹³Uni-Insubria, Varse, Italy

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Background: While little information is available concerning the molecular landscape and prognostic factors of pulmonary large cell neuroendocrine carcinomas (LCNECs) even less is known about combined LCNECs (co-LCNECs). These are LCNECs in which a part of the whole tumor (any extent) is composed of a non-neuroendocrine component, most often adenocarcinoma (ADK) or squamous cell carcinoma (SqCC). Revision of a large multicenter series of LCNECs enabled us to extrapolate co-LCNECs and describe their clinical-pathological and prognostic features.

Design: Surgical specimens of 148 LCNECs were centrally reviewed with combined features identified. Morphology, immunohistochemical (Ki-67, Napsin A, p-40, TTF-1, CD44, OTP, SSTR2A, SSTR5, mASH1, p53, Rb1, MDM2) and genomic (*TP53, RB1, ATM, JAK2, KRAS, STK11*) findings were studied and correlated with Overall Survival (OS).

Results: LCNEC diagnosis was confirmed in 123 patients (Table1): 75 pure LCNECs, 48 co-LCNECs (37 combined with ADK; 8 with SqCC) and 3 LCNECs showing only Napsin A positivity but no morphologic evidence of ADK. Applying Ki-67 labeling index cut off at 55% for the NE component, lesions were subdivided into two subgroups: co-LCNECs-A Ki-67 <55% (15 combined with ADK and 3 with SCC) and co-LCNECs-B Ki-67 ≥55% (25 combined with ADK and 5 with SqCC). Napsin A and Alcian blue expression were statistically more frequent in LCNECs-A (P=0.01; P=0.004) compared to LCNECs-B. Median OS for all LCNECs was 1.4 years; statistical differences in OS (HR 1.96, 95% CI 1.30-2.95, P <0.001) were observed between pure (median OS 1.3 years) and co-LCNECs (median OS 1.8 years). This difference in OS was maintained between pure LCNEC-A and co-LCNEC-A (P <0.05) but disappeared for LCNEC-B. At univariate analysis variables associated (P <0.05) with poor OS were p40 expression and absence of combined features, while variables associated with longer OS were tumor peripheral location, Napsin A and Alcian blue staining. At multivariable analysis, only tumor location and Napsin A remained significantly associated with OS, after adjustment for Ki-67 and study center.

With regards to molecular analyses, 26 cases (54,2 %) of the co-LCNEC group were studied (see Table 1). No single next-generation sequencing marker was statistically associated with OS.

Table 1. Next-generation sequencing (NGS) analysis*

	All LCNEC§	Combined LCNEC	Pure LCNEC	P-value
	(123)	(48)	(75)	
NGS analysis#	66 (100)	26 (100)	40 (100)	
WT	11 (16.7)	5 (19.2)	6 (15.0)	
Mutated	55 (83.3)	21 (80.8)	34 (85.0)	0.74
TP53		, ,		
WT	27 (40.9)	12 (46.2)	15 (37.5)	
Mutated	39 (59.1)	14 (53.8)	25 (62.5)	0.61
RB1				
WT	42 (63.6)	17 (65.4)	25 (62.5)	
Mutated	24 (36.4)	9 (34.6)	15 (37.5)	1.00
ATM				
WT	60 (90.1)	23 (88.5)	37 (92.5)	
Mutated	6 (9.1)	3 (11.5)	3 (7.5)	0.67
JAK2				
WT	64 (97.0)	25 (96.2)	39 (97.5)	
Mutated	2 (3.0)	1 (3.8)	1 (2 .5)	1.00
KRAS				
WT	54 (81.8)	21 (80.8)	33 (82.5)	
Mutated	12 (18.2)	5 (19.2)	7 (17.5)	1.00
STK11		·		
WT	63 (95.5)	25 (96.2)	38 (95.0)	
Mutated	3 (4.5)	1 (3.8)	2 (5.0)	1.00

^{*} ION AmpliSeg Cancer Hot Spot Panel v2, Thermo Fisher Scientific

#66 LCNEC patients out of 123 were studied: 26 co-LCNECs and 40 pure LCNECs.

Conclusions: The identification and morphologic characterization of combined features in LCNECs as well as the application of Ki-67 cut off at 55% contribute in predicting clinical outcome of pure-LCNEC and co-LNEC patients.

1942 CTNNB1 Mutations and Co-Mutations in Non-Small Cell Lung Cancer

Andres Mindiola Romero1. Laura Tafe2

¹Dartmouth Hitchcock Medical Center, Lebanon, NH, ²Dartmouth-Hitchcock Medical Center, Lebanon, NH

Disclosures: Andres Mindiola Romero: None; Laura Tafe: None

Background: CTNNB1 encodes for β-catenin, which is a member in the Wnt signal transduction pathway required for proliferation, survival and differentiation of different epithelial cells. Mutation of CTNNB1 causes constitutional changes in the β-catenin protein that impedes its degradation, leading to an uncontrolled proliferation of the mutated cell. CTNNB1 exon 3 hot-spot mutations are described in various tumor types and, for instance, in endometrial cancer, are associated with high risk of disease recurrence. The role of CTNNB1, frequency and type of co-mutations has not been well characterized in non-small cell lung carcinomas (NSCLC).

Design: Between 2013-2018, lung cancer samples from 855 patients were sequenced on the Ion Torrent PGM with the 50 gene AmpliSeq Cancer Hotspot Panel v2. Our in-house sequencing database was searched to identify patients with *CTNNB1* mutations; co-mutations in genes commonly altered in NSCLC were also recorded.

Results: Thirteen patients (1.5%) with *CTNNB1* mutations (p.S37F (5) (one also with p.D32Y), p.S37C (3), and p.S33C, p.D32H, p.G34E, p.D32V, p.D32N, in one patient each) were identified. All tumors were adenocarcinoma histology. Five patients underwent lobectomy and the predominant histologic patterns were solid (2), papillary (2) and micropapillary (1). The patients' age ranged from 54-82, at the time of diagnosis, and eight were female (61.5%). All patients were current (2) or former (11) smokers. Five patients presented with stage IV disease, two with stage III, two with stage III, and four with stage I disease. Six are deceased and seven are alive with disease. Co-

[§] The present study was performed exclusively on the 123 surgical specimens confirmed after expert pathologist revision as LCNECs out of the 148 originally collected specimens; in more details the excluded 25/148 ones were reclassified as: 8 non neuroendocrine Large Cell Carcinomas, 3 Atypical Carcinoids, 7 Squamous Carcinomas and 7 Small Cells Neuroendocrine Carcinomas

mutations were identified in all but one case and consisted of: BRAF (1 – V600E), EGFR (3 – ex19 deletions), KRAS (4), PIK3CA (3), and TP53 (3). In two patients, EGFR, PIK3CA and CTNNB1 mutations were co-occurring.

Conclusions: We identified *CTNNB1* mutations in 1.5% of lung adenocarcinomas, which, in our population, were associated with frequent co-mutations. Additional clinicopathologic data will be aggregated from our internal patients and available databases (cbioportal) to better understand the clinical implications of *CTNNB1* mutations in NSCLC. Co-mutation with *EGFR* may be a mechanism of primary resistance to EGFR TKIs. In addition, *CTNNB1* mutations make patients eligible for newer small-molecule TKIs (e.g. TTKi).

1943 Immunohistochemical Evaluation of Orthopedia Homeobox (OTP) Expression on Tumors and Normal Tissues from Various Organs

Mohammad Mohammad¹, Jianhui Shi², Aihua Li³, Haiyan Liu², Fan Lin²

¹Geisinger Commonwealth School of Medicine, Danville, PA, ²Geisinger Medical Center, Danville, PA, ³Burlingame, CA

Disclosures: Mohammad Mohammad: None; Jianhui Shi: None; Aihua Li: *Employee*, Epitomics, an abcam company; Haiyan Liu: None; Fan Lin: None

Background: Recent study demonstrated that orthopedia homeobox (OTP) was a highly sensitive and specific marker for diagnosing pulmonary carcinoids (PCs) (Nonaka D et al., Am J Surg Pathol 2016; 40: 738-744). In this study, we focused on the evaluation of OTP expression in a large series of non-neuroendocrine carcinomas and tumors from various organs.

Design: Immunohistochemical analysis of OTP (rabbit monoclonal antibody, EP490, Abcam) was performed on 1233 cases of tumor and normal tissues on tissue microarray (TMA) sections, including mesothelioma (N=18), lung neuroendocrine carcinoma (CA) (N=24), lung adenocarcinoma (ADC) (N=88), lung squamous cell CA (N=66), papillary thyroid CA (N=47), ENT squamous cell CA (N=28), breast fibroadenoma (N=20), pancreatic neuroendocrine tumor (N=33), pancreatic ADC (N=43), adrenal pheochromocytoma (N=14), endometrial CA FIGO II (N=59), ovary papillary serous CA (N=41), clear cell CA of uterus and ovary (N=22), melanoma (N=32), skin neuroendocrine CA (N=27), invasive urothelial CA (N=43), bladder small cell carcinoma (bladder SCCs) (N=24), prostatic ADC (N=38), germ cell tumors (N=60), colonic ADC (N=164), angiosarcoma (N=12), papillary renal cell carcinoma (RCC) (N=33), clear cell RCC, low grade (N=79), clear cell RCC, high grade (N=51), hepatocellular CA (N=47), liver metastatic neuroendocrine CA (N=18), glioblastoma multiforme (GBM) (N=23), mantle cell lymphoma (N=13) and hairy cell leukemia (N=1). Samples from normal tissue included pancreas (N=13), rectum/appendix/colon (N=39) and ilium/duodenum/stomach (N=13). Nuclear expression for OTP was interpreted as negative (<5% tumor cells stained), 1+ (<25%), 2+ (26-50%), 3+ (51-75%), and 4+ (>75%).

Results: Six of 10 (60%) PCs were diffusely positive for OTP. One of 33 (3%) pancreatic NETs, 2 of 24 (8%) bladder SCCs, and 1 metastatic NET in the liver (negative for TTF1 and no lung mass identified) were positive for OTP. All other tumors and normal tissues were negative. Representative cases are shown in Figure 1.

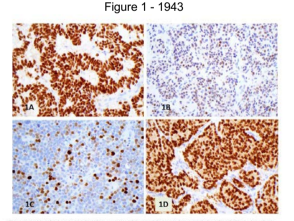


Figure 1 showing representative cases with positive nuclear staining for OTP. Note that pulmonary carcinoid (1A), pancreatic NET (1B), small cell CA of the bladder (1C), and a metastatic NET in the liver which was negative for TTF1 and positive for PR (no lung/chest mass identified on a CT scan).

Conclusions: Our data demonstrated that OTP expression was only rarely identified in non-pulmonary neuroendocrine tumors/carcinomas, which further validated the previous report of OTP to be a highly specific marker for diagnosing PCs. The diagnostic sensitivity for PCs in this study appears to be lower than the previous report, which is probably due to the small number of cases included. Caution should be taken because rare pancreatic NETs and bladder SCCs can be positive for OTP.

1944 HLA-1 Loss is Common in Non-Small Cell Lung Cancer and May Contribute to Failure of Immune Checkpoint Therapy

Margaret Moore¹, Edward Stelow²

¹University of Virginia, Charlottesville, VA, ²University of Virginia Health System, Charlottesville, VA

Disclosures: Margaret Moore: None; Edward Stelow: None

Background: Quantification of PD-L1/PD-1 expression in non-small cell lung cancer (NSCLC) is not always predictive of efficacy of immune checkpoint inhibitor therapy, and response to these agents remains limited to a minority of patients. Human Leukocyte Antigen 1 (HLA-1) participates in presentation of aberrant peptide antigens, enabling cytotoxic T cells to recognize and destroy tumor cells. Correspondingly, loss of HLA-1 expression precludes immune recognition, and this loss is one proposed mechanism of treatment failure of immunotherapy. We hereby examine the expression of PD-L1 and HLA-1 in a large cohort of NSCLCs to determine the patterns and frequency of expression of these two markers.

Design: NSCLC resection specimens from 2011-2014 classified as adenocarcinoma (AC) or squamous cell carcinoma (SCC) were identified. Tumor microarrays with 1.0 mm diameter cores were constructed, with 2 sections of neoplastic tissue and 1 section of uninvolved lung for each case. Immunohistochemistry for PD-L1 (22C3) and HLA-1 was performed. Membranous tumoral PD-L1 staining was semi-quantitatively scored as <1%, 1-50%, and >50%. HLA-1 was scored as intact, partial (clonal) loss, or complete loss with total absence of staining.

Results: 131 cases were identified, with 73 ACs and 58 SCCs. 75 cases (57%) demonstrated diffuse staining for HLA-1; 16 (12%) demonstrated clonal loss in a subset of cells; and 40 (31%) demonstrated complete loss of expression. HLA-1 was retained in all cores of non-neoplastic lung. HLA-1 expression did not significantly differ by gender, smoking history, age, histosubtype, primary tumor size, node status, or grade of differentiation. PD-L1 expression was not significantly associated with HLA-1 expression, and there was no difference in HLA-1 expression in cases with robust PD-L1 expression (>50%) compared to those with focal expression (1-50%). All combinations of HLA-1/PD-L1 expression were observed (HLA-1+/PD-L1+, HLA-1-/PD-L1+, HLA-1-/PD-L1-, HLA-1-/PD-L1-).

Characteristic: N	HLA-1: Complete Loss N=40	HLA-1: Partial Loss N=16	HLA-1: Intact N=75	p-Value
Sex Female: 57 Male: 74	21 (37%) 19 (26%)	4 (7%) 12 (16%)	32 (56%) 43 (58%)	0.17
Smoking History Smoker: 123 Nonsmoker: 8	38 (31%) 2 (25%)	16 (13%) 0 (0%)	69 (56%) 6 (75%)	0.67
Mean Age at Diagnosis	64	65	68	0.13
Histology Adenocarcinoma: 73 Squamous Cell Carcinoma:58	20 (27%) 20 (34%)	7 (10%) 9 (16%)	46 (63%) 29 (50%)	0.30
Primary Tumor Size (cm)	3.4	2.8	2.9	0.44
Node Status N0: 95 N1 and N2: 36	29 (31%) 11 (31%)	13 (14%) 3 (8%)	53 (56%) 22 (61%)	0.73
Grade Well/Moderately Differentiated: 90 Poorly Differentiated: 41	27 (30%) 13 (32%)	9 (10%) 7 (17%)	53 (59%) 22 (54%)	0.53
PD-L1 Positive (≥1%): 72 Negative (<1%): 59	17 (24%) 23 (39%)	10 (14%) 6 (10%)	45 (63%) 30 (51%)	0.16

Conclusions: HLA-1 loss is common in NSCLC and can readily be assessed by immunohistochemical methods. In this study, there is no association between PD-L1 expression and HLA-1 expression. Given that HLA-1 expression may be a determinant of response to immunotherapeutic agents targeting the PD-1/PD-L1 axis, it may be considered as an adjunct immunohistochemical marker prior to initiation of immune checkpoint therapy or in instances of treatment failure.

1945 Histopathologic Features in Interstitial Lung Disease Associated with Short Telomere Syndromes Urooba Nadeem¹, Aliya Husain², Jane Churpek³, Ayodeji Adegunsoye², Mary Strek⁴, Simone Feurstein² 1 University of Chicago Hospital, Chicago, IL, 2 The University of Chicago, Chicago, IL, 3 The University of Wisconsin School of Medicine and Public Health, Madison, WI, 4 University of Chicago Medicine, Chicago, IL

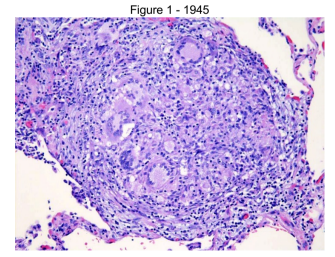
Disclosures: Urooba Nadeem: None; Aliya Husain: None; Jane Churpek: None; Ayodeji Adegunsoye: *Speaker*, Boehringer Ingelheim; Mary Strek: None; Simone Feurstein: None

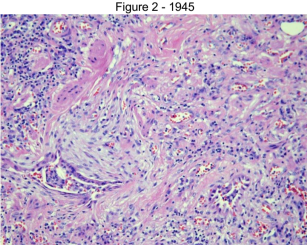
Background: Short telomere syndromes (STS) are multisystem accelerated aging syndromes caused by inheritable gene mutations in telomere maintenance genes; frequent manifestations include bone marrow failure and interstitial lung disease (ILD). Insufficient information is available about histologic findings and patterns of ILD in individuals with STS and whether these vary by specific inherited STS gene. Our study aims to describe the morphology of lung disease in patients with STS.

Design: Probands enrolled in the Inherited Hematologic Disorders Registry at our hospital with a personal or familial history of pulmonary fibrosis underwent genetic testing via targeted genomic capture and next-generation sequencing (NGS) of 13 STS genes. Available specimens(n-8), including wedge resections(n-6) and lung explants (n-2), were analyzed for 15 histopathological features (Table 1).

Results: Eight patients with known STS with ILD had available histology (M=7, F=1 average age at diagnosis 56.3 years). Pathognomonic findings of usual interstitial pneumonia (UIP) were seen in all patients. One patient with two different mutations in the telomere-associated genes showed features of both UIP and non-specific interstitial pneumonia (NSIP). There was associated lymphoplasmacytic infiltration which was mild in 50% and extensive in the remainder. Other histologic findings were variable, e.g. non-necrotizing granulomas and upper lobe predominance are noted in 50% of the patients. Pulmonary hypertension was seen in all of the patients except one who had a very early disease.

Histologic findings	% of patients (n=8)
Pattern of ILD	UIP alone 87.5%
	UIP and NSIP 12.5%
Fibrosis	100%
Honeycombing	87.5%
Fibroblastic foci	100%
Bronchial metaplasia	100%
Granulomata	50%
Upper lobe predominance	50%
Lymphoplasmacytic infiltrates	100% (50% mild, 50% severe)
Lymphoid aggregates	87.5%
Pleuropulmonary fibroelastosis	12.5%
Pulmonary hypertension	87.5%
Submucosal fibrosis	87.5%
Pleural fibrosis and thickening	100%
Eosinophils	100%
Reactive atypia	50%





Conclusions: STS associated lung disease frequently shows UIP with additional findings including inflammatory (lymphoplasmacytic infiltrate) and hypersensitivity pneumonia-like (granulomas and upper lobe predominance) features. Consideration should be given to testing for STS in patients with UIP with these histologic findings. It is important to identify this cohort of patients because these patients may have a more rapid progression of their ILD and an increased toxicity from immunosuppressive drugs especially in post-transplant setting. If a pathogenic gene mutation is identified, first-degree family members (especially siblings) should be informed about disease monitoring options and knowing the risks of future health issues.

1946 Utility of HEG1 Immunohistochemistry for Diagnosis of Malignant Mesothelioma

Julia Naso¹, Shoutaro Tsuji², Andrew Churg³

¹University of British Columbia, Vancouver, BC, ²Kanagawa Cancer Center Research Institute, Yokohama, Japan, ³University of British Columbia/Vancouver General Hospital, Vancouver, BC

Disclosures: Julia Naso: None; Shoutaro Tsuji: None; Andrew Churg: None

Background: Immunohistochemistry is important to help differentiate malignant mesothelioma from other malignancies. However, the sensitivity of mesothelial markers is particularly limited for sarcomatoid mesothelioma. A HEG homologue 1 (HEG1) antibody was recently proposed to be a specific marker for malignant mesothelioma.

Design: The SKM9-2 monoclonal HEG1 antibody was used in immunohistochemistry on tissue microarrays containing malignant mesotheliomas (n=66), non-small cell lung carcinoma (NSCLC, n=153) and high grade serous (HGS) ovarian carcinoma (n=17). Results were compared to immunohistochemistry using the mesothelial markers calretinin, D2-40, CK5/6 and WT1.

Results: HEG1 immunohistochemistry showed intense membranous staining in 50 (94%) of 53 epithelioid mesotheliomas and 2 (15%) of 13 sarcomatoid mesotheliomas. Immunostaining tended to be diffuse in epithelioid mesotheliomas and focal in sarcomatoid mesotheliomas. Correlation of immunohistochemistry results between HEG1 and other mesothelial markers was moderate, with correlation coefficients ranging from 0.52 (D2-40) to 0.77 (CK5/6). Compared to HEG1, other tested mesothelial markers had similar sensitivities for epithelioid mesothelioma (92-96%) and sarcomatoid mesothelioma (15-31%). All NSCLC cases were negative for HEG1 (adenocarcinoma n=53, squamous cell carcinoma n=60, large cell carcinoma n=13, sarcomatoid carcinoma n=21). Focal HEG1 immunostaining was present in 5 (29%) of 17 HGS ovarian carcinomas.

Conclusions: HEG1 positivity is 100% specific for epithelioid mesothelioma versus NSCLC, but cross reactivity with HGS ovarian carcinoma is a potential confounder. Sensitivity of HEG1 immunohistochemistry for epithelioid and sarcomatoid mesothelioma is 94% and 15%, respectively. HEG1 immunohistochemistry is potentially useful for differentiating mesothelioma from pulmonary malignancies, but the utility for sarcomatoid mesothelioma is limited by low sensitivity.

1947 DLL3 Expression in Small Cell Lung Carcinomas (SCLC)

Patricia Odashiro¹, Gloria Tomarchio², Andréanne Gagné³, Michèle Orain⁴, Patrice Desmeules⁵, Philippe Joubert⁶

¹Université Laval, Boischatel, QC, ²Università degli Studi UMG, Catanzaro, Catanzaro, Italy, ³Laval University, Quebec City, QC, ⁴Institut Universitaire de Cardiologie et de Pneumologie de Quebec Research Center, Quebec, QC, ⁵IUCPQ - Quebec Heart and Lung Institute, Quebec, QC

Disclosures: Patricia Odashiro: None; Gloria Tomarchio: None; Andréanne Gagné: None; Michèle Orain: None; Patrice Desmeules: *Grant or Research Support*, Pfizer Canada; *Grant or Research Support*, Novartis; Philippe Joubert: None

Background: Introduction: Lung cancer is the deadliest cancer in North America. Neuroendocrine tumors account for 20-25% of lung malignancies. These comprise typical and atypical carcinoid tumors and high grade neuroendocrine carcinomas, including large cell neuroendocrine carcinomas and small cell lung carcinomas (SCLC). The latter present at an advanced stage in most of the cases, rendering them ineligible for surgical treatment. Unfortunately, currently available chemotherapy treatments are inefficient and highly toxic. In recent years, a strategy using an antibody targeting delta-like protein-3 (DLL3) as an entryway to expose SCLC tumor cells to a toxic substrate has been used with success. The effectiveness of these molecules seems closely related to the amount of DLL3 expressed by the tumoral cells. However, the prevalence of DLL3 expression, as well as the heterogeneity of expression in the same tumor and the corresponding metastases has not been fully evaluated. Objective: To evaluate the immunohistochemical (IHC) expression of DLL3 in SCLC.

Design: Material and Methods: 39 cases of SCLC were retrieved from our institution biobank. In 35 cases, the 2 most representatives FFPE blocks of the tumor were retrieved for IHC testing, and in 4 cases, 1 block. The staining was performed on the Ventana BenchMark Ultra PLC with the DLL3 sp347 clone. The evaluation of DLL3 staining was done by a pathology's resident on digitized slides. The cases were classified for DLL3 expression according to 4 different thresholds: <1%, 1-49%, 50-74%,> 75%.

Results: Results: The prevalence of DLL3 expression in SCLC is shown in Table 01.

Table 01. Expression of DLL3 in SCLC:

Threshold of DLL3 expression	n	Mean %
<1%	3	7,7
1-49%	9	23,1
50-74%	2	5,1
>75%	25	64,1
TOTAL	39	100

Conclusions: Conclusion: Our results demonstrate a high prevalence of DLL3 expression in SCLCs. Our data confirm the relevance of performing IHC validation and clinical-pathological correlation studies with the aim of better characterizing the DLL3 expression threshold as a predictive biomarker for DLL3 targeted agents.

1948 Independent and Combined Effects of CYP and GST Genotypes as Risk Modifiers in Non-Small Cell Lung Cancer

Anumesh Pathak¹, Surya Kant², Rahul Pandey¹, Vandana Tiwari¹, Nuzhat Husain³

¹Dr Ram Manohar Lohia Institute of Medical Sciences, Lucknow, Uttar Pradesh, India, ²Respiratory Medicine, King George Medical University, Lucknow, Uttar Pradesh, India, ³Dr Ram Manohar Lohia Institute of Medical Sciences, Lucknow, India

Disclosures: Rahul Pandey: None; Nuzhat Husain: None; Nuzhat Husain: None

Background: Genetic polymorphisms in key genes encoding enzymes involved in the bio-activation (Cytochrome p450 (CYP)) or detoxification (Glutathione S-transferase (GST)) of environmental carcinogens including tobacco specific nitrosamines are potential lung cancer risk factors. The frequencies of these variants and consequently their effects vary across ethnicities. The interactive effects with CYP-GST combination along with smoking have not been documented. Five single nucleotide variants and two homologous deletion variants associated with CYP and GST genes and their interactive effects with smoking in Non-Small Cell Lung Cancer (NSCLC) were investigated

Design: The case control study comprised 244 cases of histologically diagnosed NSCLC and 224 healthy controls. IEC approval and informed consent were obtained. Mean age of cases was 56.51±11.21+ 2SD, male: female ratio was 4.19, 91 were non-smokers and 153 smokers subdivided into 20 (47.7%), 20-40 (38.7%) and 40(13.6%) in pack-years. Histological subtypes included 180 adenocarcinoma (ADC), 57 squamous cell carcinoma (SCC) and 09 adeno-squamous cases. Majority were diagnosed in late stages with only 5 in stage II, 78.2% had lymph-node metastasis and 139 cases had distant metastasis. DNA was isolated from whole blood and genetic polymorphism analyses were determined by polymerase chain reaction coupled with restriction fragment length polymorphism followed by agarose and poly acrylamide gel electrophoresis for genetic variants. All the statistical analyses were performed with Graph Pad Instat Version 3.05 and SPSS version 16(Chicago, USA).

Results: Significantly high risk of NSCLC and subtype ADC was associated with variant CYP2A6, GSTT1 and GSTM1 (Table 1). Multiplicative interaction with higher risks between combinations of the variant genotypes of CYP and GST was evident- GSTM1/CYP2A6 (0del./variant)OR12.51(CI;5.11-30.38,p<0.001), GSTT1/ CYP2A6(0del/variant) OR6.59(CI,2.69-16.16,p<0.001), GSTM1/CYP2A13(0del variant)OR 14.48(CI;4.82(2.23-10.48,p<0.001) and GSTT1/ CYP2A13 (0del/variant)OR6.11(CI,2.73-13.60,p<0.001). Polymorphism in CYP and GST gene and smoking interaction, stratified for quantity, carried a modest increase in risk susceptibility to NSCLC. No correlation with stage, nodal status or presence of metastasis was observed

Table-1: Frequency of CYP and GSTs Gene Family Genotypes and Risk of NSCLC and Histological Subtypes

Genotypic Variant	wt/wt (%)	Variants#/ mutants (%)	OR#(95%CI)	p value	
CYP1A1 chr 15 p13 (rs1	7861094(T/C)): N	lissense Variant	•	•	
Controls(N=175)	34.29	65.71	1.7(1.11-2.27)	0.01 [*]	
Cases(N=217)	23.04	65.96			
SCC(N=55)	21.82	78.18	1.87(0.91-3.81)	0.1	
ADC(N=162)	23.46	76.54	1.7(1.05-2.74)	0.03*	
CYP1B1 chr 2,p13 (rs10					
Controls(N=163)	28.22	71.78	1.3(0.81-2.1)	0.3	
Cases(N=187)	22.99	77.01			
SCC(N=44)	20.45	79.55	1.52(0.68-3.43)	0.4	
ADC(N=143)	23.77	76.23	1.26(0.75-2.10)	0.5	
CYP2A6 chr 19,p13 (rs ²					
Controls(N=169)	37.87	62.13	4.2(2.4-7.2)	<0.001*	
Cases(N=182)	12.64	87.36			
SCC(N=45)	13.33	86.67	3.96(1.58-9.88)	0.003*	
ADC(N=137)	12.40	87.59	4.3(2.37-7.80)	<0.001*	
CYP2A13 chr 20 p13 (rs		Intron Variant			
Controls(N=161)	44.19	55.91	1.7(1.06-2.58)	0.03*	
CASES(N=177)	32.20	67.80			
SCC(N=49)	32.65	67.35	1.62(0.83-3.19)	0.2	
ADC(N=128)	46.88	53.12	0.89(0.56-1.42)	0.7	
GSTP1 chr 11q13(rs 169)5(A/G)) Ile105Va	/ Missense Variant			
Controls(N=197)	37.6	62.94	1.3(0.85-1.91)	0.3	
Cases(225)	31.55	68.44			
SCC(N=62)	33.87	66.13	1.15(0.63-2.09)	0.7	
ADC(N=163)	30.67	69.33	1.33(0.85-20.68)	0.2	
GSTT1 chr 22q11.2(226	637) homologou	us recombination(delet			
Controls(N=194)	92.78	7.22	3.9(2.07-7.26)	<0.001*	
Cases(N=220)	76.82	23.18			
SCC(N=57)	84.21	15.79	2.41(0.98-5.90)	0.08	
ADC(N=163)	74.23	25.77	4.46(2.34-8.53)	<0.001*	
		gous recombination(d			
Controls(N=212)	93.39	6.61	4.5(2.42-8.36)	<0.001*	
Cases(N=232)	75.86	24.14			
SCC(N=64)	76.56	23.44	4.3(1.95-9.56)	0.003*	
ADC(N=167)	74.85	25.15	4.7(2.49-9.05)	<0.001*	

Abbreviations: OR: odds ratio; CI: confidence interval; ADC: Adenocarcinoma; SCC: Squamous cell carcinoma; N:Total number; wt/wt: wild genotype; Variants: Hetrozygous+Mutant genotype; * p< 0.05 was considered statistically significant.p-values were derived from Chisquare test. All p-values are two-sided. Control wt/wt and cases wt/wt form reference category for calculating OR (95%CI) and p-value.

Conclusions: Functionally relevant polymorphisms in CYP and GST genes with gene-gene and gene-environment interactions play a significant role in modifying the susceptibility to NSCLC in population of Indian ethnicity.

1949 Intratumor Heterogeneity of Ki-67 is a Powerful and Histology-Independent Resource to Dissect Clinical Outcome of Lung Neuroendocrine Neoplasms by Means of Artificial Intelligence Tools

Giuseppe Pelosi¹, Matteo Bulloni², Mauro Papotti³, Jasna Metovic⁴, Gabriella Fontanini⁵, Linda Pattini⁶

¹University of Milan, Milan, Lombardia, Italy, ²Politecnico di Milano, Casalpusterlengo, Lombardy, Italy, ³University of Turin, Torino, Torino, Italy, ⁴Department of Oncology, Turin, Piedmont Region, Italy, ⁵Department of Surgical, Medical, Molecular Pathology and Critical Area, University of Pisa, Pisa, Italy, ⁶Politecnico di Milano, Milan, Lombardia, Italy

Disclosures: Giuseppe Pelosi: None; Matteo Bulloni: None; Mauro Papotti: None; Jasna Metovic: None; Gabriella Fontanini: None; Linda Pattini: None

Background: Little is known about the biological role of intratumor heterogeneity of the proliferation marker Ki-67 in predicting clinical outcome in lung neuroendocrine neoplasms (Lung-NENs), which encompass typical carcinoid (TC), atypical carcinoid (AC), large cell neuroendocrine carcinoma (LCNEC) and small cell lung carcinoma (SCLC) according to the 2015 WHO classification. Whether artificial intelligence tools may be useful, it is unclear.

Design: Thirty consecutive surgically resected Lung-NENs comprising 10 TC, 14 AC and six LCNEC, all with long-term follow-up, were immunohistochemically stained for Ki-67 and scanned at 40x (NanoZoomer XR, Hamamatsu, Japan). A tailored algorithm was constructed to recognize all Ki-67-stained tumor cells and the obtained patterns were described using spatial statistics, graph modeling, fractality and Shannon entropy parameters. A Support Vector Machine classifier with polynomial kernel was then trained, employing the 5 parameters that resulted most informative out of the 620 initially computed, to distinguish dead (true positive) from alive (true negative) patients.

Results: Over 100 repetitions of 5-fold cross-validation, the model averaged 84.3% diagnostic accuracy in the prediction of ultimate clinical outcome (dead vs alive) in the 30 Lung-NEN patients under evaluation, which resulted to be independent of WHO classification. The corresponding values of sensitivity, specificity, PPV and NPV were 73.6%, 89.6%, 78.2%, and 87.4%, respectively.

Conclusions: The intratumor heterogeneity of Ki-67 is a powerful and histology-independent resource to unravel clinical outcome of Lung-NENs by using machine learning algorithms.

1950 Utility of Methylthioadenosine Physphorylase (MTAP), 5-Hydroxymethylcystosine (5-hmc) and BAP1 Immunocytochemistry (ICC) in Cytology Specimens for the Diagnosis of Malignant Mesothelioma (MM)

Roshan Raza¹, William Travis¹, Prasad Adusumilli¹, Marina Asher¹, Darren Buonocore¹, Jason Chang¹, Denise Frosina¹, Achim Jungbluth¹, Irina Linkov¹, Natasha Rekhtman¹, Marjorie Zauderer¹, Marc Ladanyi¹, Jennifer Sauter¹

**Memorial Sloan Kettering Cancer Center, New York, NY*

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Background: Cytologic diagnosis of MM is challenging since atypia in mesothelial cells is not specific for malignancy and there is no architecture to assess for invasion. Loss of BAP1 expression by ICC and homozygous deletion of p16/CDKN2a by FISH are specific but not sensitive for MM. Co-deletion of MTAP occurs in most MM with p16/CDKN2A deletions and can be detected by ICC, which has advantages over FISH. Recently, 5-hmc has been reported to show 92% sensitivity and 100% specificity for distinguishing MM from benign mesothelial proliferations. To our knowledge, these 3 markers have not yet been studied in combination in cytology. Herein, we determine the sensitivities and specificities of BAP1, MTAP and 5-hmC ICC for the diagnosis of MM in cytology.

Design: ICC with MTAP, 5-hmC and BAP1 was performed on all available MM cytology specimens from 2017-present and 20 benign specimens with reactive mesothelial cells on cell block. ICC was scored as nuclear loss of BAP1 expression, loss or marked reduction of cytoplasmic MTAP expression, loss of nuclear expression of 5-hmc in at least 50% of tumor cells or non-contributory (NC) in cases without internal positive control. All available clinical next generation sequencing (NGS) data were collected.

Results: Cytology specimens (9 fluids, 3 FNA, 1 touch preparation) from 13 patients with diagnosis of MM confirmed by histology (11 epithelioid, 2 biphasic) contained adequate tumor cellularity on formalin-fixed cell blocks for study inclusion. All 20 cases with benign mesothelial cells showed retained BAP1 and MTAP expression and retained or <50% loss of 5-hmc expression. MM in 7, 3 and 6 patients showed BAP1 loss, MTAP loss and/or ≥50% loss of 5-hmc expression (sensitivities of 58, 23 and 46%), respectively. BAP1 ICC was NC in one MM specimen. While sensitivity of MTAP was low, MTAP loss was seen in 2 MM with retained BAP1 and either retained or <50% loss of 5-hmc expression. Combined sensitivity of all 3 markers was 77%. CDKN2A deletions were detected by NGS in all 3 MM with MTAP loss by ICC. MTAP ICC was retained in 2 MM with CDKN2A deletions detected by NGS.

Patient #	Specimen type	Site	Subtype	BAP1	MTAP	5-hmc	NGS: BAP1 mutation or CDKN2A deletion
1	Fluid	Pleural	Epithelioid	Lost	Retained	≥50% loss	BAP1
2	FNA	Lymph node	Biphasic	NC	Lost	Retained	BAP1, CDNK2A
3	Fluid	Pleural	Epithelioid	Retained	Retained	<50% loss	N/A
4	Fluid	Pleural	Epithelioid	Retained	Retained	≥50% loss	N/A
5	Fluid	Pleural	Epithelioid	Lost	Retained	≥50% loss	BAP1
6	FNA	Chest wall	Epithelioid	Retained	Retained	<50% loss	CDKN2A
7	FNA	Supra- umbilical mass	Epithelioid	Lost	Retained	<50% loss	No
8	Drainage	Abdomen	Epithelioid	Retained	Lost	<50% loss	CDNK2A
9	Touch preparation	Lung	Epithelioid	Lost	Lost	≥50% loss	BAP1, CDNK2A
10	Fluid	Pleural	Epithelioid	Lost	Retained	≥50% loss	BAP1, CDNK2A
11	Fluid	Pleural	Epithelioid	Retained	Retained	<50% loss	No
12	Fluid	Pleural	Epithelioid	Lost	Retained	Retained	N/A
13	Washing	Pelvic	Biphasic	Lost	Retained	≥50% loss	N/A

Abbreviations: FNA, fine needle aspiration; n, number; N/A, not available; NC, non-contributory; NGS, next generation sequencing

Conclusions: In this small cohort, the individual sensitivities of each marker for MM are low, but the combined sensitivity of all 3 markers is 77%, suggesting the use of these markers in combination or in an algorithmic fashion may support the diagnosis MM on cytology. Furthermore, in a subset of cases with retained BAP1, MTAP may be of benefit in reaching a diagnosis of MM.

1951 Association of ERBB4 Mutations with Advanced-Stage and TP53 Mutations in Lung Carcinoma

Rongqin Ren¹, Konstantin Shilo², Weiqiang Zhao², Matthew Avenarius³, Jason Garee³, Sean Caruthers², Dan Jones²
¹The Ohio State University Wexner Medical Center, Westerville, OH, ²The Ohio State University, Columbus, OH, ³The Ohio State University Wexner Medical Center, Columbus, OH

Disclosures: Rongqin Ren: None; Konstantin Shilo: None; Weiqiang Zhao: None; Matthew Avenarius: None; Jason Garee: None; Sean Caruthers: None; Dan Jones: None

Background: Signaling in the epidermal growth factor receptor (EGFR) family of receptor tyrosine kinases (HERs) is triggered by ligand-binding of the extracellular domain (ECD) followed by homo- and heterodimerization among HERs resulting in trans- and autophosphorylation of the tyrosine kinase domain (TKD). Targetable oncogenic mutations in lung cancer in *EGFR/HER1* and *ERBB2/HER2* have been shown to alter signaling crosstalk whereas *ERBB4/HER4* have been less-studied. Here, we assess the clinicopathologic and molecular features of *ERBB4*-mutated lung carcinoma.

Design: 1526 lung cancers over a 3-year period [~99% non-small cell lung carcinoma (NSCLC)] were assessed by a 50-gene amplicon-based next-generation sequencing (NGS) panel. Exons 3-9 (ECD) and exons 15 and 23 (TKD hotspots) of *ERBB4*, exons 19, 20 and 21 of *ERBB2* and exons 3,7,15 and 18-21 of *EGFR* were analyzed. The panel also included other genes in lung cancer including *AKT1*, *BRAF, KRAS, NOTCH1, PIK3CA, and TP53*.

Results: *ERBB4* mutation was detected in 27 cases (1.8%), compared to 15 cases with *ERBB2* mutations (1%) and 170 cases with *EGFR* mutations (11.1%). The tumors comprised of adenocarcinoma (17/27, 63.0%) and squamous cell carcinoma (6/27, 22.2%), majority presenting as stage IV disease (15/27, 55.6%). Missense mutations were predominant (22/27, 81.5%), with 19 of them located in ECD and 3 in TKD, with 5 nonsense mutations; the previously characterized R106C/H and L789R changes were not seen. A high rate of accompanying *TP53* mutations was noted (20/27, 74.1%) which was higher than the overall *TP53* mutation rate in this series (49.2%, p = .03, Fisher's exact), with *KRAS* mutation also commonly seen (8/27, 29.6%) (See Table). Two of three TKD-mutated cases also had concurrent activating *EGFR* mutations (an uncommon M600I exon 15 variant and a typical exon 19 in-frame deletion). No co-occurring *ERBB2* mutations were noted. *ERBB4* was the lone mutation detected in only 4 tumors.

The Distribution Of TP53 And KRAS Mutations In ERBB4 Positive Carcinomas

	TP53			
KRAS		+	-	
	+	3/27 (11.1%)	3/27 (11.1%)	
	-	17/27 (63.0%)	4/27 (14.8%)	

Conclusions: *ERBB4* mutation is typically seen in advanced-stage lung cancer, with a range of variants noted, mostly located in the ECD. The high frequency of co-occurring *TP53* mutations in *ERBB4*-mutated carcinoma may be associated with more aggressive clinical course or a *TP53*-related mutator phenotype. Unlike for *EGFR- and ERBB2*-mutated tumors, *ERBB4 and KRAS* are not mutually exclusive. The co-occurrence of *EGFR* mutations in a few cases with *ERBB4* TKD mutations suggests oncogenic signaling cross-talk can rarely occur and may be targetable in those scenarios.

1952 The Impact of Intratumoral Glucocorticoid Synthesis Through 11β Hydroxysteroid Dehydrogenase 1 on the Intratumoral Immune Microenvironment in Non-Small Cell Lung Carcinoma: A New Marker for Predicting Response to Immune Checkpoint Blockade Therapy in Non-Small Cell Lung Carcinoma

Ryoko Saito¹, Yasuhiro Miki², Jiro Abe³, Chihiro Inoue⁴, Ikuro Sato⁵, Hironobu Sasano⁶
¹Tohoku University Graduate School of Medicine, Sendai-Shi Aoba-Ku, Miyagi-ken, Japan, ²Disaster Ob/Gyn, Int Res Inst of Disaster Sci, Tohoku Univ, Sendai, Miyagi, Japan, ³Department of Thoracic Surgery, Miyagi Cancer Center, Natori, Miyagi, Japan, ⁴Tohoku University Hospital, Sendai, Miyagi, Japan, ⁵Department of Pathology, Miyagi Cancer Center, Natori, Miyagi, Japan, ⁶Tohoku University, Sendai-shi, Miyagi-ken, Japan

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Background: The status of the intratumoral immune microenvironment is important to guarantee the effect of immune checkpoint (IC) blockade therapy, which has been broadly used in patients with non-small cell lung carcinoma (NSCLC). Glucocorticoid (GC) is a hormone well-known to act strongly on the immune system. Therefore, we examined the correlation between intratumorally synthesized GC through 11β hydroxysteroid dehydrogenase (HSD) 1 and the immune microenvironment in NSCLC.

Design: We evaluated 125 surgical specimens from patients with NSCLC (95 adenocarcinoma and 30 squamous cell carcinoma), assessing mainly the immunoreactivity for 11βHSD1 and 11βHSD2 and the levels of tumor-infiltrating lymphocytes (TILs) and CD3- or CD8-positive T cells. Furthermore, we examined the correlations between 11βHSD1 immunoreactivity and the therapeutic efficacy of IC blockade therapy using nine biopsy specimens from patients with NSCLC. Subsequently, we explored the mechanisms of GC effects on the intratumoral immune microenvironment, focusing on cytokines.

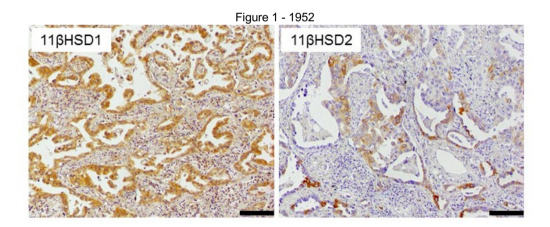
Results: 11βHSD1 immunoreactivity was significantly inversely correlated with the numbers of intratumoral TILs, CD3-positive T cells, and CD8-positive positive T cells. Additionally, we found 11βHSD1 immunoreactivity tended to be inversely correlated with the efficacy of the IC blockade therapy. According to the in vitro study, GC reduced the expression of cytokines such as IL-8 and IL-6, resulting in an inhibition of monocytes migration. Furthermore, production of cortisol, active GC, was confirmed in the cell lines expressing 11βHSD1.

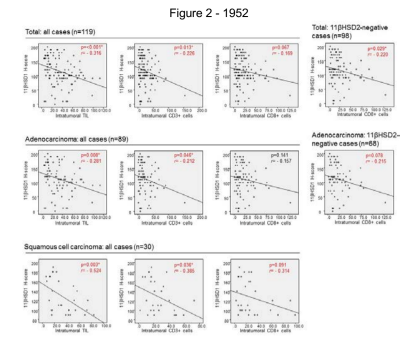
Table: The correlation between immunoreactivity of 11BHSD1 and therapeutic efficacy of IC blockade therapy

	The percentage of strongly stained cells for 11βHSD1				
	<70% (n=3) ≥70% (n=6) p value				
PR (n=5)	3	2	0.058		
PD (n=4)	0	4			

^{*}p value <0.05, significant.

11βHSD1: 11β hydroxysteroid dehydrogenase 1, PR: Partial remission, PD: Progressive disease.





Conclusions: This is the first study demonstrating the significant inhibitory effects of intratumorally synthesized GC through 11β HSD1 on tissue immune microenvironment in NSCLC, and the possible correlation of intratumoral 11β HSD1 status with the efficacy of IC blockade therapy. Our results provided new insights into the therapeutic strategies and the efficacy prediction of IC blockade therapy.

1953 Comparison of Metastatic Malignant Mesothelioma to Primary Site: A Multi-Institutional Analysis

Jefree Schulte¹, Aliya Husain¹, Yin Hung², Mari Mino-Kenudson², Andrew Nicholson³, Yu Zhi Zhang⁴, Jeffrey Mueller⁵

¹The University of Chicago, Chicago, IL, ²Massachusetts General Hospital, Boston, MA, ³Royal Brompton Hospital, London,

United Kingdom, ⁴Royal Brompton & Harefield NHS Trust, London, United Kingdom, ⁵University of Chicago Medical Center,

Chicago, IL

Disclosures: Jefree Schulte: None; Aliya Husain: None; Yin Hung: None; Mari Mino-Kenudson: None; Andrew Nicholson: None; Yu Zhi Zhang: None; Jeffrey Mueller: None

Background: Although malignant mesothelioma (MM) typically presents with locoregional disease, it does have the capability to metastasize. Albeit uncommon, a metastatic site may be the first diagnostic specimen in a patient presenting with MM. Recent studies highlight the prognostic significance of histopathologic parameters (histologic subtype, nuclear grade [NG], presence of necrosis) in MM. These prognostic factors may be useful to the treating physician in crafting treatment plans. This study seeks to compare these parameters between metastatic and primary sites in MM.

Design: Paired cases of pleural MM with metastatic and primary sites were identified from the pathology archives at the participating institutions with review of H&E stained sections. Histologic subtype was noted in all cases. For epithelioid MM, NG (1, 2, or 3, as previously described in the literature by Kadota et. al.) and necrosis were determined.

Results: 60 paired cases were identified with the primary site subtype comprised of 36 epithelioid, 24 biphasic, and 0 sarcomatoid cases. 57 of 60 (95%) metastases showed epithelioid morphology; 3 (5%) metastases were biphasic. The positive predictive value of epithelioid subtype at metastatic site was 63%, sensitivity 100%, and specificity 88%. NG and the presence or absence of necrosis at metastatic sites were not correlative with primary site histologic subtype (p=0.34 and p=0.78, respectively). Thirty-three pairs of metastases and primaries with epithelioid morphology were graded; 3 metastases were too small to grade. Pairs were more likely to show a higher NG at primary rather than metastatic sites (p<0.01) with 6 of 8 (75%) metastatic sites with NG 1 showing NG 2 at primary site, and 3 of 16 (19%) metastatic sites with NG 2 showing NG 3 at primary site. All metastases with NG 3 had NG 3 at primary site. There were no pairs with lower NG at primary site than metastatic site. The presence or absence of necrosis at metastatic site was not predictive of the presence or absence of necrosis at primary site (p=0.39).

Conclusions: The application of pathologic parameters to metastatic MM may not accurately predict the parameters at primary site. While sensitive, epithelioid morphology at metastatic site is not specific. Metastatic sites may underestimate nuclear grade at the primary site. The presence of necrosis in a metastatic site may not be predictive of necrosis at the primary site. Biopsy of the primary site may be required to more accurately classify the tumor.

1954 Immunohistochemical Biomarkers in Biphasic Malignant Mesothelioma: A Multi-Institutional Analysis

Jefree Schulte¹, David Chapel², Kenzo Hiroshima³, Yin Hung⁴, Sonja Klebe⁵, Kenneth Lee⁶, Mari Mino-Kenudson⁴, Prodipto Pal⁷, Ming Tsao⁸, Aliya Husain¹

¹The University of Chicago, Chicago, IL, ²Brigham and Women's Hospital, Boston, MA, ³Chiba University Graduate School of Medicine, Chiba, Chiba, Japan, ⁴Massachusetts General Hospital, Boston, MA, ⁵Flinders University and SA Pathology, Bedford Park, SA, Australia, ⁶Concord Repatriation General Hospital, Sydney, NSW, Australia, ⁷University Health Network, Mississauga, ON, ⁸Princess Margaret Cancer Centre, Toronto, ON

Disclosures: Jefree Schulte: None; David Chapel: None; Kenzo Hiroshima: None; Yin Hung: None; Sonja Klebe: None; Kenneth Lee: None; Mari Mino-Kenudson: None; Prodipto Pal: None; Ming Tsao: None; Aliya Husain: None

Background: Recent advancements in immunohistochemistry (IHC) in malignant mesothelioma (MM) include nuclear loss of BRCA-associated protein 1 (BAP1) and cytoplasmic loss of methylthioadenosine phosphorylase (MTAP) as specific markers of malignancy. GATA3 is described as a useful marker in distinguishing sarcomatoid MM and sarcomatoid carcinoma of the lung. This study explores the immunoreactivity of these markers in biphasic (BP) MM.

Design: Cases of BPMM were collected from 6 institutions. Histologic diagnosis and IHC for BAP1 and GATA3 were performed at each institution. A subset of cases was studied utilizing MTAP IHC at one institution. The immunoreactivity (retained or lost for BAP1 and MTAP; positive or negative for GATA3) was recorded for epithelioid (E) and sarcomatoid (S) components. The S component of a small subset was graded as low (LG) or high grade (HG); survival analysis was performed on this subset.

Results: Of 100 cases of BPMM, 94 were tested for BAP1. Seventy-eight (83%) showed concordant retention (n=45) or loss (n=33) in E and S components. Sixteen (17%) were discrepant: E lost/S retained (n=12) or E retained/S lost (n=4). Eighty-six cases were tested for GATA3. Seventy (81%) were concordantly positive (n=51) or negative (n=19) in E and S components. Sixteen (19%) were discrepant: E positive/S negative (n=8) or E negative/S positive (n=8). Twenty-six cases were tested for MTAP. Sixteen (62%) showed concordant retention (n=12) or loss (n=4) in E and S components. Ten (38%) were discrepant: E lost/S retained; no cases of E retained/S lost. Thirty-one cases were graded (24 LG, 7 HG). Median survival for LG and HG was 18.3 and 17.7 months, respectively (p=0.39). Seven (32%) and 8 (38%) of LG cases showed discrepant staining for BAP1 and MTAP, respectively. No HG case was discrepant for BAP1; 2 (40%) were discrepant for MTAP, which was similar to LG cases (p=0.66).

	E retained/S retained	E lost/ S retained	E retained/S lost	E lost/S lost
BAP1	45	12	4	33
low grade	13	7	0	2
high grade	4	0	0	1
	E positive/S positive	E negative/ S positive	E positive/S negative	E negative/S negative
GATA3	51	8	8	19
	E retained/S retained	E lost/ S retained	E retained/S lost	E lost/S lost
MTAP	12	10	0	4
low grade	10	8	0	3
high grade	2	2	0	1

Conclusions: Although the majority of BPMM showed concordance between E and S components, a subset showed discrepant staining. Although not statistically significant, discrepancy was more prevalent in BPMM with LG S component. Cases with discrepant BAP1 or MTAP (E lost/S retained) or LG morphology, may represent two tumor clones or E MM with reactive stroma; further study is needed. GATA3 was positive in the majority of BPMM and may remain a useful marker for S MM.

1955 Mutational Analysis in Pulmonary Adenocarcinomas with PD-L1 Expression

Reza Setoodeh¹, Ann Walts¹, Elias Makhoul¹, Alberto Marchevsky² ¹Cedars-Sinai Medical Center, Los Angeles, CA, ²Cedars-Sinai Medical Center, West Hollywood, CA

Disclosures: Reza Setoodeh: None; Ann Walts: None; Elias Makhoul: None; Alberto Marchevsky: None

Background: Immunotherapy has dramatically changed the treatment landscape of various malignancies including lung adenocarcinomas. The 2019 NCCN guidelines initially recommended single agent immunotherapy as a first line treatment option for advanced lung adenocarcinoma with PD-L1 expression levels of 50% or greater. However, given the recent data suggest that PD-L1 monotherapy is less effective in patients with EGFR or ALK gene alterations, the NCCN panel recently deleted the recommendation for subsequent immunotherapy in these patients.

Design: We retrospectively analyzed 124 cases of lung adenocarcinoma with PD-L1 expression (>1% expression) using PD-L1 IHC 22C3 PharmDx test and correlated PD-L1 expression with the presence of EGFR mutations or ALK gene rearrangement in the 120 cases where all 3 tests were performed. PD-L1 expression levels were subcategorized as low expression level (1%-49% of the tumor cells express PD-L1) and high expression level (50%-100% of the tumor cells express PD-L1).

Results: High expression levels of PD-L1 (50%-100%) were observed in 40% (50/124) of the cases. Among the 5 (4%) tumors that harbored ALK gene rearrangement, 4 showed high PD-L1 expression. Among the 31 (26%) tumors with detected EGFR mutations, only 6 showed high PD-L1 expression. These two alterations were mutually exclusive. Results are summarized in the table 1.

PD-L1 expression level	Total Cases	EGFR mutations	ALK rearrangement	EGFR or ALK positive
Low (1%-49% expression)	74/124 (60%)	25/120 (20.8%)	1/120 (0.8%)	26/120 (21.6%)
High (50%-100% expression)	50/124 (40%)	6/120 (5%)	4/120 (3.3%)	10/120 (8.3%)
All PD-L1 expressors (1%-100%)	120 cases	31/120 (25.8%)	5/120 (4.1%)	36/120(30%)
with molecular and FISH results				

Conclusions: In our study cohort 40% (50/124) of lung adenocarcinomas with PD-L1 expression showed high (50%-100%) expression levels. 20% of these high PD-L1 expressors were positive for either EGFR or ALK alterations: 8% (4/50) harbored ALK gene rearrangement and 12% (6/50) showed EGFR mutations. We corroborated the previously reported association of ALK gene rearrangement with high expression levels, and EGFR mutations with low expression levels of PD-L1 in lung adenocarcinomas that express PD-L1. Our study showed a substantial number (30%) of lung adenocarcinomas that express (>1%) PDL-1 have EGFR mutations or ALK gene rearrangement, suggesting that it is important to consider the results of these tests simultaneously in order to be able to stratify patients according to current NCCN therapeutic guidelines.

1956 Heterogeneity of PD-L1 Expression in Non-Small-Cell Lung Cancer

Xuxia Shen¹, Yuan Li²

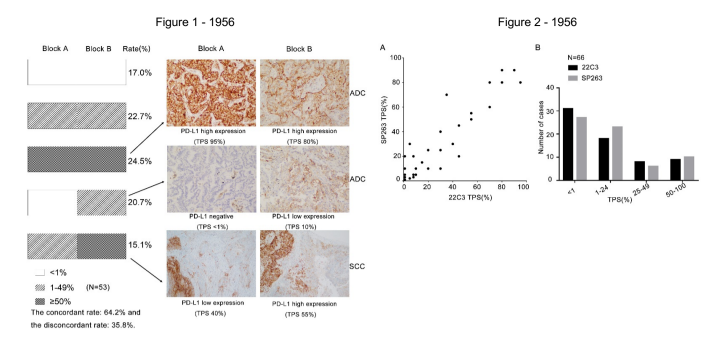
¹Fudan University Shanghai Cancer Center; Shanghai Medical College, Fudan University, Shanghai, China, ²Fudan University Shanghai Cancer Center, Shanghai, China

Disclosures: Xuxia Shen: None; Yuan Li: None

Background: PD-L1 is a predictive marker of anti-PD-1/PD-L1 therapies for non-small cell lung cancer (NSCLC). Heterogeneous PD-L1 expression may cause dilemmas in anti-PD-1/PD-L1 therapies when faced with discrepant biomarker results. Our aim was to comprehensively analyze the heterogeneity of PD-L1 expression defined as intratumoral area, paired samples and clones of anti-PD-L1 antibody to optimize tumor sampling and improve its accuracy.

Design: We selected 1002 NSCLC surgically resected specimens, 54 cell block and 73 biopsy specimens. We analyzed the associations of PD-L1 expression with histopathological characteristics, assessed the heterogeneity between paired cell block and biopsy samples (n=54), paired biopsy and resected samples (n=19), paired two blocks of the same resected sample (n=53), paired primary and metastatic lesions (n=29), and compared the consistency of clones of PD-L1 antibody (22C3 and SP263, n=66).

Results: High PD-L1 expression (TPS≥50%) was observed in 10.2% (102/1002) of the resected NSCLC, including in 7.2% (61/852) of adenocarcinoma(ADC), and 27.3% (41/150) of squamous cell carcinoma(SCC). High PD-L1 expression was significantly associated with solid (41/138, 29.7%)/ micropapillary (7/28, 25.0%) ADC and non-keratinizing SCC (31.0%,40/129)(p<0.05). A total of 13.0% (7/54) PD-L1 discordance cases were observed between paired cell block and biopsy specimens (kappa=0.773). The cell block overestimated or underestimated PD-L1 status in 5 of 54 (9.3%) and 2 of 54 (3.7%) of the paired biopsy specimens (p=0.644). A total of 31.6% (6/19) PD-L1 discordance cases were observed between paired biopsy and resected samples (kappa=0.521). The biopsy specimen overestimated or underestimated PD-L1 status in 4 of 19 (21.1%) and 2 of 19 (10.5%) of the matched resected specimens (p=0.368). A total of 19 of 53(35.8%) PD-L1 discordance cases were observed between different paraffin blocks of the same resected specimens (kappa=0.455). A total of 27.6% (8/29) PD-L1 discordance cases were observed between primary and matched metastatic lesions(kappa=0.550). High concordance of 92.4% (61/66) was observed for PD-L1 expression between clone 22C3 and SP263(kappa=0.877).



Conclusions: We found significant heterogeneity of PD-L1 expression in different subtype of ADCs and SCCs, in primary and metastatic lesions and in different tissue samples. Such discrepancies were mainly due to intratumoral heterogeneity of PD-L1 expression and association with histopathological features.

1957 Diffuse Alveolar Injury with Delayed Epithelialization (DAIDE) in Previously Healthy Patients with Idiopathic Acute Respiratory Distress Syndrome After Trimethoprim-Sulfamethoxazole Exposure Angela Shih¹, Martin Taylor², Jenna Miller³, Jennifer Goldman³, Mari Mino-Kenudson¹

¹Massachusetts General Hospital, Boston, MA, ²Boston, MA, ³Children's Mercy Hospital, Kansas City, MO

Disclosures: Angela Shih: None; Martin Taylor: None; Jenna Miller: None; Jennifer Goldman: None; Mari Mino-Kenudson: None

Background: Recently, we identified two young adults with idiopathic acute respiratory distress syndrome (ARDS) histologically characterized by diffuse alveolar injury with marked alveolar denudation, a novel pattern termed DAIDE. Both patients were exposed to trimethoprim-sulfamethoxazole (TMP-SMX). Here, we report four additional patients with DAIDE following TMP-SMX exposure and detail the features of all six patients.

Design: The original 2 patients were identified through routine surgical pathology, and an additional 4 patients were identified in the consultation archives. DAIDE was identified on surgical biopsy (n=4), autopsy (n=1), or both (n=1). Six patients with DAD on surgical biopsy were used for comparison. Clinical information was obtained from the medical record. H&E and IHC for AE1.3/CAM5.2, CD68, and CK5/6 were reviewed.

Results: Six otherwise healthy patients (M:F=1:5, median age: 21 years; age range: 10-37 years) initially presented with URI symptoms that rapidly progressed to ARDS; extensive infectious and rheumatologic workups were negative. No patients had a history of vaping. All patients were intubated and placed on ECMO for a median of 93 days (range: 1 to 190 days). Surgical biopsy was performed in 5 patients (6 days before to 10 days after intubation), and autopsy was performed in 2 patients (12 and 466 days after intubation). All 5 surgical biopsies showed marked alveolar denudation with histiocytes replacing alveolar lining and nested peribronchiolar metaplasia, consistent with DAIDE. Hyaline membranes were rare when present (n=3). The 2 autopsies showed patchy alveolar-filling fibrosis and reepithelialization of most alveoli, with focal areas of alveolar denudation and histiocytic lining, suggestive of partially resolving DAIDE. In comparison, control surgical DAD cases showed prominent hyaline membranes, only focal alveolar denudation, and no significant histiocytic lining. On follow-up, DAIDE patients underwent bilateral lung transplant (n=2); overall, 3 patients are deceased (n=3).

Conclusions: DAIDE appears to be a novel variant of DAD with a rapid and severe clinical course involving previously healthy patients and is characterized by extensive alveolar denudation with a lining replaced by histiocytes and a paucity of hyaline membranes. Although

our patients share a history of TMP-SMX exposure, the role of TMP-SMX in DAIDE is currently unclear. Nonetheless, patients with DAIDE appear to have a poor prognosis.

1958 Correlation of MTAP Immunohistochemistry with CDKN2A and MTAP Genomic Alterations in Malignant Mesothelioma

Lynette Sholl¹, Adrian Dubuc¹, Jason Hornick², David Chapel¹

¹Brigham and Women's Hospital, Boston, MA. ²Brigham and Women's Hospital, Harvard Medical School, Boston, MA

Disclosures: Lynette Sholl: *Advisory Board Member*, LOXO Oncology; Adrian Dubuc: None; Jason Hornick: *Consultant*, Eli Lilly; *Consultant*, Epizyme; David Chapel: None

Background: Immunohistochemistry (IHC) for methylthioadenosine phosphorylase (MTAP) was recently shown to be a specific marker of *CDKN2A* homozygous deletion and thus for diagnosis of malignant mesothelioma (MM). However, direct comparisons of MTAP IHC to *MTAP* gene alterations are lacking.

Design: MTAP IHC (clone 42-T, Santa Cruz) was performed on 34 MM. *MTAP* and *CDKN2A* genomic alterations were examined for each tumor: 28 tumors were tested by hybrid capture next-generation sequencing (NGS), 10 by single nucleotide polymorphism microarray, and 4 by both methods.

Results: The cohort included 20 epithelioid, 11 biphasic, and 3 sarcomatoid MM. Four tumors showed homozygous *MTAP* deletion, 14 heterozygous *MTAP* deletion, and 16 neutral *MTAP* copy number. No *MTAP* mutations were identified. All 18 cases with homozygous or heterozygous *MTAP* deletion also had a *CDKN2A* deletion, whereas 4 cases with homozygous or heterozygous *CDKN2A* deletion had neutral *MTAP* copy number.

CDKN2A copy number, MTAP copy number, and MTAP IHC were not significantly related to histotype in this cohort. By IHC, MTAP was completely lost in 13 tumors. MTAP expression was at least partially retained in 21 tumors, of which 7 showed diminished MTAP staining relative to internal controls and 7 showed discrete foci of retained and lost MTAP expression (heterogeneous staining). No tumors with neutral CDKN2A copy number showed MTAP loss by IHC. Similarly, all 16 tumors with neutral MTAP copy number retained MTAP by IHC (although 3 had diminished staining and 5 had heterogeneous staining). Of 22 tumors with homozygous or heterozygous CDKN2A deletion, 13 showed MTAP loss by IHC. Of 14 tumors with heterozygous MTAP deletion, MTAP expression was lost in 10 and retained but diminished in 4. Of 4 tumors with homozygous MTAP deletion, MTAP expression was lost in 3 and only focally retained in 1.

MTAP Copy Number vs MTAP IHC				
	MTAP Copy Number			
	Neutral	Homozygous Deletion		
IHC retained	13	0	1 (very focal)	
IHC retained (diminished)	3	4	0	
IHC lost	0	10	3	
CDKN2A Copy Number vs N	ITAP IHC			
	CDKN2A Copy Number			
	Neutral	Heterozygous Deletion	Homozygous Deletion	
IHC retained	10	2	2	
IHC retained (diminished)	2	2	3	
IHC lost	0	10	3	

Conclusions: *CDKN2A* and *MTAP* gene copy number status is usually but not always correlated. MTAP IHC correlates well with *MTAP* copy number in tumors with normal copy number or homozygous deletion. Tumors with heterozygous *MTAP* deletion generally show MTAP loss by IHC but may show diminished staining. *MTAP* copy number did not consistently predict heterogeneous or diminished MTAP expression. Additional studies will be needed to establish the diagnostic specificity of heterogeneous MTAP expression, which may reflect clonal heterogeneity or epigenetic mechanisms of gene silencing.

Pulmonary Neuroendocrine Cell Hyperplasia (PNECH); 10-Year Experience at a Single Institution Snehal Sonawane¹, Daniel Dresser², Swati Mehrotra³, Gerald Camren³, Bradford Bemiss³, Stefan Pambuccian³, Vijayalakshmi Ananthanarayanan³

¹LUMC, Lombard, IL, ²Loyola University Medical Center, Chicago, IL, ³Loyola University Medical Center, Maywood, IL

Disclosures: Snehal Sonawane: None; Swati Mehrotra: None; Bradford Bemiss: None; Stefan Pambuccian: None; Vijayalakshmi Ananthanarayanan: *Advisory Board Member*, Boehringer Ingelheim

Background: The entity diffuse idiopathic neuroendocrine hyperplasia (DIPNECH) is a clinical syndrome defined by the WHO as a generalized proliferation of neuroendocrine cells either confined to the bronchial mucosa or resulting in formation of carcinoid tumorlets or tumors. When thus defined, DIPNECH is a diagnostic entity seen in clinically symptomatic patients presenting with histologic alterations of neuroendocrine proliferations combined with radiological features of airtrapping and multiple bilateral nodules. The significance of incidentally discovered neuroendocrine cell hyperplasia in surgically resected specimens and its correlation to radiological and clinical features is not known. This study aims to characterize the clinicopathological and radiological patterns identified in patients with histologically identified diffuse neuroendocrine cell hyperplasia (NECH).

Design: We searched the pathology database starting Jan, 2009 till Sept, 2019. for the for the combination of the following keywords "diffuse" and "neuroendocrine cell hyperplasia (NECH)" and "lung" and identified ten cases that met the search criteria. The radiological data was reviewed by one thoracic radiologist in a blinded fashion. Clinical, radiological and pathological features of these patients are depicted in Table 1.

Results: There was female predominance in our study (9/10) and all patients were above the age of 45 years. 70% (7/10) of the NECH cases were diagnosed incidentally during imaging workup or postoperative surveillance for other malignancies. The remaining 3 cases were biopsied for presentation of respiratory symptoms and or radiological finding of interstitial lung disease. Radiologically, bilateral involvement of lung was seen in 9 out of 10 cases with air trapping in 2 cases. The radiological diagnosis of DIPNECH could be made only in three cases, one of which was incidentally discovered (Case 5). Nine cases showed presence of carcinoid tumor or tumorlet along with NECH and 4 cases showed granulomas along with NECH.

		C	haracteristics of the patients at the time of diagnosis	of DIPNECH			
Case number	Age	Sex	Clinical History/ Reason for investigation	Radiology findings, Bilateral(BL)/ unilateral (UL); Single (S)/ Multiple (M) nodules	Carcinoid present/ absent	Granuloma present/ absent	
1	49	F	Shortness of breath and cough with abnormal chest radiology	BL-M	Present	Present	
2	81	F	B-cell lymphoproliferative disorder; follow up	BL-M	Present	Present	
3	52	M	H/o Kartagener's syndrome, underwent transplant	UL-M	Present	Absent	
4	61	F	Fever, cough, shortness of breath, Chronic obstructive pulmonary disease	BL-M	Present	Absent	
5	56	F	MEN -1 syndrome and gastrinoma; follow up	BL-M	Present	Absent	
6	73	F	Dyspnea, Shortness of breath, Suspicious for Interstitial lung disease	BL-M	Absent	Present	
7	66	F	Adenocarcinoma lung; follow up	BL-M	Present	Absent	
8	75	F	Breast nodule follow up	BL-M	Present	Absent	
9	63	F	Chronic cough, carcinoma breast follow up	BL-M	Present	Present	
10	73	F	Carcinoma breast follow up	BL-M	Present	Absent	

Conclusions: In conclusion, in this study, vast majority of patients with NECH were incidentally discovered during surveillance and follow up imaging for an unrelated malignancy or nodules. None of these incidentally discovered patients were clinically symptomatic. The criteria for evaluation of NECH in incidentally discovered asymptomatic patients is not well defined and necessitates further investigation and appropriate follow-up for an evolving interstitial lung disease.

1960 Correlation of Gamma-Delta T-Lymphocytes and Tumor-Associated Neutrophils with Clinicopathologic, Immune, and Molecular Features in Resected Lung Adenocarcinomas

Elisabeth Tabb¹, Daniel Shepherd¹, Mingjuan Zhang¹, Marina Kem¹, Mari Mino-Kenudson¹, Yin Hung¹ ¹Massachusetts General Hospital, Boston, MA

Disclosures: Elisabeth Tabb: None; Daniel Shepherd: None; Mingjuan Zhang: None; Marina Kem: None; Mari Mino-Kenudson: None; Yin Hung: None

Background: Recent studies have implicated local microbiota in activating gamma-delta T-lymphocytes (gdTL) and subsequently inducing neutrophilic infiltration to promote oncogenesis in murine models of *KRAS*-mutant lung adenocarcinoma. This study aimed to examine the translational relevance of these preclinical findings by evaluating gdTL and tumor-associated neutrophils (TAN) in human lung adenocarcinomas, including those with and without *KRAS* mutations.

Design: We quantified the numbers of gdTL and TAN using immunohistochemistry for T cell receptor-gamma chain and myeloperoxidase, respectively, from 3 high-power fields (HPF; each 0.24 mm²) of viable tumor areas on tissue microarray (duplicate of 2-mm core) sections of 236 human lung adenocarcinomas resected in 2010-2012. High and low gdTL or TAN were defined as above or below the respective median. Data were correlated with demographics, histologic features (AJCC 8th edition), immune parameters (tumoral PD-L1 expression, CD8+ tumor-infiltrating lymphocytes [TIL]), molecular alterations (using a multiplex-PCR based assay), and outcome via chi-square or logrank tests as appropriate (significance: p<0.05).

Results: gdTL ranged 0-177 (median 3.3) per HPF (in 228 tumors with evaluable data); whereas TAN ranged 0-140 (median 1.3) per HPF (in 223 tumors with evaluable data). The numbers of gdTL and TAN correlated with each other (p<0.0001). Both high gdTL and high TAN were associated with smoking history (p<0.01; p<0.05), solid/high-grade acinar histologic patterns (p<0.0001; p<0.001), the presence of tumor necrosis (p<0.05; p<0.001), and elevated CD8+ TIL (p<0.001; p<0.05). High TAN - but not high gdTL - was also associated with greater total tumor size (p<0.01), invasive size (p<0.05), advanced stages 3-4 (p<0.05), and worse progression-free survival (P<0.05), with a trend toward worse overall survival (P=0.05). However, gdTL and TAN were not associated with tumoral PD-L1 expression or molecular alterations including *KRAS* or *EGFR*.

Conclusions: In human lung adenocarcinomas, we identified correlations of gdTL and TAN with smoking history, aggressive histology, and elevated CD8+ TIL, but no associations with mutation status. TAN also appeared to be associated with worse patient outcome. Increased gdTL and TAN may reflect aggressive tumor biology; our data also suggests that their effects may be more general and not restricted to *KRAS*-mutant tumors.

1961 Association Between PD-L1 Expression and c-Met Alterations in Non-Small Cell Lung Carcinoma Troy Tenney¹, Gabriel Sica², Debra Saxe³, Taofeek Owonikoko³, Suresh Ramalingam³, Linsheng Zhang² 1 Emory University, Atlanta, GA, 2 Emory University Hospital, Atlanta, GA, 3 Emory University School of Medicine, Atlanta, GA

Disclosures: Troy Tenney: None; Gabriel Sica: None; Debra Saxe: None; Taofeek Owonikoko: None; Linsheng Zhang: None

Background: PD-L1 expression in non-small cell lung cancer (NSCLC) is used as a biomarker to treat patients with PD1 blockade therapy. PD-L1 expression may be related to tumor stromal interactions guided by underlying genotypic/phenotypic characteristics of a tumor and tumor antigenicity. Increasing evidence has shown that c-Met pathway activation in coordination with interferon gamma can lead to PD-L1 upregulation. In order to determine whether the underlying genomic characteristics of a tumor, including c-Met status is associated with PD-L1 status, we analyzed lung NSCLC that had been genomically characterized and correlated with PD-L1 status.

Design: Immunohistochemical staining for PD-L1 was performed with clone 22C3 on a Dako Autostainer and scored as no expression (<1%), low expression (1-49%) or high expression (50% or greater). Flourescent in situ hybridization was performed for ALK, RET and ROS1 (break apart probes) and c-Met amplification/polysomy. Mutations were detected by next generation sequencing of a 26 gene panel (CMP26, based on Illumina TST26) or multiplex hotspot mutation assay (SNaPshot, 10 gene panel).

Results: We evaluated 103 lung NSCLC (85 adenocarcinomas/favor adenocarcinoma, 7 squamous cell carcinomas, 8 not otherwise specified, 2 adenosquamous and 1 combined large cell neuroendocrine carcinoma with adenocarcinoma). No PD-L1 expression was seen in 34% (N=35) while PD-L1 expression was seen in 66% (N=68) with low expression in 25% (N=26) and high expression in 41% (N=42). Among the no expression cohort there was a slight increase in EGFR mutated patients as compared to PD-L1 expression cohort (23% vs. 13%) while similar frequencies in KRAS mutations were seen in both cohorts (no expression 40%; PD-L1 expression 38%). c-Met alterations in the form of amplification, high polysomy and mutations (exon 14 skip mutations and juxtamembrane mutations) were associated with PD-L1 positive status (no PD-L1 expression 9% (n=3) vs expression 25% (n=17)), p=0.05), especially when there is high PD-L1 expression (29% (n=12); p=0.03). Only in the no PD-L1 expression category was concurrent EGFR mutation and c-Met amplification/polysomy observed (2 of 3).

Conclusions: Higher PD-L1 expression is associated with c-Met amplification, polysomy and mutations. These results indicate that patients with abnormalities in c-Met may benefit from combined inhibition of PD-1 pathway and c-Met pathway.

1962 Loss of MTAP by Immunohistochemistry is Common in Pulmonary Sarcomatoid Carcinoma as well as Sarcomatoid Mesothelioma

Simone Terra¹, Anja Roden¹, Eunhee Yi¹, Marie-Christine Aubry¹, Jennifer Boland¹ Mayo Clinic, Rochester, MN

Disclosures: Simone Terra: None; Anja Roden: None; Eunhee Yi: None; Marie-Christine Aubry: None; Jennifer Boland: None

Background: Differentiating malignant pleural mesothelioma from reactive mesothelial processes can be quite challenging. Ancillary tests such as BAP1 immunohistochemisry (IHC) and p16 fluorescence in situ hybridization (FISH) are very helpful tools to aid in this distinction. IHC for MTAP has recently been proposed as an effective surrogate marker for p16 FISH, and it is an attractive alternative test due to shorter turn-around time. There is little data regarding the specificity of MTAP IHC for mesothelioma, or whether it may be useful to distinguish mesothelioma from other entities in the differential diagnosis. While there are many reliable markers to distinguish epithelioid mesothelioma from adenocarcinoma, this is not true of sarcomatoid mesothelioma, which can be very difficult to distinguish from sarcomatoid carcinoma. The goal of this study was to determine if MTAP loss is present in pulmonary sarcomatoid carcinoma or only in sarcomatoid mesothelioma.

Design: Well-characterized cases of sarcomatoid carcinoma (n=35) and sarcomatoid mesothelioma (n=62) were included; diagnoses were confirmed by two thoracic pathologists with incorporation of immunophenotype, clinical and radiographic features. Each case was stained for MTAP (clone 2G4) and BAP1 (clone C-4). Successful staining was confirmed by presence of internal positive control for both stains.

Results: Loss of MTAP expression by IHC was observed in 18 of 35 pulmonary sarcomatoid carcinomas (51%); 32 of these cases also had successful BAP1 staining performed, which was retained in all cases. MTAP expression was lost in 38 of 62 sarcomatoid mesotheliomas (61%); BAP1 was successful in all 62 cases, and showed loss in 6 (10%). In the 6 cases of sarcomatoid mesothelioma with BAP1 loss, 5 also had loss of MTAP, while MTAP expression was retained in 1 case.

Conclusions: Loss of MTAP expression by IHC is common in pulmonary sarcomatoid carcinoma, present in half of cases. This may reflect homozygous p16 deletion, which has been described in a few cases of sarcomatoid carcinoma studied by FISH analysis. This rate is similar to what is observed in sarcomatoid mesothelioma (61%). Therefore, MTAP loss is not specific for mesothelioma, and this stain is not useful to distinguish between these two malignancies. MTAP loss is more common than BAP1 loss in the setting of sarcomatoid mesothelioma (61% vs 10%, respectively).

1963 Novel Technique for Mutation-Specific Monoclonal Antibody Generation: MET Exon 14 Skipping in Lung Adenocarcinomas

Basile Tessier-Cloutier¹, Andrea Beharry², Deepu Alex³, Parneet Cheema², Stephen Yip¹, Brandon Sheffield²

¹University of British Columbia, Vancouver, BC, ²William Osler Health System, Brampton, ON, ³BC Cancer, Vancouver, BC

Disclosures: Basile Tessier-Cloutier: None; Andrea Beharry: None; Deepu Alex: None; Parneet Cheema: None; Stephen Yip: Advisory Board Member, Bayer; Advisory Board Member, Pfizer; Advisory Board Member, Roche; Speaker, Roche; Brandon Sheffield: Grant or Research Support, Pfizer; Advisory Board Member, Novartis; Consultant, Roche; Grant or Research Support, Astra Zeneca; Grant or Research Support, Boehringer Ingelheim

Background: *MET*exon 14 skipping (*MET*ex14) mutations present in 4% of lung adenocarcinoma is now becoming an important alteration to test for targeted therapy, similarly to *ALK*. The only commercially available way to test for *MET*ex14 mutations is through next generation sequencing. There is a need for a faster and more available method to be used for the detection and validation of *MET*ex14 mutations but traditional animal base monoclonal antibody (mAbs) techniques are slow and difficult to scale. A newer, faster and animal-free approach using instead B-cells cloning to generate *in vitro*recombinant antibodies (rAbs) is increasingly popular. Here we compare a novel rAbs technique to the more traditional mAbs generation approach in developing a mutation-specific monoclonal immunohistochemistry (IHC) to *MET*ex14 mutation.

Design: Using the same amino acid sequence overlapping the fusion of *MET*exon 13 and 15, we generated a total of 72 antibody clones: 68 rAbs and 4 rabbit-based mAbs. The clones were validated by enzyme-linked immunosorbent assay (ELISA) and IHC using a combination of synthetic peptides, *MET*ex14 mutated cell line (H596) and archival lung adenocarcinoma tissue with *MET*ex14 mutation.

Results: Of the 68 rAbs screened for affinity by ELISA, 37 were retained for IHC validation along with all 4 mAbs. Using the METex14 mutated cell line, strong (3/3 staining intensity) diffuse (100% tumor cells staining) membranous staining was achieved in 4 of the rAbs. Seven other rAbs had weak-to-intermediate (1-2/3 staining intensity) non-diffuse (5-75% tumor cell staining) membranous staining and the remaining 26 rAbs showed no membranous staining. The best mAbs clone only showed focal (5%) weak-to-intermediate staining with substantial background staining and the other 3 mAbs were completely negative.

Conclusions: The rAbs technique was an effective approach to generate METex14 mutation-specific IHC clones. It can be scaled up more readily as opposed to the traditional animal-based hybridoma technique, and as a result, it increases its rate of success and decreases

cost. This technique might allow for easier transition of mutation-based biomarkers to IHC and improve turnaround time and access for predictive tests in oncology. The top 4 rAbs are currently being tested on an extended cohort of lung carcinoma tissue.

1964 Reproducibility and Accuracy of Intra-Operative Assessment on Tumor Spread Through Air Spaces (STAS) in T1 Lung Adenocarcinomas

Julian Villalba¹, Tréah May Sayo², Yin Hung², Amy Ly², Angela Shih², Marina Kem², Lida Hariri³, Mari Mino-Kenudson²

¹Boston, MA, ²Massachusetts General Hospital, Boston, MA, ³Massachusetts General Hospital, Harvard Medical School, Boston, MA

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Background: STAS has been associated with worse prognosis in patients with early-stage lung adenocarcinoma (ADC), particularly those with sublobar resection. While intra-operative assessment of STAS in frozen section (FS) has been advocated by thoracic surgeons to guide management, information on the reproducibility and accuracy of FS for diagnosing STAS is limited to date.

Design: We evaluated 73 T1 lung ADC resections previously assessed by intra-operative consultation (IC) and with ample non-neoplastic lung parenchyma in FS slides. Five pulmonary pathologists blinded to clinicopathologic data independently and separately reviewed paired FS and FS permanent (FP) slides in each case evaluating for STAS (presence, absence, or equivocal) and processing artifacts per published criteria (Kadota et al. JTO 2015). Separately, a consensus panel of 3 pathologists reviewed all the slides in permanents to reach a final integrated assessment for STAS on each case and recorded additional parameters, including the number of tumor clusters identified as STAS. Diagnostic accuracy was determined by comparing the most prevalent diagnosis achieved by >3 observers with the final integrated diagnosis. Performance characteristics including interobserver agreement (IOA), positive predictive value (PPV), and negative predictive value (NPV) were determined.

Results: STAS was identified by the consensus panel in 11% of 28 low-grade, 32% of 19 intermediate-grade and 89% of 26 high-grade tumors. The Fleiss kappa value for the STAS diagnosis among the 5 observers in FS slides(0.30 ± 0.03 [k±SE]) was lower than that in FP slides(0.41 ± 0.03), but both increased to 0.63 ± 0.05 and 0.82 ± 0.05 , after removal of cases considered as equivocal by any observer. Detecting STAS by FS showed high specificity (95%) and PPV (88%), but low sensitivity (46%) and NPV (70%). Processing artifacts were frequently noted by most observers, present in 82% of STAS-positive, 87.5% of equivocal, and 46% of STAS-negative FS slides. IOA in FS was higher in low-grade tumors (p=0.008), STAS-negative cases (p<0.001), those without artifacts (p=0.03), and STAS-positive FS slides with >10 STAS clusters (p=0.04).

Conclusions: FS is highly specific but not sensitive for the diagnosis of STAS in Stage I lung ADC. IOA on STAS is only fair in FS, partly due to concurrent artifacts, raising concern on implementation of IC for STAS assessment. Multi-Institutional studies aimed to establish standardized criteria for STAS and to improve intraoperative techniques are warranted

1965 Ki67 Proliferative Index Has Limited Use in Pulmonary Large Cell Neuroendocrine Carcinoma

Ann Walts¹, James Mirocha¹, Alberto Marchevsky²

¹Cedars-Sinai Medical Center, Los Angeles, CA, ²Cedars-Sinai Medical Center, West Hollywood, CA

Disclosures: Ann Walts: None; James Mirocha: None; Alberto Marchevsky: None

Background: A recent paper suggested that Ki67 proliferative index (Ki67%) can help stratify pulmonary typical carcinoids into grades 1 and 2 lesions, but not distinguish them from atypical carcinoids (Mod Pathol 2018). There is limited information about Ki67% values in pulmonary large cell neuroendocrine carcinomas (LCNEC). The 2015 World Health Organization (WHO) describes Ki67% as 40%-80% in LCNEC but provides no specific references.

Design: Sections from 77 consecutively resected LCNEC diagnosed by WHO criteria were stained with Ki67 antibody (Ventana Inc, Tucson AZ). Ki67% was measured using Aperio ScanScope AT Turbo, eSlide Manager and ImageScope software (Leica Biosystems, Vista CA) as previously reported. Cases were stratified into 6 classes by 10% Ki67 increments. Using the Kaplan-Meier method and SAS version 9.4 software (Cary, NC), overall (OS) and disease-free survivals (DFS) were compared by AJCC (8th ed) stage, by 6 Ki67% classes, and with Ki67% cut-points at 20% and 40%.

Results: The 42 males and 35 females ranged from 46 to 87 years of age at diagnosis. The tumors ranged from 0.9 cm to 11.5 cm. AJCC (8th ed) stage groups included IA (n=33), IB (n=7), IIA (n-2), IIB (n=16), IIIA (n=16), and IIIB (n=3). 32 (41.6%) LCNEC had a combined adenocarcinoma component. Ki67% was measured only in the NEC component. The system measured Ki67% positivity using 4072 to 44533 tumor nuclei per case (mean 16610 ±8039). Ki67% ranged from 1% to 64% (mean 26%; median 26%). Surprisingly, only 16 (21%) tumors had Ki67% ≥ 40%. OS ranged from 1 to 298 months (median 25 mos). DFS ranged from 1 to 276 months (median 9 mos). OS and DFS differed across AJCC stage (log-rank P=0.045 and P=0.029, respectively). However, neither OS nor DFS significantly correlated with

Ki67% when 6 classes were used (log-rank P=0.82 and P=0.21, respectively) or when 2 classes were used with either 20% Ki67% (log-rank P=0.95 and P=0.53) or 40% Ki67% (log-rank P=0.78 and P=0.91) as cut-point.

Conclusions: Our findings do not support the use of 40% as the minimum Ki67% in LCNEC as suggested by WHO or the use of 20% as the minimum Ki67% for LCNEC as described for diagnosing entero-pancreatic neuroendocrine carcinoma.

1966 Expression of Insulinoma-Associated 1 (INSM1) in Non-Small Cell Lung Cancers: A Diagnostic Pitfall for Neuroendocrine Tumors

Minhua Wang¹, Deepika Kumar², Rita Abi-Raad³, Adebowale Adeniran⁴, Guoping Cai⁵

¹Yale School of Medicine, Guilford, CT, ²Yale New Haven Hospital, New Haven, CT, ³Yale University School of Medicine, New Haven, CT, ⁴Yale University School of Medicine, Seymour, CT, ⁵Yale University, Wallingford, CT

Disclosures: Minhua Wang: None; Deepika Kumar: None; Rita Abi-Raad: None; Adebowale Adeniran: None; Guoping Cai: None

Background: Insulinoma-associated 1 (INSM1) has recently been reported as a highly sensitive and specific marker of pulmonary neuroendocrine tumors. It has also been noticed that INSM1 expression can be seen, although uncommonly, in non-neuroendocrine tumors. The aim of this study was to evaluate the expression of INSM1 in non-small cell cancers (NSCLCs) to avoid diagnostic pitfall.

Design: A cohort of 164 NSCLCs were selected and tissue microarrays were constructed, including 97 adenocarcinomas (AdC), 51 squamous carcinomas (SqCC), 16 other non-small cell carcinomas. INSM1 was stained and evaluated based on the intensity of nuclear staining and percentage of nuclear staining cells. Nuclear immunoreactivity was scored as 0, 1+, 2+ and 3+, representing no staining, weak staining, moderate staining, and strong staining, respectively, and the percentage was categorized as focal (<10% tumor cells), patchy (10-50% tumor cells), and diffuse (>50% tumor cells). No staining (0) and weak staining (1+) were classified as negative; focal moderate staining (2+) as equivocal; patchy and diffuse moderate staining (2+) and strong staining (3+) as positive. (Figure 1)

Results: Among 164 cases, INSM1 was positive in 16 cases (9.8%) and equivocal in 14 cases (8.5%). Of 97 adenocarcinomas, INSM1 was positive in 6 cases (6.2%) and equivocal in 9 cases (9.3%). The positive AdC cases included 1 case with focal 3+ nuclear staining, 2 cases with diffuse 2+ staining, and 3 cases with patchy 2+ staining. Of 51 squamous carcinomas, INSM1 was positive in 9 cases (17.6%) and equivocal in 6 cases (11.8%). The positive SqCC cases included 1 case with patchy 3+ staining, 2 cases with diffuse 2+ staining and 6 cases with patchy 2+ staining. Focal 3+ nuclear staining was seen in 1 of 16 other NSCLC cases. (Table 1)

Diagnosis	Case (n)	Positive	(n=16, 9.	Equivocal (n=14, 8.5%)	Negative (n=134, 81.7%)			
		3+ Diffuse	3+ Patchy	3+ Focal	2+ Diffuse	2+ Patchy	2+ Focal	0 and 1+
Adenocarcinoma	97	0	0	1	2	3	9	82
Squamous cell carcinoma	51	0	1	0	2	6	5	37
Other non-small cell carcinoma	16	0	0	1	0	0	0	15

Table 1. INSM1 expression in 164 non-small cell Lung cancers



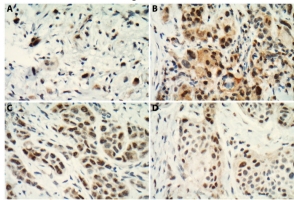


Figure 1 Representative micrographs show nuclear staining of INSM1 in non-small cell lung cancers. (A) Focal 3+ staining in a squamous carcinoma (SqCC); (B) Patchy 3+ staining in a SqCC; (C) Diffuse 2+ staining in an adenocarcinoma; (D) Patchy 2+ staining in a SqCC.

Conclusions: Our study demonstrates that INSM1 is expressed in a subset of NSCLCs and suggest that caution must be exercised in interpreting INSM1 staining, especially with limited sample such as biopsy and cell block sections. Although INSM1 is useful for the diagnosis of neuroendocrine tumors, it should not be used as a stand-alone marker in differentiating primary lung tumors.

1967 LINC00520 Promotes EGFR-TKIs Acquired Resistance in Non-Small Cell Lung Carcinoma

Yue Wang¹, Yuan Li¹

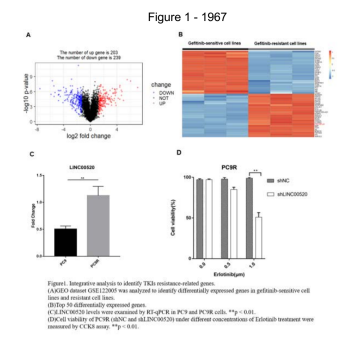
¹Fudan University Shanghai Cancer Center, Shanghai, China

Disclosures: Yue Wang: None; Yuan Li: None

Background: *EGFR* tyrosine kinase inhibitors (TKIs) therapy is a validated approach in the treatment of *EGFR*-mutated non-small cell lung carcinoma (NSCLC), but resistance universally develops and it has become a major obstacle in prolonging the survival of patients. More novel molecular biomarkers are still urgently required to elucidate the underlying mechanisms of resistance. This study aimed to investigate the role of *LINC00520* in the acquired resistance of NSCLC to *EGFR*-TKIs.

Design: Gene expression profiles from GEO dataset were analyzed to identify the genes associated with *EGFR*-TKIs resistance. *EGFR*-mutated NSCLC cell line PC9 was cultured with gefitinib for more than 6 months to acquire gefitinib-resistance, which was designated as PC9R. The expression patterns of *LINC00520* were characterized using reverse transcription quantitative polymerase chain reaction (RT-qPCR), and lentiviral vectors were used to infect cells to regulate the expression. Cytotoxicity of *EGFR*-TKIs on infected cells was determined by cell counting kit-8 (CCK-8). Survival follow-up time of 948 NSCLC samples from TCGA dataset were enrolled in this study. In addition, Statistical analysis was mainly performed by R programming language and GraphPad Prism 7.0 (GraphPad).

Results: LINC00520 is highly expressed in gefitinib-resistant cell line PC9R relative to PC9 (p<0.05). Inhibiting LINC00520 with lentivirus vectors induces apoptosis in PC9R. LINC00520 could promote cell proliferation and induce resistance. Statistics from TCGA dataset demonstrate there is no significant difference in LINC00520 expression between LUAD tissues (483) and normal tissues (347), but the expression level in LUSC tissues (486) is much higher than normal (338). More interestingly, higher LINC00520 TPM implies a worse survival rate in LUAD (DFS: HR=1.4, p =0.043, OS: HR=1.4, p =0.028), while this phenomenon does not appear in LUSC (DFS: HR=1.1, p =0.7, OS: HR=0.86, p =0.28).



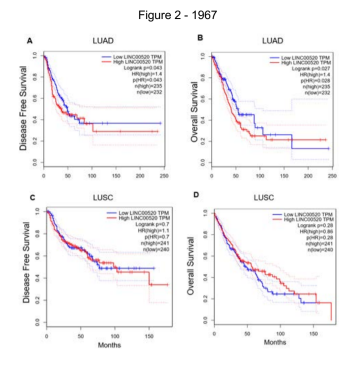


Figure 2. Kaplan-Meier survival curves for Disease Free Survival and Overall Survival in NSCLC (A)DFS and (B)OS of LUAD patients according to LINC00520 expression.

(C)DFS and (D)OS of LUSC patients according to LINC00520 expression.

Conclusions: *LINC00520* is involved in acquired resistance of *EGFR*-TKIs in NSCLC. It may serve as a predictor and a potential therapeutic target for *EGFR*-TKIs resistance.

1968 Comparison of Four Proposed Architectural Grading Systems at Predicting Recurrence in a Cohort of 278 Stage I Lung Adenocarcinoma

Ilyas Yambayev¹, Artem Shevstov¹, Paulo Moreira², Kei Suzuki¹, Travis Sullivan³, Kimberly Rieger-Christ³, Eric Burks⁴
¹Boston University, Boston Medical Center, Boston, MA, ²Boston Medical Center, Boston, MA, ³Lahey Hospital & Medical Center, Burlington, MA, ⁴Boston University Mallory Pathology Associates, Boston, MA

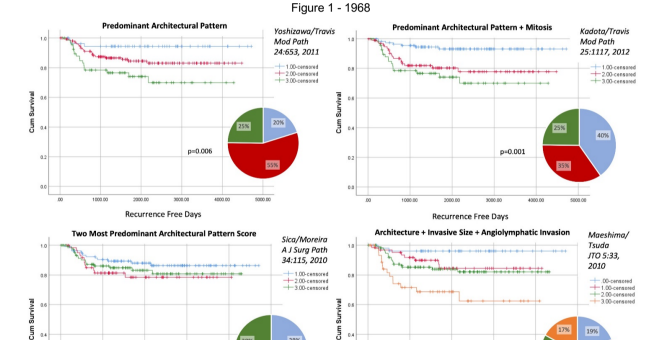
Disclosures: Ilyas Yambayev: None; Artem Shevstov: None; Paulo Moreira: None; Kei Suzuki: None; Travis Sullivan: None; Kimberly Rieger-Christ: None; Eric Burks: None

Background: Lung cancer is the most common cause of cancer death worldwide. Screening by LDCT is expected to increase the frequency of early-stage NSCLC of which lung adenocarcinoma (LUAD) is the most common subtype. Although the assignment of predominant histologic subtypes is now recommended, there remains no widely accepted prognostically relevant grading system. Several grading systems have been proposed however there has been no direct comparison of these grading systems in an independent cohort. Here we compare several previously published architecturally based grading systems in a large cohort of stage I LUAD.

Design: H&E slides were reviewed from stage I LUAD resection specimens form a multi-institutional cohort of 278 patients diagnosed between 2005-2015. The staging was reassigned using AJCC 8th edition after determining the invasive size and assessing for pleural invasion. Comprehensive histologic subtyping in 5% increments was performed along with mitotic figure counts and assessment for angiolymphatic invasion. This data was applied to compare recurrence-free survival rates using 4 published grading systems. The demographic, smoking status and stage characteristics are summarized in Table 1.

Results: Figure 1 shows the Kaplan Meier curves for 4 published grading systems. Predominant architectural pattern assignment alone was prognostically significant in stratifying patients into low (lepidic), intermediate (acinar/papillary), and high (solid/micropapillary) grades (p=0.006). The combined scoring system of the two most predominant histologic patterns proposed by Sica et. al. proved not to be prognostically significant in our cohort. The addition of mitotic rate to predominant architectural pattern proposed by Kadota et. al. successfully shifted a large fraction of architecturally intermediate tumors into the low-risk category. The addition of angiolymphatic invasion to tumors with at least 30% combined high-grade patterns (micropapillary, cribriform, and solid) as proposed by Maeshima et. al. allowed for the detection of a small but significant high-risk recurrence group.

Age	68 (62-74)
Median (Q1-Q3)	
Race	204 (74)
White (%)	49 (18)
Black (%)	11 (4)
Asian (%)	14 (5)
Other (%)	
Gender	106 (38)
Male (%)	172 (62)
Female (%)	
Smoking Status	139 (50)
Current smoker (%)	107 (39)
Former smoker (%)	28 (10)
Never smoker (%)	4 (1)
Unknown (%)	
AJCC 8 th ed. Stage	101 (36)
IA1 (pT1a)	121 (44)
IA2 (pT1b)	35 (13)
IA2 (pT1c)	21 (7)
IB (pT2a)	
Tumor Size	1.7 (1.2-2.4)
Total Size	1.3 (0.8-1.8)
[median (Q1-Q3), cm]	
Invasive Size	
[median (Q1-Q3), cm]	



Conclusions: Predominant architectural pattern assignment alone is a valuable grading system. The addition of mitotic grade and angiolymphatic invasion allows for further refinement to identify higher proportions of low-risk and a small but significant subset of very high-risk LUAD which might aid in the precision clinical management of early-stage LUAD.

p=0.339

Recurrence Free Davs

17%

p=0.001

Recurrence Free Days

19%

1969 UV Genomic Signature Classifies Lung Melanomas of Unknown Primary as Metastases from Occult Cutaneous Melanomas

Chen Yang¹, Francisco Sanchez-Vega¹, Alexander Shoushtari¹, Marc Ladanyi¹, William Travis¹, Klaus Busam², Natasha Rekhtman¹

¹Memorial Sloan Kettering Cancer Center, New York, NY, ²Chatham, NJ

Disclosures: Chen Yang: None; Francisco Sanchez-Vega: None; Alexander Shoushtari: *Grant or Research Support*, Bristol Myers Squibb; *Advisory Board Member*, Bristol Myers Squibb; *Advisory Board Member*, Immunocore; *Consultant*, Castle Biosciences; Marc Ladanyi: None; William Travis: None; Klaus Busam: None; Natasha Rekhtman: None

Background: We have encountered a group of patients with melanomas involving the lung in the absence of a clinically known primary melanoma elsewhere. A subset of patients presented with solitary large tumors. While primary pulmonary melanomas (PPM) is a category included in the thoracic WHO classification, given the absence of normal melanocytes in the lung its existence has been questioned. Herein we investigate genomic profiles of melanomas of unknown primary origin involving the lung. In particular, we sought to determine whether UV genomic signature - a characteristic feature of most cutaneous melanomas - is present in such tumors.

Design: Cases of melanomas involving the lung with no known primary elsewhere were identified retrospectively. The clinicopathologic characteristics of each case were annotated. All cases included in the study underwent targeted DNA next-generation sequencing (NGS) interrogating up to 468 cancer genes. Genomic signatures were analyzed based on a method described by Alexandrov LB et al. (*Nature* 2013; 500:415-421).

Results: Ten NGS-profiled melanomas involving the lung were identified. Five patients had solitary lung lesions with the median size of 5.2 cm (range 2.6 to 10.1 cm). Of those, 3 tumors were endo/peri-bronchial, thus meeting the suggested criteria for PPM. Hilar nodes were involved in 2 patients, and 8 tumors had epithelioid morphology resembling non-small cell carcinoma. No evidence of primary melanomas was found for any patients on clinical follow-up (median 28 months; range 5 to 42 months). Genomic testing revealed the following driver mutations commonly found in melanomas: *BRAF* (n=6), *NRAS* (n=1) and *KIT* (n=1). Genomic signature analysis was feasible for 8 cases harboring >20 mutations required for reliable analysis, including 4 patients with solitary masses. This revealed the presence of a dominant UV signature in all cases. In contrast, none of the primary lung carcinomas tested by the same method (n = 255) had a UV signature.

Conclusions: The consistent presence of a UV signature provides strong support for an occult or regressed cutaneous origin of melanomas involving the lung, and argues against the concept of PPM. Clinical presentation as solitary large (reaching >10 cm) masses occasionally with hilar adenopathy and epithelioid morphology may closely mimic primary lung carcinomas both clinicoradiologically and pathologically, representing a major potential diagnostic pitfall.

1970 Multifocal Invasive Mucinous Adenocarcinomas Involving Different Lobes are Clonally Related and Represent Intrapulmonary Spread as Defined by Molecular Profiling

Soo-Ryum Yang¹, Jason Chang², Charles Leduc³, Kay See Tan², Snjezana Dogan², William Travis², Maria Arcila², Marc Ladanyi², Natasha Rekhtman²

¹Department of Pathology, Memorial Sloan Kettering Cancer Center, New York, NY, ²Memorial Sloan Kettering Cancer Center, New York, NY, ³University of Montreal Hospital Centre, Montreal, QC

Disclosures: Soo-Ryum Yang: *Consultant*, Invitae; Jason Chang: None; Charles Leduc: None; Kay See Tan: None; Snjezana Dogan: None; William Travis: None; Maria Arcila: *Speaker*, biocartis; *Speaker*, Invivoscribe; Marc Ladanyi: None; Natasha Rekhtman: None

Background: Pulmonary invasive mucinous adenocarcinoma (IMA) commonly presents as a multifocal disease. It is widely recognized that diffuse 'pneumonic-type' IMA represents aerogenous spread of a single tumor. However, IMAs may also present as discrete nodules in different lobes, raising the possibility of separate primary tumors. Here, we explored the clonal relationship of IMAs involving different lobes using comparative molecular profiling.

Design: We performed a retrospective review of patients with IMAs who had genomic analysis performed on tumors in different lobes. Molecular assays included DNA-based targeted next-generation sequencing (NGS) for 410-468 cancer genes combined with RNA-based NGS fusion assay (Archer) and non-NGS panels for a subset of cases. Tumor clonal relationships were assessed by comparing somatic alterations between the separate tumor sites.

Results: Twenty-one patients with genomically-profiled IMAs involving contralateral (n=19) or ipsilateral different lobes (n=2) were identified. In most patients (n=14), tumors had discrete nodular presentation. Second IMA presented metachronously in 11 patients with a mean latency of 4.2 years. Notably, in 3 patients, contralateral spread manifested ≥8 years (up to 11 years) after initial tumor resection. Genomic analysis was performed on 2 separate IMAs in 19 patients and 3 separate IMAs in 2 patients, resulting in a total of 44 genotyped tumors. Comparative genomic analysis revealed that tumors in all patients shared matching driver alterations including *KRAS* (n=17), *NRG1* (n=2), *ERBB2* (n=1) and *BRAF* (n=1). In addition, in tumor pairs profiled by NGS and Archer, other shared

alterations were identified for a total of 2 – 5 shared alterations per pair (mean of 3 shared alterations). The probability of chance co-occurrence for a full set of shared alterations was 1 x 10⁻⁶ to 4 x 10⁻³².

Conclusions: Molecular profiling supports that multifocal IMAs involving different lobes represent intrapulmonary spread of a single tumor rather than separate primary tumors, including tumors presenting contralaterally after a remarkably long latency (>8 years). Overall, these findings reinforce the unique biology and clinical behavior of IMAs, and draw a sharp distinction with multifocal non-mucinous lung adenocarcinomas, which recent molecular studies confirm to represent predominantly separate primary tumors.

1971 Comparative Characterization of Tumor-Infiltrating Immune Cell Subsets Related to PD-L1 and B7-H3 expression in Non-Small Cell Lung Cancer

Diana Yim¹, Jaemoon Koh¹, Sehui Kim², Seung Geun Song¹, Young A Kim³, Yoon Kyung Jeon¹, Doo Hyun Chung¹

¹Seoul National University College of Medicine/Hospital, Seoul, Korea, Republic of South Korea, ²Seoul, Korea, Republic of South Korea, ³Seoul Metropolitan Government-Seoul National University Borame Medical Center, Seoul, Korea, Republic of South Korea

Disclosures: Diana Yim: None; Jaemoon Koh: None; Sehui Kim: None; Seung Geun Song: None; Young A Kim: None; Yoon Kyung Jeon: None

Background: PD-L1 and B7-H3 are known as immune checkpoint molecules inhibiting anti-tumor immune responses. However, the different tumor microenvironment, especially tumor-infiltrating immune cell subsets, related to PD-L1 and B7-H3 expression remain unclear. We addressed these issues in human non-small cell lung cancer (NSCLC).

Design: PD-L1 and B7-H3 expression in tumor cells were evaluated using immunohistochemistry. Composition of tumor-infiltrating immune cells, including lymphoid cells, macrophages and dendritic cells, was analyzed using flow cytometry for fresh tissues from a prospective cohort of 71 patients with NSCLC and was compared according to PD-L1 and B7-H3 expression status.

Results: In NSCLC, tumoral PD-L1 and B7-H3 expression both positively correlated with the absolute numbers of panimmune cells, CD3+ T cells, CD8+ T cells, plasmacytoid dendritic cells (DCs) and the percentage of CD8+/CD3+ T cells. Tumoral PD-L1 and B7-H3 expression both negatively correlated with the percentage of CD4+/CD3+ T cells. Tumoral PD-L1 expression also positively correlated with the absolute numbers of regulatory T cells (Treg), DCs, myeloid DCs (mDC), CD1c+ DCs, plasmacytoid DCs (pDC), and the percentage of M1/MQ macrophages, DCs/CD3-CD45+ cells, DCs/immune cells and negatively correlated with the absolute number of M1 macrophages and the percentage of M2/MQ macrophages. However, B7-H3 expression showed no significant correlation with these molecules. B7-H3 expression was rather positively correlated with the absolute number of CD141+ DCs and the percentage of MQ/CD3-CD45+ cells.

Conclusions: NSCLCs with high PD-L1 expression showed different composition of tumor-infiltrating immune cells from NSCLCs with high B7-H3 expression. These results suggest that PD-L1 and B7-H3 may influence different immune cells and form different tumor microenvironment, and may be useful therapeutic targets in NSCLC.

1972 Homologous Recombination Defects and Lung Carcinomas--Mutational Spectrum of ATM, BRCA1 and BRCA2-altered Lung Carcinomas

Ju-Yoon Yoon¹, Jason Rosenbaum²

¹Perelman School of Medicine at the University of Pennsylvania, Philadelphia, PA, ²UPenn, Center for Personalized Diagnostics, Philadelphia, PA

Disclosures: Ju-Yoon Yoon: None; Jason Rosenbaum: None

Background: Defects in the homologous recombination (HR) DNA repair pathway, a subset of which are related to germline or somatic mutations in BRCA1, BRCA2, ATM, have been shown to be a targetable phenotype, where survival benefit from PARP inhibition has been demonstrated in some carcinomas. Although variants in HR genes have been reported in lung cancer, their frequencies and clinical significance have not been well characterized.

Design: We examined the local cohort of patients with the referral diagnosis of "lung cancer", sequenced by the in-house 152 gene massively parallel sequencing (MPS, also known as next-generation sequencing or NGS) assay. All alterations were filtered and reviewed for disease-associated variants. For *ATM*, our assessment was limited to p.V2424G, a variant with the highest penetrance among the ATM variants, associated with increased breast cancer risks at levels comparable to disease-associated *BRCA1/2* variants.

Results: 1,473 cases were successfully sequenced by the in-house solid MPS assay, among which 23 patients (1.6%) were found to harbor disease-associated *BRCA1* (14/23) or *BRCA2* (9/23) alterations, with variant allele fractions (VAFs) ranging 4-69%. These cancers were mostly adenocarcinomas (17/23, with 5 carcinoma NOS and 1 SCC). 10/23 harbored *KRAS* hotspot mutations, and no cases

harbored *EGFR* mutations. 6 lung cancers with *ATM* p.V2424G were identified, with generally higher VAFs (55-92%). 2/6 cases with *ATM* p.V2424G also harbored the *EGFR* p.L858R variant, while 2 cases also harbored *KRAS* codon 12 mutations. Interestingly, the two *KRAS/ATM*-mutated cases were from a single patient, with different *KRAS* mutations.

Conclusions: Disease-associated *BRCA1/2* variants are rare in lung carcinomas, and many cases were associated with *KRAS* hotspot variants, suggesting that *BRCA1/2* mutations may be somatic in origin, likely in the setting of significant smoking history. In contrast, while *ATM* p.V2424G is equally rare, the genomic context and VAFs of the *ATM* variants suggest possible germline events. Assessment of other genes in the HR pathway is currently underway.

1973 Differential Diagnostic Values of SATB2 in Primary Pulmonary Enteric Adenocarcinoma and Pulmonary Metastases of Colorectal Carcinoma

Jingping Yuan¹, Huihua He², Lin Xiong², Li Xu²

¹Wuhan, Hubei, China, ²Renmin Hospital of Wuhan University, Wuhan. Hubei, China

Disclosures: Jingping Yuan: None

Background: Pulmonary enteric adenocarcinoma (PEA) is a rare histologic type of lung adenocarcinoma. PEA is composed mainly of tall columnar cells arranged in an irregular acinar or cribriform pattern with extensive central necrosis, closely resembling the appearance of intestinal epithelial and colorectal carcinomas under the microscope. Immunohistochemically, PEA is usually positive for CK7. However, some cases lack CK7 expression and are positive for intestinal differentiation markers, such as CDX2, villin, and CK20. For these reasons, it is difficult to distinguish between PEA and pulmonary metastases of colorectal carcinoma (MCRC), so new identification methods need to be explored. SATB2 expression is tissue-specific, and the only epithelial cells expressing this protein in adult tissue are the glandular cells lining the lower gastrointestinal (GI) tract. The sensitivity of SATB2 in colorectal adenocarcinoma reaches 80%–97%, and their low expression in primary pulmonary tumors. Therefore, this study investigated differential diagnostic values of SATB2 in PEA and MCRC.

Design: According to the WHO primary PEA diagnostic criteria, the cases of lung adenocarcinoma were collected from patients being treated at the Renmin Hospital of Wuhan University from 2015.1-2019.9 were screened. The specimens were independently reviewed by two pathologists, and immunohistochemical staining of lung adenocarcinoma markers (CK7, TTF-1, and NapsinA) and intestinal cancer markers (CK20, CDX2, and villin) was performed to aid identification. Finally, after excluding possible colorectal cancer metastasis by carefully analyzing the clinical histories and imaging examinations, we recruited 51 primary PEA specimens and 17 MCRC specimens for study. The sensitivity and specificity of immunomarkers SATB2, CK7, TTF-1, NapsinA, CK20, CDX2 and villin for distinguishing PEA from MCRC are evaluated.

Results: The expression rates of SATB2 in PEA and MCRC were 0.00% (0/51) and 100.00% (17/17), respectively. The sensitivity of SATB2-, CK7+, TTF-1+, NapsinA+, CK20-, CDX2- and villin- for distinguishing PEA from MCRC were 100.00%, 98.04%, 49.02%, 45.10%, 66.67%, 58.82%, 47.06%, respectively. The specificity of SATB2-, CK7+, TTF-1+, NapsinA+, CK20-, CDX2- and villin- for distinguishing PEA from MCRC were 100.00%, 88.24%, 88.24%, 100.00%, 82.35%, 88.24%, 88.24%, respectively.

Table 1: The diagnostic efficiency of the single immunomarkers in PEA							
Indicator	PEA (n=51)	MCRC (n=17)	sensitivity (%)	specificity (%)			
SATB2-	51	0	100	100			
CK 7+	50	2	98.04	88.24			
TTF-1+	25	2	49.02	88.24			
NapsinA+	23	0	45.1	100			
CK20-	34	3	66.67	82.35			
CDX2-	30	2	58.82	88.24			
villin-	24	2	47.06	88.24			

Figure 1 - 1973

Figure 2 - 1973

Conclusions: Our study shows that the sensitivity and specificity of SATB2, which can all reach 100%, is much higher than those of common lung adenocarcinoma immunomarkers (TTF-1, NapsinA) and intestinal cancer immunomarkers (CK20, CDX2 and villin). SATB2 can be viewed as the best immunomarkers for distinguishing PEA from MCRC. The diagnostic value of CK7 is slightly inferior to SATB2, the results of CK7 can be used as a reference for differential diagnosis of SATB2.

1974 MET exon 14 Variants in Non-Small Cell Lung Carcinoma: Prevalence, Clinicopathologic and Molecular Features

Lisi Yuan¹, David Bosler²

¹Cleveland Clinic, Cleveland, OH, ²Cleveland Clinic, Beachwood, OH

Disclosures: Lisi Yuan: None; David Bosler: None

Background: Somatic *MET* exon 14 skipping mutations (*MET* ex14) are targetable driver mutations for non-small cell lung cancer (NSCLC), responsive to *MET* inhibitors. This study seeks to further characterize the clinicopathological features and mutational profile of *MET* ex14 variant NSCLC.

Design: Retrospective review of all *MET* ex14 tested NSCLC. Genomic DNA was extracted from FFPE or FNA. Testing for selected *BRAF*, *EGFR*, *HER2*, *KRAS*, *MET* mutations was performed using a clinically validated NGS assay, the Cancer Hotspot Panel v1 (Thermo Fisher, Waltham, MA) customized to include *MET* exon 14 splice site variants, followed by MiSeq sequencing (Illumina, San Diego). Variants were classified as significant (Tier1/2) or variants of uncertain significance (VUS) per 2017 AMP/ASCO/CAP Joint Consensus Guidelines. PD-L1 expression was assessed by immunohistochemistry.

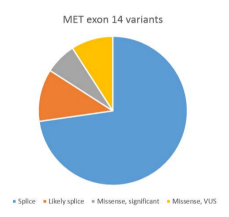
Results: Of 2296 NSCLCs tested between 2017-7/2019, *MET* ex14 variants were present in 44 (1.9%). A recurring VUS not expected to impact exon 14 splicing seen in 41 cases (c.2975C>T (p.Thr992lle) was excluded from analysis. In positive cases, median age was 76 (59% men; 41% women), and 46.7% were FNA specimens. 32 of 44 variants were MET exon 14 skipping (previously reported and/or involve the canonical recognition site), while the other 12 mutations were significant missense (3) or VUS (9). Of 9 VUS, 5 were adjacent to the canonical splice site and likely to impact splicing, and 4 were missense variants. Average allele fraction was 30.2. Four cases had concomitant mutations (3=KRAS, 1=EGFR). Of 35 cases with known clinical staging, stage 1-2=20(57%), stage 3=3 (9%), and stage 4=12(34%). Of 19 resected NSCLSs, histological types and growth pattern included 7 lepidic predominant, 6 acinar predominant, 2 micropapillary predominant, 1 solid predominant, 1 sarcomatoid, and 2 adenosquamous. PD-L1 expression in 27 cases is shown in Table 1.

Table 1.

Prevalence of MET exon 14 mutations			1.9%						
Total patients	44 patients: 26 mer	44 patients: 26 men, 18 women							
	Mean age: 76 years								
Clinical stage	Total 35 cases								
available	Stage 1 and 2: 20 c	ases (57%)	Stage 3: 3 cases (9%)		Stage 4: 12 cases (34%)				
Tumor resected	Total 19 cases								
	Histologic type and	growth pattern							
	Lepidic pattern	Acinar pattern	Micropapillary Solid predom		Solid	Adeno-		Sarcomatoid	
	predominant	predominant			predomin	nant	squamous		
			predominant						
	7	6	2		1		2	1	
PD-L1	Total 27 cases, pos	itive in 22 cases (8	32%)					·	
expression	0%	1%-4	49%:			>50%			
	5 cases (18%)	11 ca	cases (41%)			11 cases (41%)			
	Stage 1-3		PD-L1 <50%		Sta	ge 4			
	PD-L1 <50%	PD-L1 > 50%			%	PD-L1 > 50%			
	12 cases (67%)	6 cases (33%)	2 cases (33%)		3%)	4 cases (67%)			

Figure 1 - 1974

Figure 1



Conclusions: Most MET variants identified in our cohort (73%) are MET ex14 skipping. Another 11% likely result in exon 14 skipping, while the other 16% are missense variants presumably unrelated to splicing. The prevalence of MET ex14 variants is lower than previously reported (1.9% vs 3%), and a large percentage of tumors has lower clinical stage and less aggressive pathologic features, both possibly reflecting sampling differences attributed to universal testing of NSCLC at our institution rather than testing of only advanced disease.

1975 Diagnosis of Lung Biopsies with the 2015 WHO Criteria and Detection of Sensitizing Mutations: A Single-Institution Experience with 5032 Cases

Yupeng Zeng¹, Ying Ding¹, Liuyang Xu¹, Boya Zhai¹, Xiang Zhang¹, Qiaoyun Ge¹, Jiao Li¹, Qiyuan Song¹, Xiao Li¹, Zhihong Zhang¹

¹Jiangsu Province Hospital, Naniing, Jiangsu, China

Disclosures: Liuyang Xu: None; Zhihong Zhang: None

Background: The 2015 WHO classification of lung tumors provided the first specialized classification for small biopsies. This article aimed to apply the newest classification to reclassify a group of small lung biopsies and analyze their status of the main driver mutations.

Design: 5032 cases of small lung biopsies (bronchoscopic, needle, or core biopsies) were selected, which ranged from 2015 to 2018. We applied the newest classification to reclassify them and analyzed the relationship between the diagnostic subtypes of these biopsy specimens and the mutation rates of EGFR and ALK.

Results: The numbers of small lung biopsies each year during 2015-2018 were respectively 1068, 1299, 1511 and 1154. There were 3280 men and 1752 women, ranging in age from 11 to 93 years (median 63 years). The most common diagnosis was primary lung cancer (3130,

62.2%), followed by inflammatory lesion (1326, 26.4%), uncertain case (350, 6.9%), metastatic tumor (165, 3.3%), primary nonepithelial malignant tumor (36, 0.7%), and benign tumor (25, 0.5%). Among the primary lung cancer, the dominant type was adenocarcinoma (1421, 28.2%), followed by NSCC, favor adenocarcinoma (501, 10.0%), squamous cell carcinoma (368, 7.3%), and NSCC, favor squamous cell carcinoma (360, 7.2%). The tests of the main driver mutations using ARMS-PCR technology demonstrated that EGFR was positive in 56.1% (499/889, in adenocarcinoma and NSCC, favor adenocarcinoma), ALK in 5.7% (12/211, in NSCC), and ROS1 in 0.9% (2/211, in NSCC). 898 NSCC specimens went through a immunohistochemical (IHC) test for ALK (D5F3) and 38 of them were positive (4.2%). Moreover, EGFR mutations were enriched in adenocarcinoma (64.1%) compared to NSCC, favor adenocarcinoma (34.1%) (P<0.0001). On the contrary, the fraction of adenocarcinoma with ALK mutations was notably lower than in NSCC, favor adenocarcinoma (4.2% vs 8.4%, P=0.021).

Conclusions: The criteria for small lung biopsies proposed by the 2015 classification of lung tumors should be applied to pathologists' daily work. It can improve the diagnostic efficiency and quality of small lung biopsies and assist oncologists in accurately understanding the pathologic diagnosis. In this way, accurate treatment and improved prognosis are more available to the patients.

1976 Differential Diagnosis of Multiple Primary and Intra-Lung Metastasis of Lung Cancer by Multiple Gene Detection

Zhihong Zhang¹, Liuyang Xu¹, Xiao Li¹, Yupeng Zeng¹, Pan Ji¹, Shuaishuai Zhuo¹ ¹Jiangsu Province Hospital, Nanjing, Jiangsu, China

Disclosures: Zhihong Zhang: None; Liuyang Xu: None; Shuaishuai Zhuo: None

Background: To study the differential diagnosis of multi-focal lung cancer and lung cancer with pulmonary metastasis by detecting the different lesions of the same patient. To explore the differences in prognosis between MPLC and IM, and to explore the factors affecting the prognosis of multi-focal lung cancer and the tumor heterogeneity of multi-focal lung cancer in combination with histopathology and molecular biology.

Design: Fifty patients with multi-focal lung cancer were screened, and the relevant clinical information was noted; the patients were diagnosed by ACCP standard. Mutations of the lesions were detected by ARMS-PCR, and the detected genes included EGFR, ALK, ROS1, MET, KRAS, RET, HER-2, BRAF, NRAS and PIK3CA. The results of genetic testing were compared with those of ACCP standard diagnosis.

Results: We analyzed a total of 101 tumors from 50 patients. Classification based on gene testing contradicted the clinicopathologic diagnosis in 10 (20%) of the comparisons, identifying independent primaries in 6 cases diagnosed as metastasis and metastases in 4 cases diagnosed as independent primaries. Another 7(14%) tumor pairings were assigned an "equivocal" result based on gene testing. The results of gene testing of the remaining 33(66%) tumor pairings were consistent with the clinicopathologic diagnosis. The mutant heat map indicated that IM patients have a higher rate of mutation consistency than MPLC patients. The difference of prognosis between patients with mutations and those with wild-type genes patients was statistically significant (P=0.002). The difference of prognosis between patients with IM patients with MPLC and those with IM was statistically significant (P=0.038). The difference of prognosis between patients who had different condition was statistically significant (P=0.038).

Conclusions: Multi-gene detection of multi-focal lung cancer has a certain auxiliary effect on the differential diagnosis of multiple primary lung cancer and lung cancer with pulmonary metastasis, which can complement the clinical standards, but also has some limitations.

1977 Clinicopathological Features of Insertion Mutations in Kinase Domain of EGFR and ERBB2 in Non-Small-Cell Lung Cancer

Ruiying Zhao¹, Chan Xiang¹, Lei Zhu¹, Jinchen Shao¹, Jikai Zhao¹, Lianying Guo¹, Shengji Ma¹, Anbo Yu¹, Yuchen Han² Shanghai Chest Hospital, Shanghai Jiao Tong University, Shanghai, China, ²Shanghai, China

Disclosures: Ruiying Zhao: None

Background: While in-frame insertion mutations in the kinase domain of *ERBB* (*EGFR* and *ERBB2*) are known as driver mutations of non-small-cell lung cancer (NSCLC), the clinicopathological features and their role in tumor behavior have not been fully identified.

Design: We detected 4050 NSCLC formalin-fixed paraffin-embedded (FFPE) tissues using a DNA capture-based NGS assay. Afterwards, the insertions in the kinase domain of *ERBB* were identified by real-time PCR. The clinicopathological features were analyzed, and the invasion and metastasis behavior of tumor were studied.

Results: In total, 8.4% (339/4045) of the samples were detected to harbor an in-frame insertion in the kinase domain of *EGFR* and *ERBB2* (*ERBB* insertion), among which 191 insertions were in exon 20 of *ERBB2*, 141 insertions were in exon 20 of *EGFR*,

and 7 insertions were in exon 19 of *EGFR*. In *ERBB2*, 12 insertion types were revealed, among which the hotspot mutations were p.A775_G776insYVMA (71.7%, 137/191) and p.G776delinsVC (15.2%,29/191). By compassion, the insertions in *EGFR* were really disperse, as 42 insertions were detected, among which p.A767_V769dup (23.6%, 35/148) and p.S768_D770dup (20.9%, 31/148) were most frequent. 94.4% (320/339) of *ERBB* insertions were found in adenocarcinoma, among which surgical samples were more common than small biopsies (10.1% vs 7.3%, P=0.016). Among adenocarcinoma, female have higher frequencies of *ERBB* insertions than male (11.3% vs 6.8%, P<0.001). The median age of *EGFR* insertion carrier, *ERBB2* insertion carrier and non-*ERBB* insertion carrier was 54-, 49- and 62-year old respectively (P<0.001). Compared with invasive adenocarcinoma (IA) (5.6%), adenocarcinoma in situ (AIS) (28.4%) and minimally invasive adenocarcinoma (MIA) (20.0%) were more likely to harbor an *ERBB* insertion (P<0.001). The pleural invasion frequency of *EGFR* insertion carrier, *ERBB2* insertion carrier and non-*ERBB* insertion carrier was 26.7%, 16.1% and 26.7% respectively (P>0.05), while the lymph node metastasis rate was 8.3%, 21.7% and 14.9% respectively (P>0.05).

Conclusions: The insertion mutations in kinase domain of *EGFR* and *ERBB2* were more common in younger, female and adenocarcinoma patients. AIS and MIA were more frequent to harbor an *ERBB* insertion than IA, which suggest *ERBB* insertion may be related to the evolution of adenocarcinoma. *ERBB2* insertion carrier tend to have a lower pleural invasion rate while a higher lymph node metastasis rate.

1978 Correlation between PD-L1 Expression and Clinicopathological and Molecular Characteristics of Non-Small Cell Lung Cancer: A Large Scale Multi-Centric Real-World Study of Chinese Cohort Qiang Zheng¹, Yan Huang², Xin Zeng³, Xiaoyan Chen⁴, Chunyan Wu⁵, Shihong Shao⁶, Xiaosong Guan⁷, Yuan Li⁸

¹Fudan University Shanghai Cancer Center; Shanghai Medical College, Fudan University, Shanghai, China, ²Department of Pathology, Tongji University Shanghai Pulmonary Hospital, Shanghai, China, ³The First Affiliated Hospital of Guangzhou Medical University, Guangzhou, Guangdong, China, ⁴Ruijin Hospital of Shanghai Jiaotong University, Shanghai, China, ⁵Tongji University Shanghai Pulmonary Hospital, Shanghai, China, ⁶The First Affiliated Hospital of Qingdao University, Qingdao, Shandong

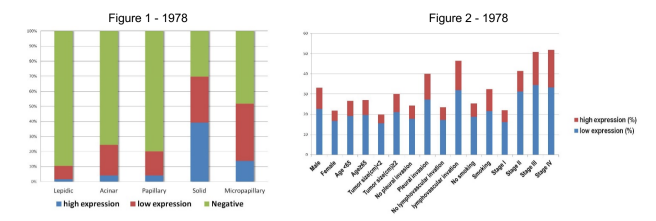
Province, China, ⁷Shanghai Histomed Diagnostic Center, Shanghai, China, ⁸Shanghai Cancer Hospital, Shanghai, China

Disclosures: Qiang Zheng: None; Xin Zeng: None; Xiaoyan Chen: None; Shihong Shao: None

Background: Programmed cell death ligand-1 (PD-L1) is a predictive marker of anti-PD-1/PD-L1 immune therapies for non-small cell lung cancer (NSCLC). The definite relationship between PD-L1 expression and clinicopathological, molecular profiles of NSCLC in Chinese population is remains unclear.

Design: A total of 6126 NSCLC specimens were enrolled from 6 centers in China. We analyzed PD-L1 (22C3) expression by immunohistochemistry on Dako Autostainer Link 48 platform. The status of *EGFR* was defined by RT-PCR or NGS in 2382 samples and *ALK* was tested by IHC, FISH or NGS in 1716 samples.

Results: High PD-L1 expression was observed in 11.9% of resected 4471 NSCLC samples, including in 9.0% of invasive adenocarcinoma (ADC, n=3507), and 25.0% of squamous cell carcinoma (SqCC, n=768). Clinically, PD-L1 high expression was tightly associated with smoking(p=0.001), advanced stage, larger tumor size, pleural invasion, and lymphovascular invasion (p < 0.001). Histopatholoically, PD-L1 high expression was more prevalent in aggressive histologic subtypes included solid, micropapillary, and cribriform subtype (p < 0.001). PD-L1 high expression was more frequent in EGFR-wild type than in mutant type (12.9% vs. 4.7%, p < 0.001). Furthermore, PD-L1 high expression was more prevalent in rare EGFR mutant types than in common mutations (42.1% vs. 20.8%, p=0.031). Besides, PD-L1 high expression was more frequently identified in ALK fusion cases (14.6% vs. 6.5%, p=0.001). A total of 1665 small biopsy cases included 1454 primary specimens and 211 metastatic specimens. The prevalence of PD-L1 high expression in surgical samples was much lower than in primary biopsy samples. Among of them, PD-L1 high expression was also prevalent in EGFR-wild type than in mutant type (32.4% vs 15.0%, p < 0.001). PD-L1 high expression was also more frequently identified in ALK-positive biopsy cases (32.1% vs. 19.2%, p=0.029), which was identical with surgical specimens. We found that high PD-L1 expression was more prevalent in metastatic specimens than in primary biopsy specimens (30.8% vs. 21.8%). In metastatic ADC specimens, the rate of high PD-L1 expression was greater than in primary ADC biopsy samples (33.7% vs. 21%).



Conclusions: This study gave an panoramic view on PD-L1 expression and clinicopathological profiles based on the largest Chinese NSCLC cohort. The discrepancy of PD-L1 expression between surgically resected specimens and biopsy specimens and metastatic lesions may result from inter/intra-tumoral heterogeneity.

1979 Centralized Population-Based Testing of PD-L1 Expression in Non-Small Cell Carcinoma of Lung

Chen Zhou¹, Gang Wang², Cheng-Han Lee³, Tadaaki Hiruki⁴, Diana Ionescu⁵

¹University of British Columbia, Vancouver, BC, ²BC Cancer Vancouver Centre, Vancouver, BC, ³Vancouver, BC, ⁴British Columbia Cancer Agency, University of British Columbia, Vancouver, BC, ⁵UBC at BC Cancer, Vancouver, BC

Disclosures: Chen Zhou: None; Gang Wang: None; Cheng-Han Lee: None

Background: Accurate assessment of PD-L1 expression is critical for selection of patients of non-small cell lung cancer (NSCLC) for immunotherapy with PD-L1/PD-1 inhibitors. However, only limited reports of PD-L1expression in population of NSCLC in North America are available. This study reports PD-L1 expression level in a large patient population of NSCLC in Canada.

Design: PD-L1 testing of NSCLC was performed for patients from the whole province of British Columbia, Canada in a centralized provincial pathology laboratory at BC Cancer, Vancouver Centre. The test used Dako PD-L1 IHC 22C3 PharmDx and Dako Autostainer Link 48 immunostainer, as it was FDA approved companion diagnostic test for pembrolizumab. PD-L1 protein expression is determined by using Tumor Proportion Score (TPS), which is the percentage of viable tumor cells showing partial or complete membrane staining at any intensity.

Results: From January 2017 to March 2018, 1,716 NSCLC was tested, which included 1,301(75.8%) adenocarcinoma, 284 (16.6%) squamous cell carcinoma and 131 (7.6%) NSCLC, NOS. PD-L1 expression level in adenocarcinoma was similar to that in squamous cell carcinoma (p>0.05), but was significantly higher than in NSCLC, NOS (p<0.05, Table 1). The overall percentage of high PD-L1 expression (TPS³50%) was 39%. The high PD-L1 expression (TPS³50%) was found in 42% distant metastases, 45% mediastinal lymph nodes, and 35% lung primary tumors. The differences in the distribution of high PD-L1 expression among distant metastatic sites, mediastinal lymph nodes metastases and primary sites had statistical significance (c2=11.8, p<0.01).

	PD-L1, <1%	PD-L1, 1-49%	PD-L1, 350%	Total
Adenocarcinoma	450 (35%)	337(25%)	514(40%)	1,301
Squamous cell	89(31%)	86(30%)	109(39%)	284
carcinoma				
NSCLC, NOS	44(34%)	46(35%)	41(31%)	131
Total	583(34%)	469(27%)	664(39%)	1,716

Table 1: PD-L1 expression level in different types of NSCLC

Conclusions: Using FDA approved Dako PD-L1 IHC 22C3 phamaDx assay, we found that adenocarcinoma had a similar PD-L1 expression level to squamous cell carcinoma but significant higher expression than NSCLC, NOS. The percentage of high PD-L1 expression (TPS³50%) was significantly higher in mediastinal lymph nodes and distant metastatic sites than in primary sites, which suggests that PD-L1 testing of metastatic NSCLC could identify more patients eligible for immunotherapy.

1980 Is STAS Assessment on Frozen Sections Reliable?

Fang Zhou¹, Julian Villalba², Treah May Sayo³, Navneet Narula⁴, Mari Mino-Kenudson³, Andre Moreira⁵

¹NYU School of Medicine, New York, NY, ²Boston, MA, ³Massachusetts General Hospital, Boston, MA, ⁴NYU Langone Health, New York, NY, ⁵New York Langone Health, New York, NY

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Background: Spread through air spaces (STAS) has been reported to be associated with a worse prognosis in adenocarcinoma of lung. Recently it has been proposed that STAS be reported on frozen sections (FS) as an indication for more aggressive surgery (lobectomy vs sublobar resection). We undertook this study to evaluate the reliability of STAS assessment on FS compared to FS controls (FSC) and non-frozen remaining tumor (RT).

Design: Cases of adenocarcinoma that had FS of the tumor were identified retrospectively from two institutions. For each case, the following was recorded: presence(+)/absence(-) of STAS on FS, FSC, and RT; and % of tumor patterns: lepidic(L), acinar(A), papillary(P), micropapillary(M), solid(S).

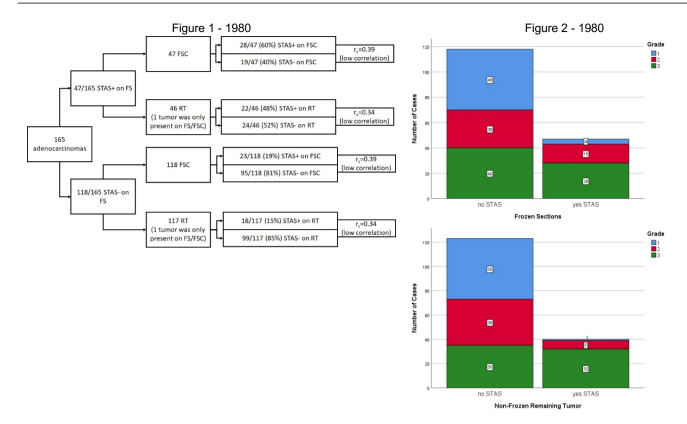
Grades were defined as follows: G1=L predominant with A/P G2=A/P predominant with <20% M/S G3=M/S predominant or ≥20% M/S

Cross-tabulations and Spearman's correlations (r_s) were performed in SPSS (see Table). r_s 0.3-<0.5 = low correlation.

Results: 165 cases were found. In 2 cases the tumor was only present on FS/FSC slides. STAS+ was present on FS in 47/165(28%), of which 28/47(60%) had STAS+ on FSC (r_s =0.39) and 22/46(48%) had STAS+ on RT (r_s =0.34) (1 tumor was only present on FS/FSC). Of the 40 STAS+ cases on RT, 18/40(45%) did not have STAS (STAS-) on FS (r_s =0.34). 118/165 of cases were STAS- on FS; of these, 18/117(15%) were STAS+ on RT (r_s =0.34) (1 tumor was only present on FS/FSC). Fig 1.

Of the 47 cases with STAS+ on FS, 4(15%) were G1, 15(32%) were G2, and 28(60%) were G3 (r_s =0.30). Of the 15 G2 cases with STAS+ on FS, 7 had 10% to <20% high grade pattern (M/S). Of the 28 G3 cases with STAS+ on FS, 13 had \geq 30% high grade pattern. Of the 40 cases with STAS+ on RT, 1(2.5%) case was G1, 7(18%) cases were G2, and 32(80%) cases were G3 (r_s =0.46). Fig 2.

Size of correlation			Interpretation			
coefficie	ent (r _s)					
0.5 to 1.0	0		Moderate to			
0.03 to <	0.5		Low correlat			
0 to <0.3			Negligible co			
	SC (r _s =0.39)					
			FS		Total	
			no STAS	yes STAS		
FSC	no STAS	Count	95	19	114	
		% within FS	80.5%	40.4%	69.1%	
	ves STAS	Count	23	28	51	
	,	% within FS	19.5%	59.6%	30.9%	
Total	I	Count	118	47	165	
		% within FS	100.0%	100.0%	100.0%	
FS vs R	T (r _s =0.34)	70				
			FS		Total	
			no STAS	yes STAS	1	
RT	no STAS	Count	99	24	123	
		% within FS	84.6%	52.2%	75.5%	
	yes STAS	Count	18	22	40	
	,== =	% within FS	15.4%	47.8%	24.5%	
Total	1	Count	117	46	163	
10.01		% within FS	100.0%	100.0%	100.0%	
FSC vs	RT (r _s =0.42)	70 17101111110	100.070		100.070	
	(13 0.72)		FSC		Total	
			no STAS	yes STAS	- I Jiai	
RT	no STAS	Count	98	25	123	
IXI	110 3173	% within FSC	87.5%	49.0%	75.5%	
	yes STAS	Count	14	26	40	
	yes STAS	% within FSC	12.5%	51.0%	24.5%	
Total		Count	112.576	51.0 %	163	
		% within FSC	100.0%	100.0%	100.0%	
Grade v	Grade vs FS (r _s =0.30)			100.076	100.076	
Grade V	3 1 3 (1 _s -0.30)		grade			Total
			G1	G2	G3	Total
Frozen	no STAS	Count	48	30	40	118
section	110 0170	% within grade	92.3%	66.7%	58.8%	71.5%
36611011	yes STAS	Count	92.576	15	28	47
	yes o i Ao	% within grade	7.7%	33.3%	41.2%	28.5%
Total	1	Count	52	45	68	165
iolai		% within grade	100.0%	100.0%	100.0%	100.0%
Grade v	s FSC (r _s =0.39		100.0% 100.0% 100.0%			100.070
Sidde V	5 . 55 (1s-0.55	'I	grade	Total		
			G1	G2	G3	
FSC	no STAS	Count	49	30	35	114
. 55		% within grade	94.2%	66.7%	51.5%	69.1%
	yes STAS	Count	3	15	33	51
	300 OTAO	% within grade	5.8%	33.3%	48.5%	30.9%
Total	l	Count	52			165
% within grade		100.0%	100.0%	100.0%	100.0%	
Grade v	s RT (r _s =0.46)	70 Within grade	100.0 /6	100.076	100.070	100.070
J. 340 V			grade			Total
			G1	G2	G3	
RT	no STAS	Count	50	38	35	123
13.1	100170	% within grade	98.0%	84.4%	52.2%	75.5%
	yes STAS	Count	1	7	32.270	40
	700 0170	% within grade	2.0%	15.6%	47.8%	24.5%
Total	<u> </u>	Count	51	45	67	163
iolai		% within grade	100.0%	100.0%	100.0%	100.0%
		70 WILLIAM GLAUE	100.070	100.070	100.070	100.070



Conclusions: The correlations among FS, FSC, and RT are low. FS STAS+ cases remain STAS+ only in 60% of FSC and 48% of RT. STAS shows higher correlation with grade on RT than it does on FS. These results show a lack of reliability in the assessment of STAS on FS, and do not support the proposal of reporting STAS in FS to make intraoperative clinical decisions, as doing so may subject patients to unnecessarily aggressive surgery.